Study of nuclear properties with muonic atoms

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Abstract Muons are a fascinating probe to study nuclear properties. Muonic atoms can easily be formed by stopping negative muons inside a material. The muon is subsequently captured by the nucleus and, due to its much higher mass compared to the electron, orbits the nucleus at very small distances. During this atomic capture process, the muon emits characteristic X-rays during its cascade down to the ground state. The energies of these X-rays reveal the muonic energy level scheme, from which properties like the nuclear charge radius or its quadrupole moment can be extracted. While almost all stable elements have been examined using muons, probing highly radioactive atoms has so far not been possible. The muX experiment has developed a technique based on transfer reaction inside a high-pressure hydrogen/deuterium gas cell to examine targets available only in microgram quantities.

1 Preface

This article is based on a lecture given at the 2019 summer school of the University of Pisa titled “Rewriting Nuclear Physics Textbooks: one more step forward” [1]. The intent of this write-up is to give a broad and general introduction to the topic (with plenty of references for the interested student to dig even deeper) and to give an example of the current progress of the field. Therefore, in terms of experiments, we focus almost completely on our own muX experiment, which marks the current edge of what is possible with muonic atom spectroscopy.

2 Introduction

Muons are fascinating particles with experiments being conducted in the context of particle, nuclear and atomic physics [2]. Additionally, also applied research is possible by measuring the spin precession and dynamics of muons inside materials through the $\mu$SR technique [3] thus probing the internal magnetic fields of the sample under study.

Muons are so-called leptons—such as electrons—and classified into the second family within the Standard Model of particle physics [4]. The properties of the muon have been extensively studied over the past 84 years since its discovery in 1936 [5,6]. Its lifetime [7].
for example, is the best measured lifetime of any unstable particle and searches for rare decays of the muon violating lepton family number have reached unprecedented sensitivity [8]. With a mass of 105.66 MeV/c², they are about 207 times heavier than an electron. Their lifetime of 2.197 μs is short, but still long enough in order to allow the large range of experiments alluded to above. Muons are also at the center of several current discrepancies between the Standard Model of particle physics and experimental results: (i) the discrepancy between the measured anomalous magnetic moment and its prediction [9], (ii) discrepancies in the rare decays of B mesons involving muons [10], and (iii) the discrepancy of the measured proton radius between the experiments using a muon as a probe compared to the ones involving electronic probes [11]. It is thus currently an exciting time to pursue experiments involving muons and who knows what new secrets these fascinating particles will reveal.

Muons are constantly being produced in the upper atmosphere through the interaction of cosmic rays with the remaining air. Due to their high energy, a large fraction of them reaches all the way to the ground and even penetrates deep into the rock [12,13]. As such these cosmic muons are part of the natural background. However, with a flux at sea level of around 1 muon/cm²/min the intensity is too low to perform precision experiments. For this reason, the interactions happening in the upper atmosphere are replicated through the bombardment of a target (typically a low-Z material such as graphite) with a proton beam from an accelerator. The muons are not produced directly, but follow from the decays of pions created in the following interactions of the primary proton with the target nucleons:

\[
\begin{align*}
p + p &\rightarrow p + n + \pi^+ \\
p + p &\rightarrow p + p + \pi^0 \\
p + p &\rightarrow d + \pi^+ \\
p + n &\rightarrow p + n + \pi^0 \\
p + n &\rightarrow p + p + \pi^- \\
p + n &\rightarrow n + n + \pi^+ \\
p + n &\rightarrow d + \pi^0
\end{align*}
\]

(1)

These are only the single pion production channels. For higher proton energies, higher multiplicity pion production is possible.

Figure 1 shows the HIPA proton accelerator complex at the Paul Scherrer Institute (PSI) in Switzerland [14] where the muX experiment described in Sect. 4 takes place. Its 590 MeV proton beam reaches a world-leading power of 1.4 MW. Two target stations are available for muon production serving a total of seven beam lines for particle physics and materials science. The highest intensities available at a low momentum of 28 MeV/c (very suitable for precision physics as these muons can easily be stopped) are \(5 \times 10^8 \mu^+/s\) and \(7 \times 10^6 \mu^-/s\) [15] many orders of magnitude more than what is available from cosmic rays.

