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I. Analysis of Visible Spectra

Naval Weapons Support Center Applied Sciences Department Crane, Indiana 47522

6 October 1975

FINAL REPORT for Period 1 July 1973 to 1 July 1975

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Prepared for Commander Naval Air Systems Command (Code AIR-31OC) Washington, D. C. 20361



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Visible spectra of flares conta ning magnesium as a fuel and lithium, sodium, potassium, rubidium and cesi i nitrates as the oxidizers are presented. Atomic and molecular emissions are identified and assigned to electronic transitions. The emission characteristics of the atomic emitters are the same as those observed in other studies. Several emission bands, possibly from diatomic alkali metal species, are observed and tentatively assigned.

The intensity of the 518 nm magnesium line is seen to decrease by two orders of magnitude when the oxidizer is changed from lithium nitrate to cesium nitrate. This change could not be easily explained on the basis of calculated equilibrium temperatures and species concentration

Recommendations for further work are discussed.

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## INTRODUCTION

Pyrotechnic compositions used for the production of visible light, either for illumination or for signaling purposes, rely on emission from atomic and molecular species formed as reaction products from the combustion of magnesium and a solid oxidizer, usually an alkali or alkaline earth metal nitrate or perchlorate.<sup>1</sup> The most widely studied system is the magnesium-sodium nitrate flare which is used for illuminating purposes. It has been shown that most of the visible light produced by this flare comes from the broadened sodium resonance lines centered at 589 nm.\* During the course of the studies to determine the processes occurring in the magnesium-sodium nitrate system, additional information was obtained on the visible and near-infrared spectra of pyrotechnic compositions containing barium and strontium salts and the other alkali metal nitrates; namely, lithium, potassium, cesium and rubidium.<sup>5,6</sup> It was determined that the use of any alkali or alkaline earth metal nitrate other than sodium nitrate for illumination purposes was unsatisfactory. The use of alkali metal nitrates for colored signals was also considered unsatisfactory in comparison to the alkaline earth compounds currently used. It was apparently for these reasons that no attempt was made to further investigate the spectra of the magnesium/alkali metal nitrate flares in any detail. The spectra were presented as either direct reproductions of the spectrograph film<sup>5</sup> or rapid scanning spectrometer data.<sup>6</sup> In neither case was any detailed attempt made to discuss or to analyze the spectra.

Since the alkali metal nitrates and perchlorates find wide use in pyrotechnics, it appears reasonable to more completely evaluate the emission characteristics of flares containing magnesium and an alkali metal-containing species. This is the first in a series of reports which will attempt to characterize the processes occurring in flares containing lithium, sodium, potassium, rubidium and cesium nitrates as oxidizers. This first report will describe the visible emission spectra of these flares.

#### EXPERIMENTAL

## Initial Calculations

To completely accomplish the objectives of the work, a series of experimental flares were made which were designed to allow investigations of the emission spectra of high and low concentrations of alkali metal emitters and the emission spectra of combinations of emitters. An initial series of calculations was done using the NASA thermodynamics program<sup>7</sup> to optimize the magnesium, alkali metal nitrate, binder formulations to give the maximum adiabatic temperature. The organic binder level was fixed at five percent and the fuel to oxidizer ratio was varied until the maximum temperature was computed. The final formulas used in this study are given in Table 1. Table 2 contains the computed maximum temperatures and the important species concentrations for the formulas. As can be seen from Table 2, the adiabatic temperatures decrease in order Li>Na>K>Rb>Cs as would be predicted.

An examination of Table 1 reveals that formulations 7307, 7308, 7312 and 7313 contain mixtures of different alkali metal nitrates. The purpose of these formulations was two-fold. The major reason for doing this was to provide high and low concentrations of the alkali metals (on the order of 100:1) for spectral measurements in the resonance line region of the spectrum. The results of this study will be published in Part III of this series of reports. The second reason was to allow an examination of the visible spectrum for possible alkali metal diatomic molecular emission.

