Simulations of sawtooth-wave adiabatic passage with losses

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The results of simulations of cooling based on sawtooth-wave adiabatic passage (SWAP) are presented, including the possibility of population leaking to states outside of the cycling transition. The amount of population leaking can be substantially suppressed compared to Doppler cooling, which could be useful for systems that are difficult to repump back to the cycling transition. The suppression of the leaked population was more effective when simulating the slowing of a beam than in cooling a thermal distribution. As expected, calculations of the leaked population versus branching ratio of spontaneous emission show that the suppression is more effective for narrow linewidth transitions. In this limit, using SWAP to slow a beam may be worth pursuing even when the branching ratio out of the cycling transition is greater than 10%.

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I. INTRODUCTION

The ability to laser cool atoms [1,2] has enabled the exploration of the collective behavior in atomic gases, ultracold scattering, and other effects. Methods used to laser cool atoms, as well as some variations, have been used for molecules [3] and nano and microscale objects [4]. However, not all atoms, molecules, or nanoscale objects can be effectively laser-cooled with known techniques, which motivates the search for new cooling methods.

The authors of Ref. [5] described a method for laser cooling called the sawtooth-wave adiabatic passage (SWAP) based on chirping counterpropagating light waves. This method was proposed for cooling narrow linewidth transitions. The basic idea is that the Doppler shift of the counterpropagating light waves leads to a stimulated absorption from the beam opposite the atom’s velocity (due to the blue shift of that beam) followed by a stimulated emission from the beam propagating in the direction of the atom’s velocity (due to the red shift of that beam). If the linewidth is narrow, the spontaneous emission that occurs between the stimulated absorption and emission will be small. This leads to a stimulated emission that occurs between the stimulated absorption from the beam and the red shift of that beam. If the linewidth is narrow, the spontaneous emission that occurs between the stimulated absorption and emission will be small. This leads to a stimulated emission that occurs between the stimulated absorption from the beam and the red shift of that beam.

The authors of Ref. [6] described in more detail the simple theoretical model for this process and presented the results from a simple theoretical model supported both the possibility of cooling and the interpretation of the mechanism.

The authors of Ref. [7] performed SWAP cooling of Dy using a transition at 626 nm with a 136-kHz linewidth. This transition is $\sim$20 times broader than that in the demonstration on Sr. The authors of Ref. [8] described experimental results for SWAP cooling of Sr in a magneto-optical trap. These studies [5–9] showed that SWAP gives a more rapid reduction of the kinetic energy compared to Doppler cooling, although the final temperature that can be reached is several times hotter than Doppler cooling. Finally, as first suggested in Ref. [5], SWAP cooling is promising for molecules because the loss of population to states outside of the cycling transition can be reduced due to the relative suppression of spontaneous emission. The authors of Ref. [9] calculated SWAP cooling for diatomic molecules in a magnetic field and did find a reduction in the population into leakage channels.

In this paper, we perform calculations for a more simplified case than Ref. [9] to understand the role of leakage during SWAP cooling. For the model below, there is only one leak state and it cannot be connected to the excited state by a laser transition. While not quite as realistic as Ref. [9], this model allows for a more transparent interpretation of the results of the cooling when population can leak from the cycling transition. In particular, we present results on the performance of SWAP cooling as a function of branching ratio $B$ into the leak state for various spontaneous decay rates.

SWAP cooling for small $B$ is important for molecules, but larger branching ratios might also be interesting. Part of our motivation was to determine whether SWAP is worth pursuing when this branching ratio is greater than 1/2. For example, laser cooling of antihydrogen, $\bar{H}$, on the 1S–2P transition was predicted [10] to give an average final energy $E_f \simeq 30$ mK compared to a Doppler temperature of $\sim 2$ mK. Because of the long lifetime of the 2S state, one could attempt SWAP cooling on the 2S–3P or 2S–4P transition to obtain colder $\bar{H}$ which would improve, for example, the measurement of the 1S–2S transition [11]. Unfortunately, the simulations below suggest that the short lifetime of these states and the unfavorable branching ratio preclude cooling $\bar{H}$ by SWAP.

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The results presented below suggest that SWAP with leaks to other states is more effective for slowing a beam of particles than for cooling. As expected, the quality of the SWAP cooling increases as the spontaneous decay rate decreases. However, for very narrow lines, SWAP might be useful even for branching ratios, $B > 0.1$.  

