

Efficient 2^3S positronium production by stimulated decay from the 3^3P level

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We investigate experimentally the possibility of enhancing the production of 2^3S positronium atoms by driving the 1^3S-3^3P and 3^3P-2^3S transitions, overcoming the natural branching ratio limitation of spontaneous decay from 3^3P to 2^3S . The decay of 3^3P positronium atoms toward the 2^3S level has been efficiently stimulated by a 1312.2 nm broadband IR laser pulse. The dependence of the stimulating transition efficiency on the intensity of the IR pulse has been measured to find the optimal enhancement conditions. A maximum relative increase of $\times(3.1 \pm 1.0)$ in the 2^3S production efficiency, with respect to the case in which only spontaneous decay is present, was obtained.

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I. INTRODUCTION

Positronium (Ps) is the neutral matter-antimatter bound state of an electron (e^-) and a positron (e^+). Ps has two distinct ground states: the singlet 1^1S (para-Ps), annihilating into two γ -rays with a lifetime of 0.125 ns, and the triplet 1^3S (ortho-Ps), annihilating into three γ -rays with a lifetime of 142 ns [1]. Ps, being a purely leptonic two-body system, is well-known for offering an ideal testing ground for high-precision quantum electrodynamics (QED) calculations [2].

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Among the many precision experiments, the most accurate were recently conducted using two-photon Doppler-free laser spectroscopy of the 1^3S-2^3S transition [3]. The 2^3S level has an extended lifetime of 1142 ns in vacuum. This is due to its optical metastability: single-photon radiative decays to 1^3S are prohibited by the electric dipole selection rules, and the reduced overlap between the positron and the electron wave functions increases its annihilation lifetime by a factor of 8 [4]. On top of its high-precision spectroscopy applications, 2^3S Ps is one of the few notable candidate systems being considered for measuring the gravitational interaction between matter and antimatter [5], together with Ps in long-lived Rydberg states [6,7], antihydrogen [8–10], and muonium [11]. Moreover, as pointed out in [12], the metastable 2^3S Ps is scarcely sensitive to stray electric fields and is a good candidate for atom interferometry, provided that a beam with sufficiently low divergence and high intensity is demonstrated [5]. Furthermore, an intense source of polarized 2^3S atoms has been recently shown to be of extreme usefulness to achieve Bose-Einstein condensation of Ps [13].

2^3S Ps sources have been demonstrated via rf transition from laser-excited 2^3P Ps in a weak magnetic field [14], via two-photon Doppler-free 1^3S-2^3S laser excitation [15,16] and more recently via single-photon excitation of 1^3S to 2^3P in a rapidly switching electric field [12] and via single-photon excitation of 1^3S to 3^3P with radiative decay to 2^3S in an electric field [17] and in the absence of an electric field [18]. This last method in particular showed that it is possible to build an almost monochromatic 2^3S Ps source with a selected and tunable velocity distribution in the 10^4 ms $^{-1}$ range, with an overall efficiency between 0.7% and 1.4% according to the selected velocity [18].

In the present work, following this experimental line, we investigate the possibility of stimulating the 3^3P-2^3S transition to increase the overall 2^3S production efficiency. Indeed, laser-excited 3^3P Ps can spontaneously decay radiatively to 2^3S via the dipole-allowed 3^3P-2^3S transition (rate $A_{23} = 1.1 \times 10^7$ s $^{-1}$) with $\sim 10\%$ measured branching efficiency [17,18], limited by the competition with the more efficient spontaneous decay channel 3^3P-1^3S ($A_{13} = 8.4 \times 10^7$ s $^{-1}$). Here we demonstrate that it is possible to increase the 3^3P-2^3S transition rate, and thus the branching efficiency of the 2^3S decay, through stimulated emission.

II. EXPERIMENTAL METHODS

A straightforward way to stimulate the 3^3P-2^3S transition consists in introducing a synchronized broadband IR laser pulse at 1312.2 nm [5] on top of the pulsed UV laser at ~ 205 nm used for 1^3S-3^3P excitation [19]. In the present experiment, a 1312.2 nm laser pulse has been obtained from the same optical setup producing the UV beam, which is described in [20,21]. In this setup, a sequence of nonlinear optical conversion processes are used to generate the 205 nm wavelength, and in particular an optical parametric generation (OPG) crystal yields as a byproduct a broadband, amplified idler beam at 1314 nm. Both 205 and 1314 nm wavelengths could be tuned by varying the temperature set point of the OPG crystal around its nominal working point of $T_1 = 175.6^\circ$ (Fig. 1). This temperature value, chosen for the UV wave-

