OPTICAL CONVERSION PROCESSES

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FOREWORD

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The overall goal of this experimental program is to identify scalable techniques that efficiently convert existing high power UV lasers to lasers operating at longer wavelengths in the visible. Two nonlinear optical conversion techniques that we have considered are: stimulated Raman and parametric conversion involving the KrF laser (248 nm).

The objective of this contract is to suggest likely acceptor atoms for each technique which will thereby allow us to evaluate some of the key technical
issues involved. These include: the production of receptor candidates in the gas phase (typically refractory metals), the volumetric removal of the lower laser level in the stimulated Raman approach to prevent "bottle necking" and allow recycling of the atoms during the laser pulse, and the consideration of overall system efficiency and scalability to high power.
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I. INTRODUCTION

The overall goal of this combined experimental and theoretical program is to successfully and efficiently convert using scalable techniques the output of a high power KrF laser into longer wavelengths so as to vastly improve its propagation characteristics.

Since the first reported lasing of an inert gas halogen laser, a number of similar systems have demonstrated lasing characteristics. Operating at various wavelengths, with different efficiencies, a major class of electronic transition lasers came into existence. Recently, analogous mercury halide compounds showing similar formation kinetics have been shown to lase in the visible, \(1, 2\) albeit in high temperature (\(\sim 275^\circ\)C) cells (see Figure 1). However, the most efficient laser reported to date in this group is the KrF laser operating at 248 nm. It has also produced the highest energy outputs reported utilizing e-beam pumping and e-beam controlled discharge pumping and has a demonstrated capability for being scaled to high average power. In certain applications, especially those requiring transmission through the atmosphere, its short wavelength severely limits its usefulness. This limitation in propagation at short wavelengths arises due to absorption by atmospheric ozone and to Rayleigh scattering which increases as \(\lambda^{-4}\) as the wavelength gets shorter. Ozone absorption is severe for wavelengths \(\sim 3000 \AA\).

Figure 1  Demonstrated Potentially Scalable Electronic Transition Lasers
Figure 2 shows vertical transmission from a height of 3 km as a function of wavelength. Also plotted are quantum efficiency of conversion from KrF wavelengths and the total percentage transmission of converted KrF radiation. From the figure, it is apparent that to efficiently utilize KrF laser radiation, its conversion wavelength should be between 340 and 400 nm to maximize its atmospheric transmission with minimal loss from quantum yield considerations. Xenon fluoride lasers, while possessing a more attractive wavelength for propagation, have not yet demonstrated the combined efficiency and energy density comparable to KrF. Any optical conversion scheme for altering the wavelength of KrF laser radiation to the 340 to 400 nm wavelength range could have higher overall efficiency than the XeF laser if the photon conversion efficiency is $\geq 40\%$. Such efficiency for conversion is a reasonable goal for the program we are discussing here. For supporting evidence, one can look over the past year at a number of milestones that have been reported relevant to the optical conversion of UV excimer lasers. With regard to overall conversion efficiency, an XeF laser has been converted, at near unit photon conversion efficiency, using barium vapor.\(^3\) Also KrF conversion to a series of UV-visible lines due to 6 Stokes and 2 anti-Stokes transitions in high pressure molecular hydrogen was reported showing good overall conversion efficiency.\(^4\) Also, a number of acceptor candidates identified as potential converters for KrF have been produced in the gas phase using scalable techniques.\(^5\) In view of the above, it seems reasonable and important to pursue scalable techniques.

Figure 2  Propagation of Converted KrF Photons
that could efficiently convert KrF laser output to longer wavelengths. Two non-linear optical conversion techniques that we have considered to achieve this goal are stimulated Raman and parametric conversion processes.

For the stimulated Raman process, phenomenologically, the acceptor atom can be thought of as absorbing as incident KrF photon thereby making a transition to an excited virtual state and then, with the emission of a Raman photon at longer wavelengths, proceeding to a level near the ground (initial) state. Through collisions with an efficient quenching gas, it can return to the initial state for subsequent re-excitation by the KrF laser field, i.e., exhibit high efficiency by recycling the metal atoms. The Raman process is enhanced when the virtual state is close to a real state.

Another method of "down conversion" to lower energy, longer wavelength photons applicable to UV laser light is parametric down conversion. In this process, conversion is achieved by the utilization of the non-linear properties of the medium (the acceptor atom or molecules). Here an atom in state 0 upon exposure to KrF laser light of frequency $\nu_1$ goes to a virtual state 1 and re-emits three photons of frequencies $\nu_2$, $\nu_3$ and $\nu_4$ such that $\nu_1 = \nu_2 + \nu_3 + \nu_4$. At the end of this process, the atom returns to its initial state entirely by optical transitions. Once again, if the various atomic transitions ($\nu_1$, $\nu_2$, $\nu_3$, and $\nu_4$) in the acceptor are allowed and the dipole moments are large, near resonant effects enhance the overall process such that efficient down-conversion should be likely.

At AERL during the current reporting period, theoretical and experimental research have been carried out on potentially efficient scalable schemes for converting KrF photons to longer wavelengths. By theoretical calculations, we have identified a number of promising candidates to convert
the KrF laser radiation to longer wavelengths using parametric processes (see Table 1). This adds to the stimulated Raman candidates reported in Ref. 5. Experimentally, we have modified a commercial KrF laser (Tachisto Corp., Needham, MA) to provide a focused output beam of nearly a GW/cm² and used this laser to convert to wavelengths near 300 nm using atomic iron as the acceptor candidate. (6) The iron was produced using scalable techniques in densities sufficiently high to provide single pass amplified spontaneous emission when exposed to a field of KrF photons from an untuned laser source. In addition, we were able to demonstrate laser action in the organo-metallic precursor (i.e., the Fe(CO)₅). This represents a likely laser pumped photodissociation process followed by inversion of a photofragment produced in the initial step. The results of these combined theoretical and experimental efforts are summarized in the following sections of this report.

