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Absolute frequency measurement of the $^{1}\text{S}_0 - ^{3}\text{P}_0$ transition of $^{171}\text{Yb}$

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Abstract
We report the absolute frequency measurement of the unperturbed transition $^{1}\text{S}_0 - ^{3}\text{P}_0$ at 578 nm in $^{171}\text{Yb}$ realized in an optical lattice frequency standard relative to a cryogenic caesium fountain. The measurement result is $518\,295\,836\,590\,863.59(31)$ Hz with a relative standard uncertainty of $5.9 \times 10^{-16}$. This value is in agreement with the ytterbium frequency recommended as a secondary representation of the second in the International System of Units.

Keywords: optical lattice clock, SI second, frequency metrology

(Some figures may appear in colour only in the online journal)

1. Introduction

Time and frequency are the physical quantities that can be measured with the lowest uncertainty so far and atomic frequency standards based on the microwave hyperfine transition of the caesium ground state realize the second in the International System of Units (SI). The fractional uncertainties of the today’s best Cs fountains are in the low $10^{-16}$ region [1–3] while optical frequency standards are now surpassing the caesium standards both in terms of accuracy and stability. Optical frequency standards using various atomic species and reference transitions [4–13] are being developed around the world also motivated by the prospect of a redefinition of the SI second based on an optical transition [14]. Optical frequency standards are of interest as highly sensitive probes for fundamental physics such as relativistic geodesy [15–17], tests of relativity [18], quantum simulations [19], and searches for the variation of fundamental constants [9, 20, 21].

In optical lattice clocks thousands of neutral atoms are trapped in an optical potential at the light-induced Stark shift cancellation (magic) wavelength [22]. Optical lattice frequency standards have now demonstrated systematic fractional uncertainties at the $10^{-17}–10^{-18}$ level [5] and also an unprecedented frequency stability [23] opening the path for frequency comparisons beyond the uncertainty of the realization of the SI units in hundreds of seconds [24]. In particular, different groups developed ytterbium optical lattice frequency standards worldwide [23–26] and the spin and angular-momentum forbidden transition $^{1}\text{S}_0 - ^{3}\text{P}_0$ of ytterbium 171 at 578 nm is recommended as a secondary representation of the SI second by the International Committee for Weights and Measures (CIPM). Absolute frequency measurement of this transition has been performed relative to the realization of the second with caesium standards at the National Institute of Standards and Technology (NIST) [8], at the National Metrology Institute of Japan (NMIJ) [27, 28], and at the Korea Research Institute of Standards and Science (KRISS) [26]. Optical frequency ratios between ytterbium and strontium lattice standards were performed at NMIJ [25] and RIKEN [24, 29].
We designed and operated an optical lattice frequency standard based on neutral ytterbium atoms. We report the frequency measurement of the unperturbed clock transition $^1S_0 - ^3P_0$ of $^{171}$Yb against the cryogenic Cs fountain ITCsF2 [1] with a total fractional uncertainty of $5.9 \times 10^{-16}$. The systematic uncertainty of the Yb optical lattice frequency standard is below that of the Cs fountain as well as the statistic uncertainty. Absolute frequency measurements of optical frequency standards are important to show confidence in their reproducibility and are at the base of a possible redefinition of the SI second based on an optical transition.

The paper is organized as follows: in section 2 we describe the realization of the Yb lattice frequency standard. In section 3 we report on the study of systematic frequency shifts and obtain the uncertainty budget for the Yb frequency standard. This is followed in section 4 by a description of the absolute frequency measurement. We finally report and discuss the results.

2. Ytterbium lattice frequency standard

Figure 1 shows the energy levels relevant for the cooling, trapping, and state manipulation of ultracold $^{171}$Yb atoms and for the operation of the frequency standard. $^{171}$Yb is fermionic with nuclear spin $I = 1/2$.

The clock transition $^1S_0 - ^3P_0$ is at 578 nm and has a natural linewidth $< 10$ mHz [30]. The magic wavelength where the light-induced scalar Stark shift cancels for the transition is 759 nm. A bias magnetic field splits the clock transition in two resonances from each $m_F = \pm 1/2$ state. We interrogate with a linearly-polarized probe the two $\pi$-transitions $m_F = +1/2 - m_F = +1/2$ and $m_F = -1/2 - m_F = -1/2$ of which we take the average. By averaging over the two $\pi$-transitions we cancel the first-order Zeeman shift and the vector components of the light shift induced by the lattice [8]. The tensor components of the light shift is zero for spin 1/2 atoms. Furthermore, the atomic ensemble can be prepared in a single spin state enhancing the spectroscopic signal and reducing the shift due to cold collisions [31].

Ytterbium atoms can be cooled to microkelvin temperatures in a two stages magneto-optical-trap (MOT) exploiting the strong $^1S_0 - ^3P_0$ transition at 399 nm (linewidth 29 MHz) and the weaker $^1S_0 - ^3P_0$ transition at 556 nm (linewidth 182 kHz). Then the atoms are loaded in a one-dimensional optical lattice engineered by a retro-reflected laser beam at the magic wavelength 759 nm. The atoms are prepared in a single spin state using an atomic pumping scheme on the $^1S_0 - ^3P_0$ transition. We probe the clock transition using Rabi spectroscopy with a 60 ms pulse at 578 nm. A repumper at 1389 nm resonant with the $^3P_0 - ^3D_0$ transition can be used to pump atoms out of the excited state $^3P_0$ to the ground state $^1S_0$. Finally, a normalized state detection is performed measuring the 399 nm fluorescence with a photomultiplier tube (PMT). The total duration of a single spectroscopy sequence is 310 ms.