II. THEORETICAL MODEL

All of the calculations used are based on the model introduced in Ref. [6], with a few modifications, which will be explicitly noted. The system is an atom with center-of-mass motion constrained to one dimension and with three internal states instead of the two internal states of Ref. [6]. The translational motion is represented by momentum eigenstates which are stepped in units of the photon momentum, $\hbar k$ with $k$ the photon wave number. The two internal states treated in Ref. [6] are a ground state $|g\rangle$ and an excited state $|e\rangle$. To incorporate the possibility of population leaking to other states, the calculations below include a third state $|l\rangle$. For simplicity, the counterpropagating lasers cause transitions between the ground and excited states, but do not connect the excited and leak states. The state of the atom is specified by its internal state and its momentum state, which is given in multiples of the photon momentum. For example, $|e, -6\hbar k\rangle$ represents the atom being in the excited state and in the $-6\hbar$ momentum state. For conciseness, we will use the symbol $n_p$ to refer to the momentum state; in the example, $n_p = -6$.

The master equation

$$\frac{d\hat{\rho}(t)}{dt} = \frac{1}{i\hbar} [\hat{H}(t), \hat{\rho}(t)] + \hat{\mathcal{L}}(\hat{\rho})$$

(1)

determines the evolution of the atom through the time dependence of the density matrix $\hat{\rho}$. The time-dependent Hamiltonian $\hat{H}(t)$ leads to a coherent evolution of the system from the stimulated absorption or emission of a photon and the associated recoil. The Lindblad superoperator $\hat{\mathcal{L}}(\hat{\rho})$ models the incoherent evolution from the spontaneous emission and its associated recoil. Within the rotating wave approximation, the time-dependent Hamiltonian is

$$\hat{H}(t) = \frac{\hat{p}^2}{2M} - \frac{\hbar}{2} \Delta(t) \hat{\sigma}_z + \frac{\hbar}{2} \Omega_s(t) \cos(kz) \hat{\sigma}_x,$$

(2)

where $\Delta(t)$ is the time-dependent detuning $\hat{\sigma}_z = |e\rangle\langle e| - |g\rangle\langle g|$, $\Omega_s$ is the standing wave Rabi frequency, the $W(t)$ is a window function not used in Ref. [6] but defined below, $\hat{\sigma}_z = |e\rangle\langle g| + |g\rangle\langle e|$, and

$$\cos(kz) n_p \hbar k = \frac{1}{2} (n_p - 1) \hbar k + \frac{1}{2} (n_p + 1) \hbar k.$$  

(3)

The time-dependent detuning has a sawtooth profile centered on the Doppler-free transition frequency that goes from $-\Delta_s/2$ to $\Delta_s/2$ with linear ramp $\alpha$: $\Delta(t) = -\Delta_s/2 + \alpha t$ with $t = 0$ defining the start of the ramp. The duration of the ramp $T_s = \Delta_s/\alpha$ will be used below.

For $\hat{H}(t)$, the only difference from Ref. [6] is that we use a windowing function to turn the standing wave on and off. The windowing function reduces the ringing that results from instantaneous changes in the detuning by smoothly going to 0 at the beginning and end of the ramp. In all of the calculations, we used $W(t) = \exp[-36(t - T_s/2)^2/(T_s/2)^2]$ where $0 \leq t \leq T_s$, but almost any smooth function which is flat during the middle part of the ramp will lead to similar results. In all of the calculations, we chose an exaggerated duration for the SWAP compared to previous calculations (e.g., Ref. [6]) to account for the window function and to leave a small amount in $|e\rangle$ at $T_s$; the population left in the excited state at $T_s$ decays to the ground and leak states with their respective branching ratio which would decrease the effectiveness of SWAP in decreasing leakage out of the $g,e$ cycle. We chose $\Delta_s = 240\omega_0$ with $\omega_0$, the recoil energy although using values 1/2 this size did not strongly affect the results.

