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Original
Ytterbium optical lattice clock at INRIM / Pizzocaro, Marco; Bregolin, Filippo; Milani, Gianmaria; Rauf, Benjamin; Thoumany, Pierre; Costanzo, Giovanni Antonio; Levi, Filippo; Calonico, Davide. - (2015), pp. 300-303. ((Intervento presentato al convegno 2015 Joint Conference of the IEEE International Frequency Control Symposium and the European Frequency and Time Forum, FCS 2015 tenutosi a Colorado Convention Center (CCC), usa nel 2015.

Availability:
This version is available at: 11583/2683401 since: 2017-09-29T10:54:01Z

Publisher:
Institute of Electrical and Electronics Engineers Inc.

Published
DOI:10.1109/FCS.2015.7138846

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Ytterbium optical lattice clock at INRIM

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Abstract— We present an optical lattice clock based on ytterbium \(^{171}\)Yb atoms developed in the laboratories of INRIM. In the experiment, we cool and trap ytterbium atoms in a two stage magneto-optical trap (MOT) (at 399 nm and 556 nm for the first and second stage, respectively). Atoms are then transferred in a horizontal, one-dimensional optical lattice at the magic wavelength (759 nm). Here the clock transition at 578 nm is probed by a laser stabilized on an ultra-stable cavity. We describe the generation of all the laser sources, the physic package and the operation of the clock. Lasers at 399 nm, 556 nm and 578 nm are obtained, with different techniques, using non-linear crystals starting from infrared sources. The clock laser is stabilized using a high finesse notched ULE cavity. The lattice is made with a titanium-sapphire laser. The aluminum vacuum chamber is designed for wide optical access and its temperature is measured by 8 thermistors for blackbody shift evaluation. Our system allows for fast loading of the lattice with \(1 \times 10^9\) atoms trapped in the lattice in 250 ms. We obtained preliminary spectroscopy results and we locked the clock laser to the atomic line. Future perspectives are discussed.

I. INTRODUCTION

Among other the clock transition \(^1S_0\)\(^-^3P_0\) at 578 nm of ytterbium \(^{171}\)Yb neutral atom is recommended as secondary representations of the SI second reflecting the measurements made at National Institute of Standards and Technology (NIST) [1], [2], at the National Metrology Institute of Japan (NMIJ) [3] and at the Korea Research Institute of Standards and Science (KRISS) [4]. Figure 1 show the Ytterbium transitions relevant for clock operations. Note that \(^{171}\)Yb is fermionic with nuclear spin \(I = 1/2\) and further hyperfine structure is present. Ytterbium is easy to cool and trap in a double stage magneto-optical trap (MOT) exploiting the strong transition at 399 nm \(^1S_0\)\(^-^1P_1\) (linewidth 29 MHz) and achieving microkelvin temperatures with the weaker transition at 556 nm \(^1S_0\)\(^-^3P_1\) (linewidth 182 kHz). Ytterbium can then be loaded in an optical lattice at the magic wavelength (759 nm). A repumper at 1389 nm resonant with the \(^1P_0\)\(^-^3D_1\) transition can be used to pump atoms from the clock state \(^3P_0\) to the ground state \(^1S_0\).

We are developing a ytterbium optical lattice clock based on \(^{171}\)Yb. The experimental setup and the laser ensemble are complete while the characterization of the clock is under way. In the following we will describe the physic package, the laser ensemble and the operation of our clock as well as first spectroscopy results and future perspectives.

II. EXPERIMENTAL SETUP

A. Physic package

Figure 2 shows a scheme of the physic package and of the vacuum chamber. The atomic source is an effusion oven at 400 °C that produces a collimated atomic beam. Atoms are trapped in a custom aluminum chamber with indium-sealed viewports, designed for wide optical access. The chamber is designed without a Zeeman slower. Instead the distance between the trapping region and the atomic source is made as short as possible to maximize the atomic flux. During operation the chamber is kept in ultra-high vacuum (pressure \(< 10^{-9}\) mbar) by two ion pumps and one non-evaporable getter pump.

The water-cooled MOT coils are outside the vacuum chamber, in the vertical direction. Three pairs of Helmholtz coils are used to compensate the stray magnetic field. Eight thermistors are placed on the aluminum vacuum chamber for blackbody shift evaluation.

