VISIBLE LASER RESEARCH
ONE-METER DEVICE

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VISIBLE LASER RESEARCH
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KrF, XeF, HgCl, Stimulated Raman Scattering, Blue-Green, Hydrogen

This final report summarizes research results obtained at Avco on a one-meter laser device. This apparatus was originally designed and constructed to be a useful scaling device for evaluation of the performance characteristics of the then recently discovered rare gas halide laser candidates. Subsequently, under this contract, a large number of technical achievements have been attained which have advanced the undere-
20. standing of various laser systems. These accomplishments have provided DARPA/ONR with very useful information for the evaluation of these lasers for a variety of mission applications.
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I. INTRODUCTION

Many contributions to the understanding of the physics and scaling limitations of rare gas halide and mercury halide lasers have been generated as part of this DARPA/ONR sponsored research program.

In carrying out this research, a sequence of experimental investigations was typically performed in the following order. New laser candidates were initially lased on a small-scaled e-beam device. These newly discovered lasers were then subsequently investigated on a small-scale, cable-driven e-beam under experimental conditions which were selected: to determine the formation processes by both e-beam and discharge pumping, to determine the quenching kinetics, to measure gain and absorption, and often to explore the effects of elevated temperature on the laser's performance characteristics. This kinetic data base was then used to develop and refine a laser code with predictive capability with regard to laser performance under power extraction. These code predictions were then tested by comparison to the laser's performance on the 1-m device that was constructed under this contract. This then allowed projections to be made for a variety of DARPA/ONR mission objectives.

Using this approach, scaling issues for a number of laser candidates were explored in detail. A complete description of the
results of these early investigations is available in prior Semi-Annual and Interim Technical Reports (Ref. 1). A brief summary highlighting these research results is listed below.

A. KrF LASER ($\lambda=249$ nm)

Krypton fluoride was the earliest rare gas halide laser that was subjected to intense investigation under this program. Small-scale experiments suggested it was efficient and volumetrically scalable. As part of this program, the following significant results were obtained:

- Important KrF* formation and quenching rates were measured.

- KrF* formation in KrF* laser mixtures was shown to occur predominantly from reactions involving KrF* as a precursor and consequently was saturable with cavity flux.

- Rate constants for three-body recombination reactions of the type

$$\text{KrF}^* + \text{Kr} + \text{Rg}' \rightarrow \text{Kr}_2\text{F}^* + \text{Rg}'$$

were calculated as a function of temperature.

- Use of these rates in a kinetics model enabled us to predict KrF* laser performance.

- Using e-beam pumping, a laser output energy of 102 J with an active laser volume of 8.5 $\lambda$ (12J/$\lambda$) at an intrinsic efficiency of 9% was obtained.

- Using e-beam controlled discharge excitation of KrF*, a laser output energy of 75 J was obtained in an active volume of 7.5 $\lambda$ (10 J/$\lambda$) at an intrinsic efficiency of 10%. The discharge enhancement ratio (discharge energy into the medium/e-beam energy absorbed in the medium) was 2.

The laser output, KrF*, and KrF*2 sidelight intensities were measured and shown to agree with model predictions.

KrF laser scaling projections were made.

KrF* and KrF*2 fluorescence was measured as a function of temperature. The results were consistent with the calculated temperature dependence of the three-body quenching of KrF* by Ar and Kr.

The KrF intrinsic laser efficiency was shown to improve with temperature.

B. XeF LASER (λ=350 nm)

Xenon fluoride subsequently became the leading rare gas halide laser candidate for continued DARPA funded research due to its longer operating wavelength, which makes it potentially useful for applications involving atmospheric propagation. The following results were obtained:

- Important XeF* formation and quenching rates were measured.
- An XeF* formation chain in neon rich mixtures was proposed and shown to be consistent with all experimental observations.
- XeF* formation was shown to proceed with unit branching in e-beam excited argon and neon rich mixtures.
- Using pure e-beam excitation, an XeF* laser output of 36 J was obtained in an active volume of 4.5 cm³ (8J/cm³) at an intrinsic efficiency of 2.6%.
- This laser efficiency was shown to be limited by the slow vibrational mixing in the upper level manifold and bottlenecking in the lower level.
- Elevated temperature was identified as a technique for improving XeF laser efficiency; subsequently its intrinsic efficiency was shown to increase by more than a factor of two at elevated temperature (~450°K).
- This increase in laser efficiency was shown to be a result of improved energy extraction brought about by an
increased rate of lower level removal and upper state vibrational mixing.