### Construction of Flares

The chemicals used in making the formulations ranged from research grade lithium, rubidium and cesium nitrate to a practical grade sodium and potassium nitrate. These oxidizers were ground and dried prior to mixing. The particle size was approximately 30 microns. The magnesium was MIL-M-382, granulation 17, which has an approximate particle size of 75 microns. The binder used was a combination of Dow Epoxy Resins<sup>®</sup>, DER 321 and DER 732 and a polyamine CX 3482.1 in the ratio DER 321 - 61.3<sup>°°</sup>, DER 732 - 26.3<sup>°</sup> and CX 3482.1 - 12.4<sup>°</sup>. This is a typical epoxy binder formulation currently in use in illuminating flares. In an effort to prevent contamination, the mixing, screening and drying apparatus was washed and dried between each formula change.

# TABLE 1. FLARE FORMULATIONS

Group 7307

Group 7308

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Group 7311

Magnesium Lithium Nitrate Rubidium Nitrate Binder	43.2% 51.1 0.7 5.0	Magnesium Potassium Nitrate Binder	35.2% 59.8 5.0
Dilliger	••••		

# Group 7312

Magnesium	39.6%	Magnesium	43.2%
Sodium Nitrate	54.9	Lithium Nitrate	51.05
Lithium Nitrate	0.5	Cesium Nitrate	0.75
Binder	5.0	Binder	5.0
DINGE	010	••••••	

Group 7309

# Group 7313

Magnesium Rubidium Nitrate Binder	25.3% 69.7 5.0	Magnesium Lithium Nitrate Sodium Nitrate Potassium Nitrate Binder	43.2% 50.65 0.55 0.60 5.0
DINUCT	5.0	Potassium Nitrate Binder	0. 5.

# Group 7310

Magnesium	19.8%
Cesium Nitrate	75.2
Binder	5.0

Group #	7307	7308	7309	7310	7311	7312	7312
Max [Metal]	Li	Na	Rb	Cs	к	Li	Li
T(°K)	3081	3073	3008	2569	3056	3083	3083
[Li]*	.130	.0012		*-		.1321	.1304
[Na]		.1882		**	• =	** **	.0018
[K]					.1858		.0016
[Rb]	.0008	••	.192			<b>48 10</b>	
[Cs]	~ ~		***	.1943		. 0009	
[Mg0]	.0569	.0550	.0352	.0269	.0489	.0579	.0521
[H~O(s)]	. 3807	.3644	.3454	. 3274	.3612	.3781	.3762
[//9]	.0849	.0787	.0538	.0544	.0775	.0857	.0245

# TABLE 2. CALCULATED TEMPERATURES AND CONCENTRATIONS

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\*Concentrations are given in mule fraction.

The flares were pressed into paper tubes at 10,000 pounds dead load. Fifty grams of composition were used for each candle. The finished size was 3.3 cm in diameter and 5.1 cm in length. Five grams of ignition composition were pressed on the surface of each candle.

#### Experimental Procedure

The flares were burned face up at a distance of 6.0 meters from the spectrographs. Visible spectra from 0.40 - 0.70um were recorded on a Bausch and Lomb 1 5 m grating spectrograph. This, instrument has a 450 line/mm grating giving a dispersion of 15 Å /mm in the first order on the photographic film. Kodak Linagraph Shellburst film was used for recording these spectra. Spectra were also taken in the region of the resonance lines using a 1 m Spex spectrograph and Kodak I-N plates. The Spex instrument has a 1200 line/mm grating and gives a first order dispersion of 8 Å/mm. Spectra of a 200 watt quartz-iodine lamp were recorded on the sare plates to provide a film response correction. In these experime ts the flares were masked in such a way that the spectrographs were observing a position in the flame seven centimeters above the surface of the candle.

The spectra were scanned and digitized on an Optronics \$2000 dentiometer. Points were taken at approximately 0.60 nm intervals. The resulting digital output was processed by standard procedures and the film density converted to radiant and luminous power readings.

