## Atom-at-a-time resonance ionization spectroscopy of nobelium

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Experimental study of atomic spectra traditionally required optical spectroscopy of a primordial isotope for each element, providing a benchmark in understanding the atomic structure and revealing how relativistic effects increasingly impact the binding energy of orbital electrons of the heaviest elements. Such studies have been conducted for most elements<sup>1</sup> and theoretical modelling can be performed to high precision<sup>2,3</sup>. Today, no tabulated spectral lines exist for the transfermium el-10 ements of an atomic number Z>100. These ra-11 dioactive elements are produced in nuclear fusion 12 reactions at rates of a few atoms per second at 13 most and must be studied 'live' immediately fol-14 lowing their production, which so far precluded 15 their optical spectroscopy. Here, we report on ra-16 dioactive decay-detected resonance ionization spec-17 troscopy on the element nobelium (No), the 102nd 18 element in the periodic table, in which we identi-19 fied the ground-state transition  ${}^{1}S_{0} \rightarrow {}^{1}P_{1}$ . Com-20 bined with data from an observed Rydberg series, 21 an upper-limit for the ionization potential (IP) is 22 obtained. These accurate results provide a partic-23 ular challenge for state-of-the-art relativistic many-24 body calculations 5,6,7,8 addressing quantum electro-25 dynamic effects. The present work opens the door 26 for high-precision measurements of various atomic 27 and nuclear properties also of elements heavier than 28 nobelium and motivates future theoretical work. 29

Since the establishment of the actinide elements in the 30 periodic table, great efforts have been undertaken to in-31 vestigate their atomic spectra<sup>9</sup>. The prevailing strategy 32 included deducing many of the atomic properties from a 33 detailed knowledge of the electronic configuration, which 34 is subject to strong relativistic effects. This approach was 35 driving large optical spectroscopy research programs for 36 many years, which yielded detailed insight into the atomic 37 structure of elements up to einsteinium<sup>1</sup> including precise 38 values for their IP. The heaviest element for which optical 30 spectroscopy has hitherto been reported is fermium with 40 Z = 100 (ref. 10). A sample of about 10 pg of <sup>255</sup>Fm with a 41

half-life of  $T_{1/2} = 20.1$  h facilitated resonance laser ionization spectroscopy experiments<sup>11,12</sup>. The transfermium elements, in contrast, neither occur naturally on earth nor can macroscopic samples be synthesized. These elements are typically produced at large accelerator facilities by heavyion induced fusion-evaporation reactions, which only yield low rates. Some of their atomic properties had been accessible in aqueous-phase and gas-phase chemical studies<sup>13,14</sup>. Only recently, the first ionization potential was experimentally determined for lawrencium (Z = 103, Lr) applying surface ionization techniques<sup>15</sup>.

Advancing optical studies to the region of the transfermium elements calls for fast spectroscopy techniques of extreme sensitivity<sup>4</sup>. Challenges in this research field include producing such atoms, separating them from undesired species, which are inevitably co-produced, manipulating and detecting them within seconds, to mention but a few. In our experiments we applied a RAdiation Detected Resonance Ionization Spectroscopy (RADRIS) technique  $^{16,17}$ , using a two-step photoionization process  $^{18}$  for ionizing nobelium atoms (Z = 102). Similar techniques had previously been employed for laser spectroscopy of shortlived americium fission isomers at production rates as low as ten per second<sup>19,20</sup>. For such an approach to become feasible for elements like nobelium, requires an extensive search for atomic transitions within a spectral range proposed by state-of-the-art model calculations. Unambiguous identification of the atoms is achieved via their unique radioactive decay fingerprint  $^{21}$ .

The isotope <sup>254</sup>No with a half-life  $T_{1/2} = 51.2 \,\mathrm{s}$  was chosen for the first laser spectroscopy experiments. This isotope was produced in the two-neutron evaporation channel of the complete-fusion reaction <sup>48</sup>Ca+<sup>208</sup>Pb. The fusion products, emerging from a thin <sup>208</sup>Pb target, were separated in-flight from the intense <sup>48</sup>Ca primary beam by the Separator for Heavy Ion reaction Products (SHIP)<sup>22</sup> at the GSI Helmholtzzentrum für Schwerionenforschung. About four <sup>254</sup>No ions per second were implanted in a buffer-gas stopping cell installed in the SHIP's focal plane<sup>4</sup>. They were thermalized in 95 mbar high-purity argon gas and accumulated and neutralized on a tantalum catcher filament (see