Figure 2 displays the experimental setup for atomic spectroscopy. The atomic source is an effusion oven at 400 °C that produces a collimated atomic beam. The distance between the atoms source and the trapping chamber is kept minimal (21 cm) to increase the atomic flux without implementing a Zeeman-slower. The trapping region is a custom aluminum chamber designed for wide optical access with indium-sealed viewports. Ten platinum resistance thermometers are distributed outside the trapping chamber for precise evaluation of the black-body shift. Standard stainless-steel fittings are used for the oven and for the main chamber pumping. Two ion pumps and one non-evaporable getter pump maintains the ultra-high vacuum (pressure $< 10^{-9}$ mbar). A pair of anti-Helmholtz coils generate the magnetic gradient for the MOT. These are vertically arrayed outside the vacuum chamber and are water-cooled by a chiller. Three pairs of Helmholtz coils are compensating the stray magnetic field. Furthermore, all lasers are spatially separated from the vacuum chamber and are delivered to the atoms by polarization-maintaining optical fibers.

2.1. First stage MOT at 399 nm

Atoms are trapped from the atomic beam in the first stage MOT using the strong $^1S_0(F = 1/2) - ^3P_0(F = 3/2)$ transition at 399 nm. The 399 nm radiation is obtained by cavity-enhanced second harmonic generation (SHG) from an amplified, external-cavity diode laser at 798 nm with an output power up to 3 W. The nonlinear medium is a lithium-triborate (LBO) crystal [32]. The diode laser is fiber coupled for spatial mode cleaning and we generate up to 400 mW of light at 399 nm with an input power of 1 W into the enhancing cavity. The laser is locked to the $^1S_0 - ^3P_0$ transition through transverse spectroscopy of an auxiliary ytterbium atomic beam.

The six MOT beams add up to about 30 mW of light power with an $1/e^2$ radius of 1 cm and a detuning of $-20$ MHz. The magnetic field gradient is 0.35 T m$^{-1}$ along the z-axis. The atomic beam is slowed down by a counter-propagating 399 nm
light beam (slower beam) which has a power of about 35 mW and a detuning from resonance of −360 MHz. The leaking field of the MOT coils is exploited to slow the atoms, avoiding the use of an external spatially-varying magnetic field. With the slower beam, we can capture up to $4 \times 10^7$ $^{171}$Yb atoms. Approximately $2 \times 10^5$ atoms are trapped and cooled down to about 1 mK in 50 ms during the clock cycle.

### 2.2. Second stage MOT at 556 nm

The second stage MOT relies on the narrower $^3S^o_0(F = 1/2) - ^3P^o_3(F = 3/2)$ transition at 556 nm. The 556 nm radiation is generated by SHG of an amplified 1112 nm Yb-doped fiber laser in single pass through a periodically-poled potassium titanyl phosphate (PPKTP) crystal. About 10 mW of 556 nm radiation are obtained from 1 W of infrared light. The laser frequency is locked to a 10 cm stable cavity made of Corning ultra-low expansion glass (ULE). The 556 nm MOT beams have a total power of 2 mW, a 1/e 2 radius of 0.5 cm and are kept on during the first MOT stage. The second stage 556 nm MOT is loaded from the 399 nm MOT with an efficiency up to the 70% when the 399 nm beams are turned off. In 60 ms we apply 3 substages at different frequency, intensity, and magnetic field gradient to maximize the fraction of atoms transferred into the lattice. The first substage (30 ms) is designed for maximum atoms transfer between the two MOTs using a magnetic field gradient of 0.13 T m$^{-1}$ and maximum beam intensity. The second substage (20 ms) decreases the atomic temperature down to 10 μK moving the frequency closer to resonance, reducing the intensity 10 times, and increasing the magnetic field gradient to 0.2 T/m. The last substage (10 ms and 0.23 T/m) provides an enhanced transfer efficiency (typically 7%) of the atoms into the red-detuned optical lattice.

### 2.3. Optical lattice at 759 nm

The optical lattice at the magic wavelength 759 nm is provided by a titanium-sapphire laser with an initial output power of about 2 W. The optical lattice is formed by a focused beam tilted from the horizontal plane by 60 mrad and retro-reflected by a curved mirror. The light is delivered using a dedicated fiber and is focused to a 1/e 2 waist radius of 45 μm. The recoil energy from a lattice photon with wavenumber $k_L$, the reduced Planck constant, and $m$ the mass of the atom.