## RESULTS

This first set of experiments was designed only to obtain the visible spectra of flares containing magnesium, an alkali metal nitrate, and a bi: 'er. No attempt was made to measure the absolute value of radiant or luminous power. Relative powers in the visible can be obtained by integrating the spectra over all wavelengths from 0.46 - 0.70 um. The relative radiant powers are Li Na>K-Cs-Rb and the ratio was 1:0.04. The burning rates on all the compositions were on the order of 0.084 cm/s.

The visible radiant power spectra obtained on the Bausch and Lomb 1.5 m spectrograph for flare groups 7307 through 7313 are shown in Figs. 1-7. Each of these spectra was selected as being











FIG. 3. Group 7311 - High [K] - Numbered peaks correspond to following emitters and wavelengths in nm: (1) K, 495.1, 495.6, 496.5, (2) K, 508.4, 509.9, (3) K, 509.7, 511.2, (4) K, 532.2, 534.0, (5) K, 534.3, 536.0, (6)  $K_2$ , 570, (7) K, 578.3, 580.2, 581.3, 583.2, (8) Na, 589.3.













FIG. 6. Group 7312 - High [Li]/Low [Cs] - Numbered peaks correspond to following emitters and wavelengths in nm: (1) Cs, 455.5, (2) Li, 460.3, (3) MgO, 500.7, (4) Mg, 518.4, (5) Na, 589.3, (6) Cs, 601.0, (7) Li, 610.4, (8) Cs, 621.3, (9) Li, 670.8, (10) Cs, 672.4.





representative of a large number of spectra obtained for each group. The spectral distributions are essentially unchanged within a particular formulation group. To avoid confusion and to show as much of the important structure as possible, each figure is normalized to a value of one at its maximum. The relative magnitudes of the maxima are 1.0:1.0:0.02:0.03:0.04 for Li, Na, K, Rb and Cs, respectively.

No inconsistencies with visual observations were observed. The flares containing a high lithium concentration, e.g. group 7307, appeared red to pink. Those containing high sodium appeared yellow-white. The flares with cesium and rubidium appeared white and the potassium flares tended to be purple. Visual estimations of brightness were in agreement with the relative radiant power measurements presented earlier.

#### DISCUSSION

#### Assignment of the Spectra

#### Lithium

The electronic structure of lithium is 1s<sup>2</sup>2s<sup>1</sup> making the ground state a 2s. The following assignments are made for the lithium spectrum shown in Fig. 1. The flare burned to get this spectrum was formulation 7307, high lithium - low rubidium concentration. Reading from short to long wavelength, the peak at 460.29 nm is assigned to the 4d+2p transition in lithium. The next three peaks are MgO bands, B+X transition, at 498.59, 499.67 and 500.7 nm and are assigned to the ?, 2; 1,1 and 0,0 transitions, respectively. Any information gained from these bands and other features associated with MgO is uncertain due to a lack of basic information regarding potential energy curves, ground state assignments, etc. This region will be observed in more detail and the results given in another report. The next observable feature is the series of three lines at 516.73, 517.27 and 518.36 nm and these are assigned to the 4s+3p transition of atomic magnesium. The next major feature is the sodium resonance doublet at 589 nm. This sodium emission is strong but the sodium is present only as an impurity. The strong peak at 610.36 nm is assigned to the  $3d \rightarrow$ 2p transition of atomic lithium. The two small peaks at 620 and 630 nm are rubidium and will be discussed later.

The most striking feature and the feature which contributes most of the visible energy is the broadened lithium resonance line centered at 670.78 nm, the  $2p \rightarrow 2s$  transition. The broadening at half-height is slightly greater than 20 nm. A further discussion of the alkali metal resonance line broadening will be presented in another report. The sharp line located at 672.34 nm is due to a cesium impurity and its assignment will be discussed in conjunction with the cesium analysis.

There were no unexpected features observed in the magnesiumlithium nitrate flare.

## Sodium

Although a great deal has been written about the sodium nitrate flare, we will analyze sodium in the same way as the other alkali metal nitrates in order to provide complete documentation. Much of what is written here is certainly redundant.