The Lindblad superoperator is somewhat more complicated than in Ref. [6] due to the branching ratio to the leak state. We will use $\gamma$ for the total decay rate of the excited state and $B$ as the branching ratio to the leak state $|l\rangle$, which is not part of the cycling transition; $1 - B$ is the branching ratio to the ground state $|g\rangle$. The Lindblad superoperator is given by

$$\mathcal{L}(\hat{\rho}) = \frac{\gamma}{2} \left( |e\rangle\langle e| \hat{\rho} + \hat{\rho} |e\rangle\langle e| - 2 \frac{3}{4} \hat{\rho}_s + \frac{1}{4} e^{ikz} \hat{\rho}_s e^{-ikz} + \frac{1}{4} e^{-ikz} \hat{\rho}_s e^{ikz} \right),$$

(4)

where

$$\hat{\rho}_s = (1 - B) \hat{\sigma}_g^+ \hat{\sigma}_g^+ + B \hat{\sigma}_l^- \hat{\sigma}_l^-,$$

(5)

with $\hat{\sigma}_g^- = |g\rangle\langle e| = \hat{\sigma}_g^+ \hat{\sigma}_g^+ \hat{\sigma}_l^- = |l\rangle\langle e| = \hat{\sigma}_l^+ \hat{\sigma}_l^+ \hat{\sigma}_l^-$. The Lindblad superoperator of Ref. [6] results when $B = 0$.

A nontrivial issue is how to treat the density matrix when going from the end of one SWAP to the beginning of the next. For the calculations below, we made choices for this treatment that assumes there is a sufficient delay between SWAPs that the population left in the excited state due to nonadiabaticity or due to $\gamma \neq 0$ will decay to 0. In a mathematical sense, this means $t \rightarrow \infty$, but, in practice, most runs finished with less than 10% population in the excited state at time $T_s$ so that adding $3 - 4/\gamma$ between SWAPs would decrease the excited population to a negligible amount. Between SWAPs we allocated the excited state population between the $|g\rangle$ and $|l\rangle$ states using the branching ratio of Eq. (5) and the momentum population was allocated by adding the population in $|e\rangle n_p$ into $n_p$ with fraction 3/5 and $n_p \pm 1$ with fraction 1/5 each. Lastly, the delay between SWAPs can lead to coherences between the momentum states of $|g\rangle$ that can affect the steady-state populations. Because we were uncertain about the experimental possibilities for coherences between multiple SWAP cycles, we set the coherence terms in the density matrix to 0 after each SWAP cycle.

III. BASIC PARAMETERS

As noted in Ref. [6], the dynamics of this master equation can be scaled, which means the outcomes are determined by scaled parameters. We will define the parameters in terms of the total decay rate of the excited state, with the scaled parameters being denoted by an over-tilde. The important
parameters are $\tilde{\Omega}_r = \Omega_r / \gamma$, $\tilde{\alpha} = \alpha / \gamma^2$, $\tilde{\Delta}_r = \Delta_r / \gamma$, and $\tilde{\omega}_r = \omega_r / \gamma$, where $\hbar \omega_r = (\hbar k)^2 / (2M)$ is the recoil energy.

As discussed in Ref. [6], there are dimensionless ratios that are important for the effectiveness of SWAP. The adiabaticity parameter $\kappa = \tilde{\Omega}_0^2 / \tilde{\alpha}$ determines whether the $g \leftrightarrow e$ transition is adiabatic or diabatic. At the Landau-Zener level, the probability for an adiabatic transition is $P = 1 - \exp(-\pi \kappa / 2)$. The range of the sweep has to be large enough to contain the Doppler-shifted resonances plus a bit extra to accommodate the transients at the beginning and end of the sweep: $\Delta_r > 4k|\nu|$. The amount of time spent in the excited state should be much less than the lifetime of the excited state: $\tilde{\omega}_r / 2 < |k \nu - 2\omega_r|$. Finally, the resonances should be separated: $\Omega_r/\gamma < \omega_c$.

IV. RESULTS AND DISCUSSION

The results from three different cases are presented in this section. The case where the population is confined to a cycling transition is briefly treated before the more complicated three-state system. For the purpose of obtaining an absolute energy in some plots, we fixed the wavelength to be $\lambda = 689.5$ nm and kept $\hbar \omega_0$ fixed at 0.36 $\mu$K times Boltzmann’s constant $k_B$.

The authors of Refs. [5–9] suggested that SWAP would be useful for the case where there is a leak in the cycling transition to other states. The idea is that the stimulated emission step would suppress the spontaneous emission into the leak state(s) $|l\rangle$. The results below give an indication of how effective this might be.

A. No leak, $B = 0$: Steady state versus $\tilde{\omega}_r$

Before treating cases where population can leak into state $|l\rangle$, it is worthwhile to examine how the excited state linewidth affects the effectiveness of the SWAP procedure. We did this by investigating the steady-state temperature reached using SWAP, as a function of the scaled recoil frequency $\tilde{\omega}_r = \omega_r / \gamma$. SWAP was proposed as a method for cooling atoms with narrow linewidths. This corresponds to larger-scaled recoil energies.

