Chemical Experiments with Superheavy Elements

Andreas Türler*

Abstract: Unnoticed by many chemists, the Periodic Table of the Elements has been extended significantly in the last couple of years and the 7th period has very recently been completed with eka-Rn (element 118) currently being the heaviest element whose synthesis has been reported. These ‘superheavy’ elements (also called trans-actinides with atomic number ≥ 104 (Rf)) have been artificially synthesized in fusion reactions at accelerators in minute quantities of a few single atoms. In addition, all isotopes of the transactinide elements are radioactive and decay with rather short half-lives. Nevertheless, it has been possible in some cases to investigate experimentally chemical properties of transactinide elements and even synthesize simple compounds. The experimental investigation of superheavy elements is especially intriguing, since theoretical calculations predict significant deviations from periodic trends due to the influence of strong relativistic effects. In this contribution first experiments with hassium (Hs, atomic number 108), copernicium (Cn, atomic number 112) and element 114 (eka-Pb) are reviewed.

Keywords: Relativistic effects · Superheavy elements · Transactinides

1. Introduction

Where does the Periodic Table of the Elements end? What is the heaviest element in the Periodic System? Are there still undiscovered ones to be found in nature? Is there an 8th period and how many elements will it contain? Will we need to introduce the superactinides starting with atomic number 121? Will the current principles governing the groups and periods of the Periodic Table still be valid for the heaviest elements and what defines the place an element occupies in the Periodic System? These intricate questions are the topic of current research in nuclear chemistry. Fig. 1 shows a modern Periodic Table.

The nuclide $^{244}$Pu, with a radioactive half-life of 81.2 million years could still be present on earth as a primordial element or be of cosmic origin due to explosive stellar nucleosynthesis, for example, from a nearby supernova that occurred after the formation of our solar system. Indeed, a concentration of about 2500 atoms $^{244}$Pu per gram of Bastnaesite, a mineral highly enriched in rare earth elements, was reported nearly 40 years ago.[1] However, a recent search for $^{244}$Pu in Bastnaesite from same mine using accelerator mass spectrometry remained negative with a detection limit of less than 550 atoms per gram.[2]

Thus, all known elements heavier than Pu are man-made. The last element that can be produced in weighable quantities is fermium with atomic number 100. It can be produced by breeding in high-flux nuclear reactors and quantities of few nanograms have been isolated and supplied for chemical experiments, i.e. for laser spectroscopic measurements of the atomic level structure.[4] All elements heavier than fermium have so far been produced only at accelerators in heavy ion fusion reactions. Up to element lawrencium (Z = 103), the identification of a new element and the assignment of the atomic number were made based on chemical properties. The position at which a newly discovered radioactive species was eluted from a cation-exchange column in $\alpha$-hydroxysobutyratesolution was indicative of its ionic radius and thus its position within the group of actinides.[1]

The element with atomic number 101, mendeleievium (Md), was the first for which only single atoms could be produced and analyzed. In an experiment about 10$^7$ atoms of $^{257}$Es were bombarded with $\alpha$-particles (He$^{\alpha}$ ions) to synthesize single atoms of Md. These were subsequently identified as single atoms after chemical isolation by their characteristic radioactive decay.[5] This was the first of a long series of ‘one-atom-at-a-time’ chemistry experiments. In this case the classical mass action laws and the definitions of chemical equilibrium and partition coefficient are no longer valid. The question arose whether one single atom actually displays chemical properties and how typical these properties are compared to bulk properties and how they can be measured. It can be shown that partition coefficients can be defined for single atoms, if one replaces concentrations by probabilities to find a single atom in a given state, i.e. in the gas phase or adsorbed to a surface in gas-phase chemistry, or in the aqueous phase vs. the organic phase in a liquid–liquid extraction step.[6] As will be discussed later, it is indeed possible to perform chemical experiments with only one single atom or molecule at a time and to quantitatively measure thermochromatic properties, distribution coefficients, and so on. Such experiments need to be designed in such a manner that the atom under investigation has the possibility to probe its state repeatedly, such as in chromatography experiments.