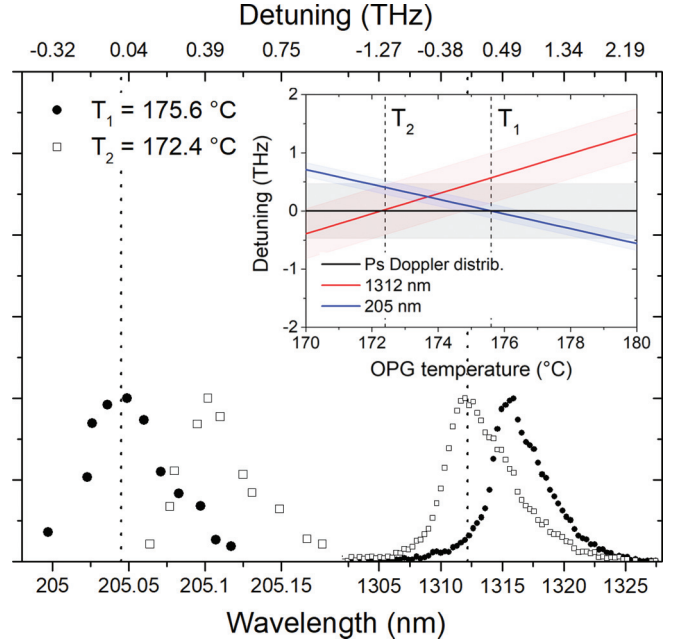


FIG. 1. Laser spectra (arbitrary units) measured in the two OPG temperature set points optimized for stimulating the $1^3S-3^3P-2^3S$ transition (white squares, 172.4°) and for the highest 1^3S-3^3P excitation efficiency (black circles, 175.6°). The dotted lines mark the transition resonances. Inset: measured laser detuning from the theoretical resonance frequency as a function of the OPG crystal temperature for the 1^3S-3^3P 205 nm laser (blue band) and the 3^3P-2^3S 1312 nm laser (red band), respectively, compared to the reference Ps Doppler distribution as measured in [19] (gray band).

length to fall on the 1^3S-3^3P resonance ($\lambda_3 = 205.045$ nm), is ineffective for a stimulated emission experiment as the bandwidth of the IR does not get to cover the 1312.2 nm wavelength with sufficient optical power to efficiently stimulate the 3^3P-2^3S transition. Hence, a different set point $T_2 = 172.4^\circ$ was selected to gain enough power at 1312.2 nm, while maintaining an acceptable detuning $\delta\nu_3$ of the UV beam from the 1^3S-3^3P resonance frequency, to minimize the unavoidable reduction in excitation efficiency. The set-point setting accuracy was limited to 0.5° by the TC200 temperature controller. At T_2 , the induced 1^3S-3^3P detuning was $\delta\nu_3 \approx 410$ GHz, corresponding to 205.103 nm and 0.9σ of the Doppler distribution of our Ps source ($\sigma_v \approx 470$ GHz [19]). A reduction in the 1^3S-3^3P excitation efficiency of about $\sim 50\%$ was expected [19], due to the Doppler selection of a Ps atoms distribution with $\lambda_3\delta\nu_3 \approx 0.85 \times 10^5$ ms $^{-1}$ average velocity in the direction parallel to the laser beam.

A direct way to observe an enhancement in the 2^3S signal due to the action of the stimulating laser is to compare the Ps annihilation time distribution with the UV laser only to that measured with both UV and IR pulses. The experimental methodology was the same as that used in previous works [17–19]. Bursts of 10^7 e^+ , 7 ns in time length [22], were guided by a 25 mT magnetic field, focused by an electric field of about 300 V cm $^{-1}$ and implanted at 3.3 keV in a circular spot of ~ 3 mm full width at half-maximum (FWHM) into a e^+ /Ps converter held at room temperature. The converter is constituted by a Si(111) p -type crystal with nanochannels