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Methods section). For a short time during every measure- 141 ment cycle, the incoming flux of <sup>254</sup>No ions was turned off 142 84 and the adsorbed nobelium atoms were evaporated from the 143 85 filament by heating this briefly to a temperature of about 1,350 K. Nobelium atoms in the ground state  $5f^{14}7s^{2}$  <sup>1</sup>S<sub>0</sub> <sup>145</sup> residing in laser beam paths in the vicinity of the filament 88 will undergo element-selective ionization in a two-step ex-80 citation scheme as shown in the inset of Fig. 1a. The ion-90 ization proceeded by resonantly exciting the singlet state 91  $5f^{14}7s7p^{1}P_{1}$  with ultraviolet (UV) light from a tunable dye 92 laser  $(\lambda_1)$ , followed by a second excitation into the contin-93 uum beyond the IP with UV light in the wavelength range  $349-353 \,\mathrm{nm} \,(\lambda_2)$  from a more powerful excimer laser. In-95 duced photoions are subsequently guided by electrostatic QF fields to a silicon detector where the characteristic alpha 97 decay of  $^{254}$ No is detected  $^{23}$ . 98

A spectral range from 28,887 to  $33,191 \,\mathrm{cm}^{-1}$  was 99 chosen to locate the  ${}^{1}S_{0} \rightarrow {}^{1}P_{1}$  atomic transition 100 in nobelium (Methods). Latest predictions based on 101 Multi-Configuration-Dirac-Fock (MCDF) and Relativistic-102 Coupled-Cluster calculations <sup>5,6,7,8</sup> suggest wavenumber val-103 ues clustering between 29,256 and 31,709 cm<sup>-1</sup> (Table 1a) 104 for the excited state  ${}^{1}P_{1}$ . The sufficient treatment of cor-105 relation effects by the different approaches decisively im-106 proved the picture given by the first MCDF calculations  $^{24}$ 107 focusing on quantum electrodynamic effects. Nonetheless, 108 about 6,110 laser scan steps needed to be conducted be-109 fore discovering this elusive first-step resonance. In Fig. 1a 110 (full triangles) we show the observed resonance in terms of 111 normalized alpha-decay count rates. A relatively high laser 112 power was initially used, which was favorable for an efficient 113 level search leading to a power-broadened transition. It is 114 centered at a wavenumber of  $\overline{\nu}_1'\,=\,29,961.457^{+0.041}_{-0.007}\,{\rm cm}^{-1}$ 115 as determined from a subsequent narrowband scan using an 116 intra-cavity etalon, see Fig. 1a (bullets). The dominant part 117 in the quoted uncertainties originates from the pressure shift 118 (Extended Data Table 1) resulting in a 1.4-parts-per-million 119 relative precision, which is far more precise than from mod-120 ern theories, and hence provides a powerful benchmark. Ta-121 ble 1a gives a comparison of the experimental result for the 122 wavenumber of the  ${}^{1}P_{1}$  state with theoretical predictions. 123

Having located the resonance in <sup>254</sup>No the spectral reso-124 lution was increased in order to measure the isotope depen-125 dence of the  $^1\mathrm{S}_0 \rightarrow \ ^1\mathrm{P}_1$  transition energy by reducing the 126 laser intensity. In Fig. 1b we show the measurement on the 127 same transition in  ${}^{252}$ No (T<sub>1/2</sub> = 2.4 s), produced under similar conditions using a  ${}^{206}$ Pb target. Successful spec-128 129 troscopy was achieved with implantation rates lower than 130 one atom per second delivered to the buffer-gas stopping cell 131 indicating the high efficiency of our setup (Methods). We 132 observed a signal displacement, the isotope shift, of about 133  $0.32 \,\mathrm{cm}^{-1}$ , which is due to a difference in nuclear size and 134 shape and will be published in a separate paper focussing 135 on the nuclear structure aspects of our work. 136

To further confirm the identification of the  ${}^{1}P_{1}$  state, 137 the saturation characteristics of the first-step resonance 138 has been measured (Fig. 2). Saturation was observed at 139 rather low photon fluxes, which is a clear indication for a 140

sizable dipole-transition amplitude. We obtained a corresponding Einstein coefficient for spontaneous emission of  $\dot{A}_{ki} = 4.2^{+2.6}_{-2.8} \times 10^8 \,\mathrm{s}^{-1}$  by fitting a rate-equation model (P.C., H.B. et al., manuscript in preparation) to the saturation data. This value is in agreement with various theoretical predictions (Table 1a) supporting that the short-lived  $5f^{14}7s7p^{1}P_{1}$  atomic state was observed.