B. Laser ensemble

The 399 nm radiation is obtained by second harmonic generation (SHG) from a 798 nm titanium-sapphire (Ti:sapphire)
laser using a lithium triborate (LBO) crystal in an enhancement cavity [5]. The Ti:sapphire laser has an output power of 1.1 W pumped by a 8 W solid state pump laser at 532 nm. Up to 0.9 W has been obtained but a typical output of 0.5 W at 399 nm is used for the experiment. The frequency can be locked to the atomic resonance of any ytterbium isotopes by transverse spectroscopy on an auxiliary atomic beam. Polarization-maintaining optical fibers deliver the 399 nm light to the atoms for the first stage MOT, the slower beam and the detection probe beam.

The 556 nm radiation is obtained by SHG from 1112 nm amplified, ytterbium doped fiber laser using a single-pass periodically-poled potassium titanyl phosphate (PPKTP) crystal. Typically 10 mW of 556 nm light are obtained starting from 1.0 W of infrared light. The frequency of the 556 nm laser is locked to the resonance of a Corning Ultra Low Expansion glass (ULE) cavity, with acousto-optic modulators (AOMs) bridging the gap to the frequency of 171 Yb. The green laser is sent to the second stage MOT by polarization-maintaining optical fiber.

The lattice is made by a Ti:sapphire laser pumped by a solid state pump laser at 10 W. An AOM is used as an optical isolator and for power-stabilization. Typically 1 W of light is sent to the atoms using a polarization-maintaining optical fiber.

The clock laser at 578 nm is obtained by sum frequency generation (SFG) in a waveguide periodically-poled lithium niobate (PPLN) crystal using an erbium fiber laser at 1030 nm and a neodymium-doped yttrium aluminium garnet (Nd:YAG) laser at 1319 nm [6]. The output power is typically 4 mW. The frequency is stabilized using the Pound-Drever-Hall technique on a 10 cm ultra-stable cavity made by ULE, with fused-silica mirrors and ULE rings. The temperature of the cavity is stabilized by a double stage control with Peltier elements to the point of zero coefficient of thermal expansion of ULE. The temperature control is a digital implementation of the powerful Active Disturbance Rejection Control (ADRC) technique [7]. The clock light is delivered to the cavity, to the atoms and to a fiber comb by compensated optical fiber links.

A solid state laser at 578 nm is under development. The radiation is obtained using a co-doped LiLuF₄: Dy₃⁺Tb₃⁺ fluoride crystal pumped by 450 nm diode laser. [8]. This new compact system with its unique direct emission at 578 nm could replace the current laser in the future.

A pigtail distributed feedback laser at 1389 nm is used as repumper from the clock state to the ground state. It has an output on the atoms of 10 mW and a frequency variation < 100 MHz. The power is enough to power-broadened the ytterbium line to 300 MHz so that active frequency stabilization is not needed.

C. Clock cycle

The clock experimental setup is sketched in fig. 3. First 171 Yb atoms are trapped in a 399 nm MOT from the atomic beam. The six laser beams have a total power of about 30 mW with a 1/e radius of 1 cm and a detuning of −20 MHz. The magnetic field gradient is 0.4 T/m along the vertical axis. A seventh beam, with a power of 50 mW and a detuning of −360 MHz is focused counter-propagating to the atomic beam and acts as a slower. There is no dedicated magnetic field to make a Zeeman slower but we exploit the leaking field of the MOT coils. With the slower beam, we can capture up to $4 \times 10^7$ 171 Yb atoms. For clock operations, we capture typically $1 \times 10^6$ atoms in 150 ms of 399 nm MOT.

The second stage 556 nm MOT is loaded from the 399 nm MOT with an efficiency up to the 70 % simply turning off the 399 nm beams. The 556 nm beams have a total power of 2 mW, a 1/e radius of 0.5 cm and are left on during the first stage. In 60 ms we apply 3 stages at different frequency, intensity and magnetic field gradient to maximize the fraction of atoms transferred in the lattice. The first stage (30 ms) is tuned to maximize the number of atoms in the 556 nm MOT and has a magnetic field gradient of 0.25 T/m. The second stage (20 ms) the frequency is brought closer to resonance and the magnetic field reduced to 0.18 T/m to minimize the atoms temperature, that is reduced to 10 µK. The lifetime of this stage is 3 s. In the third stage (10 ms) the magnetic field gradient is increased back to 0.25 T/m and the frequency is tweaked to maximize the number of atoms in the lattice.