- The change in output spectra (from \( \lambda = 353 \) to \( \lambda = 351 \text{nm} \)) at elevated temperatures was shown to be attributable to the increase in lower level removal and a change in upper state rotational distribution.

- The energy loss due to the formation of the \( \text{XeF}^* (C) \) state and excited triatomics in \( \text{XeF} \) laser mixtures was shown to be saturable by laser cavity flux.

- The rates of lower state (\( \text{XeF}(X) \)) removal by vibrational mixing and dissociation were calculated for various vibrational states and temperatures.

- The various effects of increased temperature in \( \text{XeF} \) laser kinetics were incorporated into the comprehensive laser model.

- An \( \text{XeF} \) laser specific energy of 14 J/\( \ell \) was achieved at 3 amagats (4.7 J/\( \ell \)-Amg) with 5\% intrinsic efficiency.

- The observed laser performance was shown to be consistent with model predictions.

C. BLUE-GREEN LASER STUDIES

Research has also been conducted to explore efficient blue-green laser candidates as to their suitability for strategic laser communication applications. There have been two studies directly pursued under this program. The first involved HgC\( \ell \), whereas more recent research has been carried out exploring the technique of stimulated Raman scattering of \( \text{XeF} \) radiation in molecular hydrogen.

1. HgC\( \ell \) Laser (\( \lambda = 588 \text{ nm} \))

The initial laser demonstrations were carried out using direct e-beam excitation, where the efficiency was found to be low predominantly due to the low HgC\( \ell \)* production efficiency by direct
e-beam pumping. Under this program, alternative excitation techniques (e-beam discharge) were considered. The results obtained were as follows:

- E-beam controlled discharge experiments were carried out in Hg/Cl₂/Ar mixtures with no pre-reaction between mercury and chlorine.

- The measured small signal gain and fluorescence efficiency under optimum laser conditions suggested poor HgCl₂ formation from two of the three Hg* metastable sub-levels (i.e., 3P₀, 3P₁) as a result of "harpooning" reactions with Cl₂.

- Active medium photoabsorption of Hg/Cl₂/Ar mixtures near the laser wavelength was measured using a probe laser and found to be large (0.4 to 0.8%/cm) under typical operating conditions.

- A HgCl intrinsic laser efficiency of 3% was obtained with a discharge enhancement ratio of 11.

- The results of these HgCl₂* laser experiments were compared with model calculations with good agreement.

2. Raman Conversion of XeF

With regard to the stimulated Raman scattering experimental program, a brief list of the major accomplishments is included below.

- In small-scale experiments, 66% power conversion efficiencies of XeF pump radiation to first Stokes was observed.

- In large-scale experiments, using a 1-m laser as a pump, over 60% energy conversion of XeF to Raman output was observed.

A complete description of our Raman approach, the experimental techniques employed, and the results obtained under this contract are the main subject of this final report.
II. BLUE-GREEN CONVERSION OF XeF USING STIMULATED RAMAN SCATTERING TECHNIQUES

A. BACKGROUND

Stimulated Raman scattering (SRS) involves an inelastic scattering mechanism in which a photon incident on a atom or molecule causes the atom or molecule to undergo a change in internal energy and emit a photon that is shifted in wavelength from the incident photon by that exact change in internal energy. In the case of an atom, the internal energy is in the form of an electronic transition (see Figure 1). For a molecule, the change in internal energy can be vibrational or rotational.

The electronic Raman process can achieve large wavelength shifts in a single step but, specifically for this blue-green application, utilizing efficient, ultraviolet excimer lasers as pumps, the scattering atoms are high-temperature metal vapors (e.g., XeC\(_2\) shifted lead). These approaches thereby suffer from the technology problems associated with the production of the atom source. The molecular Raman process, on the other hand, utilizes a room temperature, chemically inert gas, but suffers from a small Stokes shift which is identical to the vibrational spacing. Since hydrogen has the largest vibrational energy
Figure 1  Optical Conversion by Stimulated Raman Scattering
spacing, it is a leading contender to shift to the blue-green in a minimum number of steps using the various ultraviolet excimer lasers available as possible pumps (see Figure 2).