Muonic atoms are atoms where a negative muon has been captured by the nucleus. Due to its 207 times higher mass compared to the electrons, the muon orbits the nucleus at a 207 times closer distance and experiences 207 times higher binding energies (neglecting in both cases the finite size of the nucleus). Some of the electrons are usually kicked out during the capture process, but are often very quickly refilled and are generally not perturbed by the presence of the muon (and vice versa). However, the negative muon effectively shields one of the protons such that for the electrons the nucleus of charge \(Z\) appears like a nucleus of charge \(Z - 1\). The details about the nucleus that can be extracted by this atomic capture and subsequent cascade down to the muonic ground state through a method called muonic atom spectroscopy is the topic of next Sect. 3. The muX experiment that was built to exploit this method and extend it to target materials that are only available in microgram quantities (such as highly radioactive targets) is described in Sect. 4.
This paper only describes the knowledge that can be obtained on the nucleus through the study of the muonic X-rays emitted in the process of the atomic capture. However, much more nuclear physics is in principle involved in the whole interaction of the negative muon with the nucleus over its complete lifetime [16]. Once the muon reaches the ground state, the overlap of the muonic wavefunction with the nucleus is so large—as a matter of fact for high-Z elements the muon spends most of its time within the nucleus—that the nuclear capture channel Eq. (3) becomes dominant over the ordinary decay of the muon Eq. (2).

\[
\begin{align*}
\mu^- & \rightarrow e^- \nu_\mu \bar{\nu}_e \\
\mu^- + p & \rightarrow n + \nu_\mu
\end{align*}
\]

As shown in Fig. 2, the total lifetime for negative muons in medium and high-Z elements is reduced to below 100 ns with only a few percent of the muons still undergoing ordinary muon decay. After this nuclear capture, the resulting daughter nucleus of charge \(Z - 1\) can be highly excited and most frequently returns to its ground state through the emission of neutrons and characteristic gamma rays. The study of these gamma rays can reveal details of the nuclear structure of the daughter nucleus—see, e.g., Ref. [17] where this technique has been employed to study nuclei relevant for neutrinoless double-beta decay.

### 3 Muonic atom spectroscopy

A muonic atom is an atom in which a negatively charged muon is captured by a nucleus. To form such an atom, a target material is bombarded with a negative muon beam. The muons suffer successive kinetic energy losses by their interaction with the outer atomic electrons in the target. When the muon is sufficiently slowed down to have similar velocity as the atomic electrons, it is attracted by the Coulomb field of the nucleus and forms a muonic atom. The capture is followed by a cascade of the muon to the ground state of the atom with the emission of Auger electrons and X-rays. Since the energy of the muonic X-rays is
determined by the exact details of the interaction of the muon with the nucleus, muonic atom spectroscopy is a sensitive tool to probe the ground-state properties of the nucleus.

Muonic atom spectroscopy is not the only tool to probe these fundamental properties of a nucleus. Two additional methods are the use of elastic electron scattering or optical spectroscopy [19]. While both electron scattering and muonic atom spectroscopy are able to measure absolute properties, optical spectroscopy generally can only deliver changes in these properties between isotopes due to the difficulty to calculate multi-electron systems with the required precision. However, a large amount of data has been collected by optical spectroscopy especially at radioactive ion beam facilities delivering information on long isotopic chains (see, e.g., Refs. [20,21]). Such data allow us to test and widen our current understanding of nuclear structure—and generally nuclear theory—by comparing the corresponding calculations of these properties to the measurements. Some recent examples showing the links between such measurements and nuclear theory can, for example, be found in Refs. [22–25].

3.1 Muonic cascade

The muon is captured at a high principal quantum number $n \sim 14$ [26]. The initial population of the levels is not well known but believed to approximately follow the $\sim (2l + 1)$ distribution (with $l$ being the quantum number of the orbital angular momentum). From this short-lived state, the muon starts its cascade down to the ground state ($1s$) of the atom. The energy of the muonic transitions at the beginning of the cascade, where the muonic levels are relatively dense, are small but still much larger than the typical binding energies of the electrons. Hence, the cascade of the muon initially proceeds through the release of Auger electrons. For lower transitions below $n \sim 6$ (where the energy gaps are much larger), the emission of muonic X-rays become the dominant de-excitation mechanism. These X-rays are characteristic for each element, and their energy varies from a few keV to a few MeV. A simplified scheme of the muon cascade is illustrated in Fig. 3. The most intense transitions are the $(n, l = n - 1) \rightarrow (n - 1, l = n - 2)$ transitions such as the $4f \rightarrow 3d$, $3d \rightarrow 2p$, $2p \rightarrow 1s$, etc. Figure 4
Fig. 3  Simplified scheme of the muonic atom levels for a point-like nucleus. The arrows indicate the transitions of the muon through the different atomic levels.

shows the energy spectrum of iron as measured with germanium detectors illustrating the muonic X-ray transitions. The whole cascade is very fast taking typically much less than a nanosecond. As described above, in the ground state of the muonic atom, the muon will disappear either via the free muon decay with a lifetime of $2.2 \times 10^{-6}$ s or through the nuclear capture process.