produced via electrochemical etching and oxidized in air at 100° for 2 h [23]. Ps formed in the converter outdiffuses back into vacuum through the nanochannels with an efficiency up to 35% losing a fraction of its formation energy by collisions with the channel walls [24]. A fraction of the emitted cloud was subsequently conveyed to 2^3S either by the 205 nm UV beam alone (i.e., through spontaneous decay from 3^3P , as in [17,18]) or by a combination of the UV and IR laser pulses (i.e., through stimulated decay from 3^3P). Both beams were linearly polarized perpendicularly to the target, with a nearly Gaussian temporal profile with a FWHM of 1.5 ns (UV) and 4.0 ns (IR), and a nearly Gaussian spectral profile with bandwidths $\sigma^{\text{UV}} \approx 120$ GHz (UV) and $\sigma^{\text{IR}} \approx 440$ GHz (IR, Fig. 1). The energy of the two pulses was (53 ± 5) μJ for the UV and (405 ± 10) μJ for the IR (at the entrance viewport of the experimental chamber). The UV spot was nearly Gaussian with (7.0 ± 0.7) mm FWHM both in the horizontal and vertical directions, while the IR spot was uniform in power and slightly astigmatic, $(12.9 \pm 0.5) \times (17.2 \pm 0.5)$ mm FWHM in the horizontal and vertical directions, due to a geometrical cut of an optical element. Two spurious light backgrounds at 532 and 894 nm (with energies ≈ 80 and ≈ 100 μJ , respectively) were found superimposed on the 1312 nm beam with the same spot size, caused by the nonideality of the dichroic mirrors used to separate the beams. Photoionization of 2^3S or 3^3P due to these backgrounds was negligible due to their low intensities and the small photoionization cross sections (10^{-16} – 10^{-17} cm^2). As the presence of an electric field would have shortened considerably the optical lifetime of 2^3S Ps [18], a fast HV switch with a rise time < 15 ns was used to disable the guiding electric field of the focusing electrode [see Fig. 2(a)] ~ 5 ns after e^+ implantation [18], such that the field was negligible when the excitation lasers were shot (~ 20 ns after e^+ implantation).

The time distribution of the annihilation γ rays due to the implanted e^+ and decaying or annihilating Ps was measured with a 20×25 mm lead tungstate (PbWO_4) scintillator [25] coupled to a Hamamatsu R11265-100 photomultiplier tube (PMT), placed 40 mm above the Ps converter. The signal from the PMT was 50%-50% split and digitized using two channels of a HD4096 Teledyne LeCroy 2.5 GS/s oscilloscope set at high (100 mV/div) and low (1 mV/div) gains to further extend the linear dynamic range of the digitizer. The data of the two channels were joined to form the so-called single-shot positronium annihilation lifetime spectroscopy (SSPALS) spectra [26], whose average is proportional to the amount of Ps/ e^+ annihilating per unit time. The total lifetime of 2^3S Ps in the absence of an electric field (1142 ns) is longer than that of 1^3S (142 ns) and of all other populated sublevels in the $n = 1-3$ manifolds. Thus, an increase in the delayed annihilations (either due to in-flight or to pick-off annihilations as the Ps atoms hit the chamber walls) for $t \gg 142$ ns (t being the time elapsed from e^+ implantation) can be directly related to the amount of 2^3S Ps (as in [18]). Relative time-dependent variations between two SSPALS spectra families with different laser configurations were quantified using the parameter $S(t) = [A_1(t) - A_2(t)]/A_1(t)$, where $A_1(t)$, $A_2(t)$ are the (averaged) integrated areas below single SSPALS shots of the two families in a selected time window centered in t . In this definition, $A_1(t)$ has the role of the reference area. An