Control of the trapping conditions, thus of the light-induced frequency shifts on the clock transition, requires locking of the lattice frequency at the megahertz level and intensity stabilization. The titanium-sapphire laser has a linewidth below 20 kHz and its frequency is measured with an optical frequency comb. The slow drift of the laser is compensated by implementing a digital lock on the beat-note signal between the laser and the frequency comb. The intensity and thus the lattice trap depth is stabilized by acting on the diffraction efficiency of an acousto-optic modulator (AOM) placed before the fiber. Finally linear polarization of the lattice light is achieved by a Glan-Thompson polarizer placed in front of the input window.

### 2.4. Single spin state preparation

Spin-polarization of the ground state is achieved using optical pumping on the $^3S^o_0(F = 1/2) - ^3P^o_3(F = 3/2)$ transition at 556 nm. A linearly-polarized 556 nm pulse and a bias magnetic field of 0.4 mT are applied along the vertical axis for 1 ms. After spin-polarization we wait 30 ms to let magnetic transients decay.

### 2.5. Clock transition spectroscopy

Spin-polarization of the ground state is achieved using optical pumping on the $^3S^o_0(F = 1/2) - ^3P^o_3(F = 3/2)$ transition at 556 nm. A linearly-polarized 556 nm pulse and a bias magnetic field of 0.4 mT are applied along the vertical axis for 1 ms. After spin-polarization we wait 30 ms to let magnetic transients decay.

The 578 nm ultrastable laser is generated by sum-frequency generation (SFG) in a waveguide periodically-poled lithium...
niobate (PPLN) crystal combining a 1319 nm neodymium-doped yttrium aluminum garnet (Nd:YAG) laser with an erbium fiber laser at 1030 nm [33]. The total power obtained at 578 nm is about 7 mW.

The frequency of the 578 nm laser is actively stabilized via the Pound–Drever–Hall (PDH) method on a 10 cm ultrastable cavity made in ULE featuring fused-silica mirrors with ULE compensation rings. The cavity is suspended horizontally on 4 points in a vibration-insensitive configuration and held in an antiacoustic chamber on top of a passive seismic-isolation platform. A two-stage control using Peltier elements keeps the cavity at the temperature where the thermal expansion coefficient of ULE vanishes (about 28 °C for our cavity). The temperature servo is a digital implementation of the active disturbance rejection control (ADRC) technique [34]. The laser is distributed to the ultrastable cavity, to the vacuum chamber for spectroscopy, and to the fiber-based optical comb by polarization-maintaining optical fibers. All fiber links are actively phase-stabilized using AOMs. From measurements with similar cavities we expect this laser to have a flicker frequency instability less than $3 \times 10^{-15}$ at 1 s of integration time [33]. The cavity drift is less than 0.1 Hz s$^{-1}$ and we use a feed-forward correction on a synthesizer to cancel it.

Spectroscopy of the clock transition is achieved with a 578 nm laser beam with a 1/e$^2$ waist radius of 200 µm collinear to the lattice with an incident power of 70 mW. The lattice retro-reflector is anti-reflection coated at 578 nm with a reflectivity of <1% at this wavelength. The probe linear polarization is aligned with the lattice polarization and to the vertical bias magnetic field using the same Glan-Thompson polarizer. The frequency of the probe light is controlled by a double-pass AOM placed before the fiber link to the atoms (clock AOM in figure 2). Probe intensity is actively stabilized. During the interrogation a 0.14 mT magnetic field is applied leading to a 290 Hz Zeeman shift of the two $\pi$-transitions from the unperturbed one (figure 3). Pulsing of the clock laser is achieved by detuning the applied frequency by 1 MHz. A $\pi$ pulse is applied for a Rabi time of 60 ms. Mechanical shutters stop all other laser beams during spectroscopy.

The resulting excitation fraction is then destructively detected and normalized applying three resonant pulses at 399 nm and by collecting the atomic fluorescence with the PMT. The detection beam has a power of 2.5 mW, a waist of 0.5 mm and it strongly saturates the $^3S_0 - ^3P_1$ transition. For each pulse the PMT signal is digitally integrated for 2 ms. The first pulse measures the population in the $^3S_0$ state. The second one reveals the background signal from scattered light and excited hot background gas atoms. A third 399 nm pulse then reveals the atoms that had been excited during the Rabi spectroscopy after repumping the atom population of the $^3P_0$ state over the short-lived $^3D_1$ level back to the $^3S_0$. For this task we use a commercial pigtail distributed feedback laser emitting at 1389 nm. The 10 mW output beam of the repumper laser is focused on the atoms thus broadening the $^3P_0 - ^3D_1$ transition to 300 MHz. Active frequency stabilization is not required and the population transfer efficiency is 90% after 6 ms. We use the fluorescence from the three pulses to digitally calculate the normalized excitation fraction and the total number of atoms.

The clock transition is observed with a typical linewidth of 20 Hz (see for example figure 4). The linewidth is limited by excessive clock-laser frequency noise as the seismic noise on the ultrastable cavity is not properly cancelled by the vibration-insensitive supports and by the passive isolation platform.