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<td>&quot;</td>
<td>~364</td>
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<tr>
<td>Mercury</td>
<td>&quot;</td>
<td>~376</td>
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<td>Lithium</td>
<td>&quot;</td>
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II. THEORY

A. INTRODUCTION

The stimulated Raman process and the calculations performed for
the ARPA program on optical conversion were discussed in detail in the last
semi-annual report. In this report, calculations performed on the four-
wave parametric down conversion will be discussed. In the parametric down
conversion process, an atom in state 0 interacting with the KrF laser light
of frequency \( \nu_1 \) goes to a virtual state 1 and emits three photons of fre-
quencies \( \nu_2, \nu_3 \) and \( \nu_4 \) such that \( \nu_1 = \nu_2 + \nu_3 + \nu_4 \) (Figure 3). At the
end of this process the atom returns to the initial state entirely by optical
transitions. None of the electromagnetic energy in the parametric process
is absorbed by an ideal medium in contradistinction to the stimulated Raman
Stokes process in which the atom is left in an excited state at the end of the
nonlinear process. Thus the problem of heat removal for a large scale con-
verter is less severe in the parametric approach.

An interesting candidate atom for the parametric down conversion of
KrF laser radiation is thallium which will be used as a concrete example in
the following discussion. In the down-conversion process, the photon at fre-
quency \( \nu_2 \) is initially produced by spontaneous and stimulated Raman scat-
tering. Since there are usually several channels available for stimulated
Raman scattering, one should first calculate the Raman cross section for the
different possible final states of the atom and choose the one that is most
probable for determining \( \nu_2 \). The frequencies \( \nu_3 \) and \( \nu_4 \) are then chosen
Figure 3 Schematic of the Parametric Down Conversion Process

\[ \chi^{(3)} = \frac{N \mu_{01} \mu_{12} \mu_{23} \mu_{30}}{(\Delta E_1)(\Delta E_2)(\Delta E_3)} \]

DIFFERENCE MIXING \( \nu_4 = \nu_1 - \nu_2 - \nu_3 \)
to maximize the nonlinear susceptibility, $\chi^{(3)}$, subject to phase matching condition. In Section II.B, the stimulated Raman cross section is calculated for the three possible final states of a thallium atom irradiated with a KrF laser. In Section II.C, the dispersion of the four waves is calculated and phase matching is discussed. The nonlinear susceptibility, $\chi^{(3)}$, for the fourwave process is calculated in Section II.D, while in Section II.E, the coupled set of nonlinear equations for the field amplitudes is solved and the resulting conversion efficiency is calculated.

**B. STIMULATED RAMAN CROSS SECTION**

The energy levels of thallium atom are shown in Figure 4. Spontaneous or stimulated Raman scattering of KrF laser radiation of energy $\sim 40225 \text{ cm}^{-1}$ by an atom in the ground state can only occur to the $6p^2P_{3/2}$, $7p^2P_{1/2}$ or $7p^2P_{3/2}$ levels. Transitions to other states are forbidden by parity selection rules. In this section, we shall calculate the stimulated Raman cross section for these three possible final states. The calculation of nonlinear susceptibilities is usually a tedious exercise. The susceptibility formulae contain products of matrix elements of the dipole operator $\vec{Q} = e \vec{r}$, of the form $< \gamma J M | \vec{Q} | \gamma' J' M' >$ which connect the initial and final atomic states through a chain of intermediate (virtual) states. Yuratich and Hanna\(^{(7)}\) have used Racah algebra to sum over the intermediate state $M$ values and write the nth order nonlinear susceptibility in a form that can be factorized into two parts, in one of which the relation between the fields is explicitly displayed; the other contains the physics of the atom in the form of reduced matrix elements (which can be related to oscillator strengths) and $6j$ symbols.

In what follows, we shall use the symbols of Ref.\(^{(7)}\) as much as possible.

Figure 4  Energy Levels in Thallium
The Raman susceptibility $\chi_r$ per atom is given by

$$\chi_r = \frac{i}{6} \frac{\rho_{aa} - \rho_{cc}}{\hbar^3 \epsilon_o} \sum K \Theta^{(K)}$$

$$\times \left[ \sum_{\gamma_2 J_2} \left\{ \frac{1}{\gamma_2 J_2} \frac{1}{\gamma J} \right\} < \gamma_1 J_1 || Q || \gamma_2 J_2 > < \gamma_2 J_2 || Q || \gamma J > \right]$$

$$\times \left[ \frac{1}{\gamma_2 J_2} \frac{1}{\gamma J} - \omega_p + \frac{(-1)^K}{\gamma_2 J_2} \frac{1}{\gamma J} \omega_s \right]^2$$

$$< \gamma J || Q || \gamma' J' >^2 = \frac{3 \hbar e^2}{2m} \frac{(2J + 1)}{\hbar \gamma J - \gamma' J'}$$

In the above, $\rho_{aa}$ and $\rho_{cc}$ represent the initial and final state fractional populations, $2J + 1$ and $2J_1 + 1$ their corresponding degeneracies whereas $\gamma_2 J_2$ represents the intermediate states; $\Omega_{\gamma_2 J_2} = (E_{\gamma_2 J_2} - E_{\gamma J})/\hbar$, $\omega_p$ and $\omega_s$ are the pump and Raman frequencies respectively; $< \gamma J || Q || \gamma' J' >$ is the reduced matrix element and $\Gamma_{ca}$ is the damping term between levels $c$ and $a$. The factor $\Theta^{(K)}$ contains all the information about the angular dependence between the pump and Raman fields. Even though the summation in Eq. (1) extends over all the intermediate states, there are only three dominant terms in the case of thallium: those in which the intermediate states are $8s^2 S_{1/2}$, $6d^2 D_{3/2}$ and $7d^2 D_{3/2}$. The value of the reduced matrix elements is obtained through their relationship to the oscillator strength:

$$< \gamma J || Q || \gamma' J' >^2 = \frac{3 \hbar e^2}{2m} \frac{(2J + 1)}{\hbar \gamma J - \gamma' J'}$$

The oscillator strengths for $p$ to $s$ and $p$ to $d$ transitions have been calculated by Anderson et al. (8) and have also been experimentally determined by

The signs of the matrix elements are determined from the work of Vriens who used a combination of experimental and theoretical results to deduce these for indium and thallium. The angular dependence factor \( \Theta^{(K)} \) for the Raman process has been calculated by Yuratich and Hanna. For our purposes, we shall assume that both the pump and Raman fields are linearly polarized and for this case, we have

\[
\begin{align*}
\theta^{(0)} &= \frac{1}{3} \cos^2 \beta \\
\theta^{(1)} &= \frac{1}{2} \sin^2 \beta \\
\theta^{(2)} &= \frac{1}{6} (3 + \cos^2 \beta)
\end{align*}
\]

where \( \beta \) is the angle between the pump and Raman fields.