The electronic structure of sodium is  $1s^22s^22p^63s^1$  making the ground state energy level of the sodium atom a 3s. The spectrum to be assigned is the one given in Fig. 2; the flare formula is 7308, high sodium - low lithium concentrations. The first apparent peak is the lithium peak at 460.29 nm described previously. The next series of peaks are the MgO bands at 500.7 nm. In addition, there is a sodium peak superimposed on these bands at 498.28 nm. This peak is due to the 5d+3p transition. The next two lines are the sodium doublet at 514.91 and 515.36 nm and are due to the 6s+3p transition. The next three lines are the magnesium lines at 516.73, 517.27 and 518.36 nm discussed previously. There is next an apparently continuous band-type feature at 530 nm and this will be discussed later.

The major feature in the sodium spectrum is the broadened resonance line continuum which has been well described both experimentally and theoretically by Douda.<sup>2,3</sup> The emission is centered at 589.3 nm and is due to the  $3p \rightarrow 3s$  transition. The halfwidth of the broadened lines is approximately 50 nm.

Superimposed on the short wavelength side of the continuum is a sodium doublet at 568.27 and 568.82 nm. These lines are due to the  $4d \rightarrow 3p$  transition. Superimposed on the long wavelength side are the lithium lines at 610.36 nm and 670.79 nm and another sodium doublet at 615.42 and 616.07 nm due to the  $5s \rightarrow 3p$  transition.

Returning to the band feature at 530 nm, we will make tentative assignments only as to possible molecular emitters and delay any detailed discussion to the second report in this series. The emission at 530 nm appears to be completely continuous and has previously been assigned to a NaMg molecular interaction.<sup>10</sup> The "shoulder" which appears at 550 nm on the wing of the broadened sodium resonance line has been assigned both to a sodium-nitrogen satellite interaction and to diatomic sodium.<sup>6,11</sup> The actual assignment remains in question and will be discussed at length in the next report.

#### Potassium

Potassium salts find heavy use in pyrotechnics as oxidizers probably because of their relative inexpensiveness and lack of emission in the visible spectral region. The fact that potassium does have significant visible emission may account in part for the current inability to obtain high purity colors in signals which incorporate it as an oxidizer.

The electronic structure of potassium is ls<sup>2</sup>2s<sup>2</sup>2p<sup>6</sup>3s<sup>2</sup>3p<sup>6</sup>4s<sup>1</sup> making its ground electronic state a 4s. The assignments are made for Fig. 3, [lare formula 731], consisting of magnesium, potassium nitrate and binder only. In terms of specific line emission, the potassium spectrum is relatively simple. Barely distinguishable at approximately 490 nm are three potassium lines assigned as 495.08, 495.60 and 496.50 nm. These lines are a result of the transitions  $8d \rightarrow 4p$ ,  $10s \rightarrow 4p$  and  $8d \rightarrow 4p$ . The next series of lines are two doublets at 508.42, 509.92, 509.71 and 511.20 nm and are assigned to the transitions  $9s \rightarrow 4p$  and  $7d \rightarrow 4p$ . Two more potassium doublets are observed at 532.23, 533.97 and 534.30, 535.95 nm and are due to the  $8s \rightarrow 4p$  and  $6d \rightarrow 4p$  transitions. A fourth set of potassium doublets is observed at 578.26, 580.20 and 581.25, 583.21 nm and are assigned to the 7s, 4p and 5d, 4p transition. The sodium doublet at 589 nm is due to an impurity in the mix. Quantitative analysis of a similar composition showed the concentration of the sodium impurity to be less than 1 ppm. This illustrates the excellent emission properties of atomic sodium. It is possible that some of this emission comes from the paper tube. Similar work using slightly different formulas showed no reduction of the sodium emission when burned without tubes<sup>12</sup> so the case is probably not the origin of the emission.