To gain a better understanding of the optical trapping conditions we performed broadband spectroscopy and observed the sideband spectrum (figure 5). From the spectroscopy results and by fitting the spectrum we extracted the trap depth $U_0$ as the longitudinal motional frequency $\nu_z$ is given by $h\nu_z = 2E_R\sqrt{|U_0/E_R|}$. Furthermore the longitudinal temperature of the atoms $T_z$ can be measured from the ratio $b_{n\pi}$ between the excitation fraction of the blue and red sideband $k_BT_z = h\nu_z/b_{n\pi}$ [35], where $k_B$ is the Boltzmann constant. For example figure 5 shows a trap depth of $284(3)E_R$ and a temperature of 7.0(14) µK. Typically we operate the optical trap at a depth of 200 $E_R$ at a temperature of 7(3) µK.

**Figure 3.** Measured $^3S_0 - ^3P_1$ transition in presence of an external magnetic field of $B = 0.14$ mT and $\pi$-polarized light with the atoms polarized in the $m_F = +1/2$ state (green dots) or $m_F = -1/2$ state (blue squares). Solid lines are lorentzian fits of the resonances. We observed a 98% single spin state purity.

**Figure 4.** Spectroscopy of a single magnetic component of the $^3S_0 - ^3P_1$ transition with spin-polarized atoms. Blue points are experimental data from a single scan of the resonance. The green line is a lorentzian fit with a linewidth of 18 Hz.
2.3. Lattice light shifts

The induced lattice light shift can be written as [24, 36]:

\[ \Delta \nu_{\text{ls}} = \left( a \Delta \nu + \frac{3}{4} d(2n^2 + 2n + 1) \left( \frac{U_L}{E_r} \right) - d \left( \frac{U_L}{E_r} \right)^2 \right) \]

\[ + (a \Delta \nu - b) \left( n + \frac{1}{2} \left( \frac{U_L}{E_r} \right)^{1/2} + d(2n + 1) \left( \frac{U_L}{E_r} \right)^{3/2} \right) \]

where \( \Delta \nu \) is the lattice frequency shift from the electric-dipole (E1) magic wavelength \( \nu_E \), and \( U_L = \xi U_0 \) is the effective trap depth that takes into account the reduced laser intensity seen by the atoms. The coefficient \( a \) is the linear slope of the polarizability per lattice frequency, while \( b \) is a multipolar polarizability, and \( d \) the hyperpolarizability coefficient. \( n \) is the average motional state occupation in the lattice.

We determine the linear part of the shift and the magic wavelength by interleaving two clock cycles operating at different lattice intensities. Experimentally we measure the

2.6. Clock operation

The clock laser is locked to the average frequency of the two \( m_F = \pm 1/2 \) components removing the first-order Zeeman shift as well as the vector light shift. We alternatively interrogate the side of each transition to derive two error signals and we use an independent lock for each. In total four interrogations are performed: two for the \( m_F = +1/2 \) state and two for the \( m_F = -1/2 \) state set at the transitions sides. Addressing the four points is controlled by frequency shifting the clock AOM (figure 2), controlling also the Rabi spectroscopy by pulsing the clock laser frequency. A computer program sets and records the frequencies of the two transitions. We then post-process the data to extract the average and splitting of the two resonances.

To evaluate systematic shifts we interleave two clock sequences for a total of four independent locks. We then post-process the data to measure the frequency difference between the two interleaved states interpolating at common timetags. Figure 6 shows an example of the instability of an interleaved measurement for a total measurement time of 22.5h. The instability is compatible with white frequency noise as \( 2.3 \times 10^{-14}\sqrt{\tau/\text{fs}} \) and is limited by the same cavity noise affecting the linewidth.

3. Systematic frequency shifts

Table 1 shows the uncertainty budget for the \(^{171}\text{Yb} \) optical lattice clock. The fractional uncertainty for the measurement reported here is \( 1.5 \times 10^{-16} \). The contributions of the systematic effects will be discussed below.

3.1. Lattice light shifts

The induced lattice light shift can be written as [24, 36]:

\[ \Delta \nu_{\text{ls}} = \left( a \Delta \nu + \frac{3}{4} d(2n^2 + 2n + 1) \left( \frac{U_L}{E_r} \right) - d \left( \frac{U_L}{E_r} \right)^2 \right) \]

\[ + (a \Delta \nu - b) \left( n + \frac{1}{2} \left( \frac{U_L}{E_r} \right)^{1/2} + d(2n + 1) \left( \frac{U_L}{E_r} \right)^{3/2} \right) \]

\[ \Delta \nu_{\text{ls}} = \left( a \Delta \nu + \frac{3}{4} d(2n^2 + 2n + 1) \left( \frac{U_L}{E_r} \right) - d \left( \frac{U_L}{E_r} \right)^2 \right) \]

\[ + (a \Delta \nu - b) \left( n + \frac{1}{2} \left( \frac{U_L}{E_r} \right)^{1/2} + d(2n + 1) \left( \frac{U_L}{E_r} \right)^{3/2} \right) \]

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frequency difference between \( U_0 = U_L \) and \( U_0 = U_H \) and calculate the resulting shift at \( U_0 = U_L \) only assuming a linear shift. We then correct these values and calculate the hyperpolarizability and multipolar shifts using equation (1) and the values \( b = -0.68(71) \) mHz and \( d = -1.9(8) \) µHz known for Yb [24, 37].