The stimulated Raman emission cross section \( \sigma_{\text{SRE}} \) i.e., related to the Raman susceptibility \( \chi_r \) by

\[
\sigma_{\text{SRE}} = -\frac{12 \pi^2 i \epsilon_0}{\lambda_r} \left( \frac{8 \pi x 10^7}{c} \right) I_p \chi_r
\]

where \( I_p \) is pump intensity in watts/cm\(^2\) and \( \lambda_r \) is the wavelength of the Raman radiation. The calculated \( \sigma_{\text{SRE}} \) at the three Raman wavelengths 3083 Å, 1.649 μ and 1.975 μ (when the atom goes respectively to \( 6p^2P_{3/2} \), \( 7p^2P_{1/2} \) and \( 7p^2P_{3/2} \) states) are respectively

\[
\begin{align*}
\sigma_{\text{SRE}}(6p^2P_{3/2}) &= 1.72 \times 10^{-28} \frac{I_p}{\Gamma_{ca}}, \text{ cm}^2 \\
\sigma_{\text{SRE}}(7p^2P_{1/2}) &= 3.40 \times 10^{-27} \frac{I_p}{\Gamma_{ca}}, \text{ cm}^2 \\
\sigma_{\text{SRE}}(7p^2P_{3/2}) &= 1.38 \times 10^{-26} \frac{I_p}{\Gamma_{ca}}, \text{ cm}^2
\end{align*}
\]

where $\bar{c}_{\text{ca}}$ is in cm$^{-1}$. The line width is taken to be the same in all three cases since the dominant width is the pumping laser width ($\sim 100$ cm$^{-1}$), natural and doppler widths being much less than 1 cm$^{-1}$. The largest stimulated emission cross-section is for the $6p^2P_{1/2} \rightarrow 7p^2P_{3/2}$ transition and thus $v_2$ is expected to be $\sim 5064$ cm$^{-1}$.

C. PHASE MATCHING FOR THE FOUR-WAVE PROCESS

In any parametric process where the atomic state of the system remains unchanged during the nonlinear optical conversion process, both momentum and energy of the photons have to be conserved. In the four wave down conversion, this means that efficient down conversion will be achieved only if the $\mathbf{K}$ vectors of the three converted waves add up to the $\mathbf{K}$ vector of the KrF pump wave. Since we are dealing with near resonant down conversion it may be possible to phase match either intrinsically or by the addition of buffer gas in the right proportion. The former method is to be preferred if it is possible since it reduces the complexity of generating the right phase matching condition.

In the case of intrinsic phase matching, again two possibilities exist. In the first case, the frequency $v_4$ is adjusted to produce the correct amount of dispersion so that all waves propagate collinearly. This may be possible because of the large dispersion of the electromagnetic wave at $v_4$ near an atomic resonance. In the second case, the intrinsic phase matching can be achieved by having the wave vectors $\mathbf{k}_2$, $\mathbf{k}_3$, and $\mathbf{k}_4$ corresponding to the frequencies $v_2$, $v_3$ and $v_4$ respectively propagate at slight angles to each other such that their vectorial sum adds up to the pump wavevector $\mathbf{k}_1$. Obviously, this can be done only when the algebraic sum of $k_2$, $k_3$ and $k_4$ is larger than $k_1$. In a lossy medium the wave number $k$ is larger than
the free space wave number at frequencies slightly below a resonant frequency. In the case of KrF laser down conversion in thallium angle phase matching should therefore be possible for \( \nu_4 \) chosen to be less than the

\[ 6p^2P_{1/2} \to 7s^2S_{1/2} \]

transition energy (26477 cm\(^{-1}\)).

An important point that is to be considered in phase matching is the real absorption of waves generated at \( \nu_4 \) by the medium. The closer \( \nu_4 \) is to a resonance, the larger is the absorption; \( \chi^{(3)} \) is also larger near a resonance. However, the absorption coefficient in the wings of a Lorentzian line decreases, as \((\Delta \varepsilon)^{-2}\), while \( \chi^{(3)} \) decreases as \((\Delta \varepsilon)^{-1}\). Hence, \( \nu_4 \) should be chosen to minimize absorption while keeping \( \chi^{(3)} \) reasonably high. For the case of thallium, collinear intrinsic phase matching seems possible only where \( \nu_4 \) is less than 1 cm\(^{-1}\) from the \( 6p \to 7s \) resonance transition; unfortunately, absorption at this frequency is expected to be very large. It is therefore not a good operating point for the down converter. Noncollinear angle phase matching is therefore considered below for thallium.

The refractive index, \( n(\nu) \), at a frequency \( \nu \) is given by the Sellmeier equation (11)

\[
n = \frac{k c}{\omega} = 1 + \frac{R e}{2 \pi} \sum_{i,j} \frac{N_i f_{ij}}{v_{ij}^2 - \nu^2},
\]

where

- \( R e = 2.818 \times 10^{-13} \) cm
- \( f_{ij} \) = oscillator strength of the \( i-j \) transition
- \( v_{ij} \) = energy difference between \( i \) and \( j \) levels in cm\(^{-1}\)
- \( N_i \) = number density of particles in the \( i^{th} \) level

Equation (6) is only valid far from any resonance; i.e., when \( |v - v_{ij}| \) is at least 5 or 10 times the width of the line at \( v_{ij} \). Using \( f_{ij} \) values from Ref. 8, \( N_1 \) from the vapor pressure curve of Nesmeyanov\(^{(12)}\) and \( v_{ij} \) values from the tables of Moore,\(^{(13)}\) the wave number \( k \) at any frequency \( v(=\omega/2\pi c) \) can be calculated as a function of medium temperature when thallium vapor is in equilibrium with the liquid phase at that temperature.