3.2 Muonic level scheme

As only one muon orbits the nucleus at the same time and the remaining electrons are far away from the muon orbits, the muonic atom is essentially a hydrogen-like atom and the calculation of the muonic energy levels follows the steps found in many textbooks on the hydrogen atom [27–29]. The gross features of the muonic energy levels are obtained in the Bohr approximation of a point-like nucleus. This can be done by solving the Schrödinger equation assuming the electrostatic interaction between the muon and the nucleus. A more accurate description of the energy levels is given by the Dirac equation.
in natural units for a muon orbiting in the field of the nucleus is expressed as:

\[ (\alpha \cdot p_{\mu} + \beta m_{\mu}) \psi(r) = (E - V(r)) \psi(r) \]  
(4)

where \( \alpha \) and \( \beta \) are the Dirac matrices, \( p_{\mu} \) the momentum of the muon, and \( \psi(r) \) the wave function of the bound muon at position \( r \) and \( V(r) \) is the electrostatic potential generated by the nuclear charge distribution. A general expression of the nuclear potential is given by [30]

\[ V(r) = -Z\alpha \int d^3 r_N \frac{\rho(r_N)}{|r - r_N|} \]

where \( \alpha \) is the fine-structure constant, \( Z \) the nuclear charge, and \( \rho(r_N) \) denotes the static charge distribution of the nucleus.

For a point-like nucleus, the electrostatic potential takes the form \( V_C(r) = -Ze^2r^{-1} \) and the resulting energy of the muonic levels is the sum of the Bohr energy and the fine-structure shift. The fine structure originates from the coupling of the spin \( s = 1/2 \) with the orbital angular momentum of the muon \( l \). The resulting quantum number of this \( ls \)-interaction is the total angular momentum of the muon \( j \). Taking the values \( |l \pm 1/2| \), the new quantum number \( j \) lifts the degeneracy of the muonic energy states such that the \( 2p \) level splits into the \( 2p_{1/2} \) and \( 2p_{3/2} \) states (following the notation \( nl_j \)).

For the low-lying levels, the point-like nucleus is a very strong approximation which only very roughly describes the muonic energy levels. The effect due to the finite size of the nucleus is typically calculated by numerically solving the relativistic Dirac equation including the nuclear potential under the assumption of some reasonable nuclear charge density. Such a commonly used charge distribution for spherical nuclei is the two parameter Fermi distribution given as

\[ \rho(r) = \rho_0 \left\{ 1 + \exp \left( 4 \ln \frac{3}{\frac{r}{c}} \right) \right\}^{-1} \]

(6)
where $t$ is the surface thickness defined as the distance at which the charge density reduces from 90 to 10% of the density at the center of the nucleus $\rho_0$ and $c$ describes the half-density radius. Due to the mass of the muon which is $\sim 207$ times larger than the mass of the electron, the binding of the muon to the nucleus is significantly stronger than the binding of the electron. Therefore, the low-lying muonic states hugely overlap with the nuclear charge distribution. For $n \gtrsim 6$, the muon is sufficiently far away from the nucleus such that the finite size corrections are very small. In the presence of a finite nuclear density, the nuclear potential changes drastically as shown in Fig. 5. This causes a reduction of up to 50% of the binding energy of the low-lying states in heavier nuclei with respect to the point-like nucleus approximation [31].

For nuclei which suffer a shape distortion, the nuclear charge distribution is developed as a multipole expansion $\rho(r) = \rho_0(r) + \rho_2(r) Y_{20}(\theta)$ where $\rho_0(r)$ is the spherically symmetric part that determines the gross and fine structure while the second term is the non-spherical part relevant for the hyperfine splitting described below. The Fermi distribution then contains two extra parameters ($\beta, \gamma$) that define the shape deformation of the nucleus as [32]

$$\rho(r,c,t) = \rho_0 \left[ 1 + \exp \left( 4 \ln 3 \frac{r - c (1 + \beta Y_{20})}{t (1 + \beta \gamma Y_{20})} \right) \right]^{-1}. \quad (7)$$