We also observed several high-lying Rydberg states in  $^{254}$ No (Fig. 3a-c). The first excitation step was set to the resonance at  $\overline{\nu}'_1 = 29,961.457 \,\mathrm{cm}^{-1}$ , while the second step was scanned with a dye laser in the range from  $23,460 \,\mathrm{cm}^{-1}$ to  $23,503 \,\mathrm{cm}^{-1}$ . Two pronounced peaks, potentially members of the same Rydberg series, were observed. Sampling in an extended scan range revealed further peaks belonging to the same series, which so far could not be assigned unambiguously.



Figure 1: For caption see end of main text.



Figure 2: For caption see end of main text.

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Table 1:	For	caption	see	end	of	main	text.	

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<b>a</b> Atomic state $5f^{14}7s7p^{1}P_{1}$ in <sup>254</sup> No					
Method	$\overline{\nu}_1' \; (\mathrm{cm}^{-1})$	$A_{ki} (s^{-1}) \times 10^8$			
Experiment	$29,961.457^{+0.041}_{-0.007}$	$4.2^{+2.6}_{-2.8}$			
CI+all-order <sup>8</sup> <sup>#</sup>	$30,203 \pm 600$	-2.0			
MCDF <sup>7</sup>	$30,609 \pm 1100$ *	3.5			
$\rm IHFSCC^{6}$ §	$30,056\pm800$	5.0			
MCDF <sup>5</sup>	$30,650 \pm 800$	2.7			
<b>b</b> Ionization potential					
Method	$IP (cm^{-1})$				
Experiment	$52,467^{**} \le IP \le 53,757.5$				
CI+all-order <sup>8 <math>\sharp</math></sup>	$54,390 \pm 1100$				
MCDF <sup>7</sup> <sup>b</sup>	$53,701 \pm 1100$ ***				
IHFSCC <sup>6</sup> §	$53,489 \pm 800$ ***				
Extrapolation $^{25}$	$53,600\pm600$				

<sup>#</sup> Configuration interaction method combined with the linearized single-double coupled cluster method (all-order).

Multi-configuration Dirac-Fock.

 Intermediate Hamiltonian Fock-space coupled cluster.
 \* As not explicitly reported in the original paper, we assessed the error from the relative difference of the calculated and measured values for the corresponding state in ytterbium.
 \* The lower that the first original content of the corresponding state in ytterbium.

\*\* The lower limit of the IP has been estimated from theoretical calculations  $^6$ , for details see text. \*\*\* We give a conservative error, the magnitude of which is based on that

we give a constructive error, the magnitude of which is based on that of the error of the  $^1{\rm P}_1$  state wavenumber evaluated with the corresponding method (Table 1a).

In the absence of perturbations originating from the interaction with buffer gas atoms, the transition energies 158 from the excited <sup>1</sup>P<sub>1</sub> state to Rydberg states  $h\nu_2(n)$  follow 159 a trend described by the well-known Rydberg formula<sup>18</sup> 160  $h\nu_2(n) = h\nu_{lim} - R_m/(n-\delta)^2$ . Here,  $h\nu_{lim} = IP - h\nu'_1$ 161 denotes the ionization limit for the excited state, n, the 162 principal quantum number of the valence electron,  $R_m$ , the 163 reduced-mass Rydberg constant for  $^{254}\mathrm{No},$  and  $\delta,$  the quan-164 tum defect. In the first-order Ritz expansion  $^{26}$  the quan-165 tum defect can be expressed as  $\delta(n) = \delta_0 + B/(n - \delta_0)^2$ 166 with the fitting parameters  $\delta_0$  and B. In the upper panel 167 of Fig. 3d, the positions of the observed peaks are plotted 168 against their principal quantum number n. The n assign-169 ment was based on restricting the quantum defect to be 170 in the range between 0 and 1, i.e., arbitrary, as the out-171 come of the analysis remained unaffected by this choice. 172 The best fit to the data showed a convergence towards 173 the value  $\overline{\nu}_{lim} = 23,785^{+11}_{-1} \text{ cm}^{-1}$ . From higher-order corrections in the quantum defect, we estimated a maximum 174 175 systematic error of  $+10 \,\mathrm{cm}^{-1}$  for the given value. Shifts 176 in the atomic spectra due to the buffer gas pressure were 177 neglected as these are expected not to exceed  $0.5 \,\mathrm{cm}^{-1}$ . 178 However, experimental observations when delaying the non-179 resonant ionizing laser pulse, suggest a fast quenching of 180 the <sup>1</sup>P<sub>1</sub> state, induced by buffer-gas collisions, into a poten-181 tially long-lived but energetically very close atomic state. 182 Hence, the observed series is not necessarily excited from 183 the  ${}^{1}P_{1}$  state. Possible candidates are a  ${}^{3}D_{3}$  or even a 184  ${}^{3}\text{D}_{2}$  state, located  $159\,\text{cm}^{-1}$  and  $1,278\,\text{cm}^{-1}$  below the  ${}^{1}\text{P}_{1}$ 185 state according to ref. 6, respectively. We thus include 186 a lower limit for the ionization potential of nobelium as 187  $\overline{\nu}_{lim} + \overline{\nu}'_1 - 1,278 \,\mathrm{cm}^{-1} = 52,467 \,\mathrm{cm}^{-1}$  corresponding to  $6.505 \,\mathrm{eV}$ . The upper limit is  $\overline{\nu}_{lim} + \overline{\nu}'_1 = 53,757.5 \,\mathrm{cm}^{-1}$ 188 189 (6.665 eV). This is because any populated state energeti-190 cally lying above the  ${}^{1}P_{1}$  state would immediately depop-191 ulate to the then lower-lying  ${}^{1}P_{1}$  and cannot lead in this 192

