

## Collisional Losses, Decoherence, and Frequency Shifts in Optical Lattice Clocks with Bosons

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We have quantified collisional losses, decoherence and the collision shift in a one-dimensional optical lattice clock on the highly forbidden transition  $^1S_0\text{-}^3P_0$  at 698 nm with bosonic  $^{88}\text{Sr}$ . We were able to distinguish two loss channels: inelastic collisions between atoms in the upper and lower clock state and atoms in the upper clock state only. Based on the measured coefficients, we determine the operation parameters at which a 1D-lattice clock with  $^{88}\text{Sr}$  shows no degradation due to collisions on the fractional uncertainty level of  $10^{-16}$ .

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Laser spectroscopy of transitions with millihertz line-width has become the essential tool for the rapidly evolving optical clocks [1,2]. These clocks promise a realization of the base unit “second” with higher stability and accuracy than the currently employed frequency standards using a microwave transition in  $^{133}\text{Cs}$ . Optical clocks enable tests of new theories, which predict, e.g., the variation of fundamental constants [3], or they can be used for relativistic geodesy field experiments [4]. Because of long coherence times and very sensitive interrogation, optical clocks with neutral atoms are also ideally suited to explore the physics of ultracold collisions. Collision experiments provide insight into the coupling between the atoms in different electronic states at ultralow temperature. Understanding collisions of alkaline earth atoms is required for proposed quantum computing schemes [5] and to test atomic structure calculations [6].

Strontium is widely used for optical lattice clocks with both bosons and fermions [2,7–9]. At ultralow temperature, collisions of fermions are suppressed by the Pauli exclusion principle [10]. The bosonic isotope  $^{88}\text{Sr}$  offers higher natural abundance and easier laser cooling. It is therefore of interest especially for transportable setups. Collisions are suppressed if a three-dimensional lattice with an occupancy of at most one atom per site [8] is used but at the expense of increased complexity of setup and experimental sequence. The use of  $^{88}\text{Sr}$  in a 1D-lattice clock can therefore be advantageous if the effects of collisions were well understood [9,11].

We have quantified the three expected collision-induced effects in our lattice clock setup: inelastic losses, collision broadening of the transition, and a collision-induced frequency shift. We predict operation conditions of a  $^{88}\text{Sr}$  lattice clock, where collisions do not limit the fractional uncertainty on the level of  $10^{-16}$ . Additional shifts due to clock-laser induced ac Stark effect and quadratic Zeeman effect in bosonic lattice clocks are well understood and can be controlled with high precision [12].

Our experiment was described before [13]. In short, we first prepare the  $^{88}\text{Sr}$  atoms in a magneto-optical trap

(MOT) operated on the strong 461 nm transition  $^1S_0\text{-}^1P_1$ . The atoms are loaded for up to 250 ms from a Zeeman-slowed atomic beam, which is deflected into the MOT region by an optical molasses.

The atoms are further cooled in a second cooling phase within 140 ms on the 689 nm intercombination line  $^1S_0\text{-}^3P_1$  to below  $4\ \mu\text{K}$ . A horizontal 1D-optical lattice is overlapped during the cooling with the MOT. It is created by a laser beam of up to 300 mW power at the magic wavelength of 813 nm. The beam is focused to a waist of  $34\ \mu\text{m}$  and retroreflected. The waist size is confirmed via measurements of trap oscillation frequencies.

After the second phase, up to  $3 \times 10^6$  atoms are trapped in the lattice ( $\sim 1000$  atoms per site). The atom number and temperature were determined by absorption imaging after a variable expansion time. Untrapped atoms are falling out of the interrogation area within 150 ms. Then the trapped atoms are interrogated on the clock transition  $^1S_0\text{-}^3P_0$  at 698 nm. The clock laser beam of 2.5 mW maximum power is overlapped with the lattice through a dichroic mirror and focused on the atoms to a waist radius of about  $39\ \mu\text{m}$ . To induce a dipole transition matrix element on the clock transition, we apply a homogeneous magnetic field of up to 3 mT.

After excitation on the clock transition the remaining ground state atoms  $^1S_0$  are detected by recording the fluorescence during a 461 nm MOT phase and then pushed away by resonant 461 nm radiation. In the same way, excited  $^3P_0$  atoms are detected after optically pumping them to the  $^3S_1$  state from where they decay to the ground state via the  $^3P_1$  state. Atoms decaying to the  $^3P_2$  level are also repumped by a second laser (inset of Fig. 1). Photo-associative losses by the repumping light are avoided by switching off the lattice and expanding the cloud of  $^3P_0$  atoms before repumping [11].