The lattice intensity is modulated changing the diffraction efficiency of the AOM before the fiber delivering the laser to the atoms. This method preserves the frequency spectrum of the titanium-sapphire laser between high and low power. Atoms are always loaded at low power with the titanium-sapphire laser between high and low power. This split gives a continuous measurement at about 0.1 ms.

We use a fast photodiode (bandwidth 3 GHz) to check that the titanium-sapphire laser is in single mode. The broadband emission spectrum of the laser is not filtered out for the present measurement.

We measured the shift at the working point \( \nu_1 = 394.798.238 \) GHz between \( U_L = 196(4)E_r \) and \( U_H = 348(5)E_r \). The effective shift at \( U_L \) is \(-0.02(4) \) Hz.

The lattice depths, \( \xi \) and \( n \) were determined by sidebands spectroscopy (figure 5). For our trap configuration, \( \xi = 0.8(1) \) and \( n = 2.1(10) \) at \( U_L \) and \( n = 1.5(8) \) at \( U_H \). We correct the nonlinearity between \( U_H \) and \( U_L \) in the effective shift and calculate the hyperpolarizability and multipolar shifts using equation (1). These two contributions are \(-0.13(6) \) Hz and \( 0.664(32) \) Hz and sum to \(-0.07(5) \) Hz. This is the largest contribution to the Yb uncertainty and is due to the uncertainty in the multipolar and hyperpolarizability coefficients at high trap depths.

We also measured the lattice shift as a function of the lattice frequency. Figure 7 shows the scalar shift measured between a trap depth of \( U_L = 208(1)E_r \) and \( U_H = 333(2)E_r \) as a function of lattice frequency. Fitting the data with a second order Taylor expansion around the magic wavelength, the linear coefficient is \( a = -27(3) \) mHz GHz\(^{-1} \) and the quadratic coefficient is \( c = -0.16(3) \) mHz GHz\(^{-2} \) to give a shift \( c\Delta\nu^2U_L/E_r \). Uncertainty is limited by \( \xi \) and not by the statistical uncertainty in the fit.

We calculated the E1-magic frequency from the measured linear coefficient \( a \), the measured shift at the working point, and the nonlinear corrections. For our system it is 394.798.205(17) GHz. This measurement of the magic frequency and the ones reported before [8, 24] are not consistent with each other as they could depend on the spectral features of the titanium-sapphire lasers if not filtered [38]. Our data is consistent with a constant shift coming from the spectrum of the laser.

### 3.2. Density shift

In \(^{171}\)Yb optical frequency standards the density-dependent shift is dominated by p-wave cold collision [39] while s-wave scattering can arise from inhomogeneity of the Rabi excitation [40]. To evaluate the density shift we run interleaved measurements varying the number of atoms. This is modulated by changing the duration of the slower beam pulse at 399 nm during the first stage MOT. This method impacts on the atomic density without changing the trapping conditions and we assume the atomic density to be proportional to the atoms number. The trap region has a radius of 45 µm, given by the lattice waist, and is 0.5 mm long, given by the size of the second stage MOT. For our setup an atomic density of \( \rho_0 = 4 \times 10^{-14} \) m\(^{-3} \) corresponds to about 1 atom per lattice well. The absolute atom number has a fractional uncertainty of 20% but the relative density measurements have an uncertainty of about 1%. During our measurements, we varied the atomic density from \( \rho = 0.1\rho_0 \) to \( \rho = 3\rho_0 \).

The density shift depends on the details of the Rabi pulse used for spectroscopy and whether the atomic sample is spin-polarized [31]. Figure 8 shows the density shift measurement as a function of the atomic density difference interleaving between high and low number of atoms with a spin-polarized sample and a final excitation fraction (measured after the Rabi pulse) of 0.39(2), given by our choice of the locking point on the resonance. We fit the data with a linear function with no offset and measured a slope of \(-0.01(4) \) Hz/\( \rho_0 \). At a different locking point and an final excitation fraction of 0.34(2) we measured in a similar way a slope of \(-0.4(2) \) Hz/\( \rho_0 \). The dependence on excitation fraction is predicted by p-wave collisions [31]. For our conditions \( \rho = 0.7\rho_0 \) and excitation fraction 0.39(2) the density shift is \(-0.01(3) \) Hz.

### 3.3. Zeeman shift

To measure the clock transition we apply a magnetic field and the Zeeman effect split the two \( \pi \)-transition resonances between \( m_F = \pm 1/2 \) states. This split gives a continuous measurement of the magnetic field experienced by the atoms while running the experiment. The operational Zeeman shift for the two components from the center is 290.1(3) Hz corresponding to \( B = 138(1) \) µT [8]. We observed no drift of the magnetic field.
field down to 0.1 \( \mu T \) h\(^{-1}\). Taking the average frequency of the two \( m_\perp = \pm 1/2 \) magnetic components removes the first-order Zeeman shift. We calculated a quadratic Zeeman shift of \(-0.14(2) \) Hz using the quadratic Zeeman coefficient known for Yb, 7(1) Hz mT\(^{-1}\) [8].