For phase-matching the following two equations have to be satisfied:

\[
(k_4 + k_3) \cos \Theta + k_2 \cos \phi = k_1 \tag{7}
\]

and

\[
(k_4 + k_3) \sin \Theta = k_2 \sin \phi, \tag{8}
\]

where \( \Theta \) and \( \phi \) are the angles that \( \vec{k}_3 \) and \( \vec{k}_2 \) make with respect to pump field \( \vec{k}_1 \) (Figure 5). In the above, it is assumed that \( \vec{k}_3 \) is parallel to \( \vec{k}_4 \). This is not a necessary condition for phase matching, but it gives the smallest angle \( \Theta \). Also the roles of \( \vec{k}_2, \vec{k}_3 \) and \( \vec{k}_4 \) can be arbitrarily interchanged.

The initial generation of photons at \( v_2 \) takes place due to the Raman process and \( v_2 \) can come at any angle. The direction \( \Theta \) can therefore be chosen preferentially by making a cavity at frequency \( v_2 \) and making the KrF beam come at an angle to the axis of the cavity. This was the reason for choosing \( k_1 \) and \( k_2 \) noncollinear. In Figure 6, \( \phi \) is plotted as functions of output wave number \( v_4 \) at different vapor temperatures. The precise operating point on this curve is chosen by maximizing the conversion efficiency and minimizing the absorption. These will be discussed more fully in the next two sections.


Figure 5  Schematic of Phase Matching
Figure 6  Phase-Matching Angle vs Output Wavenumber in Thallium
D. THE THIRD-ORDER NONLINEAR SUSCEPTIBILITY

Having identified the likely output wavelengths, we are now in a position to calculate the third-order nonlinear susceptibility for the four wave down conversion. The third-order nonlinear susceptibility \( \chi^{(3)}(\omega_1; \omega_2, \omega_3, \omega_4) \) if given by (7)

\[
\chi^{(3)} = -\frac{1}{6 \hbar^3 \varepsilon_0} \sum_{J_1, J_2, J_3} \sum_{K} \frac{(-1)^{3J + K + J_2}}{(2J + 1)(2K + 1)^{1/2}}
\]

\[
< J \parallel Q \parallel J_1 > < J_1 \parallel Q \parallel J_2 > < J_2 \parallel Q \parallel J_3 > < J_3 \parallel Q \parallel J >
\]

\[
\left\{ \begin{array}{c} J \quad K \quad J_2 \quad \{ J_1 \quad J_3 \} \\ 1 \quad J_1 \quad 1 \quad 1 \quad J_2 \quad K \end{array} \right\} F(K)
\]

where

\[
F(K) = \frac{\left((\varepsilon_1 \times \varepsilon_2)^{(K)} \times \varepsilon_3^{(1)} \times \varepsilon_4^{(0)}\right)}{(\Omega_{JJ_1} + \omega_2 + \omega_3 + \omega_4) (\Omega_{JJ_2} + \omega_3 + \omega_4) (\Omega_{JJ_3} + \omega_4)}
\]

and \( S_T \) is the overall permutation operator indicating that the expression following it is to be summed over all permutations of the pairs \( \varepsilon_1 \omega_1, \varepsilon_2 \omega_2 \) etc. \( \varepsilon_i \) is the polarization vector for the \( i \)th wave. It is easy to show that for \( K = 1 \), \( F(K) = 0 \) if all the waves are linearly polarized and thus only the cases \( K = 0 \) and \( K = 2 \) have to be considered. If we define a frequency factor \( D^{-1}(J_1, J_2, J_3) = F(K) \) with numerators put equal to unity, it is found that

\[
F(0) = \frac{1}{3} D
\]

\[
F(2) = \frac{2}{3 \sqrt{5}} D
\]

24
It is now a simple matter to estimate $\chi^{(3)}$ for the parametric down conversion process.

The dominant terms are when $\gamma J = 6p_{1/2}$, $\gamma J_1 = 8s_{1/2}$ or $7d_{3/2}$ or $6d_{3/2}$ $\gamma J_2 = 7p_{1/2}$ or $7p_{3/2}$ or $\gamma_3 J_3 = 7s_{1/2}$. There are thus six dominant terms in the summation in Eq. (9). The only unknown in Eq. (9) then are the phases of the reduced matrix elements. These are obtained from the theoretical and experimental information about the phases of the radial matrix elements available in the literature(10, 14) and using an LS or jj - coupling scheme to relate the radial matrix elements to the reduced matrix elements. The resonant denominator $D$ in $\chi^{(3)}$ contains three terms $\Delta \nu_1$, $\Delta \nu_2$ and $\Delta \nu_3$. The value of $\Delta \nu_1$ can be readily calculated knowing the KrF wavelength and the energy level of the first intermediate state. $\Delta \nu_2$ is actually equal to the width of the Raman line initially generated and can be taken equal to the width of the KrF laser pump. $\Delta \nu_3$ is a variable such that phase matching condition can be satisfied while keeping the absorption at the frequency $\nu_4$ to a minimum. The estimated $\chi^{(3)}$ from Eq. (9) for thallium atom is given by

$$\chi^{(3)} \approx (1.7 \times 10^{-29} + 5.1 \times 10^{-26}/\Delta \nu_2)/\Delta \nu_3 \text{ e.s.u.} \quad (12)$$

where $\Delta \nu_2$ and $\Delta \nu_3$ are in cm$^{-1}$. For a KrF laser that is not narrowed, the line width is $\sim 100 \text{ cm}^{-1}$, while one whose line is narrowed with an etalon for example may be $\sim 10 \text{ cm}^{-1}$. Taking $\Delta \nu_2$ to be $100 \text{ cm}^{-1}$, we get,

$$|\chi^{(3)}| \approx 5.3 \times 10^{-28}/\Delta \nu_3, \text{ e.s.u./atom} \quad (13)$$

The absorption cross section in line center is given by

$$\sigma_o = \pi r_e f/\Delta \nu$$

(14)

where $\Delta \nu$ is the width of the line in cm$^{-1}$. For the $6p_{1/2} \rightarrow 7s_{1/2}$ transition in thallium, the natural width is $\sim 2 \times 10^{-3}$ cm$^{-1}$, the doppler width at 1800 K is $\sim 2 \times 10^{-2}$ cm$^{-1}$ and collision broadened width at 1 atm is $\sim 0.2$ cm$^{-1}$. Taking the collision broadened width for $\Delta \nu$, we find the absorption cross section in line center to be $\sim 5.5 \times 10^{-13}$ cm$^2$. Since the natural line is Lorentzian, the absorption cross section at a frequency tuned $\Delta \nu$ cm$^{-1}$ away from line center can be calculated from $\sigma_o (\Delta \nu /2 \Delta \nu)^2$. Thus for $\Delta \nu \sim 200$ cm$^{-1}$, the absorption cross section is $\sim 1.4 \times 10^{-19}$ cm$^2$. The third-order nonlinear susceptibility at this detuning frequency is about $2.6 \times 10^{-30}$ e.s.u./atom. It will be shown later that for thallium, this choice of $\Delta \nu$ produces a reasonable optimum of having large $\chi^{(3)}$ and tolerable absorption at the output frequency. In the following section, we solve the coupled nonlinear equations for the evolution of the field amplitudes taking into account absorption. The density of parametric medium necessary and its length for efficient conversion is also calculated.