We first address *inelastic collisions*. Without excitation on the clock transition, we observe a background gas-limited lifetime of the atoms in the lattice of 7–8 s. In the presence of atoms in the  $^3P_0$  state, density dependent shortening of the trap lifetime is observed. Inelastic two-

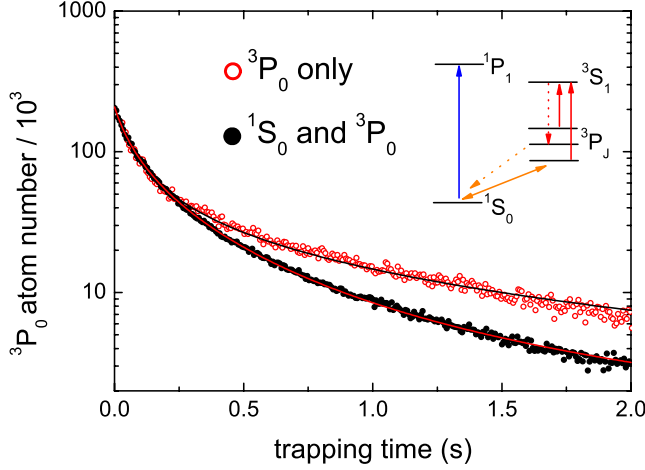


FIG. 1 (color online). Inelastic losses by collisions between  $^3P_0$  atoms (open circles), a mixture of  $^3P_0$  and  $^1S_0$  atoms (full symbols), and the results of fits (lines). The inset shows schematically the involved atomic levels. The initial number of ground state atoms is about  $1.2 \times 10^6$ .

body losses are possible by collisions of  $^3P_0$  atoms with either  $^3P_0$  or  $^1S_0$  atoms. The decay can be modeled by the coupled differential equations

$$\begin{aligned} \dot{\rho}_e &= -\Gamma\rho_e - \gamma_{ge}\rho_g\rho_e - \gamma_{ee}\rho_e^2 \\ \dot{\rho}_g &= -\Gamma\rho_g - \gamma_{ge}\rho_e\rho_g. \end{aligned} \quad (1)$$

Here,  $\rho_g$  ( $\rho_e$ ) denotes the local atomic density in the ground (excited) state,  $\Gamma$  is the inverse trap lifetime, and  $\gamma_{ge}$  ( $\gamma_{ee}$ ) is the loss coefficient for ground-excited (excited-excited) state collisions.

To distinguish experimentally between both inelastic collision channels, we first removed the ground state atoms from the lattice and observed the decay of the  $^3P_0$  population only (Fig. 1). In this case, Eqs. (1) simplify, decouple, and can be solved analytically. Integration over the spatial coordinates of a single lattice site provides an expression for its atom number:

$$N(t) = N_0 \frac{\exp(-\Gamma t)}{1 + [N_0\gamma_{ee}/(\pi^{3/2}\Gamma w_r^2 w_z)] [1 - \exp(-\Gamma t)]}. \quad (2)$$

$N_0$  denotes the initial atom number in a lattice site,  $w_r$  and  $w_z$  are the  $1/e^2$  radii of the atomic cloud in a lattice site at a given temperature and set of lattice parameters.  $\Gamma$  was fixed to the value determined before.

Many independent lattice sites are occupied with different atom numbers. We numerically sum Eq. (2) over the lattice sites. The initial axial population distribution is described by a Gaussian of  $1/e^2$  radius  $w'$  (determined from absorption images). From several data sets we determined the loss coefficient  $\gamma_{ee} = (4.0 \pm 2.5) \times$

$10^{-18} \text{ m}^3/\text{s}$ . Our uncertainty is dominated by the uncertainty of the calculation of the radii  $w_i$  and the scatter of the fit parameters. We neglect evaporation losses, since the lattice depth of  $k_B \times 45 \mu\text{K}$  is much larger than the atomic thermal energy of below  $k_B \times 4 \mu\text{K}$ . Traverso *et al.* [11] have measured a significantly larger loss coefficient in a dipole trap. The discrepancy might be attributed to effects of the dimensionality of the trap potential. When mostly the lowest axial vibrational state is populated as in our experiment, inelastic collisions are reduced [14].