There are several features in the potassium spectra which warrant attention. The first feature is the symmetrical hump centered at 570 nm. This continuous feature is tentatively assigned to diatomic potassium and will be discussed in greater detail in the next report. Another feature is the sharply increasing continuum starting at 600 nm and continuing outward. The leveling off of this feature at 650 nm may or may not be real since in this region both the film and spectrograph are rapidly losing response and the apparent leveling may be an artifact of this loss in sensitivity. No assignment has been made at this time. It is possible that this feature is the tail of the broadened potassium resonance line continuum centered at 768.2 nm. However, the broadening in potassium would have to be excessively large for this to be the case. Similarly the increased continuum observed at 425-475 nm might also be due to broadening of the 404.5 doublet in potassium, the other transition, 5p As, going to the ground state. As more data is reduced, the origins of these continuous emissions may become more apparent. These also will be discussed in the next report. There are observed transitions in diatomic potassium at 650 nm and at 406 nm.<sup>9</sup> These have been observed in absorption only.

In regard to missing features, it is interesting to note the apparent absence of the Mg and MgO emissions. These emissions actually are present. However, on the scale used in this figure, they are too small to be seen. As can be seen from Table 2, this might be due to a loss in the temperature of the system. If this is the case, conceivably the magnesium lines and the MgO band system could be used as a guide to determining the temperature distribution in a pyrotechnic flame. This will be discussed in more detail in another section of this report.

#### Rubidium

The electronic structure of rubidium is  $1s^22s^22p^33s^3p^3d^{10}4s^24p^5s^1$  making the ground state a 5s. The assignments given are for Fig. 4, flare formula 7309, consisting of magnesium, rubidium nitrate and binder. As can be seen from Fig. 4, rubidium has a large number of emission lines in the visible. At the extreme left of the figure, centered at 420.7 nm, we see the 6p+5s transition in rubidium. Unfortunately the sensitivity of the film was such that this feature was barely apparent. The two lines at 455.5 and 459.3 nm are due to a cesium impurity. In the region from 500-535 nm there are a series of several lines as yet unidentified. These are apparently narrow with little or no degrading indicating that this is probably atomic emission rather than molecular band emission. The exact wavelengths of the major lines are 515.3, 518.6, 526.4 and 532.6 nm. These lines are followed by another unidentified line at 539.4 nm.

The next line is a rubidium line at 543.15 nm and is due to the  $8d \rightarrow 5p$  transition. The next rubidium line is seen at 557.88 nm and is the transition  $9s \rightarrow 5p$ . The next line is actually two unresolved rubidium lines at 564.81 and 565.37 nm coming from the 7d+5p and  $9s \rightarrow 5p$  transitions. Next is a rubidium line at 572.45 nm and is from the 7d+5p transition. The next line is the sodium doublet at 589 nm. Skipping the band emission at 600 nm the next line is a rubidium line at 615.96 nm arising from the 8s+5p transition. The next line is a to 520.63 nm, a rubidium line from the 6d+5p transition followed by the rubidium transition at 629.83 nm. This transition is also the 6d+5p. The last two lines superimposed on a continuum are lithium at 670.7 and cesium at 672.3 nm.

The continuous emission at 600 nm, which may be molecular band emission, could be due to either diatomic rubidium or some other rubidium interaction. Further data reduction is underway and this, like the other apparent molecular species, will be discussed in the second report. The continuous emission beginning at 630 nm and continuing to cut off looks suspiciously similar to the continuum in potassium. This may be a molecular interaction, it may be only the wing of the broadened resonance line centered at 792 nm, or it may be an artifact of the spectrograph response. Further examination of the data will be presented in the second report.

Again, in this spectrum, one should note the apparent absence of the Mg and MgO emission. As in the case of potassium, the emission is present but very weak.

