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Simulating The Doppler-Free Fluorescence Spectrum For The Potassium D₁ Transitions

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Abstract:

Radiation theory (absorption, spontaneous emission, and stimulated emission) is applied to Potassium (³⁹K and ⁴¹K) to examine details of the D₁ lines, Figure 1, in the near IR at 770 nm. When examining the resonance fluorescence from two counter-propagation laser beams in a K cell, Figure 2, three prominent "Doppler-free" features—dips at the D_{1a} and D_{1b} resonances and spikes at their crossover frequencies—stand out superposed on the fluorescence background. They are examined with a detailed simulation, Figures 3 and 4, and compared to observations, Figure 5. Parametric studies of the Doppler-free features, Figures 6–8, indicate how to maximize their prominence, and thus their importance as frequency references for laboratory and atmospheric observations.

Figure 1: Hyperfine Structure of Potassium



Figure 3: Flow-chart indicating how the fluorescence is calculated. Beside each step is the symbolic representation of what is being calculated.



Figure 5: Total Fluorescence and Experimental Comparison



Figure 6: Effects of Power





An energy-level diagram of K can be described by three related models. The basic model is a two level system. The intermediate model includes electron spin, splitting the excited state into a doublet for the D_1 lines and a quartet (not shown) for the D_2 lines. The final model includes the nuclear spin of 3/2 and the associated hyperfine interaction, leading to hyperfine splitting.¹

Figure 2: Experimental setup. A CW laser beam is split in two by a half-silvered mirror. One beam is sent through a potassium cell in one direction; the other in the opposite direction. The frequency (wavelength) can be varied to examine the Doppler-free features. The detector records the fluorescence output from the four resonances.





In the above equations, a *y* subscript indicates a given transition, f_{lu} is the oscillator strength, S_y is the line strength of each transition, f_y is the offset frequency of each transition, $\Delta f_y = A_y / 2\pi$, and *Ao* is the inverse of the lifetime.

<u>Results:</u>

The simulated fluorescence response for each isotope as a function of frequency is shown in Figure 4. (At this wavelength, a frequency difference of 1.0 GHz is equivalent to a wavelength interval of 1.98 pm.) Figure 5 compares the measured and simulated responses. The main fluorescence and Doppler-free features are present in both curves. However, the calculations exhibit a higher frequency resolution, while the observations show wider shoulders near the cross over and weaker shoulders near, at least, the D_{1a} dip.

Figure 7: Effects of Temperature







(wavelength) control

Calculating The Fluorescence:

To make the simulations we modified a C++ code developed at the University of Illinois². An outline of the code is given in Figure 3 and the physical properties for potassium are given in **Table 1**³.

Table 1: Potass	sium Pr	oper	ties			
Transition		n4 to n1		n3 to n1	n4 to n2	n3 to n2
Line Strength		5		1	5	5
Spontaneous Emission Rate, A _v		Ao / 2		Ao / 6	Ao / 2	5 Ao / 2
Offsets for K39 (GHz)		0.31		0.254	-0.152	-0.208
Offsets for K41 (GHz)		0.405		0.375	0.151	0.121
			1			I
Lifetime = 1 / Ao	26.2 ns			lsotope	39K	41K
Oscillator Strength	0.339			% Occurrence	e 93.26%	6.73%
Center Wavelength	770.109	3 nm				

Figure 4: Fluorescence for each isotope



Offset Frequency (GHz)

Conclusion:

This is the first comparison for potassium of a Doppler-free simulation and observation. They show good agreement, thereby confirming the main features of the simulation. An extra dip was found in the middle of both the D_{1a} and the D_{1b} dips. The effects of parameter variation enable us to optimize the experimental set up, which will be used shortly to provide an absolute wavelength standard for a new ALO lidar system for temperature and wind observations in the mesosphere.

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