The muonic energy levels experience an additional splitting, which is known as the hyperfine splitting (HFS). This effect arises from the coupling of the total angular momentum of the muon $J^1$ with the nuclear spin $I$ and is described by a new quantum number $F = I + J$. The HFS effect is typically smaller than the fine splitting except for highly deformed nuclei where they become of the same order. The allowed transitions of the muon are given according to the selection rules of this new quantum number $F$. The HFS occurs either in deformed nuclei whose shape is not spherical or in nuclei with nonzero magnetic moment and involve the electric quadrupole (E2) and magnetic dipole (M1) interactions. The energy shift of the muonic state of total angular momentum $F$ due to the M1 and E2 interaction accounting for the distribution of the nuclear moments is given by the generalized Bohr–Weisskopf formulas as [33]

$$\Delta E(M1) = \frac{A_1}{2} \left\{ F(F + 1) - I(I + 1) - J(J + 1) \right\} \quad (8)$$

$$\Delta E(E2) = A_2 \left[ \frac{6}{4I(2I-1)J(2J-1)} \left( K(K + 1) - \frac{4}{3} I(I + 1)J(J + 1) \right) \right] \quad (9)$$

where $K = F(F + 1) - I(I + 1) - J(J + 1)$. The M1 hyperfine constant $A_1$ is determined by the magnetic interaction between the muon and the nucleus in terms of the spin and orbital magnetization. It depends on the spatial extent of the nuclear magnetic moment and the wave function and the magnetic moment of the orbiting particle. The HFS constant $A_2$ is a function of the nuclear quadrupole moment generated by the non-spherical part of the nuclear charge distribution. The quadrupole moment describes the effective shape of the nucleus taking a value of zero for spherically symmetric, positive values for prolate and negative values for oblate nuclei. Due to the muon to electron mass ratio, the magnetic moment of the muon is $\sim 207$ times smaller than the one of the electron. In muonic atoms, therefore, the quadrupole deformation is two orders of magnitude stronger than the magnetic effect contrary to the electronic atoms. However, the magnetic part of the HFS can be studied in spherical nuclei where the quadrupole interaction is eliminated.

1 While the total angular momentum is usually written as $j$ for the fine splitting, it is given as $J$ in the hyperfine splitting.
In atoms where the muon is bound to a highly deformed nucleus, the energy of several of the muonic transitions is of the same order as the excitation energy of the nucleus to states with nonzero spin. Due to the $IJ$-coupling, this leads to an admixture of ground and excited hyperfine states and a more complicated structure of the muonic energy levels. The effect is called dynamic E2 interaction and is responsible for the presence of the hyperfine splitting in even-even nuclei [34–36].

A schematic of the finite size, fine and hyperfine splitting is shown in Fig. 6. Other corrections to the muonic levels are the vacuum polarization, the Lamb shift, the nuclear polarization, the electron screening, and the anomalous magnetic moment. More details about the theory of muonic atoms can be found in, e.g., [16,19,36–41].

### 3.3 Extraction of charge radii

Profiting from the high sensitivity of the low-lying muonic transitions to the properties of the nuclear charge distribution, the charge radii of almost all stable elements have been determined employing muonic atom spectroscopy. In order to do so, the energy of the muonic levels is calculated accounting for all the relevant effects and given as a function of the charge distribution parameters. The parameters of the charge distribution can subsequently be extracted when the experimentally observed transitions are matched to the predicted function. However, typically it is the root mean square radius which is reported rather than the charge distribution parameters. The r.m.s. radius is given as $r_C = \sqrt{\langle r^2 \rangle}$, where $\langle r^2 \rangle$ is the second radial moment defined as [19]

$$\langle r^2 \rangle = \frac{1}{Z e} \int d^3 r \rho(r) r^2 .$$

The most difficult component that needs to be treated in the theoretical analysis and in the end limits, the uncertainty on the charge radius extraction is the nuclear polarization correction. This effect emerges due to the electromagnetic interaction between the muon and the nucleus exciting the system into virtual states. The effect results in a shift of the muon binding energies which for the 1s state in $^{208}$Pb is 6 keV [42]. Due to inadequate knowledge of the nuclear excited states, the uncertainty of the calculation of the nuclear polarization...
limits the experimental accuracy. Despite this limitation, very precise values for the charge radius can be obtained. For $^{208}$Pb, e.g., the charge radius was determined with a relative precision of 0.02% [43].

3.4 Extraction of quadrupole moments

The extraction of quadrupole moments from measured muonic X-rays spectra has been performed many times in the past—as example see Refs. [40,44–46]. Here, we briefly describe our extraction of the quadrupole moments for isotopically pure $^{185,187}$Re targets [47,48].