#### Cesium

The electronic structure of cesium is  $1s^22p^63s^23p^83d^{10}$ 4s<sup>2</sup>4p<sup>4</sup>4d<sup>10</sup>5s<sup>2</sup>5p<sup>6</sup>6s<sup>1</sup> making its ground electronic state a 6s. The following assignments are for Fig. 5, flare formulation 7310, composed of magnesium, cesium nitrate and binder. The two peaks at 455.54 and 459.32 nm are cesium emission lines and are assigned to the transition 7p+6s, a transition to the ground state. Visual examination of the spectrograph film shows that these lines are reversed as one would expect. The peak at 473.2 nm as well as those at 534.2, 546.5, 550.2, 563.5 and 566.3 nm are unidentified at this time. The possibility that they are molecular emissions exists but the shapes of the lines indicate that they are probably higher level atomic cesium transitions. The next line is cesium at 584.47 nm and is due to the transition 9d+6p. The next line is the sodium doublet at 589 nm. Next is a cesium line at 601.03 nm and is the Rd+6p transition. The small line is cesium at 603.41 nm and is due to a 10s.6p transition. The next line is at 621.29 nm and is the 8d+6p cesium transition. The next line is a cesium line at 635.6 nm and is due to the 9s-6p transition. This is followed by a line at 658.65 nm and is the cesium 9s.6p transition. The final line is a strong cesium line at 672.33 nm resulting from the transition 7d-6p.

In the cesium spectra, there are no unidentified band structures or continua. As a matter of note, the Mg and MgD emissions are present but weak and the underlying graybody continuum which normally has at least a tendency to increase at the longer wavelengths is absent. It should also be noted that there is no sharply increasing emission at the extreme red end of the spectrum as was observed in both potassium and rubidium.

## Mixtures

Figures 6 and 7 are the visible spectra of formula groups 7312 and 7313. Group 7312 is a high lithium - low cesium concentration formula. Group 7313 has a high lithium concentration and low sodium and potassium concentrations.

The emission features in both Figs. 6 and 7 are characteristic of lithium. The major lines are at 460.3, 610.4 and 670.8 nm. In addition, both spectra show the sodium resonance doublet at 589 nm. One should note that in Fig. 6 the sodium is present as an impurity and there is little broadening of the emission lines while in Fig. 7 the sodium concentration is about 0.2 percent and the broadening is already apparent.

In Fig. 6, the other spectral features are the Mg and MgO lines and bands and the more intense cesium emission lines. In Fig. 7, the Mg and MgO emissions are prominent features. However, the emissions from the low concentration potassium present are too weak to be shown on the scale of this figure.

It should be noted that there are no molecular emissions from diatomic or polyatomic species which might have been formed as a result of interactions between the alkali metals.

#### Summary

In an effort to summarize this mass of information and make it more useful, the wavelength and assigned transitions for the alkali metal nitrotes only are tabulated in Tables 3 to 7. If these tables are compared to the arc emission spectra of atomic alkali metals generally tabulated, <sup>13</sup> one finds virtually complete agreement in both line assignments and intensities.

#### Disappearance of Prominent Features

As one observes the spectra in Figs. 1 to 7, it is apparent that as the alkali metal nitrates are changed from lithium to desium, i.e., as the molecular weight of the oxidizer increases, the intensity of emission from the magnesium lines at 516.73, 517.27 and 518.36 nm and magnesium oxide bands at 500 nm decreases. This decrease is even greater than is apparent from the spectra since, for the sake of comparison, the powers have been normalized to a value of one at the maximum. At first glance this decrease could be attributed to a decrease in the temperature of the reaction. In an effort to correlate these two factors a simple calculation was made using the

# TABLE 3. ASSIGNMENT OF LITHIUM LINES

Wavelength (nm)	Transition
460.29	4d+2p
497.20	45+2p
610.36	3d-2p
670.78	20+25

# TABLE 4. ASSIGNMENT OF SODIUM LINES

Wavelength (	nm)	Transition				
498.28		50 °P				
514.91	€s+3p					
515.36		6s <b></b> →3p				
568.27		4d+3p				
568 52		4d+3p				
589.00	reversed and	3p+35				
589,95	broadened	3p+3s				
615.42		5s+30				
616.02	5s <b>≁</b> 3p					

# TABLE 5. ASSIGNMENT OF POTASSIUM LINES

Wavelength (nm)	Transition
495.08	8d→4p
495.60	10s+4p
496.50	8d→4p
508.42	9s→4p
509.71	7d+4p
509.92	9s+4p
511.20	7d+4p
532.23	8s+4p
533.97	8s+4p
534.30	6a+4p
535.95	6d+4p
578.26	7s+4p
580.20	7s-⊷4p
581.25	5d≁4p
583.21	5d+4p