We then investigated a mixed sample of  $^3P_0$  and  $^1S_0$  atoms. Collisions  $^3P_0 + ^1S_0$  are responsible for the different decays visible in Fig. 1. We determined the loss coefficient  $\gamma_{ge} = (5.3 \pm 1.9) \times 10^{-19} \text{ m}^3/\text{s}$  by numerical integration of Eqs. (1). Contributions to the uncertainty are again the determination of the  $w_i$  and the scatter of the fit. Careful consideration of the fitting strategy is required because losses are dominated by  $^3P_0 + ^3P_0$  collisions. Best accordance of data and fit was obtained by fitting  $\gamma_{ee}$  together with  $\gamma_{ge}$ . Nevertheless, we constrained  $\gamma_{ee}$  to the value obtained under the clean conditions of the first set of measurements. We found slightly different results for  $\gamma_{ge}$  depending on whether we fitted only the excited state population, the  $^1S_0$  atom number, or both simultaneously. We attribute this scatter to inelastic collisions that diminish the  $^3P_0$  atom number during the  $^1S_0$  atom detection. We corrected the excited state population for collisions with  $^3P_0$  atoms but neglect other types of collisions.

Different loss mechanisms are possible at the respective collision asymptotes. The asymptote  $^3P_0 + ^3P_0$  correlates at short internuclear distances to a  $^1\Sigma_g^+$  state, which can decay by fluorescence to lower molecular states. More likely as source for the losses are nonadiabatic transitions to molecular potentials dissociating to lower asymptotes. In the case of collisions at  $^1S_0 + ^3P_0$ , the latter path is very unlikely. Only one potential dissociates to the one lower asymptote  $^1S_0 + ^1S_0$  and it is very steep at the energy of the  $^1S_0 + ^3P_0$  asymptote. A direct spontaneous decay to that ground state potential is not possible due to selection rules, but at short internuclear distance complex coupling exists to molecular states that are subject to spontaneous decay [15]. This more indirect pathway can give a qualitative explanation for the significant difference between  $\gamma_{ge}$  and  $\gamma_{ee}$ . However, we cannot exclude three-body losses or processes involving a lattice photon.

As second decoherence process we identified strong *density dependent broadening* of the clock transition (see Fig. 2). To model the observed spectra we extended Eqs. (1) to a full density matrix  $\rho$  including the excitation dynamics, and integrated them numerically:

$$\dot{\rho} = -\frac{i}{\hbar}[H, \rho] + \mathcal{R}(\rho). \quad (3)$$

In the rotating wave approximation, the atom-light Hamiltonian and the relaxation matrix  $\mathcal{R}(\rho)$  become

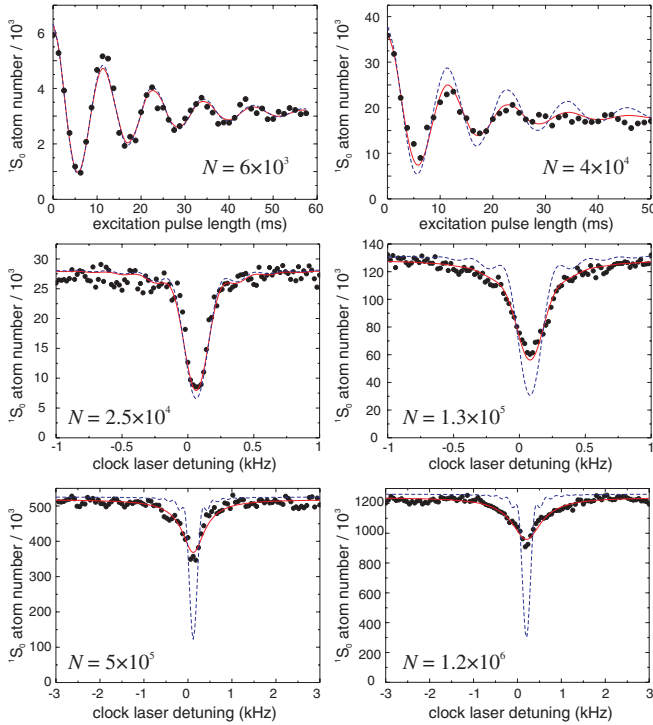


FIG. 2 (color online). Rabi oscillations (top row) and spectra of the clock transition (two lower rows) for different atom numbers  $N$ . Lines: Density matrix with dephasing constant  $\gamma_{\text{dep}}$  (red full lines) and with  $\gamma_{\text{dep}} = 0$  (blue dashed lines). The frequency axes have arbitrary offsets.