In heavy muonic atoms, there exists an intermediate domain of energy states such as $n = 5, n = 4$ where the radius of the muon orbits is larger than the nuclear size and the muon is not influenced by the presence of the surrounding atomic electrons. In this regime, the hyperfine constants $A_1$ and $A_2$ [shown in Eqs. (8) and (9)] are independent of the details of the electric quadrupole and magnetic moment distributions. Therefore, the measurement of the quadrupole energy splitting is a direct probe of the spectroscopic quadrupole moment which can be obtained by fitting the experimentally observed $5g \rightarrow 4f$ hyperfine transitions.

The $5g_{9/2} \rightarrow 4f_{7/2}$ and $5g_{7/2} \rightarrow 4f_{5/2}$ hyperfine complexes in $^{185,187}$Re are treated together with three weaker multiplets which coincide in energy, namely the $5f_{7/2} \rightarrow 4d_{5/2}$.
$5g_{7/2} \rightarrow 4f_{7/2}$ and $5f_{5/2} \rightarrow 4d_{5/2}$. Due to the nuclear spin of the rhenium isotopes in the ground state of $I = 5/2$, the hyperfine pattern consists of 76 transitions which appear as two bumps at around 360 keV. To disentangle the hyperfine lines, a very good knowledge of the spectral line shape, which describes the detector’s response, is required. The line-shape model was fitted to four background and nuclear capture lines close in energy to the hyperfine structure providing a consistent set of line-shape parameters. The obtained parameters were then fixed for each of the 76 transitions, and a global fit of the $5 \rightarrow 4$ hyperfine structure was performed. The intensities and energies of the hyperfine lines were expressed as a function of the quadrupole moment up to a quadratic term with regard to the $F = 7 \rightarrow F = 6$ transition of the $5g_{9/2} \rightarrow 4f_{7/2}$ multiplet. In this way, the quadrupole moments could be treated as parameters of the global fitting function and were determined by minimizing the overall $\chi^2$ of the fit.

The extracted nuclear spectroscopic quadrupole moment is $2.07(5) \text{ b}$ and $1.94(5) \text{ b}$ for $^{185}\text{Re}$ and $^{187}\text{Re}$, respectively. The results were compared with older literature values obtained by means of muonic X-ray spectroscopy in a natural rhenium sample [44]. A disagreement at the order of 7% was found. The discrepancy is attributed to the higher sensitivity of the isotopically pure data to the weaker transitions (neglected in the previous determination) involved in the fitting of the $5 \rightarrow 4$ hyperfine structure.

4 The muX experiment

In the following pages, we briefly describe the muX experiment which developed a new method to perform muonic atom spectroscopy with targets that are only available in microgram quantities. This enables the application of muonic atom spectroscopy to highly radioactive isotopes and elements that are only available in small quantities. The original motivation for the muX experiment is the measurement of the so far unknown charge radius of $^{226}\text{Ra}$. This is in view of an ongoing experimental effort to measure atomic parity violation in a single, trapped radium ion [49]. In order to extract the origin of the parity violation from their measured atomic transition lines and strengths, precise atomic calculations are needed, which in turn are only possible with a precise value of the charge radius as an input.

As described above, the measurement of muonic X-rays is taken by stopping a negative muon beam in a target resulting in the atomic capture of the muon and the subsequent cascade down to the ground state leading to the emission of the characteristic X-rays. A typical experimental setup contains an entrance detector providing the start signal for an incoming muon. The target is mounted behind this detector and surrounded by veto detectors in order to detect and veto the electrons from the ordinary muon decay. Typically, the entrance and the veto detectors are plastic scintillators. The detectors for the measurement of the muonic X-rays are arranged behind the veto detectors. The best choice in order to obtain good energy resolution are high-purity germanium (HPGe) detectors. Such a setup was assembled for the measurement of the stable elements $^{185}\text{Re}$ and $^{187}\text{Re}$ by the muX collaboration [48] briefly mentioned in Sect. 3.4.

However, as stated one of the main goals of the muX experiment is the measurement of the radioactive isotopes $^{226}\text{Ra}$ and other radioactive elements such as $^{248}\text{Cm}$, which are limited to an amount of only a few μg in the PSI HIPA facility. Additionally, in the case of $^{226}\text{Ra}$ the emitted gamma rates from all its daughter isotopes are so large that it would be impossible to handle much more than a few micrograms. The standard approach of directly stopping a conventional muon beam in such small target masses is not possible and a novel method for forming the muonic atoms is needed to be developed. The new approach is based on the
experience gained during the research on muon catalyzed fusion concerning the behavior of negative muons in hydrogen gases (see [50] and references therein). It relies on repeated muon transfers in a hydrogen–deuterium gas mixture and is inspired by the achievements of Abott et al. [51] and Strasser et al. [52].