E. CONVERSION EFFICIENCY

The coupled amplitude equations describing the parametric four-wave conversion process are

$$\frac{dA_1}{dz} = -\beta_1 A_2 A_3 A_4 - a_1 A_1$$

(15)

$$\frac{dA_2}{dz} = +\beta_2 A_1 A_3 A_4$$

(16)
\[
\frac{dA_3}{dz} = \beta_3 A_1 A_2 A_4 \tag{17}
\]
\[
\frac{dA_4}{dz} = \beta_4 A_1 A_2 A_3 - a_4 A_4 \tag{18}
\]

where \( A_j \) is the amplitude of the electric field of the \( j \)th wave, \( a_j \) is the linear absorption coefficient and \( \beta_j \) is the nonlinear coupling constant \( (\beta_j = 3N k_j x^{(3)}/4n_j) \). Only waves 1 and 4 are assumed to have absorption in the medium since they have resonance with the initial state of the atoms. It is not difficult to include absorption of the other two waves as well. It is convenient to rewrite the Eqs. (15) - (18) in a non-dimensional form with the help of the following substitutions:

\[
y_j = \frac{A_j}{A_{10}} \sqrt{\frac{\omega_1}{\omega_j}} \tag{19a}
\]
\[
\xi = \frac{z}{L} \tag{19b}
\]

where \( A_{10} \) is the initial amplitude of the pump wave and \( L \) is the length of the nonlinear medium. The reduced equations are,

\[
\frac{dy_1}{d\xi} = -\beta y_2 y_3 y_4 - \gamma_1 y_1 \tag{20}
\]
\[
\frac{dy_2}{d\xi} = \beta y_1 y_3 y_4 \tag{21}
\]
\[
\frac{dy_3}{d\xi} = \beta y_1 y_2 y_4 \tag{22}
\]
\[
\frac{dy_4}{d\xi} = \beta y_1 y_2 y_3 - \gamma_4 y_4 \tag{23}
\]
where

\[
\beta = \frac{12 \pi^2}{c \lambda_4} \left( \frac{\omega_2 \omega_3}{\omega_1 \omega_4} \right)^{1/2} I_{10} N L \gamma^{(3)},
\]

\[
\gamma_1 = a_1 L \text{ and } \gamma_4 = a_4 L.
\]

In the above, \(I_{10}\) is the incident intensity of the pump wave and \(N\) is the density of the nonlinear medium. \(\gamma_j^2\) gives the normalized photon density in the \(j\)th mode and the conversion efficiency is given by \(\gamma_4^2\) since \(\nu_4\) is the output frequency of interest. Equations (20) - (23) can be solved exactly analytically in the absence of losses. The initial conditions are, \(\gamma_1 = 1, \gamma_2 = \delta, \gamma_3 = \Delta\) and \(\gamma_4 = 0\) at \(\zeta = 0\). The analytical solution involves Jacobi's elliptic functions and the expression for the conversion efficiency \(\eta\) is given by

\[
\eta = \frac{\Delta \text{ sn}^2 (u, \rho)}{\Delta + 1 - \text{ sn}^2 (u, \rho)} \quad (24)
\]

where

\[
u = \beta \left[ \delta (1 + \Delta) \right]^{1/2}
\]

\[
\rho = \left[ \frac{\delta - \Delta}{\delta (1 + \Delta)} \right]^{1/2}
\]

It must be emphasized that the expression (24) is valid only for the case of no losses. Typical values of \(\Delta\) and \(\delta\) might be \(10^{-3}\). When \(\delta = \Delta\), we have \(\rho = 0\) and the Jacobian elliptic function \(\text{sn} u\) reduces to \(\sin u\). Complete conversion is obtained when \(\text{sn} u = 1\), which means \(u = \pi/2\). This implies that \(\beta\) should be equal to \(\sim \pi/2 \sqrt{\delta}\) for complete conversion.

Figure 7 shows a plot of conversion efficiency vs \(\beta\) for typical values of \(\delta\) and \(\Delta\). When we take into account the absorption of the waves, it is in general
Figure 7  Conversion Efficiency vs $\beta$, $\gamma_1 = \gamma_4 = 0$
not possible to obtain closed form analytical solutions. Equations (20) - (23) are, however, easily solved in a computer. Figures 8 to 10 show typical results of such computer runs. It is evident from the figures that absorption at $\nu_4$ affects the generated output much less than absorption of the pump wave. This is not surprising since the bulk of $\nu_4$ is generated over a small length at the very end of the nonlinear medium while absorption of $\nu_1$ takes place over the entire length of the nonlinear medium.

For $\delta \approx 10^{-3}$, we find that we need $\beta \approx 50$ for efficient parametric conversion. If one of the three generated waves is initially created by the Raman process, the corresponding resonant denominator should be replaced by the pump laser linewidth. Thus for a diffraction limited pump laser, the conversion parameter $\beta$ can be written as

$$\beta = \Lambda N P / \gamma_l$$

where $P$ is the KrF laser power in watts, $\gamma_l$ is the laser linewidth in cm$^{-1}$ and $\Lambda$ is an effective cross section in cm$^2$/w. In the case when $\Delta \nu_1$ and $\Delta \nu_3$ are very small, the expression for $\chi^{(3)}$, is dominated by a single term, and $\Lambda$ can be written as

$$\Lambda \approx \frac{9}{4 \pi^2} \frac{r e^2}{\hbar c^2} \times 10^7 \left( \frac{(gf)_{01} (gf)_{21} (gf)_{32} (gf)_{03}}{\Delta \nu_1 \Delta \nu_3} \right)^{1/2}$$