3.4. Black-body radiation shift

The black-body radiation (BBR) emitted by bodies at temperature \( T = 0 \) K surrounding the atomic cloud induces a systematic Stark shift of the clock transition frequency according to [41]

\[
\Delta \nu_{\text{BBR}} = -\frac{1}{2} \left( \alpha_1^{(0)} + \alpha_2^{(0)} \right) \langle E_0^2 \rangle_T (1 + \eta_{\text{clock}}(T)).
\] (2)

The coefficients \( \alpha_1^{(0)} \) are the static polarizabilities of the clock transition ground and excited states, \( \langle E_0^2 \rangle_T \) is the average electric field radiated by a black body at temperature \( T \), \( \eta_{\text{clock}} \) is a dynamic correction. For \(^{171}\)Yb is \( \Delta \alpha^{(0)} = (\alpha_1^{(0)} - \alpha_2^{(0)}) = 36.2612(7) \) kHz (kV cm\(^{-2}\)) and \( \eta(T = 300 \text{ K}) = 0.0179(5) \) [41, 42].

We kept into account the vacuum package and the Yb oven to evaluate the BBR shift.

The vacuum package temperature has been continuously monitored at ten different positions by platinum resistance thermometers Pt1000. Gradients are caused by the power dissipated by the MOT coils, by the temperature of the water cooling, and by the temperature of the oven. Eight Pt1000 are placed on the chamber enclosing the atom trapping region while other two are placed respectively close to the oven and far away from it. The Pt1000 calibration uncertainty is 0.2 K while the 4-wires resistance reading uncertainty is 0.06 K. The temperature spread between different thermometers lays typically within 2 K. We observed bigger inhomogeneities during transients at the start of the experiment, while changing the magnetic field gradient in the MOT coils, or increasing the oven temperature. Figure 9 shows an example of the recorded temperatures during a measurement.

We assumed a rectangular probability density function for the temperature seen by the atoms, taking as average value \( T = (T_{\text{max}} + T_{\text{min}})/2 \) and as uncertainty \( (T_{\text{max}} - T_{\text{min}})\sqrt{12} \) where \( T_{\text{max}} \) and \( T_{\text{min}} \) are the maximum and minimum temperature recorded during a single run [43]. The uncertainty coming from the temperature inhomogeneity for this measurement is 0.8 K. This approach leads to a BBR shift due to the environment temperature of \(-1.219(13) \) Hz.

The BBR emitted by the oven reaches the atom trapping region by passing through a copper vacuum-differential tube which is 15 cm long and has a 0.5 cm inner diameter. Thus the atoms are affected by both the radiation from the oven with direct line of sight and the radiation reaching the atoms after being reflected on the tube inner surface. A ray tracing analysis has been performed to calculate the effective solid angle under which the atomic cloud sees direct and reflected radiations. In the effective solid angle calculation the emissivity of the tube material is also kept into account as \( \epsilon = 0.4(2) \) and two reflections are relevant in the shift calculation.

The oven temperature of 400(10)°C has been continuously monitored using a thermocouple. The uncertainty considers the thermocouple accuracy and differences in temperature between the oven tip and the sensor position. The BBR shift contribution from the oven is then calculated to be \(-0.011(4) \) Hz.

For the first measurements in this campaign we had also a window heated at 230(5)°C with direct sight to the atoms. For the effective solid angle underlying the hot window, we modeled up to 5 reflections from stainless steel surfaces (fittings to the window) and 1 reflection from a polished aluminum surface (main chamber), with the material emissivity \( \epsilon_{\text{steel}} = 0.5(3) \) and \( \epsilon_{\text{Al}} = 0.13(10) \). The resulting shift was \(-0.12(15) \) Hz. After February 2016 we replaced the hot window with a cold intra-vacuum mirror thus removing this.

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**Figure 8.** Density shift measured interleaving between high and low atom numbers with an excitation fraction of 0.39(2). Blue points mark the experimental data with their uncertainty. The green line and the shaded region show a linear fit with its statistical uncertainty.

**Figure 9.** Example of the temperatures of the ten Pt1000 platinum resistance thermometers surrounding the vacuum chamber recorded during a single measurement run. Solid colored lines mark the temperature of the eight resistance thermometers on the trapping chamber. The black dashed line is the temperature recorded close to the atomic oven while the black dotted line is the temperature measured far away.
shift and source of uncertainty. The contribution of the hot window BBR to the final relative uncertainty for the measurement presented later is at the $5 \times 10^{-17}$ level as we averaged data with and without this shift.

3.5. Probe light shift

The interaction of the probe laser with the atoms during Rabi spectroscopy causes a Stark shift on the transition. We have calculated this shift by using the published value [8] and noticing that for a $\pi$ pulse the laser intensity scales as the inverse of the square of the Rabi time. From this rescaling it follows that the Stark shift induced by the probe laser is $0.005(18)$ Hz.

3.6. Gravitational redshift

The difference between the height of the Yb atoms and the integrated height of the Cs atoms along the free-flight trajectory is $h(\text{Yb} - \text{Cs}) = 0.15(2)$ m. The gravitational redshift for the comparison is then $1.6(2) \times 10^{-17}$.