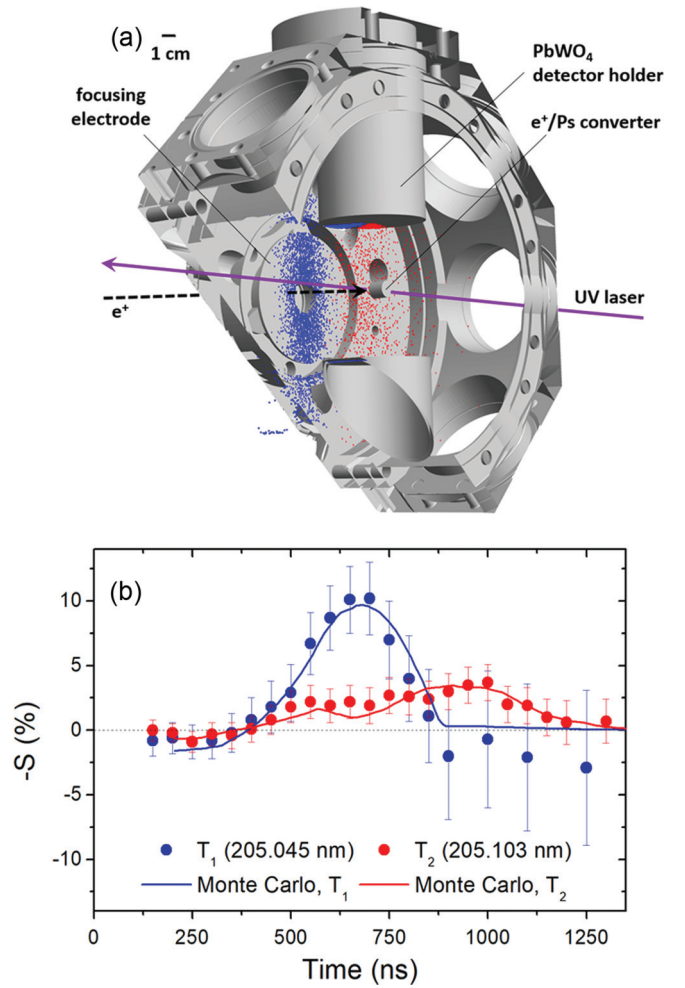


FIG. 2. (a) Distributions of Ps impact positions, shown superimposed upon the 3D drawing of the chamber walls, for the $T_1 = 175.6^\circ$ (blue circles) and $T_2 = 172.4^\circ$ (red circles) set points, emphasizing the different Doppler selection in the two detuning conditions. (b) Measurement of the annihilation time distributions of spontaneously decaying 2^3S Ps atoms from the 3^3P level without stimulated transition to 2^3S for the two temperature set points. The graph shows the curve $-S(t) = [A^{\text{UV}}(t) - A^{\text{off}}(t)]/A^{\text{off}}(t)$ (see the text). Each time distribution has been fitted with the Monte Carlo model discussed in the text (solid lines).

alternating measurement scheme between the two families was used to minimize the effect of time drifts of the experimental conditions. Contributions of eventual residual long-time drifts in the S calculation were further reduced by normalizing the shots in each family to a second-order polynomial fit of their value versus time (*detrending technique*; see [17], the Appendix).

III. RESULTS AND DISCUSSION

First, a set of reference spontaneous 2^3S Ps production measurements was acquired on the detuned OPG set point (T_2) to measure the $S(t) = [A^{\text{off}}(t) - A^{\text{UV}}(t)]/A^{\text{off}}(t)$ parameter in the absence of stimulated emission in the conditions selected for the following experiments. The measured $S(t)$ curve was compared to one acquired at the on-resonance set point (T_1)

(see Fig. 2), which was well-characterized previously [18]. A^{UV} and A^{off} correspond to the integrated averaged areas with and without the UV laser. Time windows of 300 ns width with steps of 50 ns were used to calculate $S(t)$. All measurements were obtained via alternating shots with the UV laser to the same number without it (~ 200 shots for T_1 , ~ 300 shots for T_2). The peak of 2^3S annihilations on the chamber walls is more evident in the on-resonance (T_1) measurement (between 500 and 750 ns) than in the detuned (T_2) measurement, where it is lower and smeared to longer times, in agreement with a reduced 2^3S production efficiency and longer atoms' traveled distances, because of the Doppler selection operated by the detuned UV beam that selects atoms traveling longer trajectories.

The experimental $S(t)$ curves were fitted using a previously developed Monte Carlo (MC) model [17,18]. This MC calculates the atoms' flight trajectories (starting from our Ps source velocity distribution [19]) and their excitation dynamics by simultaneously integrating the center-of-mass equations of motion and optical rate equations for the internal level dynamics. The annihilation position and time distribution for 1^3S and 2^3S Ps in flight and by collisions with the chamber walls [17] are calculated. The only fitting parameter was the $1^3\text{S}-3^3\text{P}$ excitation efficiency η_3 , while the 2^3S branching $\eta_m (=0.097)$ and quenching $\eta_q (=0.17)$ efficiencies were set according to past measurements (with the same laser delay of 20 ns; see [18]). Different Doppler selections in the two set points were now constrained to the measured UV detunings (0 GHz for T_1 and 410 GHz for T_2).

The fit (Fig. 2, solid lines) yielded $\eta_3 = (12.4 \pm 0.7)\%$ for T_1 and $\eta_3 = (5.2 \pm 0.4)\%$ for T_2 , corresponding to an $\approx 58\%$ reduction in the excitation efficiency as a consequence of the UV laser detuning, in agreement with the expected $\sim 50\%$ [19]. The annihilations' position distributions resulting from the fit [Fig. 2(a)] emphasize the difference in the atoms' flight trajectories in the chamber geometry in the two different detuning conditions.