$$\times \sum_K \left( \frac{2K + 1}{3(2J + 1)} \right)^{1/2} \begin{vmatrix} J & K & J_2 \\ 1 & J_1 & 1 \end{vmatrix} \begin{vmatrix} J & K & J_2 \\ 1 & J_1 & 1 \end{vmatrix}$$

$$x \left\{ \delta_{K,0} + 2\delta_{K,2/\sqrt{5}} \right\}$$

(26)
Figure 8  Conversion Efficiency vs $\beta$, $\gamma_1 = 0$, $\gamma_4 = 1$
Figure 9  Conversion Efficiency vs $\beta$, $\gamma_1 = 0.2$, $\gamma_4 = 1$
Figure 10  Max Conversion Efficiency vs $\gamma_4$
The quantity $\Lambda$ is an atomic parameter for a set of given frequencies. In Figure 11 a map of constant $P/\gamma_f$ is plotted in the coordinates of $\Lambda$ and $N$. Each atomic system is represented on this map by a horizontal line, whose position depends upon the density of the medium that may be produced in the gas phase at reasonable temperatures. The line of constant $P/\gamma_f$ have been plotted on the assumption that $\beta$ equals 50. Three elements are plotted on this map. These are thallium, lithium and lead. From the map it is seen that both lithium and thallium are good parametric down conversion candidates from the point of view of ease of obtaining the operating density. However, both require angle phase matching for intrinsic phase matching. Collinear phase matching conditions may be obtained by the addition of buffer vapor like mercury.

In the case of thallium an operating density of $10^{17}$ cm$^{-3}$ over a path length of 100 cm will give an integrated absorption depth of ~ 1.4 which is acceptable for a down converter. This justifies our choice of $\Delta \nu_3 \sim 200$ cm$^{-1}$ to keep $\chi^{(3)}$ reasonably large while the absorption cross section remains at a tolerable level.
Figure 11  Map of $\Delta$ vs $N$ for Certain Candidate Atoms Considered for the Parametric Down Conversion
III. EXPERIMENTS

A. INTRODUCTION

The successful demonstration of optical conversion in any laser system through stimulated Raman or Parametric conversion requires optimization of a number of variables to achieve reasonable gain, namely, overall system cross section, which is in turn influenced by the laser intensity, bandwidth and wavelength, and also the ability to produce sufficient densities of acceptor candidates to show reasonable conversion. We will describe the results and status of this program with regard to the stimulated Raman candidates in the next two sections and then describe how these systems were coupled to producing lasing in a couple of candidates in the section titled "Lasing Experiments."

B. ACCEPTOR CANDIDATE PRODUCTION

Under the early phases of this contract, the stimulated Raman and direct optical pumping cross sections for various candidates were calculated. For the stimulated Raman process, the total gain, $g_o L$, for a single pass system is equal to $N_o S R E L$, where $N$ is the density of atoms and $L$ is the active length of Raman medium. For a diffraction limited beam, the maximum value of the intensity-length product is equal to $P/\lambda_p$ where $P$ is the pump laser power and $\lambda_p$ is the pump wavelength. This, $g_o L$ can be written as,

$$g_o L = \Lambda NP/\gamma_f$$  \hspace{1cm} (28)
where

\[ \Lambda = 9.55 \times 10^{-14} \left( \frac{v}{\nu_{23}} \right) \frac{f_{12}(gf)_{32}}{\Delta v^2} \sum_{K=0}^{2} \left\{ \begin{array}{ccc} J_{3} & K & J_{1} \\ 1 & J_{2} & 1 \end{array} \right\}^2 \theta(K) \text{ cm}^2/\text{W}, \]

\( \gamma_{\ell} \) is the bandwidth in \( \text{cm}^{-1} \), and \( P \) is in watts. The quantity \( \Lambda \) is then an atomic parameter for a given pump wavelength and given pump and Raman field polarizations. In what follows, we shall assume for convenience, that both pump and Raman fields are linearly polarized and are traveling in the same direction.

In Figure 12, a plot of constant \( g_{o}L \) is presented in the coordinates of \( \Lambda \) and \( N \). Each atomic system is represented on this graph by a horizontal line, whose position depends upon the density of the medium that may be produced in the gas phase and extent represents the degree of uncertainty in experimentally achievable atom production. The lines of constant \( g_{o}L \) have been plotted on the assumption that \( P/\gamma_{\ell} \) of the pump laser equals \( 10^5 \) W cm. From this plot and the results of experiments we performed to produce the needed metal atom densities, using the techniques of flash photolysis or discharge dissociation of organo-metallics, we can conclude that for KrF down conversion, the most promising conversion systems, in terms of ease of demonstration of principle, are molecular hydrogen, atomic iron in the gas phase, and calcium vapor in a heat pipe. Although our proposal candidate, platinum, does not show as large a projected gain as does atomic iron, it has the advantage of potentially shifting the KrF output to more propagating wavelengths. One aspect many of these systems have in common is the effect of direct optical pumping as a competing process to near resonant or resonant Raman processes. It was useful,
Figure 12  Anticipated Stimulated Raman Gain of Various Accepted Candidates
therefore, to consider these effects using iron as an example, since experiments with a glow discharge showed we could produce atomic iron densities in excess of $8 \times 10^{13}$ atoms/cc using iron pentacarbonyl as the precursor. This is equivalent to a thermal temperature near $1600 \, ^\circ K$.) We were also able to generate significant densities of atomic iron by the flash photodecomposition of $\text{Fe(CO)}_5$ in a buffer of argon.

These calculations, which were summarized in our previous semiannual report, showed the anticipated gain for direct optical pumping was about 250 times greater than for the Raman process. We, therefore, had every expectation that the atomic iron system should efficiently convert the KrF pump radiation to longer wavelengths near 300 nm.