The lattice laser is delivered to the atoms by polarization-maintaining optical fiber and is focused by an achromatic lens to a waist of 45 µm. The laser is retro-reflected by a curved mirror to form a lattice with a depth of 300 recoil energies for 1 W of power. The lattice is horizontal. We trap typically up to $5 \times 10^3$ atoms in the lattice while the maximum number of atoms we trapped was $3 \times 10^4$. Lifetime of atoms in the lattice is 3.0 s.

Atoms in the lattice are probed by the clock laser at 578 nm. The clock laser is collimated with a waist of 200 µm collinear...
to the lattice and is sent to the atoms through the lattice back-reflector, that is transparent at this wavelength. The lattice and clock polarization are aligned vertically. Other than the 578 nm light, all other radiations are stopped by mechanical shutters during spectroscopy.

The spectroscopy of the atoms is performed by detecting with a photomultiplier tube the fluorescence from the atoms by 3 pulses of resonant light at 399 nm of the duration of 1 ms. The first pulse measures the atoms left in the ground state. The second pulse is used to subtract the background from scattered light and the atomic beam. After the second pulse the repumper laser at 1389 nm is used to pump back atoms from the clock state to the ground state in 12 ms. Then the third pulse detect the number of excited atoms.

Atoms can be prepared for interrogation in around 250 ms. We used clock pulses (Rabi pulses) of typically 50 ms to 100 ms. Total cycle length with detection is between 350 ms to 450 ms.

III. FIRST SPECTROSCOPY RESULTS

Figure 4 show the spectroscopic signal from $1 \times 10^4$ atoms in the lattice, while interrogated by 100 $\mu$W pulses of 578 nm light of 100 ms. From the fit of the shape of the sidebands (red line in the figure) we can deduce a trap frequency of 70 kHz (consistent with a depth of 300 recoil energies) and an atomic temperature of 5 $\mu$K.

Applying a vertical magnetic field during spectroscopy reveals the hyperfine structure of the ytterbium clock line (fig. 5). The clock line is the average of the two $\pi$ transitions, whose separation changes by 20 Hz/mT. In this figure the polarization of the clock laser was tilted respect to the vertical to show the also the $\sigma$ transitions.

We used the spectroscopy signal to lock the frequency of the clock laser to the atoms using an AOM.

IV. CONCLUSIONS

We have show the status of the ytterbium optical lattice clock at INRIM. The experimental setup is complete and we achieved first spectroscopy results. In the next step we will finish the characterization of the clock and measure its absolute frequency respect to the cryogenic fountain IT-CsF2 [9].

Moreover our ytterbium lattice clock is part of the EMRP project "International Timescales with Optical Clocks" [10], where a comparison campaign with other clocks is planned, both local and remote. The clock will be part of a proof-of-principle relativistic geodesy experiment. It will be compared to a transportable strontium clock developed at PTB moved to the Laboratoire Souterrain de Modane (LSM) in the Fréjus tunnel. The comparison will be made through a optical fiber link and with a transportable frequency comb developed at NPL at LSM. General relativity predicts a frequency shift of $10^{-16}$/m and we should measure the 1000 m elevation difference between the two clocks at the decimeter level within a few hours.

As well, the Yb clock is part of the project AQUASIM, aiming to compare INRIM’s clock to a ytterbium degenerate Fermi gas experiment at European Laboratory for Non-Linear Spectroscopy (LENS) in Florence [11]. The connection will exploit the already existing optical fiber link between the two laboratories [12] and will be useful for studies of collisions physics and quantum simulations.

ACKNOWLEDGMENT

The authors acknowledge funding from the EMRP Project SIB55-ITOC, MIUR Project PRIN2012 AQUASIM and ITN Marie Curie Project FACT. The EMRP is jointly funded by the EMRP participating countries within EURAMET and the European Union.

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