XUV frequency comb metrology on the ground state of helium

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The operation of a frequency comb at extreme ultraviolet (XUV) wavelengths based on pair-wise amplification and nonlinear upconversion to the 15th harmonic of pulses from a frequency comb laser in the near-infrared range is reported. Following a first account of the experiment [Kandula et al., Phys. Rev. Lett. 105, 063001 (2010)], an extensive review is given of the demonstration that the resulting spectrum at 51 nm is fully phase coherent and can be applied to precision metrology. The pulses are used in a scheme of direct-frequency-comb excitation of helium atoms from the ground state to the 1s4p and 1s5p 1P1 states. Laser ionization by auxiliary 1064 nm pulses is used to detect the excited state population, resulting in a cosine-like signal as a function of the repetition rate of the frequency comb with a modulation contrast of up to 55%. Analysis of the visibility of this comb structure, thereby using the helium atom as a precision phase ruler, yields an estimated timing jitter between the two upconverted comb laser pulses of 50 attoseconds, which is equivalent to a phase jitter of 0.38(6) cycles in the XUV at 51 nm. This sets a quantitative figure of merit for the operation of the XUV comb, and indicates that extension to even shorter wavelengths should be feasible. The helium metrology investigation results in transition frequencies of 5740806993(10) MHz and 5814248672(6) MHz for excitation of the 1s4p and 1s5p 1P1 states, respectively. This constitutes the first absolute frequency measurement in the XUV, attaining unprecedented accuracy in this windowless part of the electromagnetic spectrum. From the measured transition frequencies an eight-fold improved 4He ionization energy of 5945204212(6) MHz is derived. Also a new value for the 4He ground state Lamb shift is found of 41247(6) MHz. This experimental value is in agreement with recent theoretical calculations up to order mα4/5 and m2/Mα5, but with a six times higher precision, therewith providing a stringent test of quantum electrodynamics in bound two-electron systems.

I. INTRODUCTION

Atomic spectroscopy has been paramount for the discovery of the laws of physics. The ordering of spectral lines in the hydrogen atom has led to the Rydberg formula and the concept of quantization in the old Bohr mechanics. In 1947 further advance was made with the observation by Lamb and Retherford that the 2S1/2 and 2P1/2 states in hydrogen are not degenerate, but differ by 1 GHz. This result lead to the development of quantum electrodynamics (QED), which is the most precisely tested physics theory to date. Since the first measurement by Lamb and Retherford, QED contributions to energy levels in atoms are referred to as “Lamb shifts”. The theory has been elaborately tested and confirmed by experiments in various bound systems, ranging from atomic hydrogen, via one-electron heavy-ion systems, exotic atomic systems such as positronium, and muonium to one-electron molecular ions, while recently a test of the QED has also been reported in a neutral molecule. Tests have also been performed on the anomalous magnetic dipole moment of the electron (g-2), so that in fact QED can now be used to derive a new value for the fine structure constant from such a measurement.

The two main QED contributions to the energy of an atomic system are virtual photon interactions (“self energy”) and screening of the nuclear charge due to the creation (and annihilation) of electron-positron pairs (“vacuum polarization”). The magnitude of these effects depends on the considered system and its energy eigenstate, and is studied by means of precision spectroscopy. In atoms such as hydrogen and helium, the strongest QED effects are observed in the electronic ground states. For this reason spectroscopy involving the ground state is preferred. In the case of hydrogen this is achieved by inducing a two-photon transition at 2 × 243 nm from the 1S ground state to the 2S excited state, which has reached a level of precision that allows a detailed comparison between theory and experiment. This comparison is currently limited by the uncertainty in the measured proton charge radius, which in itself is not a QED phenomenon. Assuming that the QED calculations are correct, one can also use the 1S – 2S experiment in hydrogen to determine the effective proton charge radius. In this respect it is interesting to note that a recent experiment with muonic hydrogen (where the electron in hydrogen is replaced by a muon), has resulted in a proton size that differs 5σ with the CODATA value that is based mainly on hydrogen spectroscopy. The origin of this difference is still under debate.
performed by experiments using atoms with a higher nuclear charge \( Z \), as the non-trivial QED effects scale with \( Z^4 \) and higher. For this reason experiments have been performed on many different high-\( Z \) ionic species such as \( \text{Ti}^{91+} \). However, the required (very) short wavelengths for excitation, such as hard-X-rays, makes it very difficult to perform absolute frequency measurements (see e.g. \cite{22} and references therein).

Even for \( Z = 2 \) in the \( \text{He}^+ \) ion or the neutral helium atom, where QED effects are at least 16 times larger than in hydrogen, it remains difficult to perform high resolution spectroscopy. In the case of \( \text{He}^+ \) a two-photon excitation from the ground state requires 60 nm, while for neutral helium 120 nm is required to drive the \( 1s^2 \text{^1S}_0 - 1s2p \text{^1P}_1 \) two-photon transition. One-photon transitions require even shorter wavelengths, in the extreme ultraviolet (XUV), e.g. 58 nm to excite the \( 1s^2 \text{^1S}_0 - 1s2s \text{^1S}_0 \) two-photon transition. Single femtosecond laser pulses can be made sufficiently intense for convenient upconversion to XUV or even soft x-ray frequencies using HHG \cite{38}. However, as a consequence of the Fourier principle, the spectral bandwidth of such pulses is so large that it inhibits high spectral resolution. Frequency combs combine high resolution with high peak power by generating a continuous train of femtosecond pulses. In this case the spectrum of the pulse train exhibits narrow spectral components (modes) within the spectral envelope determined by the spectrum of a single pulse. The modes are equally spaced in frequency at positions given by:

\[
f_n = nf_{\text{rep}} + f_{\text{CEO}}
\]

where \( n \) denotes the integer mode number, \( f_{\text{rep}} \) is the repetition frequency of the pulses, and \( f_{\text{CEO}} \) is the carrier-envelope offset frequency. The latter relates to the pulse-to-pulse phase shift \( \Delta \phi_{\text{CE}} \) between the carrier wave and the pulse envelope (carrier-envelope phase or CEP) by

\[
\Delta \phi_{\text{CE}} = 2\pi f_{\text{CEO}}/f_{\text{rep}}.
\]

Each of the comb modes can be used as a high-resolution probe almost as if it originated from a continuous single-frequency laser \cite{39,41}. If the entire pulse train can be phase coherently upconverted, the generated harmonics of the central laser frequency should exhibit a similar spectrum with comb frequencies \( mf_{\text{rep}} + qf_{\text{CEO}} \), where \( m \) denotes an integer mode number, and \( q \) is the integer harmonic order under consideration. By amplification of a few pulses from the train, and producing low harmonics in crystals and gases, an upconverted comb structure has been demonstrated down to 125 nm \cite{42}. However, to reach wavelengths below 120 nm, HHG has to be employed requiring nonlinear interaction at higher intensities in the non-perturbative regime \cite{44}. It is well established that HHG can be phase coherent to some degree \cite{32,33,44,46}, and attempts have been made to generate low frequency combs based on upconversion of all pulses at full repetition rate \cite{47,48}. However, due to the low XUV-pulse energies the comb structure could so far not be verified in the XUV-domain for those sources. That limitation can be overcome by combining parametric amplification of two frequency comb pulses in combination with harmonic upconversion. This was recently demonstrated with direct frequency comb excitation at 51 nm in helium \cite{51}, and in this paper a full and detailed description is given of that experiment.

The article is organized as follows: in section II the measurement principle is explained, followed in section III with a description of the different parts of the experimental setup. The general measurement procedure and results are presented in section IV. In section V, part A, a discussion of all systematic effects that need to be taken into account to determine the ground state ionization potential from the measured transitions frequencies...
is given. This is followed in part B with a discussion of the timing jitter in the XUV, which can be derived from the measured Ramsey signal. In the final section VI the conclusions and an outlook are presented.