Subsequently, the 1312.2 nm IR pulse was introduced to stimulate the $3^3\text{P}-2^3\text{S}$ transition. A first measurement campaign to optimize the IR laser 2^3S production efficiency was carried out by progressively attenuating its energy from the nominal $405 \mu\text{J}$ with a set of graded neutral density filters. The idea behind this optimization search is that the desired stimulated emission from the 3^3P level competes dynamically with the repumping of this level by absorption from the 2^3S . This mainly depends on the pulse energy (maintaining the other pulse parameters, in particular the FWHM temporal profile). One expects that on our nanosecond time scale, while a low energy can induce only a low population gain on the 2^3S level, a too high energy subtracts population from that previously efficiently excited by the rising part of the pulse. The optimization measurements were acquired alternating (~ 200) shots with both UV and IR lasers to shots with UV laser only. Their associated $S(t) = [A^{\text{UV}}(t) - A^{\text{UV+IR}}(t)]/A^{\text{UV}}(t)$ reflected the relative signal changes only caused by the IR, i.e., isolating the effects of this laser pulse (resonant with the $3^3\text{P}-2^3\text{S}$ transition) on the 2^3S Ps population. The resulting $S(t)$ curves (an example of which—with 5 dB attenuation—is shown in Fig. 3, inset) exhibit an excess of annihilations in the region around 550 and 1050 ns, where the impact of 2^3S

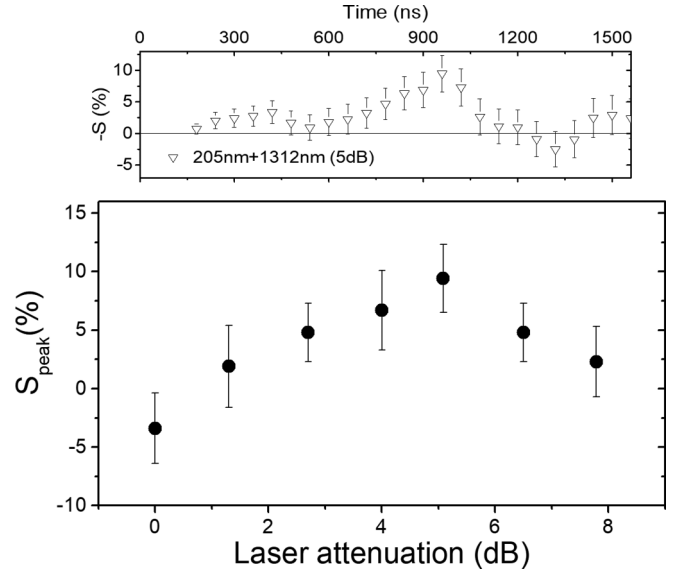


FIG. 3. Main panel: the S_{peak} value, as defined in the text, vs the attenuation in dB of the IR laser pulse starting from the maximum energy $405 \mu\text{J}$ (black points). Above: the $-S(t)$ parameter in the case of the 5 dB attenuation point.

atoms onto the chamber walls is also observed in the reference measurement (T_2 in Fig. 2). A convenient way to show the enhancement in the 2^3S production is to consider the peak amplitude of the $-S(t)$ curve, S_{peak} . The results are reported in Fig. 3 as a function of the 1312.2 nm laser pulse attenuation. The effect of the 1312.2 nm laser is compatible with zero when the beam is sent with full power. As expected, with the maximum pulse energy, a depopulation effect is observed. Then the parameter S_{peak} progressively increases as the laser energy is decreased to reach a maximum at 5 dB attenuation ($E^{\text{peak}} \approx 126 \mu\text{J}$) to slowly decrease again at even higher attenuations.

These measurements indicate that, selecting the 1312.2 nm laser pulse energy at E_{peak} , an increase of $(190 \pm 90)\%$ [corresponding to a $\times(2.9 \pm 0.9)$ gain efficiency] in the amount of produced 2^3S with respect to the reference was obtained. These findings are supported by numerical simulations based on a simple rate equations model [17], which predicts the same relative gain in the 2^3S production efficiency in such experimental conditions. The model also indicates that the same relative gain can be obtained with both UV and IR lasers set on-resonance.