context to delayed resonance ionization. In Table 1b various theoretical predictions are compared with our experimental value for the IP. This value continues the trend of increasing IP along the heaviest actinides<sup>1,27</sup> and is significantly higher than the one recently reported for the heavier element lawrencium<sup>15</sup>. In analogy to the lanthanides, the lighter homologues, this corroborates the scenario of closed 5f and 7s atomic-shells in nobelium.

In summary, laser spectroscopy of the element nobelium was successfully performed using the ultra-sensitive and highly efficient RADRIS technique. This marks a basis for future experiments, in which atomic-level energies of nobelium including the first IP can be determined with unprecedented precision. The new data provide anchor points for future theoretical work, which in turn forms an indispensable guide for such experiments. This work opens up a new horizon for laser spectroscopy in the elusive region of the heaviest elements, including those beyond nobelium, accessible with even lower yield, with different atomic structure. An example is the heaviest actinide element lawrencium, which is now within reach of such studies. Moreover, isotope shift and hyperfine structure measurements  $^{28}$ on nobelium isotopes are now feasible, forming a valuable contribution to nuclear structure studies in the region of deformed nuclei on the shore of the 'superheavy elements', elements which only exist due to nuclear shell structure. These sophisticated alternatives to established in-beam and decay spectroscopy approaches<sup>29</sup> provide complementary information on both single-particle and collective properties, alongside spin assignments, and will critically test and revise state-of-the-art nuclear models.



Figure 3: For caption see end of main text.

1. Kramida, A., Ralchenko, Y., Reader, J. & NIST- 264 Team. NIST Atomic Spectra Database version 5.3 (2016); 265 http://physics.nist.gov/asd (accessed March 2016). 266

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- 2. Eliav, E., Fritzsche, S. & Kaldor, U. Electronic structure the-267 ory of the superheavy elements. Nucl. Phys. A 944 518-550 268 (2015).269
- 3. Schwerdtfeger, P., Pašteka, L.F., Punnett, A. & Bow-270 man, P.O. Relativistic and quantum electrodynamic effects 271 in superheavy elements. Nucl. Phys. A 944 551-577 (2015). 272
- Backe, H., Lauth, W., Block, M. & Laatiaoui, M. Prospects 4. for laser spectroscopy, ion chemistry and mobility measurements of superheavy elements in buffer-gas traps. Nucl. Phys. A 944 492-517 (2015).
- 5. Indelicato, P., Santos, J.P., Boucard, S. & Desclaux, J.-P. QED and relativistic corrections in superheavy elements. Eur. Phys. J. D 45, 155-170 (2007).
- 6. Borschevsky, A. et al. Predicted spectrum of atomic nobelium. Phys. Rev. A 75, 042514 (2007).
- 7. Liu, Y., Hutton, R. & Zou, Y. Atomic structure of the superheavy element No I (Z = 102). Phys. Rev. A 76, 062503 (2007).
- 8. Dzuba, V.A., Safronova, M.S. & Safronova, U.I. Atomic 245 285 properties of superheavy elements No, Lr, and Rf. Phys. 246 Rev. A 90, 012504 (2014). 247
- Worden, E.F., Blaise, J., Fred, M., Trautmann, N. & 9. 248 Wyart, J.-F. Spectra and Electronic Structures of Free Ac-249 tinide Atoms and Ions. In The Chemistry of the Actinide and 250 Trans-actinide Elements (eds Morss, L.R., Edelstein, M., 251 Fuger, J.) vol. 3, Ch. 16.4 (Springer, 2008). 252
- 10. Sewtz, M. et al. First Observation of Atomic Levels for the 293 253 Element Fermium (Z = 100). Phys. Rev. Lett. **90**, 163002 294 254 (2003).255
- 11. Sewtz, M. et al. Resonance ionization spectroscopy of fer-256 mium (Z = 100). Spectrochim. Acta B 58, 1077-1082 (2003). 257
- 12. Backe, H. et al. Laser Spectroscopic Investigation of the El-298 258 ement Fermium (Z = 100). Hyperfine Interact. 162, 3-14 200 259 (2005).300 260
- 13. Nagame, Y., Kratz, J.V. & Schädel, M. Chemical studies 301 261 of elements with  $Z \ge 104$  in liquid phase. Nucl. Phys. A 944 262 302 614-639 (2015). 303 263