II. OVERVIEW AND PRINCIPLE OF XUV COMB GENERATION AND EXCITATION

A frequency comb is normally based on a modelocked laser producing an infinite train of pulses with fixed repetition rate and CEP-slip between consecutive pulses. The corresponding spectrum of such a pulse train consists of a comb of narrow optical modes associated with frequencies given by Eq. [1]. To convert the frequency comb to the XUV, we select only two pulses from the FCL. In this case the spectrum changes to a cosine-modulated continuum, but with the peaks of the modulation exactly at the positions of the original FCL spectrum. This “broad frequency comb” is converted in a phase-coherent manner to the XUV by amplification of the pulse pair to the millijoule level, and subsequent HHG. Once the FCL energy is transferred to the XUV by amplification of the pulse pair to the mJ level, yielding a cosine-modulated spectrum when viewed in the frequency domain is shown in Fig. 1. In the present case, the interacting fields are not separated in space, but only in time. Excitation with two (nearly) identical pulses produces a signal which is cosine-modulated according to:

\[ S(T) \propto \cos(2\pi f_{tr} T - \Delta \phi(f_{tr})) \]  

when varying \( T \) through adjustment of \( f_{rep} \) of the comb laser. In this expression \( f_{tr} \) is the transition frequency and \( \Delta \phi(f_{tr}) \) is the spectral phase difference at the transition frequency between the two pulses. Relation (3) is valid only for weak interaction, which is the case in the current experiment given an excitation probability of \( \ll 1 \) per atom.

Without HHG and in the absence of additional pulse distortions the phase shift \( \Delta \psi := \Delta \phi(f_{tr}) \) is equal to \( \Delta \phi_{CE} \). The excitation signal will exhibit maxima corresponding to those frequencies where the modes of the original comb laser come into resonance with the transition.

The spectral phase difference of the generated XUV pulse pair cannot be determined directly. Therefore we need to propagate the spectral phase difference from the frequency comb through the parametric amplifier and the HHG process into the interaction region. The phase shift \( \Delta \psi(f) \) imprinted on the pulses by the non-collinear optical parametric double-pulse amplifier (NOPCPA) is measured directly using an interferometric technique described previously \[52\]. To model the HHG process we employ a slowly varying envelope approximation described in section [V] which yields a differential XUV phase shift of the form

\[ \Delta \phi_q = q(\Delta \phi_{CE} + \Delta \psi) + \Delta \psi_q, \]  

where \( \Delta \psi \) denotes the carrier envelope phase accumulated in the NOPCPA (compare Sec. [VI]). \( q \) is the harmonic order of the resonant radiation and \( \Delta \psi_q \) is an additional phase shift due to nonlinear and transient response in the HHG process \[53, 54\]. Note that only differences in phase distortion between the two subsequent pulses from the FCL affect \( \Delta \phi_q \) and therefore the Ramsey signal. Shared distortions, such as frequency chirping due to uncompensated time-independent dispersion, have no influence on the outcome of this experiment.

The frequency accuracy of the method scales with the period of the modulation \( f_{rep} \) (here equal to 100 – 185 MHz) rather than the spectral width of the individual pulses (about 7 THz in the XUV at the 15th harmonic). An error \( \delta \) in the value of \( \Delta \phi_q \) leads to a frequency error in the spectroscopy result of \( \Delta f = \delta/(2\pi T) \). Of the two components contributing to this error, \( \Delta \phi(f) \) can be expected to be independent of \( T \), while the transients contained in \( \Delta \psi_q \) decay with increasing \( T \). Therefore \( \Delta f \) decreases at least with \( 1/T \), leading to a higher accuracy for a longer time separation between the pulses. In practice there is still a minimum requirement on the stability and measurement accuracy of phase shift of rms 1/200th of a cycle (at the fundamental frequency of the FCL in the near infrared). This ensures that the contribution \( \delta \) to the frequency uncertainty of the measured transition frequency is small enough so that the ‘mode number’ ambiguity in the transition frequency, due to the periodicity of the signal, can be resolved with confidence.

A sketch of the XUV comb principle in the frequency domain is shown in Fig. 1. A FCL serves as a source of phase-coherent pulses. A pair of such pulses is amplified in a NOPCPA to the mJ level, yielding a cosine modulated spectrum when viewed in the frequency domain. The amplified pulses are filtered spatially with two pinholes, one placed between the second and third amplification stage, and one after compression respectively. As a result the pulse pairs show less spatial intensity and phase variation compared to the unfiltered beam.

Before the IR beam is focused in a krypton jet for HHG, the center of it is blocked by a 1 mm diameter copper disk, while the outside is clipped with an iris. The donut-mode shape is used to facilitate the separation of the driving IR field from the generated XUV emitted on axis (see also section IIID).

Phase shifts in the NOPCPA, which are not common for both pulses, and therefore change the position of the comb modes, are measured by means of spectral interference. A Mach-Zehnder-like configuration is used to interfere the original comb pulses with those that are amplified by the NOPCPA, as described in \[52\] and section IIIC.

The amplified pulses are focused a few mm in front of a krypton jet, in which two phase-locked pulses of
Frequency
Original Comb
Time
Amplified Comb
Frequency
Time
Harmonic order
XUV Comb

FIG. 1. Principle of frequency comb generation and spectroscopy in the XUV. (a) Schematic of the spectral and temporal structure of the generated light at the different stages in the experiment (left to right): sharp equidistant frequencies from the infrared FC laser, followed by double-pulse amplification resulting in a cosine modulated spectrum, and finally HHG resulting in odd harmonics of the central frequency, where each of the harmonics consist of a XUV comb in the form of a cosine-modulated spectrum with period $f_{\text{rep}}$.

FIG. 2. The excitation probability of helium at 51.5 nm on the $1s^2 \ 1S_0 - 1s5p \ 3P_1$ transition, as a function of the repetition of the frequency comb laser (lower x-axis), and the delay between the pulses (upper x-axis in attoseconds). In this example $f_{\text{CEO}}$ is locked 46.21 MHz, and a 1:5 He:Ne mixture is used for the atomic beam.

FIG. 3. Scheme of the experimental setup, including: femtosecond frequency comb laser with control electronics, non-collinear optical parametric chirped-pulse amplifier (NOPCPA), a phase measurement in a Mach-Zehnder interferometer, high harmonic generation (HHG) and a spectroscopy apparatus. Part of the pump laser output at 1064 nm is used as ionization beam.

high-order harmonics are generated to create a cosine-modulated comb spectrum in the XUV. In order to avoid direct ionization of helium with higher-order harmonics, the IR intensity in the interaction region and the harmonic medium are chosen such that the desired 15th harmonic appears at the cutoff of the HHG process. The XUV beam crosses a low-divergence beam of helium atoms perpendicularly, and excites them from the ground state to the $1snp \ 1P_1$ state ($n \in \{4, 5\}$). A pulse of 1064 nm radiation ionizes the excited atoms, which are subsequently detected in a time-of-flight mass spectrometer. Tuning of the XUV comb is accomplished by changing $f_{\text{rep}}$ of the FCL. The changing mode separation effectively causes the modes in the XUV to scan over the transition. An example for a scan of the repetition time corresponding to approximately 500 attoseconds (as) is shown in Fig. 2. The number $m$ of the mode that excites the measured transition in this experiment is in the order of 50 million. This means that the repetition rate of the fundamental FCL needs to be changed only by a few Hz in order to bring an adjacent XUV-mode $f_{m+1}$ into resonance with a helium transition ($f_m$ is assumed to be at the transition initially). As a consequence the ionization signal will be cosine modulated. In order to resolve the resulting ambiguity in the mode-number assignment, the measurement is repeated with different repetition rates, corresponding to pulse delays between 5.4 ns and 10 ns.

Besides the frequency-domain perspective, this experiment can be viewed also in the time domain. In this case it can be seen as a pump-probe experiment, which tracks the dynamics of the electronic wave function that results from mixing the ground state of helium with the excited $p$-level. The first XUV pulse brings the atom into a superposition of the ground and the excited state. This superposition results in a dipole, which oscillates at the transition frequency with an amplitude that decays according to the lifetime of the excited state. The second pulse probes this oscillation. Two extreme cases can be distinguished. If the second pulse is in phase with the dipole oscillation, the amplitude of the excited state increases, and thus its detection probability. If the phase of the second pulse is shifted by $\pi$ with respect to the dipole oscillation of the helium atom, its oscillatory movement is suppressed and the probability to find an atom in the excited state is decreased.