After the discovery of lawrencium the quest for the synthesis of ever heavier elements shifted from chemical methods to nuclear physics, since the half-lives of the newly discovered nuclides became shorter and shorter with increasing atomic number. Also, the production rates were diminishing rapidly. Even the lightest transactinide element rutherfordium with atomic number 104 can be produced at best with a rate of 2 min$^{-1}$ of accelerator beam time. Since the discoverer of a new element traditionally has the right to propose its name, the quest for new elements turned into a fierce competition between scientists of the United States and Russia during the Cold War era. The recipe for synthesis of new ele-
He recovery of another six new elements at the explored, which ultimately led to the disractor systems, a new concept could be heavy ion accelerators and physical separations of both competitors was the same. The heaviest, most neutron-rich available elements was element 106 (seaborgium, Sg) in 1974\(^{[11]}\); this competition came to an end. The technological developments were exhausted and the available projectile and target combinations did not yield any detectable heavier element.

With the advent of more powerful heavy ion accelerators and physical separator systems, a new concept could be explored, which ultimately led to the discovery of another six new elements at the Gesellschaft für Schwerionenforschung (GSI), Darmstadt, Germany.\(^{[8]}\) If the doubly magic nucleus \(^{208}\)Pb, or nearby \(^{209}\)Bi are used as target materials, and these are bombarded with heavy projectiles such as \(^{54}\)Cr, \(^{58}\)Fe, or even \(^{70}\)Zn, the break-up of the magic proton and/or neutron shell requires so much energy that the fusion products can be obtained with a much lower excitation energy of only 15–20 MeV. Thus, only one or two neutrons need to be evaporated. Still, the fusion products of such reactions are relatively neutron-deficient, meaning rather short-lived. The formed fusion products were separated from the incoming beam and from abundant nucleon transfer- and fission products with the Separator for Heavy Ion reaction Products (SHIP) and detected in a segmented silicon detector. Two observations were of importance: i) the newly synthesized elements all decayed by the emission of one or several alpha-particles to already known lighter nuclei, thus pinpointing the mass number and also the atomic number of the synthesized nucleus, ii) none of the new nuclides decayed preferentially by spontaneous fission (SF) from the ground state. This implied that these nuclei were stabilized against fission by nuclear shell effects. All the isotopes of the newly discovered elements were very short-lived, with half-lives in the range of milliseconds, and only a few atoms were produced in month-long experiments. Prospect for experimental chemical investigatons of elements beyond Sg (Z = 106) were rather dim, with the exception of element 108 (hassium) where in the alpha-decay chain of the nuclide \(^{277}\)12 the isotope \(^{290}\)Hs with a half-life of about 10 s was observed. However, only two decay chains were observed in about 50 days of beam time. Very recently, the International Union of Pure and Applied Chemistry (IUPAC) has accepted the name of copernicium (Cn), suggested by the discoverers (IUPAC) has accepted the name of copernicium (Cn), suggested by the discoverers.