From practical considerations (as to the number of shifts, the efficiency of the pump and the quantum yields involved), we identified XeF/H₂ as the preferred process. This Raman process involves a very nonresonant transition (see Figure 3) and has, therefore, a small Raman cross section (relative to the atomic case) and experiments are typically carried out at high pressure. In fact, pressure is one of the most significant parameters affecting the molecular SRS process. This is because the variation in pressure will affect both the number density of the scatterer and the spontaneous Raman linewidth. Since the gain is equal to the Raman cross section times the density times the length, i.e.,

\[ \text{GAIN} = \sigma \text{NL} \]

and the Raman cross section is inversely proportional to the Raman linewidth, a consideration of the pressure influence on the gain is important.

There are three distinct pressure regimes (Ref. 2) for the H₂ Raman linewidth (see Figure 4). At very low pressures, the linewidth is primarily determined by Doppler broadening and is independent of pressure; therefore the gain is directly proportional to the pressure. At pressures near 0.3 atm, the forward scattering Raman line width decreases due to collision narrowing effects, resulting from the frequent velocity-changing but not

HYDROGEN PROVIDES THE LARGEST MOLECULAR WAVELENGTH SHIFT (4155 cm⁻¹)

Figure 2 Blue-Green Wavelengths Accessible by Stimulating Raman Scattering (SRS) of XeF, XeCl and KrF with H₂
Figure 3 Potential Energy Curves Depicting Non-Resonant Features of XeF Pumped SRS Processes in Hydrogen
Figure 4 Pressure Effect of $H_2$ Raman Linewidth
phase-changing collisions. In this regime, the linewidth becomes inversely proportional to the density of the collision partner (i.e., $H_2$), and therefore, the gain scales as density squared. At still higher pressures ($\gtrsim 3$ atm), collision broadening effects increase the Raman linewidth and the Raman gain eventually becomes independent of pressure.

Studies of stimulated Raman scattering in hydrogen gas are numerous (Ref. 3) and a variety of pump lasers have been used in these previous experiments. In fact, stimulated Raman scattering in general has a rich literature associated with it and high conversion of pump photons to Raman output has been predicted theoretically, modeled extensively and observed experimentally (Ref. 4). Consider, for example, the model prediction in Figure 5. Here the pump laser was Ruby and the Raman scatterer was liquid CS$_2$ (Ref. 5). Through variations in the cell length, the


Figure 5  Model Predictions for Cell Length Variations and Their Influence on the Resulting Stokes Shifts for Ruby Laser Pumped CS$_2$
authors predicted and observed high conversion to arbitrary Stokes, essentially approaching unit photon conversion efficiency. In our experiments, which will be described below, we routinely converted ~60% of the energy of the pump to stimulated Raman output.

All of these literature sources and our own practical experience allowed us to confidently propose that xenon fluoride could be very efficiently shifted, using molecular hydrogen, to longer wavelengths. To insure that the desired Raman process dominated over other potential competing processes, we proposed that the shift to the blue-green from 350 nm be carried out in two distinct steps in two separate cells (see Figure 6). This was for two reasons: one involves wavelength selection by placing different scatterers in the two cells, e.g., H₂ and D₂. In this way, different wavelengths across the blue-green region can be produced. The second reason involves the necessity of controlling a significant competitive process to stimulated Raman scattering for pump photons: parametric or four-wave processes (Ref. 6). These processes can occur when waves separated by a vibrational frequency copropagate in the Raman medium (see Figure 7). Different four-wave combinations can occur as long as phase matching and energy conservation are maintained.

The phase matching requirement implies that in a dispersive medium, the four-waves may not be able to couple in the axial direction. Experimentally, what is observed is the appearance of light at various Stokes and anti-Stokes shifted wavelengths in cones around the primary beam. To reduce significant parametric processes

Figure 6  Two-step two-cell Raman conversion to the Blue/Green.
PUMP PHOTONS PRODUCE STOKES AND ANTI-STOKES PHOTONS

2 PUMP PHOTONS PRODUCE STOKES AND ANTI-STOKES PHOTONS

A STOKES AND PUMP PHOTON PRODUCE ANTI-STOKES AND 2\textsuperscript{ND} STOKES PHOTONS

Figure 7 Combination Schemes for Four-Wave Parametric Processes
from occurring, one can collinearly inject desired Stokes radiation with the pump, thereby reducing the likelihood of phase matching. In addition, since the number of potential parametric processes grows rapidly with the number of intense fields present, by restricting the conversion to a single Stokes shift in any one cell, the number of potential loss processes is reduced and control over the cell's output is enhanced.
B. EXPERIMENTAL RESULTS (U)