- 14. Türler, A., Eichler, R. & Yakushev, A. Chemical studies of elements with Z>104 in gas phase. Nucl. Phys. A 944 640-689 (2015).
- 15. Sato, T.K. et al. Measurement of the first ionization potential of lawrencium, element 103. Nature 520, 209-211 (2015).
- 16. Lauth, W. et al. Resonance Ionization Spectroscopy in a Buffer Gas Cell with Radioactive Decay Detection, Demonstrated Using <sup>208</sup>Tl. Phys. Rev. Lett. 68, 1675-1678 (1992).
- 17. Backe, H. et al. Towards optical spectroscopy of the element nobelium (Z = 102) in a buffer gas cell. Eur. Phys. J. D 45, 99-106 (2007).
- 18. Letokhov, V.S. Laser Photoionization Spectroscopy (Academic Press, 1987).
- Backe, H. et al. Isotope Shift Measurements for Superde-19. formed Fission Isomeric States. Phys. Rev. Lett. 80, 920-923 (1998).
- 20. Backe, H. et al. Stability of Superdeformation for Americium Fission Isomers as Function of the Neutron Number. Nucl. Phys. A 690, 215c-218c (2001).
- 21. Laatiaoui, M. et al. On laser spectroscopy of the element nobelium (Z = 102). Eur. Phys. J. D 68, 71-77 (2014).
- 22.Hofmann, S. & Münzenberg, G. Discovery of the heaviest elements. Rev. Mod. Phys. 72, 733-767 (2000).
- 23.Lautenschläger, F. et al. Laser spectroscopy of the heaviest elements at SHIPTRAP. Nucl. Instrum. Methods B (2016), accepted.
- Fritzsche, S. On the accuracy of valence-shell computations 24.for heavy and super-heavy elements. Eur. Phys. J. D 33, 15-21 (2005).
- 25. Sugar, J. Revised ionization energies of the neutral actinides. J. Chem. Phys. 60, 4103 (1974).
- Martin, W.C. Series formulas for the spectrum of atomic 26.sodium (Na I). J. Opt. Soc. Am. 70, 784-788 (1980).
- 27.Wendt, K., Gottwald, T., Mattolat, C. & Raeder, S. Ionization potentials of the lanthanides and actinides - towards atomic spectroscopy of super-heavy elements. Hyperfine Interact. 227, 55-67 (2014).
- 28.Campbell, P., Moore, I.D. & Pearson, M.R. Laser spectroscopy for nuclear structure physics. Prog. Part. Nucl. Phys. 86, 127-180 (2016).

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- Herzberg, R.-D. et al. Nuclear isomers in superheavy ele ments as stepping stones towards the island of stability. Na *ture* 442, 896-899 (2006).
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the experiments. F.L., P.C., H.B., S.R. and M.L. analyzed the data. M.L. wrote the manuscript with input from all authors.