4.1 Transfer reactions in a hydrogen/deuterium gas cell

Figure 7 shows a sketch of the experimental apparatus. While the details of the setup will be described in Sect. 4.2, the sketch shows that at the center of the apparatus a gas cell is located with the target mounted within its volume. The gas cell is filled with 100 bar hydrogen and a 0.25% admixture of deuterium. The following paragraphs describe the processes occurring in this gas cell that allows to perform muonic atom spectroscopy with only a few microgram of target material.

Muon capture and transfer The process of the muon transfers to the target starts with a muon entering the gas cell. Due to the high pressure in the gas, the muon is quickly stopped, captured by a hydrogen molecule and a $\mu$-molecular complex is created:

$$\text{(ab)}\mu e^*.$$  

(11)

$a$ and $b$ are hydrogen isotopes and $e$ is an electron. The prepared gas mixture contains natural hydrogen and a small admixture of deuterium. Hence, $a$ or $b$ is most likely a proton ($p$) and we can rewrite Eq. 11 as

$$\text{(pp)}\mu e^*.$$  

(12)

The main dissociation modes [53] of the $\mu$-molecular complex result in the production of muonic hydrogen $\mu p$ by either direct dissociation

$$\text{(pp)}\mu e^* \rightarrow (p\mu)^* + (pe)$$  

(13)
Fig. 8 Total and transport cross sections for the elastic $\mu d$-$H_2$ scattering in the laboratory frame. The cross sections feature a deep minimum at a kinetic energy of around 4 eV due to the Ramsauer–Townsend effect [54–56]. The kinetic energy gain due to the muon transfer from muonic hydrogen to muonic deuterium $\mu p + d \rightarrow p + \mu d$ is shown by the green line.

or by electron emission and subsequent dissociation

$$stoff( pp\mu e)^* \rightarrow (pp\mu)^* + e \rightarrow (\mu p)^* + p + e .$$

(14)

Following the formation, the muonic hydrogen de-excites to the ground state and is thermalized through multiple scatterings with the surrounding hydrogen molecules. Due to the large cross section for $\mu p$-$H_2$ scattering, only a very small number of muonic hydrogen atoms can travel far enough to reach the target.

The key to increase the distance over which the muonic atoms can travel lies in the so-called Ramsauer–Townsend effect [57] which requires a small deuterium admixture in the hydrogen gas. When $\mu p$ scatters with a deuterium atom, the muon is transferred from muonic hydrogen to muonic deuterium

$$\mu p + d \rightarrow p + \mu d^* ,$$

(15)

where $d$ is the deuteron. For the conditions used in the muX gas cell, the rate of this transfer amounts to $\sim 3.5 \times 10^6$/s [58]. The reason for this transfer is the larger binding energy of $\mu d$ due to the higher mass of a deuteron compared to a proton. From the released binding energy, the $\mu d$ atom gains a kinetic energy of 45 eV [54] which is gradually lost by subsequent scatterings.

At a kinetic energy of $\sim 4$ eV [54–56], the $\mu d$-$H_2$ scattering cross section features a deep minimum due to the Ramsauer–Townsend effect (see Fig. 8) and the hydrogen basically becomes “transparent” to the $\mu d$ atom. The distance covered by the $\mu d$ atom over its lifetime can be best described by the mean diffusion radius $R_{\text{diff}}$ [56]. As the scattering cross section is anisotropic, it is best to estimate $R_{\text{diff}}$ with the so-called transport cross section

$$\sigma_{\text{tran}} = \int d\Omega (1 - \cos \theta) \frac{d\sigma(\theta)}{d\Omega} ,$$

(16)

where $d\sigma(\theta)/d\Omega$ is the differential cross section of the $\mu d$-$H_2$ scattering process and $\theta$ is the scattering angle. $\sigma_{\text{tran}}$ is also termed momentum-transfer cross section as the average momentum transfer $\Delta p = p_{\text{in}} - p_{\text{out}}$ in the scattering process is directly proportional to $\sigma_{\text{tran}}$. The calculation shows that $R_{\text{diff}} \propto 1/\sqrt{\sigma_{\text{tran}}}$ [56], which amounts to around 5 mm for
the gas mixture used in the muX experiment (compared to the $\sim 0.5$ mm for a thermalized $\mu p$ atom in a 100 bar hydrogen atmosphere) and thus a much larger fraction of the muons is able to reach the target compared to the case of a pure hydrogen gas.