1. Short-Pulse Laser Experiments: Lumonics XeF Laser

From July 1979 to March 1980, we investigated efficient generation of Stokes shifted radiation using a commercial exciplex laser as the pump (Lumonics, TE-261-2) and molecular H<sub>2</sub> and D<sub>2</sub> as the Raman scattering media. General laser specifications as detailed by the manufacturer were found to be achievable with our particular device (e.g., it produced 75 mJ of energy per pulse with XeF<sup>*</sup> and 190 mJ for KrF<sup>*</sup> with the cavity optics provided). The beam quality associated with these achievable energies was insufficient to provide focused intensity-length products to achieve laser action in the Raman gas. We, therefore, altered the cavity optics to include Brewster windows and an unstable cavity configuration of the type described by Barker and Loree (Ref. 7). This cavity produced <20 mJ of XeF laser energy in a pulse (FWHM) of 6 ns with a beam waist 10 times the diffraction limited value. The power density achievable was measured at the focus by recording the energy transmitted through various diameter pinholes. The beam waist at the focus of a 50 cm lens was found to be near 2.5 x 10<sup>-2</sup> cm in diameter with an energy of 10 mJ. This, with the 6 ns pulselength, corresponds to an achievable power density of 3 x 10<sup>9</sup> W/cm<sup>2</sup> at the focus.

Efficient conversion however relies on an intensity-density-length product well in excess of threshold to achieve

optimum conversion of XeF pump radiation to first Stokes. To measure efficiency and to parameterize the Raman process, the experimental setup shown in Figure 8 was used. Here the output of the Lumonics was focused through a pinhole to:

1) characterize the input beam, and 2) eliminate that portion of the pump with poor beam quality. In this arrangement, a second lens refocuses the beam into the center of our high-pressure Raman cell.

The Raman cells were constructed from available stainless steel shock tube sections about 40.6 cm long with an inner diameter of 3.8 cm and a wall thickness near 1.3 cm. The polished Dynasil quartz windows were near 6 cm in diameter and ~2.5 cm thick. The windows were supported by O-rings on the high-pressure side and by gaskets on the low-pressure side. The internal volume of these cells was near 465 cm³.

Pulse energies in these experiments were measured with a Scientech Model 362 energy/power meter. Pulse shapes were measured by photodiodes (ITT, type S-5 or Hamamatsu, type S-4). The outputs from the photodiodes were recorded by a Tektronics Model 7844 dual beam oscilloscope equipped with a C-51 oscilloscope camera.

Spectra were analyzed with either a 3/4 m monochromator, 1 m spectorgraph, or an optical multichannel analyzer (Tracor Northern, Model TN-1710, equipped with a Diode Array Rapid Scan Spectrometer).
Figure 8  Schematic of Experimental Setup where L is the Lumonics Laser, a and b are 50 cm fL lens, C is the High-Pressure Cell, D is a Photodiode, F are Filters and O is a Spectral Monitoring or Energy Measuring Device
Using this arrangement and these various diagnostics, the effects of laser intensity, focal length, and H\textsubscript{2} pressure on the stimulated Raman scattering threshold, gain, Stokes production and conversion efficiency were investigated.

The pump laser intensity (at fixed focal length) was varied in two ways: by repeated firings to degrade the mix from optimum or by lowering the laser charging voltage. Using either technique, the subsequent results were identical, so the more convenient technique of varying the charging voltage was usually employed. Since these experiments were all carried out with focused geometry, a related factor to the intensity achieved (W/cm\textsuperscript{2}) is the length over which the intensity was high. A visual perspective of the various focal arrangements used is shown in Figure 9. These pictures were obtained by open shutter exposures of the focused laser light into a glass cell containing a dilute solution of Rhodamine 6 G. The focal length of \textasciitilde 135 cm geometry provided a beam waist of about 0.05 cm in diameter. Figure 10 shows a plot of the energy through this size pinhole versus distance from focus. The distance between half-power points (or the effective Rayleigh range) is about 8 cm. Since the active length of the high-pressure cell is over 40 cm, there was no problem anticipated with gain clipping and no effect on the Raman output was observed when a Raman cell of \textasciitilde 90 cm was used.
Figure 9  Focal Length Profiles
Figure 10  Energy Measured Through a 0.05 cm Pinhole vs Distance from Focus
For this experimental setup (at a maximum laser output corresponding to near $4.5 \times 10^8$ W/cm$^2$ for this relatively soft focal geometry), the dependence of forward scattered Stokes generation on H$_2$ pressure was measured (see Figure 11), using the optical multichannel analyzer (note that the intensity versus wavelength calibration had not been measured at this time). The first Stokes component appears to optimize at pressures above 15-20 atms (250 psi), whereas the second Stokes appears to peak near 25-30 atms and then decrease at higher pressures. These pressure scaling results are in agreement with the KrF/H$_2$ experiments of Loree (Ref. 8) and the gain saturation reported by Bloembergen (Ref. 9).