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 <sup>328</sup> on the online version of the paper. Correspondence and requests

on the online version of the paper. Correspondence and requests for materials should be addressed to M.L.(M.Laatiaoui@gsi.de). Figure 1: Resonance ionization signals of nobelium atoms. a, Scans over the first-step resonance in <sup>254</sup>No. The count rates are normalized to a beam current of one particle microampere, equivalent to  $6.2 \times 10^{12}$  <sup>48</sup>Ca projectiles per second. Best fits to the data on the basis of a rateequation model are indicated by solid lines. Full triangles:  $0.80 \text{ cm}^{-1}$  fit profile full width at half-maximum (FWHM); photon flux:  $1.1 \times 10^{14}$  photons/pulse/cm<sup>2</sup>; laser bandwidth:  $0.18 \text{ cm}^{-1}$ ; bullets:  $0.13 \text{ cm}^{-1}$  fit profile FWHM; photon flux:  $5.2 \times 10^{12}$  photons/pulse/cm<sup>2</sup>; laser bandwidth:  $0.04 \text{ cm}^{-1}$ . Inset: a simplified ionization scheme. b, Same as a, but for <sup>252</sup>No.  $0.36 \text{ cm}^{-1}$  fit profile FWHM; photon flux:  $1.1 \times 10^{13}$  photons/pulse/cm<sup>2</sup>; laser bandwidth:  $0.18 \text{ cm}^{-1}$ . Error bars,  $\pm 1 \text{ s.d.}$ 

Figure 2: Saturation characteristics of the first-step resonance for  $^{254}$ No. Alpha-decay count rates at the resonance peak, normalized to the maximum value, are displayed versus the photon flux of the first excitation step. Laser bandwidth:  $0.18 \text{ cm}^{-1}$ . The flux for the second step was kept at  $7.3 \times 10^{15}$  photons/pulse/cm<sup>2</sup> during this measurement. A best-fitting to the data according to a rate-equation model is also shown (solid line). All error bars indicate  $\pm 1$  s.d.

Table 1: Experimental and theoretical values for the  ${}^{1}P_{1}$  state and the ionization potential of nobelium.

Figure 3: Observed high-lying Rydberg states in nobelium. a-c, <sup>254</sup>No alpha-decay count rates, normalized to the maximum value, versus the excitation-energy equivalent wavenumber for the second-step excitation. First-step excitation:  $\overline{\nu}'_1 = 29,961.457 \,\mathrm{cm}^{-1}$ . Two selected Rydberg resonances with Lorentzian-profile fits (solid lines) to the data are shown in **a** and **b** and indicated by corresponding asterisks in **c**. **d**, Top panel: the position of 7 high-lying Rydberg states (bullets) as a function of *n* and a corresponding best fit (solid line) according to the Rydberg-Ritz formula. The residuals of the fit (full triangles) are shown in the bottom panel. Error bars,  $\pm 1$  s.d.

Extended Data Table 1: Uncertainties for the value of the  $^{254}$ No first-step resonance. The peak-position in the narrowband-scan data is extracted from a best fit based on a rate-equation model. The value for the wavelength measurement represents the accuracy of the wavelength meter in multimode-fiber operation. A conservative value for the pressure shift is taken from ref. 16.

Extended Data Figure 1: **Principle of the RADRIS technique.** Laser spectroscopy on radionuclides after their production and transmission through the velocity filter SHIP<sup>22</sup>. a, Thermalization of the fusion products in the buffer gas; b, accumulation on the catcher filament; c, reevaporation from the filament; d, two-step photoionization of neutral atoms; e, accumulation of re-ionized fusion products on the PIPS detector; f, decay detection.

## Methods

Production of <sup>252</sup>No and <sup>254</sup>No. The experiments described in this paper were carried out behind the velocity filter SHIP  $^{\rm 22}$ 332 at the linear accelerator (UNILAC) of GSI Helmholtzzen-333 trum für Schwerionenforschung in Darmstadt. The isotopes 334 <sup>252,254</sup>No were produced in the complete-fusion evaporation re-335 actions  $^{206}\mathrm{Pb}(^{48}\mathrm{Ca},\!2n)^{252}\mathrm{No}$  and  $^{208}\mathrm{Pb}(^{48}\mathrm{Ca},\!2n)^{254}\mathrm{No}$  at UNI-336 LAC beam energies of  $217\,{\rm MeV}$  with cross sections of  $515^{+80}_{-47}\,{\rm nb}$ 337 and  $2050^{+460}_{-340}$  nb (ref. 30), using lead sulfide (PbS) targets<sup>31</sup> 338 of  $460 \,\mu g/cm^2$  and  $470 \,\mu g/cm^2$  average thickness, respectively, 339 which remained constant throughout the measurements. The 340 <sup>48</sup>Ca<sup>10+</sup> beam from the UNILAC exhibited a macro-pulse struc-341 ture of 5 ms beam-on and 15 ms beam-off. Beam currents were 342 typically 0.7 particle microampere (about  $4.4 \times 10^{12}$  particles 343 per second). For laser spectroscopy experiments, the beam was 344 further chopped in accordance with user-defined measurement 345 cycles. In the case of  $^{254}$ No, the implantation rate of the fusion 346 products delivered to the experiment was repeatedly checked by 347 a retractable position-sensitive 16-strip silicon detector placed at 348 the focal plane of SHIP. 349