III. EXPERIMENTAL SETUP

The measurement setup consists of four major elements: a frequency comb laser, a non-collinear parametric amplifier, a phase-measurement system, and a vacuum apparatus for HHG and excitation of helium in an atomic beam. A schematic overview of the setup is given in Fig. 3.
A. Frequency-comb oscillator and pulse stretcher

Phase-coherent pulses are obtained from a home-built Kerr-lens mode-locked Ti:Sapphire frequency comb laser. The FCL has an adjustable repetition rate between 100 MHz and 185 MHz. Dispersion compensation in the cavity is obtained with a set of chirped mirrors, supporting a spectral bandwidth of 60 nm centered around 780 nm. The FCL is stabilized and calibrated against the signal of a rubidium clock (Stanford research PRS10), which itself is referenced to a GPS receiver so that an accuracy on the order of 10^{-11} is reached after a few seconds of averaging. Before sending the FCL pulses to the amplifier, the wavelength and bandwidth of the comb pulses is adjusted with a movable slit in the Fourier plane of a grating-based 4 − f stretcher (1200 l/mm, f = 10 cm). The bandwidth is set to 6 nm in this device, ensuring that after upconversion to the XUV only one state in helium is excited at a time. The spectral clipping and losses from the gratings in the stretcher reduce the pulse energy from about 5 nJ to 60 pJ. At the same time the added dispersion and reduced bandwidth lengthens the pulse to about 2 ps.

B. Non-collinear optical parametric amplifier

A pair of subsequent pulses obtained from the stretcher is amplified in a non-collinear optical parametric chirped-pulse amplifier based on two 5 mm long BBO (beta-barium borate) crystals. Here we present a concise description of the system, while further details can be found in [52, 55]. The amplifier operates at a repetition rate of 28 Hz, and amplifies two subsequent FC pulses to a level of typically 5 mJ each. The bandwidth of the pulses remains essentially unchanged compared to that selected by the slit of the preceding stretcher, although saturation effects in the NOPCPA result in a 'cathedral'-like spectrum (see Fig. 4). The pump light for the NOPCPA (two pulses at 532 nm, 50 ps, 80 − 100 mJ per pulse) is obtained by frequency doubling 1064 nm light from a Nd:YAG-based pump laser. A relay-imaged delay line in the pump laser is used to produce these pulse pairs with a time separation between 5.5 ns and 10 ns. This time separation is adjusted carefully for each value of \( f_{\text{rep}} \) to match the time delay between consecutive pulses from the FC laser at a few-ps level.

Electronic synchronization is employed (timing jitter less than 1 ps) so that the pump and comb laser pulses arrive at the BBO crystals at the same time. The amplification happens in three stages, the first two located in the first BBO crystal and the last (power amplifier) stage uses the second. Between the first and second crystal spatial filtering is used to reduce phase-front errors. After amplification in the NOPCPA, subsequent compression, a final spatial filter is employed (based on a 1:1 telescope with \( f = 50 \) cm lenses and an 80 micron pinhole in between, mounted in vacuum). This reduces wave-front errors and results in a Gaussian beam with a diameter of 6 mm. The pulses of 300 fs duration are essentially diffraction and Fourier-limited with an energy of 1 − 2 mJ per pulse.

By choosing parametric amplification, a high gain (10^8) is achieved while avoiding some of the transient effects that conventional laser amplifiers suffer from, such as thermal lensing and inversion depletion. The reason for this is that in NOPCPA negligible power is dissipated in the amplifier medium (a nonlinear crystal), and that the amplified (signal) wave is not distorted provided the process is perfectly phase matched (see e.g. [56]). Imperfections of the wave front of the pump-beam pulses can lead to a spatially-dependent phase mismatch in the NOPCPA, which in turn influences the phase of the amplified signal beam. Therefore special attention is paid to match the properties (wave front, pulse length, energy, diameter) of the two pump pulses so that induced phase effects in the two amplified signal pulses are as equal as possible. A Shack-Hartmann sensor was used to align their direction within several μrad and their propagation axis within 10 μm. The remaining spatial differential signal distortions are reduced by the spatial filters and finally measured interferometrically as described in the next section.

C. Phase-shift measurements in the IR

Knowledge of the carrier-envelope phase change between consecutive pulses of a repetition-frequency stabilized mode-locked laser is a prerequisite for frequency-comb metrology. Because parametric amplification influences the phase of the amplified light [52, 55], a method is needed to detect a differential phase shift between the pulses. This shift is recorded for each laser shot, using spectral interferometry with the original comb pulses as a reference in a Mach-Zehnder configuration (Fig. 5).
The phase-measurement setup is an advanced version of the one published earlier [52]. In order to deal with the demand for higher spectral resolution of the phase-measurement setup and the spatial dependence of the phase shift between the pulses, several improvements were made. The most important is a motorized iris (diameter: 2 mm), which enables spatial mapping of the differential phase within 30 seconds, by scanning it along the donut mode (see also section III D). Such a wavefront scan is made before every recording of the helium signal as it cannot be done for every individual laser shot while measuring the helium signal. However, during a recording of the helium signal the iris is opened and an average value for the shift of $\Delta \phi_q$ is monitored for each laser shot. This average is compared with the wavefront scan made just before a recording of the helium signal from which a correction for the average phase shift is calculated. In this way the effective phase difference change can be continuously monitored. The differential phase distortion along the donut-mode profile has a typical magnitude of 100 mrad, and a spatial variation of 20 – 30 mrad.

Perfect spatial overlap between the amplified and reference pulses is ensured by sending both beams through a 2 meter long large-mode volume (field diameter 20 μm) single-mode photonic fiber. Before doing so, the pulses are stretched approximately 40 times in a grating stretcher. The combination ensures that both amplified and reference beam are perfectly overlapped, while the stretching and the large mode volume of the fiber enables to increase the pulse energy for good signal-to-noise ratio without inducing self-phase modulation (SPM) in the fiber. This last aspect is particularly important because otherwise intensity differences between the pulses could induce SPM and corrupt the result of the phase shift determination.

D. High harmonic generation

The central part of the amplified and spatially filtered beam is blocked by a 1.9 mm diameter copper disk to facilitate separation of the fundamental IR light and generated XUV further downstream. High harmonics are produced by focusing this donut-shaped beam ($f = 500$ mm) a few mm in front of a pulsed krypton gas jet. The intensity at the focus is estimated to be less than $5 \times 10^{13}$ W/cm², while the local gas density is estimated at a few mbar. After the focus the beam encounters an iris of 0.8 mm diameter at 40 cm distance from the jet, which is placed in the image plane of the copper disk. The iris blocks the infrared radiation with an extinction ratio of 1 : 27, while the XUV light emitted on axis can freely propagate. Thereafter the XUV beam passes the interaction chamber (described below), and enters a normal-incidence focusing grating monochromator equipped with an electron multiplier to analyze the harmonic spectrum. We estimate that about $1 \times 10^9$ photons at the 15th harmonic at 51.5 nm are generated per laser shot. The driving intensity is chosen on purpose at such a level that the 15th harmonic is positioned exactly at the cut-off of the HHG process. As a result the next harmonic (the 17th, causing background counts due to direct ionization of helium atoms in the spectroscopy experiment downstream) is about 10 times weaker.

E. Spectroscopy chamber

In the interaction chamber the XUV double pulse intersects a low-divergence beam of helium atoms at perpendicular angle to avoid a Doppler shift. The atomic beam is generated using a pulsed valve (General Valve, backing pressure 3 bar) producing a supersonic expansion with either pure helium or a mixture with a noble gas (ratio 1 He : 5 NG). By seeding in Kr, Ar and Ne the helium velocity can be varied over a factor of 4 to investigate Doppler effects. The divergence of the atomic beam is limited to approximately 3 – 4 mrad by two skimmers: one circular skimmer of 0.3 mm diameter and one adjustable slit skimmer of 0.25 mm width to set the XUV-He beam angle. This divergence is similar to the divergence of the XUV beam ($< 2$ mrad). Directly after interaction with the XUV-pulse pair the excited state population is detected by state-selective ionization (see Fig. 5) with a
50 ps pulse at 1064 nm. The resulting helium ions are detected in a time-of-flight mass spectrometer.