In the final step, the tightly bound $\mu d$ atom easily passes through the electron cloud of the target atoms as it behaves like a small neutral particle. Once close enough to the target nucleus, the muon is transferred for the last time and forms the desired muonic atom with the target nucleus $\mu A$

$$\mu d + A \rightarrow d + \mu A^*.$$  (17)

The excited muonic atom $\mu A^*$ de-excites through emission of Auger electrons and X-rays to the ground state (compare Sect. 3), which allows the measurement of the muonic X-ray spectrum.

4.2 Setup

A sketch of the experimental setup is already shown in Fig. 7. The following paragraphs will describe in more detail the different elements.

The experiment takes place in the $\pi E1$ area of the HIPA facility at PSI. The $\pi E1$ area offers a high purity negative muon beam with a muon rate of several $10^4 \mu$/s at our chosen momentum. An overview of the $\pi E1$ area including the muX experiment is shown in Fig. 9b. The beam enters the experimental setup from the left side. The red magnets are quadrupole magnets focusing the beam onto the gas cell. The gas cell shown in Fig. 9a is surrounded by an array of HPGe detectors. While in 2017 and 2018, most of the HPGe detectors were obtained from the French/UK germanium detector loan pool [59], in 2019 the Miniball germanium detector array [60] was used for the muonic atom X-ray measurements of $^{226}$Ra and $^{248}$Cm (compare Fig. 10). While in 2017 and 2018, a total of 13 and 14 germanium crystals were operational (see also [61]), this increased in 2019 due to the use of the Miniball germanium detectors to a total of 24 operational germanium crystals. The corresponding full-energy peak efficiencies at 1.3 MeV amounted to $\sim 2\%$ for 2017 and 2018 and close to $3\%$ for 2019. All HPGe detectors require constant cooling with liquid nitrogen. For this purpose, an automated cooling system for the detector array was installed.
Incoming muons are detected by a 20 $\times$ 20 mm$^2$ large and 100 $\mu$m thin plastic scintillator. In front of this entrance, detector is a 5 mm thick plastic scintillator including a circular hole with a diameter of 15 mm. This detector serves as a collimator and veto detector at the same time. All scintillators were directly coupled to silicon photomultipliers in order to record their light output. A layer of $^{208}$Pb is placed in front of this veto detector for the continuous measurement of lead muonic X-rays. Using the precisely measured muonic X-rays of $^{208}$Pb [43] the HPGe detectors can continuously be energy calibrated. Several kHz of muons continuously impinge on this lead layer allowing to perform energy calibrations and track changes in the energy calibration with good statistics every few hours. As additional calibration means, a $^{60}$Co $\gamma$-ray source mounted next to the target, natural background lines like the 2614.5 keV $\gamma$-ray of the thorium series or accidental muonic lines from the target setup like $^{12}$C (75.3 keV) or $^{27}$Al (346.8 keV) are used. The gas cell itself is a cylindrical aluminum vessel with a length of 2 cm and a diameter of 30 mm closed toward the muon beam by an entrance window made of a 0.6 mm thin carbon fiber sheet. It is reinforced by a 4 mm thick titanium support grid with an additional 2 mm thick carbon fiber grid on top. This configuration is robust enough to withstand the pressure of 100 bar inside the gas cell. The target is mounted on a plate at the back wall of the cell opposite to the entrance window. The wall of the cell behind the target also contains the inlet/outlet for the gas allowing to fill the hydrogen/deuterium gas mixture.

4.3 Transfer optimization

The gas cell is designed to stop muons in 100 bar of the hydrogen–deuterium gas mixture at room temperature. This corresponds to $\sim$10% of the density of liquid hydrogen. The remaining parameters to optimize the process of transferring muons to a target mounted at the back wall of the gas cell are the stopping position of the muon beam and the amount of deuterium admixture to the hydrogen gas.