For these calculations, we fixed the range $\Delta_r = 240$ and set the ramp rate at $\alpha = \Omega_0^2 / 2$ (i.e., $\kappa = 2$) which gives a Landau-Zener probability of $P \approx 0.96$. The initial momentum distribution was started as a thermal distribution at low temperature (typically $k_B T \sim 10 \mu$K) and the SWAP procedure was iterated several times (typically less than 20) until the average kinetic energy reached a limiting value. After each SWAP, we allowed the decay of any population in the excited state and we set the coherence between different momentum states to be zero after each SWAP as described in Sec. II. For each $\tilde{\omega}_r$, the Rabi frequency $\tilde{\Omega}_r$ was taken to be an integer and was varied in steps of 1 to approximately find the lowest steady-state energy; see Fig. I(c) where the central point was the value used and the error bars of $\pm 1/2$ roughly indicate the uncertainty in $\tilde{\Omega}_r$ that would give the lowest steady-state energy. Different values of $\tilde{\omega}_r = \omega_r / (\hbar \gamma)$ were obtained by varying the decay rate $\gamma$.

The data in Fig. 1 show the scaled average kinetic energy in a steady state versus the scaled recoil frequency. As was found in Refs. [5–9], SWAP cooling does not achieve a temperature as low as that from Doppler cooling; for the range plotted in Fig. 1, the asymptotic temperature was more than $\sim 10X$ the Doppler temperature for every $\tilde{\omega}_r$. However, as expected from Refs. [5–9], the lowest temperature is achieved when the decay rate $\gamma$ is smallest (i.e., when $\tilde{\omega}_r$ is largest).

B. One sweep with loss

The case where population can leak to state $|l\rangle$ is somewhat more complicated because the loss grows with each SWAP and the population that is lost remains at the same energy. To understand the trends for SWAP with a leak out of the cycling transition, it is sufficient to understand the results from one SWAP.

I. Single initial velocity

In this section, the initial state has all of the population in one particular momentum state. One sweep is performed. This situation mimics that of slowing a beam of atoms. The quantities of interest are the amount of population lost into state $|l\rangle$ and the change in energy of the states. In all of the calculations the atom starts in the ground state with $p = 20 \hbar k$.

There are several parameters that control the population lost into the leak state and the energy removed during the SWAP. We chose to study how the population in the leak state and the final energy varied with the branching ratio $B$ to spontaneously decay to the leak state $|l\rangle$ for a few scaled recoil frequencies $\tilde{\omega}_r$. The $\tilde{\omega}_r$ was stepped by the factor $\sim 1.4$ to show a range of behavior. The range of the sweep $\Delta_r$ was scaled for each $\tilde{\omega}_r$ so that $\Delta_r = 240 \tilde{\omega}_r$. For each $\tilde{\omega}_r$, the $\tilde{\Omega}_r$
was varied to give the best slowing for $B = 0$ and was then fixed for all other values of the branching ratio. As can be seen in Fig. 5, there could be other choices for $\tilde{\Omega}_2$, but the general trends remain. The ramp rate was chosen so that $\alpha = \Omega_2^2/2$ (i.e., $\kappa = 2$) which gives a Landau-Zener probability of $P \simeq 0.96$. A slower ramp rate would give a larger Landau-Zener probability, but at the cost of spontaneous emission between the stimulated absorption and stimulated emission steps. After the SWAP, the excited-state population was allowed to decay as in Ref. [6].

The parameters in the simulation gave a final energy after one SWAP that ranged from 0.82 times the initial energy for $\tilde{\omega}_r = 2$ to 0.84 times the initial energy for $\tilde{\omega}_r = 0.25$. For a perfect SWAP, the momentum should go from $20\hbar k$ to $18\hbar k$, meaning the expected final energy is $(18/20)^2 \simeq 0.81$ times that of the initial energy. This indicates that the SWAP procedure is effective at slowing the atoms for $n_p \sim 20$, roughly independent of the decay rate. As an example, for $\tilde{\omega}_r = 1$, 89% of the population finishes in the $18\hbar k$ momentum state when $B = 0$.