**IV. FREQUENCY METROLOGY ON 1s^2 – 1s4p AND 1s^2 – 1s5p TRANSITIONS IN HELIUM**

The generated XUV comb has been used to measure the 1s^2 – 1s4p and 1s^2 – 1s5p transition frequencies, from which an improved value for the helium ground state binding energy is derived. Because this involves tests of many systematic effects, a hierarchy of measurements can be identified (see table [II]). The first level is that of a single recording of the helium signal as a function of the repetition rate of the frequency comb laser in the IR. To record the helium signal, the infrared FC-laser repetition frequency is scanned in steps of less than 20 mHz, resulting in changes in the time separation between the two pulses of around 1 attosecond. Each scan requires recording of 20000 – 30000 laser pulses, which takes about 15 minutes. For each laser shot, we record the ion signal, together with a series of other parameters, which are the frequency comb repetition frequency \( f_{rep} \) and offset frequency \( f_{CEO} \), the individual IR-pulse energies, energy \( E_{XUV} \) of both XUV pulses and the average amplifier phase shift. The records are then binned into typically 20 groups over a Ramsey period, based on a scaled coordinate \( u \) defined as

\[
u := \left[ q(f_{CEO} + \frac{\Delta \psi}{2\pi} f_{rep}) - f_{th} \right]/f_{rep} \tag{5}\]

where \( q \) is the harmonic order (\( q = 15 \)), \( f_{th} \) is the theoretically predicted transition frequency, and \( \Delta \psi \) is the phase shift at the peak of the envelope, which is calculated from the phase measurement and additional corrections (see section [V]). We calculate the excitation probabilities \( p(u) \) per laser shot, and averages of the other measured parameters within each bin, denoted by bars over the respective symbol from here on. The measured transition frequency \( f_{ij} = f_{th} + f_{ex} \), average background \( p_0 \), and Ramsey-fringe amplitude \( A \) are then fitting parameters in the following model:

\[
p(u) = \left( p_0 + A \cos 2\pi [u + f_{ex}/f_{rep}(u)] \right)/E_{XUV}(u) \tag{6}\]

This results in a transition frequency \( f_{ex} \) for a single scan, relative to the theoretical transition frequency \( f_{th} \). The statistical error in this fit is determined via a bootstrap method [55], which requires no model of the noise sources.

The theoretical transition frequencies \( f_{th} \), used in the fitting procedure as a reference to which the experimental value is determined, are obtained by combining recent values of the theoretical ground state energy from the literature [60] with those of the excited states [61]. Predicted theoretical frequencies are 5740860963(36) MHz for the 1s^2 1S_0 – 1s4p 1P_1 and 5814248632(36) MHz for the 1s^2 1S_0 – 1s5p 1P_1 transitions.

In Fig. 2 a typical recording of the excitation of helium by scanning the XUV comb over the 1s^2 1S_0 – 1s5p 1P_1 transition is shown. The contrast of the modulation (in this example 40%) is smaller than unity due to various effects, like Doppler broadening, frequency noise and a finite constant background of about 15% due to direct ionization with the 17th and higher harmonics. Fitting of a single recording typically shows a statistical error of 1/50th of a modulation period. Depending on the repetition rate it amounts to an uncertainty of 2 – 3 MHz, which is unprecedented in the XUV spectral region.

Several scans are then performed while changing one parameter in the setup (e.g. helium velocity) to determine the magnitude of systematic effects. Such a measurement sequence is referred to as a “series” in the following. The two most frequently performed tests (series) determine the Doppler shift (by changing the helium velocity using gas mixtures) and ionization-induced frequency shift (by varying the density in the krypton jet). Less frequently the IR-pulse ratio was also varied to test its influence on the measured transition frequency. The analysis of these tests, as well as additional experiments that have been performed to determine other systematic errors, are described in more detail in section [V]. Each of the series typically requires 4 – 6 scans. From this data we extract (inter/extrapolate) the measured transition frequency in absence of the investigated effects and a coefficient for correcting the shifted values. For example, the extrapolation to zero velocity in a Doppler-shift measurement yields a Doppler-free frequency plus the slope, i.e. the angle between XUV and atomic beam. The angle can be used to correct the Doppler shift of scans at a finite known helium velocity. A session then consist of a number (2 – 15) of these series containing in total up to 60 scans. Within each session, groups of series are selected so that each group contains at least one Doppler measurement and one ionization-effect measurement. If possible, also a pulse-ratio-variation measurement is included, otherwise a default dependence is used based on previous measurements.

Figure 3 gives an example for a data set acquired in a measurement session. The different series are displayed therein as red circles for Doppler-shift measurements, green triangles for ionization-shift determinations and blue squares for the measurement of the shift related to the pulse-intensity ratio. Note, that some measurements are used to derive both the Doppler and the ionization-related contribution. These separate determinations are used to mutually correct each other for the different tested aspects (so e.g. an ionization-series value is corrected for the Doppler shift measured just before or after, and vice versa). The error bar on each series depends on the statistical error of its single recordings, and the fit of the systematic effect. After grouping, a sequence of transitions frequencies results with an appropriate error bar. Additional corrections (such as Stark shift, recoil shift) are then applied, and the theoretical value for the energy of the excited state is taken into...
FIG. 6. A measurement session performed on the $1s^2 \, ^1S_0 - 1s5p \, ^1P_1$ transition in helium at $f_{\text{rep}} = 121$ MHz. The uncorrected frequency offset from theory $f_{\text{th}}$ is displayed on the $y$-axis. The different shapes of the points indicate measurement series of systematic effects. Doppler-measurement series are shown as red circles, ionization shift measurement series with green triangles while the blue squares denote the measurement of pulse-intensity related effects.

account. The weighted average over these values then leads to a session-level value for the ground state binding energy and an error estimate for it.

Each session is performed for a combination of one upper state in helium (either $4p$ or $5p$), with one repetition rate of the frequency comb laser. Most recordings were made of the $1s^2 \, ^1S_0 - 1s5p \, ^1P_1$ transition. As an additional check also one series was measured on the $1s^2 \, ^1S_0 - 1s4p \, ^1P_1$ transition. A weighted average over all sessions then leads to a new value for the measured transition energies.

V. DISCUSSION

The experiments discussed in this article yield a two-fold result. The phase coherence of the XUV-pulse pair allows to measure the frequency of an electronic transition in helium. In addition, the contrast of the Ramsey fringes, obtained for different repetition frequencies and helium velocities is used to investigate the coherence of the generated XUV radiation. Accordingly the following discussion is divided in two major parts: an evaluation of the measured transition frequencies and a contrast analysis of the observed interference signals.

A. Determination of the ground-state energy of helium with an XUV-frequency comb

For an absolute measurement many systematic effects have to be considered in addition to the already mentioned effects of Doppler shifts, phase shifts in the NOPCPA and in the HHG. These include recoil shifts, refractive index changes (Kerr effect) in the focusing lens for HHG and the entrance window to the vacuum setup, and the influence of the plasma generated in the HHG medium. Also AC and DC Stark effects and Zeeman shifts are considered. A new ionization energy for helium is derived from these measurements by taking the excited states ionization energy of the $4p$ and $5p$ into account. The binding energies of these states are known to an accuracy better than 20 kHz from calculations [60,61]. From the systematic studies an improved experimental value is then obtained; an error budget for this value is listed in Table[11]

To remove the ambiguity due to the periodic comb spectrum, the energy of the ground state of helium is determined by measuring the frequency of $1s^2 \, ^1S_0 - 1s5p \, ^1P_1$ transition at different repetition rates $f_{\text{rep}}$ of the FCL between 100 and 185 MHz. Additionally, the frequency of the $1s^2 \, ^1S_0 - 1s4p \, ^1P_1$ transition is measured at $f_{\text{rep}} = 148.5$ MHz as a crosscheck. The correct mode number is found by plotting the possible energies of the helium ground state against $f_{\text{rep}}$ as shown in figure[11] (similarly as implemented in [12]).

Systematic phase and frequency shifts, which occur at different stages in the experiment (like phase shifts in the NOPCPA or HHG), are measured, investigated and corrected at different stages of data analysis. In this section the magnitude and treatment of individual systematic contributions to the measured frequency is discussed.

Many error sources lead to a frequency uncertainty that is a function of the delay between the pulses in a Ramsey pulse-pair. This can be a direct influence (by phase shifts), but also indirect (e.g. Doppler shift) because the fit accuracy depends on the contrast of the Ramsey interference pattern, which in turn depends e.g. on the repetition rate, helium velocity and decay rate vs. repetition time. We take these variations into account, providing not only an error, but also an error range estimate for several contributions (see Table[11]).

Some of the considered error sources relate to differential phase shifts in the IR or XUV light field (e.g. a IR wavefront error or ionization shift), while other errors shift the measured frequency (e.g Doppler shift). For this reason the corrections and errors are given either in terms of a phase shift at the $15^{\text{th}}$ harmonic, or as a frequency shift. Phase shifts convert into different frequency shifts depending on the repetition rate of the frequency comb laser, while the frequency shifts are independent of $f_{\text{rep}}$.