By now keeping the focal length and H$_2$ pressure fixed, variations in stokes output were measured as a function of XeF pump input, (see Figure 12). With data similar to these, it is possible to calculate the small-signal gain for this single-step process. From our measurements, we found threshold to be near 2.2 mJ. The pulse width was 6 ns and the length over which significant conversion occurred was near 8 cm with a beam waist at the center of the focal region of 0.05 cm. Since it is commonly regarded that a gain times length $>30$ is necessary to achieve lasing from noise (i.e., spontaneous Raman scattering), we can calculate the small-signal gain, $g$, as follows:


$H_2$ PRESSURE VARIATION (Xe F)

X-DEPLETED PUMP

□-1st STOKES

△-2nd STOKES

Figure 11 Various Laser Output Photons vs Total $H_2$ Pressure
Figure 12 Optimized XeF Conversion to First Stokes vs Input Energy
\[ g \frac{I_{\text{Th}}}{L} \approx 30 \]

where

\[ I_{\text{Th}} = \frac{2.2 \times 10^{-3} J}{6 \times 10^{-9} s \times \pi (0.025 \text{ cm})^2} = 1.9 \times 10^{8} \frac{\text{W}}{\text{cm}^2}, \]

and \( L = 8 \text{ cm} \)

\[ g = \frac{30}{I_{\text{Th}} \frac{L}{W}} = \frac{30}{1.9 \times 10^{8} \times 8} = 2 \times 10^{-8} \text{ cm/W} \]

This value is about a factor of 5 larger than the value deduced from wavelength scaling of that reported by Bloembergen (Ref. 9) at Ruby wavelengths (see Figure 13). Since the theoretical wavelength scaling for the SRS cross section is fairly straightforward, our results suggest that Bloembergen's reported value may be too low. However, a more definitive gain measurement at the relevant wavelength (\( \lambda \approx 350 \text{ nm} \)) is required to resolve this apparent discrepancy.

The pump laser produces output at both 3511 Å and 3532 Å (see Figure 14) with \( \approx 80\% \) at 3511 Å. The total temporal pulsewidth was measured to be 6 ns (FWHM). When these spectral lines were observed independently with a monochromator, their individual pulsewidths were still near 6 ns but the 353 nm line was delayed slightly from the 351 nm line. This observation is consistent with the observations of others of higher gain in the 351 nm band at room temperature (Ref. 10). Pulse shape measurements of the first Stokes pulses were initially performed with the experimental setup using the

Figure 13. First Stokes Gain for Forward SRS in H₂. Solid line is wavelength-scaled calculation normalized to Bloembergen's result with ruby; * indicates the value deduced from this work.
Figure 14 XeF Lasing Transitions
50 cm focus (see Figure 9) at pressures of 10 atm. Characteristic pulse shapes are shown in Figure 15. The top traces show the input laser pulse with the 351, 353 nm XeF lines unresolved. The bottom traces are of the corresponding 411, 414 nm first Stokes lines. The temporal widths of these Stokes-shifted lines are ~4 ns. Similar effects were observed for the softer focus. Measurements of the pulse shape of the second Stokes component from a single cell did not show any further pulse distortion, i.e., they had essentially the same 4 ns pulsewidth as the first Stokes.