Figure 2 shows the population that leaks to the state $|l\rangle$ as a function of $B$. As expected, there is no population leak for $B = 0$, and the population in $|l\rangle$ increases with the branching ratio. For small branching ratio into the leak state $B$, the leaked population $P_l$ is proportional to $B$, but increases more slowly than linearly for larger $B$. The slope of $P_l$ for $B \sim 0$ determines the effectiveness of the SWAP procedure at small branching ratios: a smaller slope means less loss and a greater effectiveness. Although the $B \sim 0$ is important, even the extreme case $B \sim 1$ (i.e., the branching ratio of spontaneous emission into the leak state is $\sim 100\%$) has less than $1/4$ of the population leaking into state $|l\rangle$ during a SWAP for the parameters in Fig. 2. As an example, for $\tilde{\omega}_r = 1$, the $B \sim 1$ population loss is 12% per sweep. If this loss were the same for successive $n_p$, there would be approximately 70% population loss after ten SWAPs. Similarly, the $\tilde{\omega}_r = 1$, $B = 1/2$ case gives 8% loss per SWAP which is a factor of 2.6 suppression of the loss that would occur for Doppler cooling.

With one exception, there is a clear trend of decreasing population leak as $\tilde{\omega}_r$ increases. This is understandable because an increased scaled recoil energy means a smaller decay rate, which should make the SWAP procedure more effective: there is more stimulated emission back to $|g\rangle$ and less spontaneous emission which could lead to either $|g\rangle$ or $|l\rangle$. As an example, the $\tilde{\omega}_r = 1$ case [dash-dot (blue) line] used $\Omega_2 = 33$ and had $P_l \simeq B/5$ for $B \simeq 0$. For this case, the loss is $\simeq 5$ times smaller than the best case using Doppler cooling. Even the worst case shown [$\tilde{\omega}_r = 0.25$ solid (red) line] has a loss for $B \simeq 0$ that is $\simeq 3$ times smaller than the best case using Doppler cooling. The exception to the trend is the $\tilde{\omega}_r = 0.7$ [dotted (green) line]. This case also had an anomalous value for $\tilde{\Omega}_2$ compared to the trends observed for the other $\tilde{\omega}_r$. We do not know why this case is different from what we expected.

2. Thermal distribution

In this section, we present the results for the effect of one SWAP when the initial distribution of momenta is thermal. The initial distribution was chosen to be proportional to $\exp(-[n_p/14]^2)$ which corresponds to a temperature, $k_B T = 98\hbar \omega_r$. This temperature was chosen to give similar SWAP parameters to Sec. IV B 1. As with the previous section, we performed calculations for several $\tilde{\omega}_r$, the range of the sweep was $\Delta_2 = 240\hbar \omega_r$, and we varied the $\tilde{\Omega}_2$ for each $\tilde{\omega}_r$ to obtain the best cooling for $B = 0$.

There was not a substantial difference in the amount of energy removed in one SWAP for different $\tilde{\omega}_r$ at $B = 0$. This trend is similar to that in the previous section. The final average energy somewhat increased with decreasing $\tilde{\omega}_r$: the largest for $\tilde{\omega}_r = 1/4$ (approximately 0.83 times the initial average energy) and the smallest for $\tilde{\omega}_r = 2$ (approximately 0.78 times the initial average energy).

The population that leaked to state $|l\rangle$ after one SWAP is plotted in Fig. 3 for several different $\tilde{\omega}_r$. The line types

![Figure 2](image1.png)

**FIG. 2.** The probability $P_l$ for the atom to leak to the state $|l\rangle$ after one SWAP as a function of the branching ratio $B$ to the leak state. The solid (red) line is for $\tilde{\omega}_r = 0.25$, the medium dash (blue) line is for 0.35, the short dash (orange) line is for 0.5, the the dotted (green) line is for 0.7, the dash-dot (purple) line is for 1.0, the dash-dot-dot-dot (maroon) line is for 1.4, and the dash-dot-dot-dot-dot (black) line is for 2.0. For all calculations, the atom starts in the ground state with $p = 20\hbar k$.

![Figure 3](image2.png)

**FIG. 3.** The probability $P_l$ for the atom to leak to the state $|l\rangle$ after one SWAP as a function of the branching ratio $B$ to decay to the leak state $|l\rangle$. The line types are the same as Fig. 2. For all calculations, the atom starts in a thermal distribution of momenta as described in the text.
are the same as for Fig. 2. The amount of population that leaks out of the cycling transition is much larger than for the previous section where only one momentum component is initially occupied. This is because the thermal distribution has several momentum components, and the SWAP parameters were chosen to give an overall efficiency. However, parameters that work well for large momenta are not so good for small momenta and vice versa. For example, at larger momentum, the time between the stimulated absorption and stimulated emission is larger, which necessitates a larger ramp rate and larger Rabi frequency to decrease stimulated emission, while a smaller momentum requires a smaller ramp rate and Rabi frequency to get separation between the stimulated absorption and stimulated emission. The best slope for small $B$ is for $\bar{\omega}_r = 2$ and is $\sim 2$ times smaller than would result from spontaneous emission from the excited state. Unlike the previous section, the interpretation of this slope is ambiguous because it is not clear how many photons were absorbed.