In the following sections, we will discuss the effect of differential phase shifts in the amplifier, in the focusing lens and the entrance window into the HHG-chamber, and in the krypton jet in which the XUV is generated. Furthermore Doppler effects, AC and DC Stark shifts, Zeeman shift, recoil shift and the contribution to the signal from the adjacent $6p$-level will be evaluated. The discussion is subdivided according to the different stages of error analysis following Table[11].
Unequal conditions experienced by the two laser pulses can shift their relative phase $\Delta \phi_{CE}$, and therefore the offset frequency $f_{CEO}$ of the corresponding frequency comb. Because every imprecision in the carrier-envelope phase shift $\Delta \phi_{CE}$ between the IR pulses is multiplied by the harmonic order of the harmonic upconversion process, special care must be taken to determine the resulting phase shifts. The harmonic order of 15 employed in this experiment sets a high demand for the detection of differential phase shifts between the IR pulses. The setup used to measure the relative spatial-phase profile of the IR-pulse pair was described in section III C. Typical measured phase shifts are on the order of 100 mrad (both positive and negative, depending on the alignment of the pump laser and NOPCPA system). Therefore in the 15th harmonic phase shifts on the order of 1.5 rad in the XUV at 51 nm are considered. The phase-shift variation across the spatial profile of the beam is typically 20 – 30 mrad in the IR, which dominates the uncertainty in the propagation of the phase shift error to the XUV.

In a previous experiment the effective influence of the measured differential spectral phase shift $\Delta \psi(f)$ was calculated by taking a simple equal weight average of all spectral components in the pulse. However, this is an oversimplified procedure. In order to propagate IR pulse distortions to the XUV we model the HHG process using a slowly-varying-envelope approximation, neglecting depletion of the HHG medium (consistent with our operating conditions). We can then write the (complex) generated electric field at the $q^{th}$ harmonic

$$E_H(t) = f_H(A(t))e^{iq(\omega_0 t + \phi(t))} + c.c.,$$

where $A(t)$ is the slowly varying (real) envelope, $\phi(t)$ the (chirp) phase of the pulse and $f_H$ the (complex) single frequency response of the HHG medium. We take $\omega_0$ to be the $q^{th}$ subharmonic of the transition frequency $\omega_{tr}$.

The phase shift between the driving laser field and generated XUV radiation depends on the IR intensity. On a single-atom level and in the strong-field limit it can be described analytically. This model contains several contributions to the harmonic yield. The most common of these are referred to as the long and short quantum trajectories which exhibit a different intensity-dependent linear phase. By selecting only the central part of the HHG emission we ensure that only those terms that show the smallest phase coefficient (the short trajectory) contribute. The yield of the on-axis emission from the 'short' trajectory is optimized by placing the HHG medium slightly behind the focus of the IR fundamental beam. If we now consider the peak of the pulse, assume that only a single quantum trajectory contributes to the HHG field (as should be the case by design of the experiment) so that there is no net interference and therefore $f_H$ has no oscillatory behaviour, and neglect phase matching (which should be fair in a sufficiently small region around the peak amplitude), we can model the response function by:

$$f_H(A) = \alpha A^n e^{i(\beta A^2)}$$

so that the Fourier transform of the distorted pulse at the transition frequency (neglecting the rapidly oscillating counter-rotating component) reads

$$\tilde{E}_H(\omega_{tr}) \approx \int dt \left[1 + n \frac{\delta A(t)}{A(t)} + i(2\beta A(t)\delta A(t) + q \delta \phi(t))\right] f_H(A(t))e^{-iq\phi(t)}.$$ 

By adjusting the intensity difference between the two pulses we attempt to make the $\delta A$ terms vanish. Then the phase difference at the transition frequency between the pulses is given by the expression

$$\Delta \psi = \arg \left(\frac{\tilde{E}_H(\omega_{tr})}{\tilde{E}_H(\omega_{tr})}\right) \approx \int dt \frac{\delta \phi(t)}{f_H(A(t))e^{-iq\phi(t)}}.$$ 

For Fourier-limited pulses ($\phi(t) = 0$) and real $f_H$ (no intensity-induced phase effect in the HHG process) this would just be a weighted average of $\delta \phi$ with $|f_H|$ as a weight function. However, since the IR pulse is near bandwidth limited and tuned close to the transition frequency, $\phi(t)$ can only vary a small fraction of unity during the pulse. Likewise, because we use a harmonic at the cutoff (the 15th), the XUV yield has a single max-
imum of limited duration around the maximum of the IR pulse. During this time we can assume the nonlinear phase $\exp(i\beta A^2)$ to be constant to first order (in t). Therefore the phase correction required at the transition frequency in the XUV is approximately equal to:

$$\overline{\Delta \psi} \approx \frac{\int dt \delta \phi(t)|f_H(A(t))|}{\int dt |f_H(A(t))|}.$$  \hspace{1cm} (12)

Due to the localization of the XUV emission, the weighting function $f_H(A(t))$ could be replaced with any function that peaks, where the pulse does. This means that the effective phase shift in the XUV at the transition frequency can be approximated using phase information in the time domain as the XUV radiation is only generated when the IR pulse is close to its maximum intensity. We estimate $A(t)$ by assuming Fourier-limited IR pulses with the spectrum found by spectral interferometry. This should be a good approximation of the actual pulse shape in the experiment, as it leads to the highest XUV yield, and the pulse compressor was set to achieve this maximum XUV yield during the measurements. $\delta \phi(t)$ is calculated using the same assumption by just calculating the inverse Fourier transform of the two spectral-interferometry images. In a separate experiment we determined that the amount of XUV radiation at the 15th harmonic depends on the IR intensity to the 9th power, which has actually been used as the peaking weighting function for $\delta \phi(t)$.

The temporally-averaged phase shift using the reconstructed XUV intensity and temporal phase shift results in a value of $\overline{\Delta \psi}$ for each laser shot. This is then used in the helium signal fitting procedure.

We analyzed our data using equal weight spectral averaging over the measured spectral phase as well. The resulting transition frequency coincides with our more accurate model to within 1 MHz. However, for the simple spectral average of the measured phase shift, the spread in the values for the series with different repetition rates of the comb laser is significantly larger, which confirms that weighted temporal averaging models the phase shift better.

To minimize the intensity-related frequency shifts from HHG in the spectroscopy, we take care that the energy in the driving pulses is on average equal within 5%. Together with the filtered beam profile and equal temporal profile, this ensures that the intensity in the focus is equal on the few percent level. To compensate for possible intensity differences we determined the observed transition frequency as a function of IR-pulse energy difference and interpolate linearly to zero difference. First the energies of the pulses are measured with a photo-diode. The integral of the photo-diode signal is recorded with an oscilloscope for each pulse and divided by the mean value. A value for the peak intensity of a pulse is obtained from the measured energy, using again the spectral data available in the interferograms recorded for the phase measurement. We typically find a frequency shift that, linearly extrapolated, corresponds to an IR phase shift smaller than 0.2 rad for 100% relative intensity variation, which is less than 3 rad at the 15th harmonic. This interpolation also includes (and therefore minimizes) nonlinear phase shifts in the beam splitter, focusing lens and vacuum window, which are not included in the phase measurement using the Mach-Zehnder interferometer. Our relative intensity determination is accurate to within about 2% for small pulse energy differences, so that the error due to this determination can be estimated to be 60 mrad, corresponding to a frequency error of 1.5 MHz at 150 MHz repetition frequency in the XUV.

The “Amplifier phase” error in Table I combines the uncertainty of the phase measurement and wavefront deviation in the fundamental IR beam. The latter is calculated based on an average over the wavefront scan over the donut mode of the IR pulses. However, in principle there can be a systematic error for this effect at the level of a single helium signal recording, especially if the intensity profile is not perfectly homogeneous. In that case the average of the phase measured over the beam is not representative for the phase shift at the 15th harmonic. To minimize this effect, we employ tight spatial filtering just before the phase-measurement setup to reduce wave-front errors and to smoothen the intensity profile to the lowest order Gaussian. During a session the pinhole we use for spatial filtering typically starts to wear out after a few hours of operation, leading to slightly asymmetric intensity distributions over time that vary randomly for each measurement session. The pinhole was also regularly replaced and realigned, as was the NOPCPA, so that the intensity and phase variations average down significantly over the near 200 scans that were analyzed. Therefore the error due to the phase deviations and measurement accuracy is listed as a statistical error, conservatively based on the rms amplitude of the phase deviations as measured for each helium signal scan.