Once again, the possibilities to synthesize even heavier elements seemed to be exhausted. Attempts to synthesize element 113 in the reaction \(^{70}\)Zn + \(^{209}\)Bi at GSI were not successful. Nevertheless, a rather unexpected development was reported by the Florov Laboratory in Dubna, Russia. Instead of using a closed-shell target nucleus, they employed the doubly magic projectile \(^{40}\)Ca and heavy, neutron-rich actinide targets such as \(^{210}\)U or \(^{242,244}\)Pu to synthesize...
superheavy elements with atomic numbers 112 and 114 with larger production cross sections than the so-called ‘cold’ fusion reactions used at GSI. First results indicated that the nuclide $^{283}$Cn has a fantastic long half-life of about 5 min and decayed by SF, while one atom of $^{294}$114 lived longer than 30 s. Even though these results were published in renowned journals, these results could not be reproduced in independent experiments and it was feared that the group in Dubna had to face the same fiasco as the group in Berkeley, which was forced to retract an article about the alleged synthesis of element 118. In subsequent improved experiments the group in Dubna reported new decay data that was measured with a gas-filled magnetic separator system. In theses experiments several new nuclides were observed, among them $^{283}$Cn with a half-life of about 4 s and $^{294}$114 with a half-life of 2.6 s. Both nuclides decayed by alpha-particle emission and were formed with production cross sections that were 5 to 10 times higher than those observed in cold fusion reactions for Cn. So far, in two independent experiments the same decay characteristics for $^{283}$Cn were indeed observed: first in a chemistry experiment and shortly after in an experiment conducted at SHIP. Also, an experiment at Berkeley was successful in confirming decay data for the nuclides $^{284}$Cn, $^{286}$Cn, $^{288}$Cn, $^{290}$Cn, $^{292}$Cn, $^{294}$Cn and $^{296}$Cn. Unfortunately, all the decay chains ended with SF of a previously unknown nuclide, so that the connection to the known region of heavy nuclides could not be established. Also, a number of confirmation experiments have not yet been conducted due to the enormously long beam times required. Nevertheless, the community was absolutely stunned about this success. The relatively large production cross sections can only be explained by a larger survival probability of these superheavy nuclides compared to lighter ones. Indeed, already in the early sixties, the nuclear shell model predicted a so-called ‘island of stability’ at proton numbers between 114 and 126 (the exact location is still under debate) and neutron number 184. Although, the newly discovered nuclides in Dubna are more neutron deficient, it appears as if the shores of this fabled island have indeed been reached. Modern shell model calculations reproduce the observed decay properties of the heaviest nuclides rather accurately, predict the existence of heavy isotopes of elements 108 (Hs) or 110 (Ds) with half-lives approaching tens of years! Although it might not be possible to find these elements in nature, it might one day be possible to store such exotic atoms in traps and study their atomic properties in detail over an extended time period.

Is it indeed possible to study the chemical properties of single atoms or molecules and why is this interesting? As the case of the nuclear shell model demonstrated, it might be interesting to have a look at the answers that theoretical chemists give to predict the chemical properties of superheavy elements. Traditionally, the chemical properties of an unknown element and its compounds were predicted by exploiting the fundamental relationships of the physicochemical data of the elements and their compounds within the groups and the periods of the Periodic Table. However, with the increasing nuclear charges in heavy elements, the innermost electrons become more tightly bound and are thus moving with velocities which are a considerable fraction of the speed of light. Therefore they obey the laws of relativistic quantum mechanics. Due to relativistic rearrangements, which are predicted for the electronic structure of the heaviest elements, quite unexpected chemical properties are to be expected. These so-called relativistic effects are already severely influencing the chemical properties of lighter elements such as gold. The yellow color and the rather inert chemical behavior can be attributed to relativistic effects and explained by fully relativistic quantum chemical calculations. Early predictions of the chemical properties of superheavy elements were rather spectacular. One of the articles was entitled ‘Are elements 112, 114, and 118 relatively inert gases?’. Nowadays, with the use of powerful supercomputers the accuracy of quantum chemical calculations is increasing rapidly. Latest calculations come to the conclusion that element 112 (Cn) is not completely inert as expected for a noble gas, but should form a chemical bond with Au, which, however, is weaker than that of the lighter homolog Hg. Also, element 114 might be relatively inert but should interact stronger with an Au surface than Cn. Element 118 is predicted to be the first element in the row of noble gases with an electron affinity. Due to the considerable experimental difficulties in experimentally studying the chemical properties of superheavy elements, the gap between experimental observables and theoretical predictions is still large.