A final measurement campaign was conducted to directly evaluate the 2^3S branching efficiency achieved by setting the 1312.2 nm laser to the optimal energy. The set of measurements was acquired alternating (~ 200) shots with both UV and IR lasers to shots with both lasers off. The resulting $S(t) = [A^{\text{off}}(t) - A^{\text{UV+IR}}(t)]/A^{\text{off}}(t)$ curve, compared to the reference 2^3S curve obtained in the same temperature set point T_2 and in the absence of the IR enhancement laser (previously plotted in Fig. 2), is shown in Fig. 4. The 2^3S branching efficiency was evaluated by fitting $-S(t)$ with the Monte Carlo model. The fit was now performed by varying η_m while keeping all other parameters fixed to those determined in the reference measurement. The found value for the

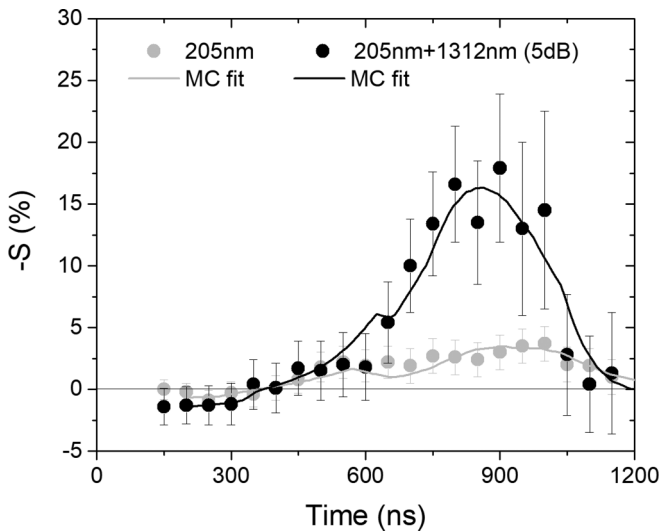


FIG. 4. The experimental curve of the $-S$ parameter ($= [A^{\text{UV+IR}}(t) - A^{\text{off}}(t)]/A^{\text{off}}(t)$) measured with the empirically determined optimal 1312.2 nm laser intensity, compared to the reference $-S$ obtained without the IR laser.

branching efficiency was $\eta_m^{\text{UV}} = 0.297 \pm 0.019$. This value, compared to the branching efficiency previously estimated from the measurement with the UV laser alone with the same laser delay ($\eta_m^{\text{UV}} = (0.097 \pm 0.027)$ [18]), leads to a $2\ ^3S$ Ps enhancement of $\times(3.1 \pm 1.0)$, in perfect agreement with the IR pulse optimization measurement.

IV. CONCLUSIONS

In conclusion, we experimentally demonstrated the possibility to efficiently stimulate the $3\ ^3P-2\ ^3S$ transition of Ps by employing a pulsed, broadband 1312.2 nm laser. The highest enhancement efficiency was found by tuning the intensity of the IR laser pulse inducing the $3\ ^3P-2\ ^3S$ transition. In these optimal conditions of our experiment, due to the pulsed excitation dynamics, the relative enhancement of the $2\ ^3S$ Ps atoms production from the excited $3\ ^3P$ population, with respect to the reference of the spontaneous emission decay, was measured to be $\times(3.1 \pm 1.0)$. This corresponds to a branching efficiency of $2\ ^3S$ production from the $3\ ^3P$ level of up to $\sim 30\%$. In any case, the overall $2\ ^3S$ Ps excitation efficiency was limited by the present technical restriction in producing both pulsed laser beams with the correct wavelengths and energies using a single optical parametric generation stage. The variation in the temperature set point of the laser generation crystal, necessary to output the correct 1312.2 nm

wavelength with sufficient energy, induced a UV detuning of about 410 GHz and a reduction in the amount of excited $3\ ^3P$ of $\approx 58\%$.

In a future realization of this experiment, the present technical limitations could be overcome by separating the two UV and IR laser lines, i.e., having independent nonlinear optical generation and amplification stages. An advantage of having the two laser wavelengths independent from each other would be to retain the full tunability characteristics of our $2\ ^3S$ source [18] while conveying $\sim 30\%$ of what could be excited in $3\ ^3P$ to $2\ ^3S$. Furthermore, if one accepts sacrificing the mentioned velocity selection, a further increase up to a factor of 5 of the overall excitation efficiency could be obtained by enlarging the UV laser bandwidth to cover efficiently the Ps Doppler profile. Finally, the first laser spectroscopy of the $3\ ^3P-2\ ^3S$ transition would become feasible. A laser system with independent UV and IR laser lines is currently in development to take full advantage of this $1\ ^3S-3\ ^3P-2\ ^3S$ stimulated excitation scheme in view of future measurements on a beam $2\ ^3S$ Ps.

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