2. Measurement series level errors

Some of the systematic effects are related to the condition of the setup during the measurement. In particular transient and alignment-related effects are investigated alternately during a measurement session, forming series of measurements, which can be used to derive and mutually correct systematic shifts.

Transient phase shifts occur when the first pulse changes the propagation conditions for the second one. In this case the condition of equal pulses is not sufficient to avoid an additional phase shift. Transient shifts that occur within the Mach-Zehnder interferometer are readily detected and corrected for. This includes shifts that are caused in the optics after the NOPCPA.

The largest transient effect is due to ionization of the HHG medium. What matters is only the ionization between the two points in time where the 15th harmonic is generated. Ionization leads to a lowering of the refractive index especially for the IR fundamental pulse,
to the effect that the phase velocity of the second pulse becomes faster than that of the first pulse. Ionization is intimately linked to the HHG efficiency and therefore cannot be avoided completely. As we vary the IR intensity (by adjusting an iris size in the IR beam path) we find that at too high intensity, the second XUV pulse can be strongly suppressed due to ionization of the medium and resulting phase-mismatching effects induced by the first pulse. In order to keep the latter to a minimum, we operate at an intensity at which a reasonable XUV yield is obtained, while simultaneously keeping the XUV pulse-energy imbalance to within less than 20% (see Fig. 7). In order to determine the remaining effect due to ionization, we change the pressure in the HHG medium (by adjusting the valve driver), thereby varying the plasma density, and record the measured transition frequency effectively for different ion densities in the HHG medium.

The simultaneously measured XUV energy is used as a relative gauge for the density of ions in the HHG interaction zone, as no direct measurement of the level of ionization of the HHG medium was available for the helium measurements. If the phase matching conditions are not severely altered due to ionization, a quadratic dependence of the XUV output on krypton density is expected, while the plasma density, and therefore the phase shift, should be proportional to the density of krypton atoms. We assume that the induced plasma density is proportional to the krypton density, and that no recombination takes place on a timescale of 10 ns.

The relation between XUV yield and ion density was verified in a separate experiment by determining the amount of ionization in the HHG process using a collector grid just below the krypton jet. The voltage applied to this grid was on the order of a few volts to avoid secondary emission. For the conditions in the experiment the relation between XUV energy and recorded number of ions is equal to a power law with an exponent of 2.1(6), in good agreement with the expected 2.0.

To obtain the ‘ionization free’ frequency we extrapolate the frequency offset vs. XUV energy (when varying Kr density) to zero XUV energy using this experimentally determined exponent (see Fig. 8). From the extrapolations we typically find shifts between 0 and 2 rad at the 15th harmonic for standard conditions, with uncertainties in the range between 0.5 and 2.0 rad. The average correction is on the order of 0.1 XUV cycles. Figure 8 shows the fitting of a typical ionization measurement series, from which a value is derived for \( f_{ex} \) at zero gas density in the HHG jet and therefore at zero XUV intensity.

We also have considered krypton atoms in the HHG medium possibly left in an excited state after the first pulse. These excited state atoms would have a different nonlinear susceptibility and therefore produce a different nonlinear phase shift. Such a shift would not be detected by reducing the gas density. However, the HHG cutoff energy for such excited state atoms would lie far below the cutoff energy of the ground state atoms, as the first excited state is about 10 eV above the ground state. Because the cutoff is already set to the 15th harmonic for ground state krypton, this radiation can not be produced by the excited Kr atoms.

The Doppler effect is monitored by varying the helium beam velocity using mixtures with noble gases. The velocity of helium in the atomic beam is measured (in a separate experiment) by monitoring the frequency shift in the helium signal for different angles between atomic and XUV beam. Helium atoms in the pulsed atomic beam have a speed of 2000(320) m/s for a pure-helium expansion, helium in neon (pressure ratio 1:5) results in a speed of 830(200) m/s, and helium in argon (pressure ratio 1:5) is slowed down to 500(250) m/s. The XUV and atomic beam can be aligned perpendicularly using the different gas mixtures within approximately 10\(^{-4}\) rad, resulting in a typical statistical error of 13 MHz for a single
the ground state energy is smaller than 0
beam angle) and the angle is kept small with changing
is included in the error budget.

that is proportional to \( \cos \phi \) (\( \phi \) being the XUV-atomic
beam angle) and the angle is kept small with changing
signs by readjusting it, the systematic Doppler shift in
the ground state energy is smaller than 0.5 MHz, which
is included in the error budget.

3. Measurement session level errors

The biggest correction at the measurement session
level comes from the recoil shift on the considered
transitions and can be calculated with high precision. It
amounts to 18.28 and 18.75 MHz for the 1s \(^1S_1 \rightarrow
up \:^1P_1(n \in 4, 5)\) transitions respectively.

As discussed in section III D, the IR beam is separated
from the XUV beam an iris placed after the HHG inter-
action zone. However, some diffracted IR light can still
reach the spectroscopy zone. Compared to the original
IR beam, the diffracted light is found to be at least 27
times lower in intensity. this light can still produce an
AC-Stark shift but only during the time of the excitation
pulses (and not in between), Therefore it shows up in our
measurement as a phase shift, which results in a different
frequency shift for different \( f_{rep} \).

The required correction is determined from a measure-
ment with the beam block removed (i.e. with full IR
intensity present in the interaction region). In this case
we find a shift of 17.5(6.0) MHz on the 1s \(^2S_0 \rightarrow 1s5p \:^1P_1\)
at 148 MHz repetition frequency. With the beam block
inserted for the regular helium measurements, a shift of
less then 0.65 MHz is expected. Converting this to an
equivalent phase shift of 14(14) mrad in the XUV allows
to correct the other measurements performed at different
repetition rates.

We have also estimated the theoretically expected fre-
quency shift at \( f_{rep} = 148 \) MHz for the same case.
The estimated average peak intensity with the beam
stop removed is about 25 GW/cm\(^2\), over a pulse dura-
tion of 300 fs. Between the two pulses the intensity is
zero. The AC-Stark shift can be calculated by taking
the time-averaged intensity \( \frac{2}{3} \) (for pulses 6.7 ns apart)
of 1 MW/cm\(^2\). It leads to a theoretical AC-Stark shift of
20 MHz, in good agreement with the experimental value,
measured with the full IR beam passing the interaction
zone.

Finally, we need to consider that the pulses for the
phase measurement are split off via a 1 mm thick beam
splitter (with fused silica as substrate material). After
this point the pulses travel through the focusing lens
(few mm BK7 glass) and enter the vacuum setup for
HHG via a Brewster window (another few mm BK7).
Any difference in intensity between the two subsequent
pulses that are used for the XUV comb generation, or any
transient effect in these glass pieces, can cause an addi-
tional phase shift which is not taken into account by the
generic phase-measurement procedure during recordings
of the helium transition. To be able to still correct for
this phase shift, a separate experiment was performed re-
moving the IR/XUV separation iris in the vacuum setup.
In this way the IR pulses could travel through the in-
teraction zone to the monochromator, where they were
coupled out through a window. Two beam splitters in-
side the monochromator vacuum chamber reduced the
pulse energy to less than 0.1% of the original energy,
thus avoiding spurious phase effects in the output win-
dow and other optics. A phase measurement was then
performed on these pulses, while varying the pulse-energy
ratio (which can be adjusted via the pump laser). This
measurement resulted in an additional correction and un-
certainty due to the optics placed between the phase-
measurement setup and HHG (beam splitter, lens and
Brewster window) of 8.5(15) mrad in the IR. This con-
tribution (multiplied by 15 to take the harmonic order
into account) is called “NL-phase shift” in the error bud-
get in Table I where NL stands for the nonlinear origin
of these differential phase shifts.

From Table I it is clear that the error contributions dif-
fere between the sessions due to different sizes of the cor-
responding data sets. Also a higher accuracy is achieved
for the lower repetition rates in part because phase er-
rors there result in smaller frequency errors. The highest
uncertainty of 25 MHz at \( f_{rep} = 185 \) MHz is based on
a very short measurement session, consisting of only one
set of Doppler and ionization series.
4. Evaluation of the He ground state energy

In the evaluation of the error in the ground state energy, the combined statistical and systematic error based on all sessions (3.7 MHz) is combined with several systematic errors represented as frequencies.