The knowledge and understanding of the chemistry of the superheavy elements, especially of the early transactinides Rf, Db, and Sg and their compounds, both experimentally and theoretically, is already quite extensive as documented by a first textbook entitled ‘The Chemistry of Superheavy Elements’ and several review articles (e.g. ref. [22]). While rapid, automated chemical separations in aqueous solution were (and are) the method of choice for detailed investigations of Rf and Db, the development of gas-phase chemical separation methods have allowed first chemical studies of heavier transactinide elements such as Sg, Bh, and Hs. The spectacular results obtained at弗洛罗夫实验室在Dubna使用$^{48}$Ca离子束和actinide targets to synthesize relatively long-lived isotopes of Cn and 114 give chemists the perspective to experimentally investigate even the heaviest known elements. Very recently an experiment to elucidate for the first time the chemical properties of Cn was successful. Here, latest results on chemical experiments with elements Hs, Cn, and 114 will be discussed.

**Experimental Techniques**

For chemical investigations, the element of interest has to be transported rapidly from the location of its production in a heavily shielded vault at the accelerator to a chemistry laboratory, or, the element has to be converted in situ to a chemical compound. Usually, the element of interest is transported attached to small aerosol particles in gas through a capillary to the chemistry laboratory. This transport can be accomplished within few seconds. Unfortunately, the transport yield is negatively affected by increasing beam intensities. The transport is unspecified, thus also undesired by-products of the nuclear reaction are transported. The harsh ionizing conditions created by the beam passing through the target recoil chamber severely limit the in situ synthesis of volatile transactinide compounds. In order to assess nuclides that are produced with few picobarn cross sections, the overall efficiency has to be as high as possible.

Luckily, chemists sometimes face very favorable and clear cut cases. Such a case constitutes the chemistry of Hs. If Hs has any resemblance to its lighter homolog Os, then it should also form tetroxides. Due to the perfect tetrahedral symmetry of OsO$_4$, this compound only interacts very weakly with non-reducing surfaces and is therefore very volatile, similar to a noble gas. In experiments with Hs, recoiling atoms were converted in situ with an admixture of O$_2$ gas to the volatile HsO$_4$. Another chemically very favorable case constitutes Cn as an expected homolog of Hg. Sufficiently volatile reaction products such as Hg and Cn can be transported directly with the flowing gas. It is well known that Hg interacts strongly with other metals such as Zn or Au. Single atoms of Hg can be quantitatively adsorbed on an Au surface by diffusion-controlled deposition from a stream of He. The isolated atoms (or
Hot topics to the cold or compact detector array registering the nuclear decay (room temperature and transported with the gas flow through a perfluoroalkoxy (PFA) capillary).

44±6°C for HsO4 and -82±7°C for OsO4. Solid lines (red and black) represent results of a simulation of the adsorption process with \( D_{\text{HSi3N4ads}} \) = 39 kJ/mol. The magenta line indicates the temperature profile (right-hand scale). The maxima of the deposition distributions were evaluated in situ (after two alpha decays) of the decay chain (including detection of a complete three-member \( \alpha \)-particle decay chain) is of the order of 55%. A similar set-up using gold-covered silicon detectors was used for the first successful chemical identification of Cn[13] and element 114.[24]

First Chemical Identification of Hassium and Copernicium

The first chemical identification of Hs[23] and Cn[13] signify the most important accomplishments of heavy element chemistry in past years. Even though only few single atoms can be obtained using the most favorable production reactions, an unambiguous identification of the nuclides 269Hs and 268Cn was achieved after chemical separation. In an experiment conducted in 2001 at GSI by an international collaboration using the IVO technique, a total of seven decay chains attributed to 269Hs were registered.[23] The registered decay sequences were in good agreement with the lower end (after two alpha decays) of the decay chains attributed in earlier experiments to the nuclide 277Cn. Thus, the chemistry experiments gave supporting evidence for the first identification of element 112 at the SHIP separator of GSI Darmstadt.[9]

In Fig. 3 the distribution of decay chains along the temperature gradient is shown in comparison with the adsorption peak of OsO4. Surprisingly, HsO4 is slightly less volatile than OsO4. Applying a microscopic model of the adsorption process first thermochemical properties of HsO4 could be deduced from only seven registered atoms! The standard adsorption enthalpy of HsO4 on a silicon nitride surface was determined as \(-\Delta H^{\text{ads}}\) (269HsO4) = 46 ± 2 kJ/mol.