Concerning the admixture of deuterium, it is necessary to balance two opposing effects. Additional deuterium increases the transfer rate from muonic hydrogen to deuterium. However, the additional deuterium decreases the mean diffusion radius of the muonic deuterium atom due to the increased $\mu d$-$D_2$ scattering. In order to determine the best gas mixture for
Fig. 11  (Left) Gold muonic X-rays as measured from a 5 μg gold target over 18.5 h. The 2p − 1s transitions in gold are the lines at 5.59 MeV, 5.75 MeV and 5.77 MeV. The calibration X-rays of 208Pb are measured at 5.45 MeV and 5.78 MeV. Moreover, an X-ray from muon catalyzed fusion is observed at 5.5 MeV. (Right) For each event in the HPGe detector the time difference (tDiff) between the signal in the HPGe and in the entrance detector was determined. This additional information is presented in the 2D spectrum and allows to differentiate uncorrelated X-rays such as from 208Pb (constant in time) from prompt X-rays (centered around tDiff=0) and X-rays stemming from muon transfers (starting at tDiff=0 and extending for a few hundred ns)

In the experiment, the number of muonic X-rays per muon is measured for a given target for different gas mixtures. It was found that a deuterium admixture of 0.1 to 0.25% is optimal.

The muon beam with a momentum of \( p \approx 28\text{MeV}/c \) results in a stopping distribution with a full width at half maximum of approximately 7 mm in the gas mixture. The stopping position, i.e., the center of the stopping distribution, can easily be tuned by changing the beam momentum. Muons with a 1 MeV/c lower momentum stop too close to the entrance window decreasing the probability of reaching the target. In contrast, muons with a 1 MeV/c higher momentum are not stopped in the gas but in the backing plate of the target resulting in the direct atomic capture in the backing plate material. As before, the number of observed muonic X-rays is measured for a given target material and the beam momentum carefully tuned to maximize the yield of these X-rays.

### 4.4 Transfer efficiency

In 2017, the muX collaboration demonstrated for the first time that the newly developed procedure transfers muons to a gold target with a mass of only 5 μg [61]. Due to the expected small transfer rate to the 5 μg target, the test started with the optimizations described in Sect. 4.3 for a 50 nm thick gold target evaporated onto a copper backing foil with a surface of \( \sim 4.9 \text{cm}^2 \) corresponding to a target mass of almost 500 μg. After reaching the optimal settings for this target, the target was replaced by a 10 nm thick gold target and the optimization procedure was repeated. The next optimization step involved a 3 nm thick target with the same surface area. The successive reduction in the target mass ensured a proper evaluation of the required measurement time and experimental settings for the 5 μg target.

The final gold target featured the mentioned mass of 5 μg corresponding to a 3 nm thick coating on a 1 cm² area. After a total measurement time of 18.5 h, the HPGe detectors revealed the muonic atom energy spectrum of the gold 2p − 1s region shown in Fig. 11a. The 2p − 1s lines of muonic 208Pb used for calibration purposes are in the same energy region. In addition, a line stemming from muon catalyzed fusion is observed. The muon catalyzed fusion is a clear indicator for a successful muon transfer from muonic hydrogen to deuterons. The corresponding two-dimensional spectrum showing in addition time information is shown in Fig. 11b. The time difference tDiff for the HPGe detector signals is calculated with respect
to the correspondingly detected muon in the entrance detector. Background signals, such as the calibration X-rays, and the desired muonic X-rays can in principle be differentiated by carefully analyzing this 2D spectrum.

The spectrum was measured with a momentum of \( p = 27.75 \text{ MeV/c} \) and a deuterium admixture of 0.25% in a 100 bar hydrogen/deuterium gas mixture at room temperature. The test showed that about 1% of the stopped muons in the gas cell were transferred to the 5 \( \mu \text{g} \) gold target [61].

5 Conclusions

Muonic atoms can be used to study nuclear properties with high sensitivity due to the close orbits of the muon around the nucleus. By measuring the energy of the characteristic X-rays emitted by the captured muon on its cascade down to the ground state—a technique termed muonic atom spectroscopy—the full muonic energy level scheme can be readily reconstructed. Two nuclear properties of interest can be extracted in that way from the measured X-ray spectrum: the absolute nuclear charge radius and the quadrupole moment of the nucleus. While the nuclear charge radius leads to overall shifts of the energy levels, the quadrupole moment leads to a hyperfine structure of the muonic transitions.

Almost all stable elements have been the subject of muonic atom spectroscopy campaigns with corresponding extractions of their charge radii and quadrupole moments (where applicable). However, no elements that can be handled only in small quantities—either due to their scarcity or radioactivity—could be probed so far. The muX experiment has developed a method based on muon transfer reactions inside a high-pressure hydrogen/deuterium gas cell to probe targets available only in microgram quantities. A first proof-of-principle with a target of 5 \( \mu \text{g} \) of gold was very successful and a nice muonic energy spectrum could be recorded. In 2019, measurements with the radioactive targets \(^{248}\text{Cm}\) and \(^{226}\text{Ra}\) were performed, the analysis of which is currently ongoing.

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