All of these data and the above discussion allow us to summarize the results of our single pass, single cell, XeF* pumped H₂ conversion efficiencies for optimized experimental conditions. Conversion efficiencies were calculated by measuring the output energy from the Raman cell through a series of spectral filters, then adjusting the recorded energy for the known transmission of the filters and finally ratioing it to that measured through an empty cell. A photon conversion efficiency of 100% is usually not achievable in these nonresonant molecular Raman systems because as the first Stokes radiation approaches high intensities, it itself acts as a pump and is converted to higher order Stokes emission. Table 1 contains a summary of the typical output and conversions observed. The power conversion efficiency is the relevant value, since the energy efficiency should
Figure 15 Temporal Characteristics of First Stokes (H₂) Spectral Lines Using the Short Focal Geometry
<table>
<thead>
<tr>
<th>Pump</th>
<th>Focus</th>
<th>$E_{in}$ (mJ)</th>
<th>$E_{out}$ (mJ)</th>
<th>Energy Eff (%)</th>
<th>Photon Eff</th>
<th>Power Eff</th>
<th>Power Photon Eff</th>
</tr>
</thead>
<tbody>
<tr>
<td>XeF</td>
<td>137 cm</td>
<td>9.6</td>
<td>4.2 $S_1$</td>
<td>44</td>
<td>61</td>
<td>66</td>
<td>76</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>~0 $S_2$</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
approach the power efficiency for sufficiently long pulse-lengths. These reported efficiencies did vary somewhat from experiment to experiment, however, the values presented here were observed routinely. (The actual highest efficiency for conversion to $S_1$ with $H_2$ in a single experimental observation was 57% energy and 86% power conversion efficiency.)

Other stimulated Raman conversion efficiencies for $H_2$ reported in the literature include 20% photon efficiency into $S_1$ pumped by a 1.06$\mu$m laser by A. Grasiuk (Ref. 11), while Komine and Stappaerts (Ref. 12), report power photon conversion efficiencies of near 50% into either $S_1$ or $S_2$, when $H_2$ is pumped by a tripled Nd:YAG laser at 355 nm with an oscillator amplifier configuration. Loree, et al. (Ref. 8), have also reported up to 70% energy depletion into various Stokes orders for KrF pumped $H_2$. All of these experiments suggest efficient conversion to longer wavelengths via stimulated Raman scattering in a single pass is readily achievable.

The next topic that we addressed in these short pulse, low energy experiments was the second step in the two-step process; namely, investigation of the conversion properties of the emerging $S_1$ from cell one, subsequently acting as a pump, to produce its $S_1$ in cell two. (see Figure 16).

The first of these experiments consisted of two-step shifting through two cells filled with hydrogen, i.e.,

Figure 16  Schematic of Experimental Setup where $C_1$ and $C_2$ are the Two High-Pressure Cells and $P_1$, $P_2$ the Photodiodes for Monitoring the Inputs to each Cell Respectively
XeF \rightarrow H_2 \rightarrow H_2$. The first cell was optimized for maximum conversion of XeF* to first Stokes, $S_1$, at 411, 414 nm. These emerging laser photons were separated from the depleted pump via a dielectric coated mirror. The depleted XeF* pump was thus rejected, while the first Stokes beam from cell one was focused into the second cell, also optimized to convert into its own first Stokes at 496 and 500 nm.

Experimentally, the same conversion efficiencies observed through the first cell were not observed in the second step. This was essentially due to the limitations in $S_1$ energies (acting as the pump) which could be delivered into the second cell through the various optical components (mirror, lens, window, etc.). We were apparently just over threshold in pump laser intensity for the second-step SRS process.

Specifically, the XeF pump laser was focused into the first cell with a focal length of about 135 cm and the cell was maintained at pressures $>40$ atm to insure good conversion. The output light (depleted XeF* and $S_1$ radiation) resulting from this optimized configuration was collected and then refocused into a second cell. A dielectric coated mirror was usually employed, behind the first cell, to reflect $>99.9\%$ of the depleted pump yet partially transmit ($\sim50\%$) the first Stokes radiation. The waist diameter and conversion length of this $S_1$ radiation were
measured as before (see Figure 17). From this figure, the effective gain length is near 5 cm and the power density at the focus is estimated as follows:

0.05 cm pinhole

\[
\frac{1.5 \times 10^{-3} \ J}{4 \times 10^{-9} \ s} \times \frac{4}{\pi (0.05)^2 \ cm^2} = 1.9 \times 10^8 \ \frac{W}{cm^2}
\]

0.04 cm pinhole

\[
\frac{0.7 \times 10^{-3} \ J}{4 \times 10^{-9} \ s} \times \frac{4}{\pi (0.04)^2 \ cm^2} = 1.4 \times 10^8 \ \frac{W}{cm^2}
\]

This represents a lower limit for the actual power density of this first Stokes laser light in that \(\geq 50\%\) of the power was lost in transmission through the optical components. When these are taken into account, the measured power density of the first Stokes compares favorably with the XeF pump (\(4.5 \times 10^8 \ W/cm^2\)) and indicates that the focusability of the emerging Raman Stokes shifted light is comparable to the pump.