Radiation Detected Resonance Ionization Spectroscopy. 350 The fusion-evaporation residues with a mean kinetic energy of 351 about 41 MeV (ref. 23) were separated from the <sup>48</sup>Ca primary 352 beam by the velocity filter SHIP and subsequently thermal-353 ized inside the optical cell, a buffer-gas stopping cell filled with 354 95 mbar argon of ultrahigh purity (99.9999%). The cell was sep-355 arated from the vacuum of the SHIP by a 3.5- $\mu$ m thick entrance 356 foil (Mylar) on a support grid, see Extended Data Fig. 1. The 357 geometrical transparency of the grid amounts to 90.3%. A sub-358 stantial fraction of the stopped fusion products remained in a 359 positive charged state and was collected during the accumula-360 tion time of every measurement cycle onto a catcher filament, a 361 tantalum wire of  $125\,\mu\mathrm{m}$  diameter. The filament was heated for 362 300 ms to a temperature of about 1,350 K, triggering the evap-363 oration of neutral nobelium atoms for subsequent two-step laser 364 ionization<sup>21</sup>. Ions produced during this process were promptly 365 guided by suitable electric fields to a particle detector, a Passi-366 vated Implanted Planar Silicon semiconductor detector (PIPS), 367 with which the characteristic alpha-decay or spontaneous fission 368 of the studied nobelium isotopes and their descendants was reg-369 istered <sup>17,21,23</sup>. The optimum accumulation time depends on the 370 half-life of the isotope under consideration and was 25 s (ref. 32) 371 in case of  $^{254}$ No with  $T_{1/2} = 51.2 \pm 0.4 \,s$  (ref. 29). The two-step 372 laser ionization took place during a 5-s time window every cycle, 373 while the primary ion beam was switched off. Correspondingly, 374 3-s-beam-on and 3-s-beam-off periods were chosen in the case 375 of the short-lived isotope<sup>33 252</sup>No ( $T_{1/2} = 2.42 \pm 0.06 \,\mathrm{s}$ ). We 376 further introduced 'waiting' cycles by interrupting the data ac-377 quisition while changing the laser frequencies. In Fig. 1a, for 378 instance, a 'waiting' cycle of 5 minutes was chosen, necessary es-379 pecially in the case of  ${}^{254}$ No, in order to minimize residual  ${}^{254}$ No 380 alpha-decay events that might lead to counts not belonging to 381 the chosen laser frequency. The absolute temperature of the fila-382 ment was monitored using a fast infrared pyrometer (LumaSense 383 Inc., IMPAC IS 6 Advanced). The temperature was adjusted 384 to the nominal value when necessary in order to prevent over-385 448 heating, which was observed to lead to increased background in 386 preparatory experiments. 387

Laser setup for nobelium spectroscopy. The extended level 388 search in nobelium was carried out using four tunable excimer 389 laser-pumped dye lasers (Lambda Physik, FL and LPD series, 390 bandwidth  $(5.5 \pm 0.5)$  GHz) and an Optical Parametric Oscil-391 lator (OPO) system (GWU-Lasertechnik, VisIR2, bandwidth 455 392

 $\approx 90 \,\mathrm{GHz}$ ) pumped by a frequency-tripled Nd:YAG laser (Continuum, Powerlite DLS 8050). Except for the latter 50-Hz system, all the other pump lasers were operated at a repetition rate of 100 Hz. The laser pulse duration was at most 18 ns (FWHM) with a jitter  $< 11 \,\mathrm{ns}$  for all of them. The dye lasers were set up to scan in the range from  $25,000\,\mathrm{cm}^{-1}$  up to  $31,000\,\mathrm{cm}^{-1}$ During the level search in nobelium, however, the scans were conducted mainly in the UV region of the optical spectrum. With a mean step size of about  $0.89 \,\mathrm{cm}^{-1}$  we thereby covered more than twice the spectral range from  $28,887 \,\mathrm{cm}^{-1}$  to  $30,530 \,\mathrm{cm}^{-1}$ . With the OPO system, operated in a frequency-mixing mode, and by choosing a scan step size of  $3 \text{ cm}^{-1}$ , adapted to the laser bandwidth, multiple scans from 30,000 up to  $33,191 \text{ cm}^{-1}$  were conducted. The laser wavelengths were continuously monitored with a wavelength meter (HighFinesse-Ångstrom, WS/7-UVU) that was calibrated to an internal neon lamp. Laser pulse energies in excess of 0.1 mJ were repeatedly measured at the optical cell for all tunable lasers. For an efficient beam time usage they were operated simultaneously in different wavelength ranges. They were synchronized with excimer laser synchronization units (Lambda Physik, LPA 97) with respect to the ionizing laser, an excimer laser (Lambda Physik, LPX220) delivering an average pulse energy of 45 mJ of broadband laser light in the wavelength range 349-353 nm at the optical cell. The total photon energy available for ionization was by far higher than all theoretical predictions and extrapolations of the IP of nobelium (Table 1b). The scans near the IP were performed by replacing the excimer ionizing laser by a tunable dye laser scanning in the blue range of the optical spectrum. Here, only two dye lasers were operated simultaneously. In addition, the dye lasers enabled the use of intra-cavity etalons and thus a narrowing of the laser bandwidth down to 1.2 GHz. The narrow resonance shown in Fig. 1a (bullets) was recorded using this option. More details on the laser systems used can be found in ref. 23.