$$\frac{H}{\hbar} = \begin{pmatrix} 0 & \Omega/2 \\ \Omega/2 & \Delta \end{pmatrix}, \quad \mathcal{R}(\rho)_{11} = -(\Gamma + \gamma_{ge}\rho_{22})\rho_{11}$$

$$\mathcal{R}(\rho)_{12} = \mathcal{R}(\rho)_{21}^*$$

$$= -[\gamma_{ge}(\rho_{11} + \rho_{22})/2 + \gamma_{ee}\rho_{22}/2 + \Gamma + L + \gamma_{\text{dep}}\rho_{11}]\rho_{12}$$

$$\mathcal{R}(\rho)_{22} = -(\Gamma + \gamma_{ge}\rho_{11} + \gamma_{ee}\rho_{22})\rho_{22}. \quad (4)$$

The Rabi frequency  $\Omega$  can be calculated from the experimental parameters [9,16], the detuning  $\Delta$  is determined from the spectrum itself. Damping of the coherences in the density matrix is introduced by the coefficient  $L$  representing an effective laser linewidth and spatially dependent Rabi frequencies. The dephasing by elastic collisions is described by  $\gamma_{\text{dep}}$ . We assume dephasing by collisions with  $^1S_0$  atoms. Different mechanisms can be imagined, but the experimental data do not allow to distinguish between them.

For the sake of fast computation, we neglected spatial variations in the excitation probability due to the lattice or the clock-laser profile. Furthermore, a rectangular population distribution of full width  $2w'$  in axial direction was assumed. We have verified in the case of the inelastic losses that the effect of this approximation is minor.

To fix the excitation dynamics we fitted observed Rabi oscillations and a scan of the clock transition without significant broadening. The fitted Rabi frequency is about 17% smaller than the one calculated from the peak clock-laser intensity and the homogeneous magnetic field, an effect we attribute to the finite Lamb-Dicke parameter. The FWHM laser linewidth was fitted to be about 36 Hz, which is consistent with a transition linewidth of 35 Hz we have observed at an excitation pulse length of 35 ms and a few thousand atoms. The linewidth of the laser was limited by problems in the vibration isolation of the reference resonator [13]. To match the observations at higher density we had to include also a dephasing coefficient of  $\gamma_{\text{dep}} = (3.2 \pm 1.0) \times 10^{-16} \text{ m}^3/\text{s}$  (see Fig. 2). The effect is stronger by a larger factor ( $10^3$ ) compared to collisions  $^1S_0 + ^3P_1$  in a 1000 K Sr gas [17], which could indicate the existence of a scattering resonance. We did not observe a temperature dependence of any collision coefficient  $\gamma$  between 2 and 4  $\mu\text{K}$ .

The third collision effect being very important for an optical clock with neutral atoms is the *frequency shift*  $\delta\nu$  due to collisions. We have measured  $\delta\nu$  with an alternating stabilization [18,19]. The clock-laser interrogates either about  $10^4$  atoms (used as reference) or a higher atom number (and thus density) in consecutive probe cycles. The density shift is revealed as the difference of the offset frequencies between the clock laser and its reference cavity for the alternated stabilization cycles. We shortened the cycle time to  $\tau_{\text{cycle}} = 200 \text{ ms}$  and detected the excited state population only. For the best performance for equal atom number, the stability of the difference relative to the optical frequency is better than  $10^{-15}$  at an averaging time  $\tau = 10 \text{ s}$  and scales as  $1/\sqrt{\tau}$ . The atom number is varied via the duration of the initial MOT stage.

The dependence of the observed frequency shift as a function of the difference of atom number in the two stabilizations is shown in Fig. 3. Error bars indicate the uncertainty of the shift derived from the Allan deviation of the records with a length of  $\sim 500 \text{ s}$ . Fluctuations of the atom number difference lead to the given uncertainties. The point at zero atom number difference indicates that no systematic shifts occur. At low density we observe a linear dependence of the shift on the atom density. We attribute the saturation at high density to the changed dynamics during the excitation due to losses and dephasing. A theoretical analysis of the excitation process including an excitation dependent frequency shift will be carried out in the future. The clock laser was stabilized by probing the atoms with a 5 ms  $\pi$  pulse detuned such that the excitation probability was reduced to about 35% after the pulse.