The lower volatility of HsO4 compared to OsO4 is not due to relativistic effects. Adsorption of closed-shell spherically symmetric molecules with inert surfaces is governed by van der Waals forces, which can be calculated using a model of molecule–slab interaction. It can be shown that the molecule–surface interaction energy is determined by the relativistic (or non-relativistic) \( R_{\text{val}} \) of the valence \( d \) orbitals, which are changing in the same manner down group 8.[25]

In 2006 and 2007 a series of experiments performed at FLNR Dubna allowed the unambiguous detection of an isotope of Cn after chemical separation.[13] These experiments were conducted with the isotope 248Cn. It decays with a half-life of about 3.8 s by alpha-particle emission to 246Ds,
which undergoes SF with a half-life of about 200 ms. Larger production rates were observed by using the indirect production path $^{242}\text{Pu}(^{48}\text{Ca},3\text{n})^{287}\text{Hg}$, which was reported in an experiment at FLNR in 2007. In this experiment, the adiabatic potential of $^{114}\text{Hg}$ was measured to be $\Delta H_{\text{ads}} = 44 \text{ kJ mol}^{-1}$. In conclusion, it can be stated that Cn is forming metallic bonds with Au which are weaker than for the homolog Hg. But Cn is chemically not as inert as a noble gas, as was initially suggested.\(^{[18]}\)

Experiments with Element 114

The next heavier element currently under investigation is element 114, which would be placed as eka-Pb into group 14 of the Periodic Table. Suitable nuclides for chemical experiments are $^{287}\text{Hg}$ ($T_{1/2} = 0.5 \text{ s}$), $^{288}\text{Cn}$ ($T_{1/2} = 0.8 \text{ s}$) or $^{289}\text{Cn}$ ($T_{1/2} = 2.6 \text{ s}$) which can be synthesized in the reaction $^{44}\text{Ca} + ^{242}\text{Pu}$ or $^{244}\text{Pu}$. In a first publication, observation of three atoms of element 114 was reported in an experiment at FLNR similar to the ones performed for Cn.\(^{[24]}\)

From these three events an adsorption enthalpy of $\Delta H_{\text{ads}}(\text{E}114) = 34 \pm 20 \text{ kJ mol}^{-1}$ was deduced for the adsorption on an Au surface. If true, this result would be very puzzling, since eka-Pb is expected to be less reactive than Cn and should thus deposit at higher temperatures than Cn. Also, theoretical calculations which predicted the adsorption of Cn sufficiently accurately, predict $\Delta H_{\text{ads}}(\text{E}114) = 68 \text{ kJ mol}^{-1}$, somewhat less volatile than Cn. Due to the fact that a background made the positive identification of element 114 events more difficult than in the case of Cn, this first result was met with some skepticism.

In order to remove this background, the very successful chemical techniques developed so far are coupled to a physical preseparator, such as a gas-filled separator. At GSI Darmstadt the dedicated gas-filled separator TASCA (Trans Actinide Separator and Chemistry Apparatus) was constructed and successfully commissioned.\(^{[26]}\)
Recoiling fusion products are separated from the incoming beam and from transfer reaction products with a dipole magnet and focused on a relatively small product beam spot by two quadrupole magnets. The separator is operated as a gas-filled separator which allows charge focusing. The separated product beam is passed through a thin window supported by a honeycomb structure into a gas volume. The windows can stand pressures well above 1 atm. These investigations were and will remain challenging. Presently, mainly the production process has to be optimized. This requires substantial investments in new, much more powerful accelerators and the development of the corresponding target technologies, so that the very precious target materials can be safely irradiated. In addition, the chemical separation procedures have to become even faster. One possibility is the use of vacuum thermochromatography, which should allow the investigation of nuclides with half-lives as short as about 100 ms. It appears today that the experimental chemical investigation of the complete 7th row of the Periodic Table will not just remain a dream, but become reality within the next decades.

Received: March 17, 2010