25

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# TABLE 6. ASSIGNMENT OF RUBIDIUM LINES

Wavelength	(nm)	Transition
420.70	reversed	6p→5s
536.26		8 <b>d</b> →5p
543,15		8 <b>d</b> →5p
557.88		9s+5p
564.81		7d→5p
565.37		9s+5p
572.45		7d+5p
615.96		8s+5p
620.63		6d≁5p
629.83		6d+5n

# TABLE 7. ASSIGNMENT OF CESIUM LINES

7p→6s

7p+6s

9d→6p

8d+6p

Wavelength (nm) Transition 455.54 reversed 459.32 reversed 584.47 601.03 603.41 621.29

10s→6p 8**d**+6p 9s+6p 635.60 9s**→**6p 658.65 7d+6p 672.33

27

1.808

Boltzmann equation to calculate the population of the upper levels which give rise to these transitions. We know that the intensity of an atomic line in emission is given by

 $I_{em}^{nm} = N_n h c v_{nm} A_{nm}$ (1)

where  $N_n$  is the population in the n<sup>th</sup> level,  $A_{nm}$  is the fraction of atoms carrying out the transition from level n to level m per second and hcv<sub>nm</sub> is the energy of a light quantum of wavenumber  $v_{nm}$ . In the calculation to be done, we will consider only a single emission line, the magnesium line at 518.36 nm. Thus, in Eqn. (1), the only quantity which changes as we change the oxidizer in the system is the population of the upper level, level n. The concentration in the upper level is given by the Boltzmann equation,

 $N_n = N_{tot} e^{-E/kT}$ (2)

where  ${\rm N}_{\rm tot}$  is the concentration of the magnesium, E is the energy associated with the 518.36 nm wavelength, k is Planck's constant and T is the temperature of the system. The results of this calculation for the systems under consideration are given in Table 8. In this table the experimentally obtained intensity of the magnesium line, I(Mg), at 518.36 nm is given relative to its maximum value, i.e., the value when lithium nitrate is used as the oxidizer. Since lithium nitrate was used in several formulas as the main oxidizer, there was a chance for comparison of the intensity of the magnesium line for precision among different spectra. For all systems chosen and for several spectra in each system exposed for different times, the agreement among the magnesium intensities was  $\pm 20\%$ . While this error is not small enough to allow for quantitative discussions of populations, it is sufficiently accurate to allow a general comparison between what is predicted by theory and what is observed experimentally. The values for the magnesium concentration, [Mg], again relative to the value when the oxidizer is lithium nitrate, and the temperatures, T, given in Table 8 were obtained from the thermodynamic calculations given in Table 2. The quantity N<sub>n</sub> is the relative population of the excited state of magnesium and is obtailed by multiplying the concentration of magnesium by the Boltzmann factor at the specified temperature. Examination of Table 8 reveals that the theoretical change in the upper state concentration, N , is only a factor of about two while the observed change in intensity, I(Mg), is over a factor of 100. Obviously then there are additional factors which one must consider in order to treat this problem.

Molecular Species	Group #	<u>T°K</u>	Boltzmann Factor	[Mg]	Nn	I(Mg)*
Li	7313	3083	1.0	1.0	1.0	1.0
Na	7308	3073	0.97	0.93	0.90	0.50
к	7311	3056	0.92	0.91	0.84	0.038
Rb	7309	3008	0.80	0.64	0.51	0.010
Cs	7310	2969	0.71	0.64	0.45	0.009

TABLE 8. INTENSITY AND CALCULATED POPULATION FOR MAGNESIUM

\*Determined experimentally. All other quantities in this Table are calculated.

The experiments performed here were not in any way designed to explain this phenomena. A detailed analysis of the observed decrease in intensity is not only beyond the scope of this paper but is probably impossible due to a lack of the necessary information. Any efforts to resolve this problem will have to be undertaken with entirely different methods and procedures.