**RADRIS efficiency.** The overall efficiency of the setup is defined as the ratio of the nobelium decay count rate measured with the PIPS detector  $A_{RIS}$  at maximum of a resonance to the implantation rate of nobelium ions delivered to the optical cell  $A_{Ion}$ , both normalized to the intensity of the primary beam. The spatial distribution of the <sup>254</sup>No ions delivered to the experiment was best described by a 2D-Gaussian distribution with  $\sigma_x = 22 \,\mathrm{mm}$  and  $\sigma_y = 5.7 \,\mathrm{mm}$  based on a measurement with the position-sensitive 16-strip silicon detector exhibiting an active area of  $80\,\mathrm{mm}\times35\,\mathrm{mm}.$  The ion implantation rate was extracted from the alpha-decay count rate  $A_{\alpha}$  measured with this detector according to  $A_{Ion} = A_{\alpha}/(\varepsilon_{\alpha} \times \varepsilon_{\Omega} \times \varepsilon_{daq})$ , with  $\varepsilon_{\alpha} = 0.9$ , the alpha-decay probability for <sup>254</sup>No,  $\varepsilon_{\Omega} = 0.55$ , the solid angle coverage for alpha decays from implanted  $^{254}$ No recoils $^{23}$ and  $\varepsilon_{dag} = 0.77$ , the efficiency for data acquisition, which was limited to the beam-off time windows during which data recording occurred. On average, decay rates in the PIPS detector of  $0.39 \pm 0.05$  and  $0.048 \pm 0.006$  per second and particle microampere  $(6.2 \times 10^{12} {}^{48}$ Ca projectiles per second) were obtained for <sup>254</sup>No and <sup>252</sup>No, respectively. With these numbers an overall efficiency of the apparatus of  $(6.4 \pm 1)\%$  and  $(3.3 \pm 1)\%$  was calculated, respectively, proving a high-efficiency of the applied spectroscopy technique. The difference in the quoted numbers is mainly due to the half-lives of the isotopes under investigation. In the spectroscopy of <sup>252</sup>No a shortest possible beam-off period of 3s was applied. Even though the measurement cycle was optimized to minimize the impact of the half-life on the overall efficiency, the spectroscopy of  $^{252}$ No turned out to be less efficient compared with  $^{254}$ No.

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- 30. Oganessian, Yu.Ts. et al. Measurements of cross sections for the fusion-evaporation reactions <sup>204,206,207,208</sup>Pb+<sup>48</sup>Ca and <sup>207</sup>Pb+<sup>34</sup>S: Decay properties of the even-even nuclides <sup>238</sup>Cf and <sup>250</sup>No. Phys. Rev. C 64, 054606 (2001).
- 31. Kindler, B. et al. Chemical compound targets for SHIP on heated carbon backings. NIMA 561, 107-111 (2006).
- 32. Laatiaoui, M. et al. Perspectives for laser spectroscopy of the element nobelium. Hyperfine Interact. 227, 69-75 (2014).
- 33. Sulignano, B. et al. Identification of a K isomer in  $^{252}$ No. Eur. Phys. J. A 33, 327-331 (2007).

Extended Data Table 1: For caption see end of main text.

Origin	Uncertainty $(\text{cm}^{-1}) \times 10^{-3}$
Fit Wavelength measurements Pressure shift	$\pm 4.2 \\ \pm 5.0 \\ + 34.2$



Extended Data Figure 1: For caption see end of main text.