The gravitational potential relative to the geoid at INRIM’s laboratories was measured in 2013 in the context of the European project ‘International timescales with optical clocks’ (ITOC) [44, 45]. The measurement was performed both by a levelling approach and by a Global Navigation Satellite System/geoid (GNSS/geoid) approach [46]. The gravitational potential at the Yb atoms position is $C(\text{Yb})_{\text{lev}} = 2336.25(35) \text{ m}^2 \text{s}^{-2}$ as determined by levelling and $C(\text{Yb})_{\text{GNSS/geoid}} = 2336.48(27) \text{ m}^2 \text{s}^{-2}$ as determined by GNSS/geoid [47] relative to the International Earth Rotation and Reference Systems Service (IERS) 2010 conventional reference potential $W_0 = 62,636,856.0(5) \text{ m}^2 \text{s}^{-2}$ [48, 49]. Geometric levelling is accurate over short distances but is susceptible to systematic errors which may accumulate over distances $>1000$ km [50]. Therefore the GNSS/geoid approach is beneficial in the comparison between different laboratories by satellite or long-haul optical fiber links. In this case the GNSS/geoid value gives a gravitational redshift on the Yb atoms of $13.47(3)$ Hz with a fractional uncertainty of $6 \times 10^{-18}$ limited by the uncertainty on the reference $W_0$.

The corresponding numbers for ITCsF2 are $C(\text{Cs})_{\text{lev}} = 2334.8(10) \text{ m}^2 \text{s}^{-2}$ and $C(\text{Cs})_{\text{GNSS/geoid}} = 2335.0(10) \text{ m}^2 \text{s}^{-2}$. Using the GNSS/geoid value the redshift for ITCsF2 is then $2.599(1) \times 10^{-14}$.

3.7 Other shifts

We measured that the phase-stabilized fiber links between the cavity and the atoms contribute a fractional uncertainty of $1 \times 10^{-18}$ after 1000 s of integration time. However the fiber link to the atoms is referenced to a mirror placed 30 cm from the lattice back-reflector, that is the reference for the lattice. We discovered some phase-noise between the two references which include noise synchronous with the clock cycle (e.g. MOT coils switching). We measured a fractional contribution of $5 \times 10^{-17}$ that we take as uncertainty coming from the Doppler shift.

A static electric field would cause a Stark shift on the clock transition [41]. Our vacuum chamber is made of aluminum and is a Faraday cage from external electric fields. However it is possible for electric charges to be present on the windows of the vacuum chamber [51]. In our setup two fused-silica windows are 25 mm away from the atoms and have a radius of 38 mm and a thickness of 12 mm. We estimated a capacitance of $0.6 \text{ pF}$ between the two surfaces and a time constant of $\sim 50$ d assuming a high resistivity of $10^{18} \text{ ohm}$ for fused silica [51]. We have operated our vacuum system for more than 3 years and possible charges present at the beginning should have decayed to give a Stark shift $< 1 \times 10^{-17}$ that we take as uncertainty.

The frequency shift due to collisions with the hot atomic beam and background gas can be related to the lifetime of the atoms in the trap [52]. From our observed lifetime of 2.7 s and the $C_{6}$ coefficients for Yb dimers [53] we assign a shift of 8 mHz that we take as uncertainty.

Tunnelling of atoms from one lattice site to the other leads to a broadening of the band structure of the lattice [54] and in our trap is suppressed by the deep potential. Given our trap depth of $U_{0} = 200 E_{r}$ and temperature of $T = 7(3) \mu \text{K}$ we calculated that 70% of the atoms occupy the lowest three band in the lattice [54]. We then take the width of the third band 2 mHz as uncertainty.

We calculated the line pulling from the lattice sidebands to be 2 mHz that we take as uncertainty. The line pulling from the $\pi$ and $\sigma$ transitions between $m_{JF} = \pm 1/2$ states is negligible because of spin-polarization of the atoms and the linear polarization of the probe light.

We measured a possible error in the servo loops taking the average of the error signal measured during the campaign that gives a fractional uncertainty of $1 \times 10^{-17}$. We switch the frequency of the AOM keeping a constant power to avoid thermal effects. We did not detect any shift up to 2 mHz by heterodyne interferometry, that we take as uncertainty. The switching of the AOM to apply the Rabi pulse introduces a phase jump in the phase-stabilization. We calculated a frequency shift of 0.5 mHz [55] that we take as an uncertainty.

4. Absolute frequency measurement

We measured the absolute frequency of the $^{171}$Yb frequency standard relative to the caesium fountain ITCsF2 from January 2016 to the end of May 2016. The comparison has been performed with a fiber frequency comb with a repetition rate of 250 MHz referenced to a hydrogen maser. The fountain ITCsF2 continuously measured the maser frequency for the whole duration of the campaign.

The light at 578 nm resonant with the cavity was sent to the frequency comb through an optical fiber link with active phase-stabilization. The fiber link to the comb contributes to an uncertainty of $1 \times 10^{-18}$ above 1000 s of integration time. Two different computers measured the frequency of the cavity-stabilized light relative to the frequency comb and the frequency difference between the cavity and the ytterbium atoms of the optical standard. The frequency comb and
the optical standard recorded data at different rates and we averaged and interpolated the data to common timestamps. Both computers were synchronized to INRIM’s Network Time Protocol (NTP) server within 3 ms. Since the maximum observed frequency drift of the cavity was 0.1 Hz s\(^{-1}\), we assume an uncertainty of \(6 \times 10^{-19}\) coming from synchronization.