The most likely explanation for the observed decreases lies not only in a discussion of temperature and concentration but also in a discussion of the distribution of the available energy released by the chemical reactions. Thermodynamic calculations have shown that the energy available from these reactions are 1.93, 1.47, 1.20 and 0.64 kcal/g for lithium, sodium, potassium and cesium nitrate reacting with magnesium, respectively.<sup>14</sup> This calculation assumed that the reaction products were at 1200°K. The reasoning behind this assumption and a method for using these numbers to obtain total luminous power has been presented earlier.<sup>15</sup> Unfortunately for purposes of this discussion total luminous power has no significance. It should also be noted that the energies presented above are for slightly different formulas; however, the trend will be the same.

As can be seen, there is significantly less energy available in the higher molecular weight nitrates. If this energy is now distributed differently among the various atomic levels, the observed factor of 100 decrease in intensity could probably be explained. For example, if in the case of cesium nitrate, significantly more of the energy were contained in the resonance line continuum, there would be less available for populating the upper states of magnesium and magnesium oxide and the observed decrease might easily be predicted. Any more detailed analysis than this simple "handwaving" argument will be delayed until more accurate data is obtained on the distribution of energy in the products of the reactions.

Other factors which might affect the ability to predict the observed decrease in intensity include (a) the inability of thermodynamics alone to predict species concentration and reaction kinetics in the system, (b) the lack of sufficient information concerning quenching processes which occur, and (c) the problem associated with observing emitting species in a region of the flame where their concentration is low and their emission not its maximum. As was pointed out previously, since these experiments were not designed to answer any of these questions, further discussion is merely speculation and will not be pursued at this time.

An understanding of these problems is definitely needed. For example, one might be attempting to make a flare that produced significant infrared radiation while minimizing the visible radiation. If one could choose additives which would decompose to form products

which would accept some or all of the excess energy of the reaction it should be possible to quench the visible emission. If these compounds could also be chosen so that their products emitted infrared radiation or transferred the energy to an infrared emitter, one would not only have eliminated the visible radiation but would also have increased the power of the infrared radiation given off.

### CONCLUSIONS AND RECOMMENDATIONS

The purpose of this work was to identify and characterize the emission from flares containing magnesium as a fuel and the various alkali metal nitrates as oxidizers. The emission characteristics of the atomic emitters are the same as those predicted by other studies. There were no anamolous effects observed. All the predicted lines were present and no new atomic transitions were identified.

Several molecular emission bands associated with alkali metal species were observed in flares which contained sodium nitrate, potassium nitrate and rubidium nitrate. These bands were tentatively assigned and further discussion was delayed to a second report.

A surprisingly marked effect was observed in the emission from atomic magnesium and magnesium oxide as the oxidizer was changed. The emission intensity from a magnesium line at 518.36 nm decreased by two orders of magnitude when the oxidizer species was changed from lithium nitrate to cesium nitrate. This change could not be easily explained on the basis of calculated temperature and species concentration.

On the basis of this work, the following recommendations for future work can be made:

1. To add to the current knowledge of spectroscopic and molecular data, high resolution spectra of magnesium-sodium nitrate flares should be taken in the region of the magnesium oxide  $B \rightarrow X$  transition. This data would provide information about lifetimes and potential energy surfaces which is not well established.

2. Information should be obtained about the distribution of energy among the various emitters possible in a pyrotechnic flame. This work would require simultaneous spectral power measurements in all regions of the spectrum from the near ultraviolet to the middle infrared. This data would allow more refined flare formulations for producing maximum infrared power, possibly in regions as far as five micrometers. The current work appears to indicate the possibility of non-thermal energy distributions, particularly in the near-infrared region of the spectrum. If the distributions are non-thermal, then new emitting species might be added to flames which would absorb the energy now emitted in the visible and re-emit this energy in the infrared, thus increasing infrared powers. 3. Information should be obtained on species concentration in the flame to provide a better understanding of predicted flare performance. The sharp, relatively uncomplicated, magnesium and magnesium oxide lines and bands would probably be an ideal choice for making concentration measurements for comparison with calculated concentrations. At the present time, we simply assume that the thermodynamics are accurate. The results of this study tend to indicate that this might not be the case and closer examinations appear to be indicated.

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