The biggest contribution to the uncertainty comes from the ionization shift model. This error contribution listed under “Ionization shift model” in Table I refers to the error introduced by the uncertainty in the power law of 2.1(6) between the XUV intensity and the amount of ionization (and therefore phase shift) in the harmonic generation region. As this power law is used to correct for the ionization-induced shifts for all measurements at the measurement series level, it results in a systematic error. This is determined by re-analyzing the helium ionization potential for a power law of 1.5 and 2.7, resulting in a variation of 4.9 MHz. As an additional test of (mostly) ionization-induced errors, we also determined the ground state energy using only data points selected within a limited range of XUV-pulse energy, relative to the average XUV energy for each scan. Tests were performed for symmetric and anti-symmetric XUV energy distributions with 5% and 15% deviation from the average XUV energy. A maximum deviation of 5 MHz (from the value based on all points) was obtained in case of an asymmetric selection of data points with an XUV energy higher than 1.15 times the average energy. This is comparable with the estimated accuracy of the ionization model as it is based on a typical XUV yield and full distribution of XUV energies.

During the excitation time interval, the extraction fields for the time-of-flight spectrometer are switched off. The estimated residual field of less than 0.5 V/cm results in a calculated DC-Stark shift on the 5p, m = 0 state of less than 20 kHz. There can be an additional field due to ions in the interaction zone left from the first excitation pulse. However, even for 1000 ions these fields are too small to cause a significant shift.

The influence of the Zeeman effect is estimated based on the magnetic field measured in the interaction region of 5 × 10⁻⁵ T. Because the excitation radiation originates from HHG, which has essentially zero yield for circular polarization, we can assume the circular component to be less than 1%. Since the ground state has no angular momentum, optical pumping cannot occur. Therefore only a weak (1%) ∆m = ±1 component has to be taken into account, leading to a Zeeman shift of less than 7 kHz.

It is vital for Ramsey excitation, that only one transition is excited at a time. Excitation of other states would contribute also with a cosine of almost the same period, but a different phase (which changes quickly with the repetition rate of the FCL). A contribution from the 1s² → 1s2p transition, which is in the vicinity of the 13th-harmonic can be excluded, as it can not be ionized with a single 1064 nm photon. But for the excitation of the 5p also the 6p level could possibly be excited.

To investigate this, the bandwidth of the XUV pulses was determined by coarse tuning of the IR central wavelength by moving the slit in the Fourier plane of the pulse stretcher, located before the parametric amplifier. As a result, the XUV comb spectrum is tuned as well, and scanned over the transitions. From the signal we subtracted a constant background signal from direct ionization due the 17th and higher harmonics. This background has been found by blocking the ionization beam. A typical scan (corrected for the background) is depicted in Fig. 10. The solid line is a Gaussian fit of both the 5p and the 6p resonance, yielding a 1/e half width of the XUV spectrum of 0.07 nm at the 15th harmonic.

From this width and the Gaussian profile we conservatively estimate a contribution to the 5p signal from the 6p resonance of less than 1%. This can cause a shift of the Ramsey pattern of at most 10 mrad (in the XUV). However, in practice the influence of this shift on the final transition frequency averages down for different repetition frequencies, so that the maximum error due to excitation of the 6p is less than 30 kHz for all measurements combined.

After incorporating all the systematic corrections, a clear coincidence can be seen between the results for different repetition rates. A new ground state energy for ⁴He of 5945204212(6) MHz is found by taking a weighted average over all measured frequencies at the coincidence location. The two most recent theoretical predictions of 5945204174(36) MHz from ⁶He and 5945204175(36) MHz from ⁶²He are in agreement with this value within the combined uncertainty of theory and experiment. (The uncertainty of 36 MHz in both theoretical values is based on estimated but yet uncalculated higher-order QED contributions.) Remarkably, very good agreement is found with the prediction of Korobov ⁶²He, who calculates 5945204223(42) MHz, employing non-relativistic quantum electrodynamics theory, in which the problematic divergences of QED are canceled at the operator level.

FIG. 10. Excitation spectrum (single pulse) from the ground state to the 1s5p ¹P₁ and 1s6p ¹P₁ levels in helium, obtained by coarse scanning of the central wavelength of the fundamental and therefore its 15th harmonic over the transitions.
However, the uncertainties in the present calculations are too large to decide, which of the theoretical approaches is better suited to calculate the energy structure of few electron atoms.

Compared with the best previous determinations using single nanosecond laser pulses \(^{24,26}\), our new value is almost an order of magnitude more accurate. Good agreement is found with the value of 5945204215(45) MHz from \(^{24}\) (based on the measured transition energy in that paper, but corrected for a 14.6 MHz recoil shift that was previously not taken into account, and using the most recent 2\(p\) state ionization energy from \(^{64}\)). However, there is a difference of nearly 3\(\sigma\) with the value of 5945204356(50) MHz from \(^{26}\).

### B. Signal contrast and phase stability of HHG

In the previous section phase information of the obtained cosine-shaped signals was used to determine the ground state ionization energy of helium. In the following we use the observed contrast of the same helium signals to investigate the phase stability of the generated XUV frequency comb. The signal visibility (ion signal modulation amplitude divided by the average signal) depends on a combination of effects.

The biggest contributions come from the line width of the observed transition, phase stability of the 15\(\text{th}\) harmonic, Doppler broadening, and the width of the IR-FCL modes. In one extreme case \((f_{\text{rep}} = 185\text{ MHz, helium seeded in argon)}\) we find a fringe contrast of 55\%, while for \(f_{\text{rep}} = 100\text{ MHz and pure helium the contrast is below 5\%}\. In Figure 12 the signal contrast is shown for several experimental conditions.

From an analysis of the observed visibility of the interference pattern as a function of the helium-beam velocity and comb repetition frequency, an estimate for the XUV comb phase jitter and of the effective width of the angular distribution of the atomic beam can be derived. We model the Ramsey pattern assuming Gaussian phase

\[
\text{Visibility} = \frac{1}{1 + f_{\text{rep}}^{2}} \cdot \frac{\text{width}}{\text{width} + \text{line width}}
\]

### Figure 11

**FIG. 11.** Measured \(^4\text{He} 1\text{S}_0\) binding energy \((\pm \sigma f_{\text{rep}})\) relative to the theoretical value of 5945204174 MHz \(^{60}\), plotted against the repetition frequency \(f_{\text{rep}}\) of the comb laser. A clear coincidence is seen at \(+37(6)\text{ MHz relative to theory.}\

### Figure 12

**FIG. 12.** Ramsey signal of helium, acquired with different experimental conditions. (a) \(f_{\text{rep}} = 185\text{ MHz, } 1s^2 1S_0 - 1s5p 1P_1\) transition, helium seeded in argon, contrast: 56\%; (b) \(f_{\text{rep}} = 148\text{ MHz, } 1s^2 1S_0 - 1s5p 1P_1\) transition, helium seeded in neon, contrast: 38\%; (c) \(f_{\text{rep}} = 121\text{ MHz, } 1s^2 1S_0 - 1s5p 1P_1\) transition, helium seeded in neon, contrast: 25\%; (d) \(f_{\text{rep}} = 148\text{ MHz, } 1s^2 1S_0 - 1s4p 1P_1\) transition, pure helium beam, contrast: 13\%. 