The comb contribution to uncertainty was assessed by measuring the same laser frequency with two independent combs and is \(1 \times 10^{-16}\), limited by the comb electronics noise at an averaging time of \(1 \times 10^4\) s. The frequency of the beat note between the cavity-stabilized laser and the comb was measured redundantly to detect cycle slips in the counter. We discarded all points where the counter readings did not agree at better than 0.2 Hz.

The cryogenic fountain ITCsF2 ran in two different atomic density regimes [1] during the campaign. In the low-density regime ITCsF2 contributes to an uncertainty of \(3.0 \times 10^{-16}\) with a stability of \(3.6 \times 10^{-13}/\sqrt{\text{Hz}}\). In the high-density regime it contributes to an uncertainty of \(4.8 \times 10^{-16}\) with a stability of \(2.5 \times 10^{-13}/\sqrt{\text{Hz}}\). The measurement of the hydrogen maser frequency relative to ITCsF2 was considered only in the windows of operation of the ytterbium standard.

We took a total of 31 different measurements for a total measurement time of the optical frequency standard of 227 h. Figure 10 shows the different measurements as a function of the modified Julian date (MJD). For each measurement the statistical uncertainty is dominated by the uncertainty of the cavity-stabilized laser. The systematic uncertainty varies for each measurement because of the different density values of both the fountain and optical frequency standard and because of the reduced BBR uncertainty contribution on the ytterbium after removing the hot window after the first 10 measurements. We applied a statistical analysis of the data based on the Gauss–Markov theorem [56, 57] that considers the correlations between the different measurements coming from the systematic shifts [58]. The measurement result is \(f_{\text{Yb}} = 518\,295\,836\,590\,863.59(31)\) Hz where the uncertainty is detailed in table 2. The reduced chi squared with 30 degrees of freedom for the average is \(\chi^2_{\text{red}} = 1.36\) and is calculated from the complete correlation matrix of the measurements [58]. This chi squared has a p-value of 10%. The total fractional uncertainty is \(5.9 \times 10^{-16}\) and it is dominated by the uncertainty of the caesium fountain. The fountain systematic uncertainty contributes for a total of \(4.0 \times 10^{-16}\). The statistical uncertainty resulted \(3.4 \times 10^{-16}\) but we increased this value to \(3.9 \times 10^{-16}\) to account for \(\chi^2_{\text{red}} > 1\). A contribution of \(1.6 \times 10^{-16}\) comes from the ytterbium optical frequency standard as explained above.

Figure 11 compares our value with previous measurements and the recommended frequency for \(^{171}\)Yb as a secondary representation of the second by caesium standards. Circles denote measures relative to the realization of the second by caesium standards.
\[ f_{\nu_C} (\text{CIPM2015}) = 429\,228\,004\,229\,873.2(2) \text{ Hz} \text{ as a secondary representation of the second with its uncertainty of } 5 \times 10^{-16} \ [59]. \]

5. Conclusions

We report the absolute frequency measurement of the unperturbed \(^{171}\text{Yb}\) with a fractional uncertainty of \(5.9 \times 10^{-16}\) performed in an optical lattice frequency standard and limited by the caesium fountain. The measurement result is \(f_{\nu_C} = 518\,295\,836\,590\,863.59(31) \text{ Hz}\) where the number in parentheses is the numerical value of the standard uncertainty referred to the corresponding last digit of the quoted result. Our measurement is independent from previous measurements of the transition and the result is in agreement with the recommended value for \(^{171}\text{Yb}\) as a secondary representation of the SI second. This measurement of the ytterbium transition is the first in Europe after previous measurements achieved in North America and Asia and is important in view of a possible redefinition of the SI second based on optical frequency standards.

We characterized the ytterbium optical frequency standard and evaluated its fractional uncertainty for the present measurement to \(1.6 \times 10^{-16}\). We expect that this uncertainty can be reduced to the \(1 \times 10^{-17}\) level in the near future. In this measurement the Yb standard is limited by the poor short-term stability due to excessive noise on the ultrastable cavity. Improving the cavity environment and the performance of the clock laser will allow us to reduce the uncertainty measured by interleaving to \(1 \times 10^{-17}\) (lattice shift and density shift). The nonlinear part of the lattice shift can be reduced by working with smaller lattice depths and by improving the measurement of multipolar and hyperpolarizability effects. The Zeeman shift can be reduced by working with smaller magnetic fields. The Doppler shift can be easily cancelled by referencing the clock laser to the lattice mirror. The implementation of a mechanical shutter in the atomic beam path will further reduce the effects of the background collisions and the BBR shift coming from the hot atomic oven. The static Stark shift can be better evaluated by placing electrodes on the vacuum chamber windows. The final limitation of the current setup is the temperature uniformity of the vacuum chamber leading to a total BBR contribution with a fractional uncertainty of \(2.5 \times 10^{-17}\). Improving the thermal isolation from environment and heat sources (e.g. the MOT coils) and waiting for thermal transient to decay will be beneficial in further reducing this contribution to \(1 \times 10^{-17}\).

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