C. PUMP LASER SOURCES

It was the original premise of this experimental approach to use a double dye laser as the source to perform conversion experiments. This system would have the advantage of being spectrally narrow ($\Delta \lambda \sim 1 \, \text{Å}$ near 248 nm) as well as tunable. After much difficulty, the vendor (Phase-R Corp.) was able to provide a tunable, flashlamp pumped dye laser utilizing Coumerin 504 dye (Exciton Corp.) and a cooled ADP crystal for frequency doubling (see Figure 13). Output was measured to be near 5 mJ in a 500 nsec pulse. This laser was found to be useful to perform absorption experiments of the organo-metallic/buffer gas photodecomposition/discharge dissociation mixtures to determine loss processes at the pump laser frequency. Referring to Figure 12, it can be seen that its output ($P/\gamma \sim 10^3$ W/cm) is not sufficient for many of these systems to be useful as a pump laser. It should be possible, however, to use it in a MOPA configuration to be amplified in a discharge laser device to provide somewhat tunable spectrally narrow, high power pulses.
Figure 13  Doubled Dye Laser System
The conversion experiments were performed using a discharge initiated KrF excimer laser (Tachisto Corp., Needham, MA). As delivered, this laser provided approximately 50 mJ of output in a 20 nsec pulse with a stated beam divergence near 4 mrad. The output is a rectangle ~15 mm x 4 mm. To achieve values of $P/\gamma$ as high as possible, it was necessary to improve the laser beam quality so it could be focused tightly. This was accomplished using the technique described by Loree et al of LASL and consisted of replacing the supplied output coupling mirror with a 50 cm plano-convex lens (suprasil). This provided an unstable cavity configuration of quite a good beam quality which focused near 50 cm from the Tachisto exit port. Typical output was near 30-40 mJ in 20 nsec which, therefore, provided approximately 1-2 MW of power. One and two shot exposures of the output were analyzed to obtain the spectral width and it was found to be near $10 \AA$ (~160 cm$^{-1}$ fwhm). This suggests that for this pump laser a reasonable value for $P/\gamma$ is $1-2 \times 10^4$ Wcm. Considerable improvement can be made by narrowing the bandwidth with subsequent increased ease of conversion demonstration (see Figure 12).

D. LASING EXPERIMENTS

1. Atomic Iron

As a check on our overall system, we opted to repeat the hydrogen conversion experiments of the LASL group using a 60 cm stainless steel shock tube section filled to near 10 atm with hydrogen. We were able to observe 4 or 5 Stokes shifted lines as well as one anti-Stokes line using a simple prism spectrograph and polaroid film for observation. Satisfied with our laser's performance and implied beam quality, we then
attempted our best candidate in terms of ease of demonstration, i.e., atomic iron. We were able to show on our initial attempts, single pass amplified spontaneous emission near 300 and 304 nm (see Figure 14) using discharge dissociation of Fe(CO)$_5$ in 50 torr of neon to provide the required atomic iron density. Subsequent spectra taken with a moderate resolution spectrograph showed that for high iron densities we were seeing output on four different lines (see Figure 15). The apparent doublet shape of the KrF laser pulse is real and is likely due to the presence of a strong absorber in the discharge cell. By superimposing some atomic mercury spectra onto these type of data, we were able to correlate our observations of wavelength with the known transitions in the atomic iron system to confirm the origin for these lasing lines. These conclusions are summarized in Figure 16. We were able to delay the time the KrF pump laser fired relative to the discharge and, thereby, probe the spectral output variation with the decaying atomic iron density. The intensities of the various lines were then probed through densitometer traces of the photographic plates. These data are plotted as intensity peak height in arbitrary units vs time on a semi-logarithmic scale (see Figure 17). From these data, it is readily seen that the 300 nm transition shows the greatest output and persists with decreasing iron density due to its higher total gain in agreement with theoretical calculations.

In addition to doing spectrally resolved measurements, we performed experiments with calibrated photodiodes to establish temporal characteristics and measure overall conversion efficiency (see

Figure 14  Atomic Iron Laser
Figure 15  Optically Pumped Atomic Iron Laser
Figure 16  Atomic Iron Lasing Transitions
Figure 17  Spectral Distribution as a f (Atomic Iron Density)
Figure 18). As can be seen from the figure, we could also measure simultaneously, through atomic absorption techniques using an iron hollow cathode, the production and decay of the various iron states produced in the discharge cell. The shot to shot output of the KrF pump laser was also monitored by an additional photodiode. Typical data are shown in Figure 19.

From the absorption information, it can be seen that the maximum density of atomic iron in the lowest spin-orbit state $^5D_4$ is observed at times near 300 µsec after the initiation of the discharge. Any direct discharge production must occur on a faster time scale (see Figure 20). This additional iron atom production beyond the off time of the discharge probably arises from energy transfer reactions and radical reactions as well as cascading of higher lying excited states of iron into the lowest energy state. From the work of Callear and Oldman (16), it is likely that the $J = 0, 1, 2$ and 3 levels of the $^5D$ state under our experimental conditions quickly (100 µsec) establish a Boltzmann distribution, then as a group decay into equilibrium with the $^5D_4$ state. The characteristic time of this latter process varies with the nature of the quenching gas present. For 50 torr of neon, it's likely to be of the order of several msec, whereas hydrogen was measured to relax these states with a rate constant of $7.4 \pm 0.7 \times 10^{-12}$ cm$^3$/sec. By adding 3 torr of hydrogen to the 50 torr of neon, we observed increased laser output at earlier times probably at 300 and 304 nm and at the expense of the 320 or 330 nm. Also, data showing the time for peak signal (see Figure 21) coincide closely with the atomic absorption data shown in Figure 19, supporting the induction time for optimum iron production.

Figure 18 Experimental Approach
Figure 19 Experimental Data
Figure 20  Discharge Cell Characteristics
Figure 21  Converted Laser Output vs Atomic Iron Density
In addition to showing conversion of KrF with atomic iron produced in a discharge, we were able to demonstrate lasing with iron produced by the photodecomposition of Fe(CO)$_5$ in a buffer of Ar. In contrast to the discharge, we saw prompt iron atom production (see Figure 22) and therefore chose to fire the laser at times near 40 $\mu$sec after the flash initiation. The flashlamp temporal behavior is such that it exhibits a fwhm of nearly 8 $\mu$sec and a time to 98% extinction of approximately 40 $\mu$sec. Since we routinely produced more iron atoms in the discharge than in the flash (e.g., $4 \times 10^{12}$ vs $3 \times 10^{11}$ atoms/cc), we observed higher conversion efficiency in the discharge method. These data suggest that for the higher iron densities, we demonstrated overall conversion efficiency of 1-2%. This is in complete agreement with our expectations. As discussed in our proposal, to produce higher conversion efficiency with these refractory candidates, one must recycle the atoms during the KrF laser pulse time scale, i.e., return the lower laser level population to the initial state through collisions with an efficient quenching gas for subsequent re-excitation by the laser field (see Figure 23). In that the pump laser used in these experiments had a 20 nsec pulse, it is difficult to demonstrate this effect. By adding sufficient quenching gas to relax on a time scale fast compared to 20 nsec, the likelihood of competing unfavorably with the spontaneous emission lifetime ($\sim 2$ nsec) is assured whereas any scaleup KrF laser system is projected to have pulse lengths of order 1 $\mu$sec and the recycling time requirement is thereby significantly reduced. We did, however, investigate the effect of some of the quenching candidates on the flash production of iron. These results are shown in Figure 24. These data are interpreted as evidence that the initial production of iron atoms is not significantly affected by the
Figure 22  Flash vs Discharge: Iron Laser Output
• MECHANISM