The momentum distribution after SWAP, Fig. 4, indicates that the atoms that leak into the state $|l\rangle$ are those with smaller energy. In this calculation, $\bar{\omega}_r = 1$ and the branching ratio back to $|g\rangle$ is $1 - B = 0.99$. For this case, 0.605% of the atoms leak to the state $|l\rangle$. The solid (red) line shows the initial thermal distribution of the atoms. The medium dashed (blue) line shows the momentum distribution after the SWAP for the atoms that finish in the $|g\rangle$ state. The increase in the distribution for small $|n_p|$ shows that there has been cooling during the SWAP, even with the loss. The short dashed (orange) line is the momentum distribution for atoms that leaked to $|l\rangle$. This distribution has been scaled so the integral would be the same as for the $|g\rangle$ state. This distribution is even more strongly peaked at small $|n_p|$ which indicates that the atoms that leak preferentially have smaller energy. This trend is plausible because choosing $\Omega_s$ to give the largest energy decrease indicates the SWAP is most efficient for the states with larger $|n_p|$.

C. Relative efficiency with loss

One possible measure of the efficiency of SWAP cooling is the change in energy divided by the population lost to the leak state. To make it dimensionless, we define it as $R \equiv \langle (KE)_0 - (KE)_f \rangle / (KE)_0 P_f$ for $B = 0.02$. One calculation was for a single initial momentum 20$\hbar k$ and the other was for a thermal distribution. For the $n_p = 20$ plot, the red X’s are when maximizing the change in $KE$ while the black *’s are when maximizing $R$.

D. Implication for $\dot{H}$

One of the motivations for the above studies was to explore whether $\dot{H}$ can be further cooled from the apparent limit of Doppler cooling in the ALPHA trap: the average final energy in the simulation was $\bar{E}_f/k_B \sim 30$ mK [10]. This simulation was based on one laser Doppler cooling on the
1S–2P transition at 121.6 nm. The idea for further cooling is to excite the \( \bar{\text{H}} \) to the metastable 2S state and attempt to laser cool on the 2S–3P or 2S–4P transition. The best case is the 2S–4P transition which has the larger photon momentum and smaller decay rate, meaning it will have the larger \( \tilde{\omega}_r \); the value \( \tilde{\omega}_r \approx 0.065 \) is smaller than any of the values simulated above. For this transition, the Doppler temperature \( T_D \approx 0.31 \) mK. Thus, the starting \( k_B T/(\hbar \tilde{\omega}_r) \) is five times that in Fig. 3 assuming \( T \sim 20 \) mK [10]. Unfortunately, the branching ratio to leak states is \( B \approx 0.88 \) for either transition. We simulated multiple SW APs until all of the population had leaked out of the cycling transition and found a decrease of 16% in the kinetic energy. This is a large decrease considering the unfavorable conditions (i.e., large branching ratio and large linewidth). However, since populating the 2S state is very difficult, the simulations strongly suggest that SW AP cooling is not worthwhile for further cooling \( \bar{\text{H}} \).

V. CONCLUSION

Results from simulations of SWAP cooling of a thermal distribution or slowing a beam were presented when a loss channel was present. For the cases investigated, there was not a large difference in the cooling or slowing versus the branching ratio \( B \) for spontaneous decay into the leak state(s) for one SWAP, and there was less population lost during one SWAP when slowing a beam compared to cooling a thermal distribution. The population lost to the leak state strongly depended on the branching ratio as well as the spontaneous decay rate of the excited state. For small branching ratios, the population lost to the leak state is proportional to the branching ratio but did not increase as rapidly for larger \( B \). For the same branching ratio, atoms with larger scaled recoil energy \( \varepsilon_r/(\hbar \gamma) \) have less population lost into the leak during each SWAP. For the cases we investigated, the population lost during each SWAP for small \( B \) was as much as five times smaller than would be lost during Doppler cooling. This confirms the suggestion in Ref. [5] that SWAP could be useful for cooling molecules. SWAP might be useful for slowing beams even for unfavorable (i.e., large) branching ratios out of the cycling transition if the scaled recoil energy is large.

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