From a linear fit through the origin (excluding the open data points in Fig. 3 because of saturation), the coefficient of the absolute collision shift as a function of the atom number  $N$  can be determined to better than 4%. The coefficient for the shift as a function of the density is

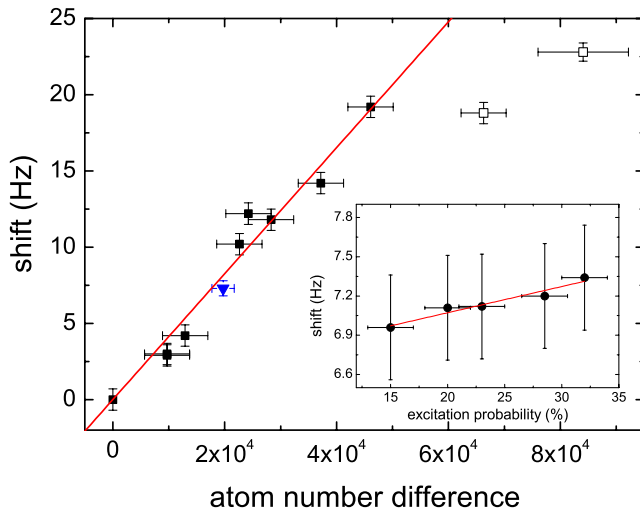


FIG. 3 (color online). Observed density dependent shift of the  $^{88}\text{Sr}$  clock transition. We assume linear scaling of density with atom number. The open symbols were excluded from the linear fit (full line, see text). The inset shows the dependence of the shift on the excitation probability with a linear fit. The triangle in the main graph represents the data in the inset. The shift was rescaled according to the trap parameters and atom temperature.

$\Delta\nu_\rho = 2\delta\nu\pi^2w'w_z^2w_z/N\lambda = (7.2 \pm 2.0) \times 10^{-17} \text{ Hz m}^3$  with an uncertainty mainly due to the density determination ( $\lambda$ : magic wavelength). A temperature dependence of the shift was not observed between 2 and 4  $\mu\text{K}$ .

The inset in Fig. 3 shows the dependence of the density shift on the excitation probability after the clock-laser pulse, which was varied by the clock-laser detuning in the stabilization cycle. We observe a weak relative influence of  $(2.9 \pm 4.5) \times 10^{-3}$  per percent excitation probability.

Having quantified three collision influences, we give guidelines for the design of a 1D-lattice clock with bosonic  $^{88}\text{Sr}$ . Assuming a typical lattice depth of  $k_B \times 10 \mu\text{K}$ , an atom temperature of 3  $\mu\text{K}$ , and an available lattice laser power of 300 mW, one could choose a lattice waist of 75  $\mu\text{m}$ . With  $w' = 280 \mu\text{m}$  and at the current level of accuracy for the density shift correction of 4%, a density shift of about 1 Hz would be tolerable to reach a fractional accuracy of  $10^{-16}$ , which is also the present uncertainty due to the blackbody shift [2]. This would limit the total atom number to about  $2 \times 10^4$  distributed over about 1400 sites, a value comparable to or larger than in present lattice clocks with  $^{87}\text{Sr}$ . The collisional broadening is then about 1.3 Hz. A new density shift measurement in the proposed lattice should yield an improved correction and allows for increasing the atom number until the collisional broadening becomes relevant. Aiming at a line width of about 10 Hz, operation with more than  $10^5$  atoms is feasible.

With a cycle time of 200 ms, the quantum projection noise limited relative stability in 1 s is  $2 \times 10^{-17}$ . To achieve this stability, however, the atom number has to be controlled to about 0.2%. At this density, losses do not distort the observed line or limit the excitation probability.

We conclude that collisions do not impose limitations for a bosonic  $^{88}\text{Sr}$  lattice clock compared to state-of-the-art ones with  $^{87}\text{Sr}$ . We expect that these newly observed and quantified decoherence mechanisms also will have impact on the design and operation of experiments with alkaline earth atoms that require long coherence times like quantum computing, millihertz linewidth lasers [20], or optical Feshbach resonances [21].

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