ORGANO-METALLIC

+ PHOTONS

↓

QUENCHER KrF LASER

M* + hν M**

• KINETICS

LOSS M + M + X → M2 + X \( \tau = 0.5 \) msec

RECYCLE M* + Q → M + Q \( \tau = 3 \) nsec

Figure 23  Production and Recycling of Metal Atoms
Figure 24  Quenching Gas Effects
presence of the surrounding gases, however, different quenchers exhibit efficiencies when acting as a third body, X, in the iron atom nucleation loss process (see Figure 23). Although these data are conveniently displayed as following exponential decay, the atoms are likely decaying according to the integrated solution for a third order process, i.e., \( \frac{1}{[Fe]} \) vs time.

These experiments are continuing with our next efforts being directed toward calcium, whose production is limited to thermal techniques, but whose output is in a good wavelength region from propagation consideration.

2. Iron Pen acarbonyl, Fe(CO)₅

During the course of our investigation of the atomic iron laser discussed in the previous section, we observed stimulated emission from the organo-metallic precursor used in these experiments. A hint of this output can be seen in our initial experiments on atomic iron (see Figure 14) in the frame marked 'laser no discharge'. By optimizing the laser cell interaction volume and Fe(CO)₅ density, we were able to generate significant output at a number of wavelengths as shown in Figure 23. Using spectrally narrow filters, we were able to account for 60% of the output as being associated with the 384 nm transition. Most of the remaining photons are associated with the 562 nm line. The other observed transitions got stronger with repeated exposure of the same gas sample to a sequence of KrF laser pulses. It is likely they are due in some way to the Fe₂(CO)₉ formed by the action of UV light on Fe(CO)₅ by the following sequence. (17)

\[
\text{Fe(CO)}_5 + h\nu \rightarrow \text{Fe(CO)}_4 + \text{CO}
\]

Figure 25  Optically Pumped Iron Pentacarbonyl Lasing Transitions
Fe(CO)$_4$ + Fe(CO)$_5$ $^{(m)}$ Fe$_2$(CO)$_9$

We did however, see emission on those lines even on the first fill, but no special precautions were taken to isolate our storage containers from the action of the room lights, etc. and therefore there is likely to be Fe$_2$(CO)$_9$ "impurities" in the iron pentacarbonyl.

Since the output showed prompt temporal behavior (see Figure 26), it is likely to be lasing from an excited state of the Fe(CO)$_5$ or a photo-fragment produced in the initial photon absorption step, i.e.,

$$h\nu (\text{KrF}) + \text{Fe(CO)}_5 \rightarrow \text{FeCO}_5^{**}$$

$$\text{Fe(CO)}_5^{**} \rightarrow \text{Fe(CO)}_5^{*} + h\nu$$

$$\text{Fe(CO)}_5^{**} \rightarrow \text{Fe(CO)}_4^{**} + \text{CO}$$

$$\text{Fe(CO)}_4^{**} \rightarrow \text{Fe(CO)}_4^{*} + h\nu$$

We did see a trend with increased iron pentacarbonyl density (see Figure 27) and this behavior can be explained by recognizing that the extinction coefficient for Fe(CO)$_5$ absorption at 248 nm is $1.7 \times 10^{-17} \text{ cm}^2$, (18) so for 0.2 torr, the distance for 50% extinction is 2.7 cm. We observed maximum output when the KrF pulse was focussed through a gas path length of approximately 4 cm. Increased converted output would be expected by reducing this to very short distances but severe damage to the optical window occurred for these cases. Clearly, pumping the iron carbonyl end-on (see Figure 20) in this matter is not optimal. We are, therefore, planning on using a cylindrical lens to transversely pump the Fe(CO)$_5$ in a dye cell configuration and look for output perpendicular to the pump laser.

Figure 26  Experimental Data
Figure 27  Variation of Output with Fe(CO)$_5$ Density
These experiments on photodissociation of encapsulated metals open a new field of candidates which do not require atom production techniques. If further research indicates efficient conversion can be achieved, the overall project would be simplified with regard to the engineering aspects of a practical device. More work in this area is continuing with emphasis being placed on transverse pumping, conversion efficiency, and new candidates.
IV. SUMMARY

With the delivery and modification of an untuned high power KrF laser, experiments began on optical conversion to longer wavelengths with various candidates generated using scalable production techniques. For ease of demonstration of principle, the candidates hydrogen, iron and calcium were identified as compatible with the available power and bandwidth of our pump laser and were therefore first considered. We lased hydrogen in the manner described in Reference 4 to calibrate our experimental setup and procedure. The next day we observed single pass amplified spontaneous emission from atomic iron produced by the discharge dissociation of Fe(CO)$_5$. In addition, we observed lasing in the organo-metallic precursor (i.e. the iron pentacarbonyl) where the inversion is from a molecular excited electronic state of the parent or a fragment produced in the photodissociation process. Experiments to evaluate this new class of conversion candidates and to demonstrate conversion in atomic calcium are continuing.

These results demonstrate that efficient conversion of KrF towards longer wavelengths is achievable using an untuned laser as the pump. Also, the technique of producing refractory metals in densities sufficiently high for single pass amplified spontaneous emission by the decomposition of organometallic compounds is clearly established.

In addition to the stimulated Raman candidates, the theoretical effort has identified a number of parametric acceptor candidates for converting KrF to more propagating wavelengths. An experimental evaluation of some of these candidates will be undertaken shortly.
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