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\(S\) is included (red square symbol at \(f\)) for \(seeded\) in \(argon\).}
TABLE I. The error budget for the measurement of the ground-state binding energy of helium. The corrections and uncertainties are given either in radians or in MHz at the 15th harmonic, depending on the type of effect (see text). The NL-phase shift denotes the differential phase shift (multiplied by 15 to account for its effect at the 15th harmonic) induced in the optics after the phase measurement but before the HHG jet (see V A 3).

| Effect                  | Correction | Systematic uncertainty | Statistical uncertainty |
|-------------------------|------------|------------------------|-------------------------|
|                         | Single recording level |                       |                         |
| Statistical fit error   | —          | —                      | typ. 50–150 mrad        |
| Amplifier phase         | 0 – 2.25 rad| —                      | 75 – 375 mrad           |
| Combined error single recording | —       | —                      | typ. 160 – 360 mrad     |
|                         | Series level |                       |                         |
| Doppler shift           | 0 – 10 MHz | see text               | 8 – 18 MHz              |
| Ionization shift        | 0 – 2 rad  | see ‘ionization shift model’ | 0.5 – 2.0 rad          |
|                         | Session level |                      |                         |
| AC-Stark shift          | 14 mrad    | 14 mrad - see text     | —                      |
| NL-phase shift          | 128 mrad   | 225 mrad               | —                      |
| Recoil shift 4p         | 18.28 MHz  | ≤ 1 MHz                | —                      |
| Recoil shift 5p         | 18.75 MHz  | ≤ 1 MHz                | —                      |
| Combined error single session | 9 – 25 MHz | —                      |                         |
|                         | Ground state evaluation |                  |                         |
| Weighted mean sessions  | —          | —                      | 3.7 MHz                 |
| Ionization shift model  | —          | —                      | 4.9 MHz                 |
| Doppler shift           | —          | —                      | 500 kHz                 |
| DC-Stark shift          | —          | —                      | < 1 kHz                 |
| Adjacent 6p-level       | —          | —                      | < 30 kHz                |
| Zeeman shift            | —          | —                      | < 7 kHz                 |
| Total error in ground state energy | —    | —                      | 6 MHz                   |

noise, while the Doppler effect is taken into account with various distributions, including Gaussian and rectangular. Both Doppler and phase-noise distribution widths are fitted in the procedure to the observed visibilities, and scans are included that were used in the frequency determination, plus additional scans that were performed on the 6p and 7p transitions. In all cases we assume a common background count rate of 15%. The resulting Doppler width and phase jitter depends on the function used for the Doppler distribution. For a Gaussian profile, we find a FWHM angular beam divergence of 2.4 mrad and 0.42 cycles XUV jitter. A rectangular distribution leads to a divergence angle of 4.3 mrad and 0.35 cycles XUV phase jitter. In both cases the model seems to have deficiencies. While in the former case the visibility seems to be systematically underestimated for slow beams, in the latter case this underestimate is less pronounced and the fit becomes worse for the fast He beams where the angular distribution is important. We therefore estimate an intermediate value of 0.38(6) cycles for the XUV phase jitter.

If we consider the contributions to the XUV comb jitter then the biggest contribution arises from the bandwidth of the modes of the FC in the IR. The bandwidth of the FC was measured in a separate experiment by beating the FC modes with a frequency doubled narrow bandwidth Er-fiber laser (NP Photonics, with a short-term line width of 5 kHz) to yield a value of 1.6 MHz FWHM, corresponding to a jitter of 1/6 of an XUV cycle.

A second contribution comes from the phase uncertainty of 47 mrad FWHM in the amplified pulses after correction for the measured phase shift (evaluated near the peak intensity of the pulse), leading to a jitter of 1/9 cycle in the XUV. For HHG we take into account that the phase of the generated harmonics is proportional to the intensity of the fundamental beam [44], with an estimated factor of $\approx 10^{-13}$ rad cm$^2$ / W, using the results presented in [64]. Combined with an NOPCPA pulse-ratio distribution FWHM of 2.5%, this gives a small expected jitter of 0.02 XUV cycles. Fluctuations of 6% in the IR pulse intensity lead to variations in the level of ionization in the HHG jet of more than 35% (as ionization scales with at least the 7th power of the IR intensity for the conditions in the experiment). If we adopt a typical average ionization correction of 0.1 XUV cycles, then this induces a phase jitter of 0.04 XUV cycles. Krypton density fluctuations (< 20%) induce less than half this value.

Statistically independent combination of these noise sources leads to a total expected phase jitter in the XUV of 0.21 cycles. This is lower, but comparable to the jitter extracted from the contrast measurements. The difference might be due to uncertainties in the exact experimental conditions, in particular the Doppler broadening and the bandwidth of the frequency comb laser modes during the experiments.

The (estimated) contributions to the jitter suggest that the contrast should hardly be influenced by intensity-induced phase noise from the HHG process. This was investigated experimentally by taking only a subset of
the data points to determine the contrast, using a selection criterion based on the measured XUV intensity. With this procedure we find that the visibility of the interference does not change (within a few percent) if we restrict the XUV intensity variation to symmetric bands of 5%, and 40% around the mean XUV energy. We therefore conclude that intensity dependent effects do not play a significant role as a source of phase noise in the XUV comb in the present experiment.

VI. CONCLUSIONS AND OUTLOOK

The metrology presented in this article represents a significant advance in the history of precision extreme ultraviolet spectroscopy. As is illustrated in the historical overview of the spectroscopy of the helium atom, depicted in Fig. 13, the theoretical and experimental accuracy of the value of the binding energy of helium improved by many orders of magnitude over the last hundred years. Experimentally the biggest progress has always been initiated by new methods, which in turn also lead to advances in the theoretical understanding. The pioneering measurement by Herzberg in 1958 based on a helium lamp and grating spectrometer was not significantly improved upon until the first laser excitation in 1993.

In this paper we have described the first absolute frequency measurement in the XUV spectral region, based on parametric amplification and harmonic upconversion of two pulses from an IR frequency comb laser. Direct frequency comb excitation in the XUV of helium from the ground state is demonstrated, leading to an almost 10-fold improved ground state ionization energy, and in good agreement with theory.

For the employed method it is vitally important to control and detect phase shifts in the amplification and harmonic upconversion process. An important aspect of direct excitation with an upconverted frequency comb is the possibility to detect systematic errors due to phase shifts between the pulses from the NOPCPA, HHG and ionization, but also from the AC-Stark effect. Uncorrected errors of this kind show up as a frequency shift that is proportional to $f_{rep}$, and can therefore be detected easily. In the current experiment no dependence on the repetition rate is observed in the measurements (see Fig. 11), indicating that the systematic effects have been taken into account properly. The accuracy of the presented method scales with $f_{rep}$, which also means that it can be improved by orders of magnitude by choosing frequency comb pulses further apart. If the pulse delay would be extended to $> 150$ ns, then the ionization shift introduced in the HHG process would in fact vanish altogether, as the time between the pulses would be sufficiently long to replace the gas in the focal region with a new sample before the second pulse arrives. This would reduce the error in the spectroscopy significantly because the biggest source of uncertainty is currently due to ionization effects in the HHG process. To fully exploit the enhanced accuracy with larger pulse delays, the sample has to be trapped and cooled to reduce Doppler and time-of-flight broadening which would otherwise wash out the ever denser signal modulation. In view of this, measurements on He$^+$ are particularly interesting. This ion has a hydrogen-like electronic configuration which can be excited from the ground state using a two-photon transition at 60 nm. Such an experiment has the potential to perform QED tests beyond what has been possible so far in atomic hydrogen.

In the present work a frequency comb laser in the XUV range is demonstrated, for the first time at wavelengths as short as 51 nm. The phase jitter of the XUV frequency comb modes is found to be 0.38(6) cycles. From the results we conclude that the observed comb jitter in the XUV is not dominated by the HHG upconversion process, but most likely by the stability of the original comb in the infrared. This form of technical noise can be considerably reduced by locking the comb laser to a stable

![FIG. 13. (color online) (a) Advance in theoretical and spectroscopic accuracy of the ground state energy of helium over the last 100 years. The shaded areas emphasize new spectroscopic tools that allowed to overcome limits of previous technologies. Red squares represent the precision of calculations. The precision obtained in experiments is indicated with blue circles. (b) Comparison of helium ground state ionization energy values obtained experimentally (blue circles) and theoretically (red squares) in the last two decades. The values are plotted with respect to the most recent theoretical result.](image)
optical reference cavity. It seems therefore feasible to extend comb generation and spectroscopy to even shorter wavelengths into the soft-X-ray region, provided that the carrier-phase noise of the fundamental and upconverted pulses is kept low enough. Further possibilities would arise if not subsequent pulses from the FC, but any two-pulse sequence could be selected for upconversion. This would increase the accuracy significantly below 1 MHz as the pulses could have a much increased time delay, thus making it possible to resolve multiple transitions with discrete Fourier transform spectroscopy techniques. In view of these possibilities, we envision applications such as QED test of He+ and highly charged ions, precision spectroscopy of simple molecules such as H₂, coherent XUV or X-ray imaging, and possibly even the emergence of X-ray nuclear clocks.

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