The variation of the conversion features with the pressure in the second cell was qualitatively similar to that measured in the single cell experiments. Figure 18 shows a plot of the output as measured with the OMA when the pressure in the second cell was varied. (These data are not directly comparable in that the sensitivity of the OMA is greater at 500 nm that at 400 nm.) The points at "zero pressure" indicate that the input consisted of both \(S_1\) and \(S_2\).
Figure 17  Energy Measured Through 0.05 or 0.04 cm Diameter Pinholes vs Distance from Focus for $S_1$ Radiation
Figure 18 Cell 2 (H₂, H₂) Pressure Variation Effect on S₁ Pump and the Generation of S₂
Threshold in the second cell was measured by monitoring the first Stokes input with a photodiode and the second Stokes output with another photodiode. The results are summarized in Figure 19. In these experiments, the first Stokes, acting as a pump in the second cell, is probably not far above threshold.

The intensity-length product for conversion of XeF*/H₂/H₂ can be calculated from these second-cell experiments. The threshold energy was near 1 mJ. The pulselength was 4 ns with a beam waist of 0.05 cm and a conversion length ~5 cm at the focus of the second cell. The intensity-length product is, therefore, near 6 x 10⁸ W/cm which is considerably less (even after wavelength scaling is accounted for in the calculation) than the value calculated from the single-cell experiments. This lowered (apparent) threshold is caused by the presence of a small amount of second Stokes radiation emitted from cell 1 and entering into cell 2. This small amount of second Stokes can act as the initial input for the conversion process and would require less amplification (i.e., a total gain significantly <30) for stimulated Raman scattering to occur.

The temporal features of the two-step conversion are depicted in Figure 20. This figure shows the XeF pump, the first Stokes output of cell 1 and the second Stokes output of cell 2, when both cells were filled with H₂.
Figure 19 Cell 2 Threshold for $S_2$ as a Function of $S_1$ Input Intensity
Figure 20  Pulse Temporal Narrowing in Successive Conversion
The pulsewidth goes from 6 ns for the XeF laser to 4 ns for \( S_1 \) to 2.5 ns (FWHM) for \( S_2 \), although the general pulse shapes remained the same.

The typical energy conversion efficiency measured in this manner was 25 to 30% into \( S_2 \) output from the \( S_1 \) input, i.e., 496 out/411 in. This corresponds to a photon conversion efficiency of 30 to 35%, a power efficiency due to pulse shortening of 40 to 48% and a photon power efficiency of 48 to 56%. The total power conversion efficiency from the XeF input into cell 1 into \( S_2 \) output from cell 2 was 29%. As discussed above, conversion efficiency characteristics of what was achieved in a single cell should be achievable in any subsequent cell for an overall conversion of \( \eta_1 \eta_2 \). Since we measured power conversion efficiencies of 66% in a single cell, overall conversion \( >40\% \) to the blue-green should be achievable, but were unobtainable here due to the low pump intensities available.

This conjecture was tested somewhat by conducting two-step experiments with KrF* as the pump. The KrF laser operates at higher output intensities and was capable of generating significant quantities of \( S_1 \) through \( D_2 \) (see Figure 21). In this case, the first cell contained 67 atm of \( D_2 \) which resulted in an output consisting mainly of depleted pump (249 nm) and \( S_1 \) at 268 nm. These laser transitions were collected and focused into the second cell.
Figure 21 Two-Step KrF/D$_2$/H$_2$ Raman Conversion
containing H₂. They each generated their own Stokes components in cell 2, but most of the output consisted of a few first Stokes orders (see Figure 21). Of particular interest is the fact that the 268 nm S₁ output of cell 1 was depleted over 70% in cell 2, with the majority of the output appearing in the sought after 308 nm line (which would be analogous to the production of 470 nm with XeF as the pump).

2. Long-Pulse Laser Experiments: 1-Meter E-beam Pumped Lasers

Included as part of this DARPA-supported program, we also investigated Raman scattering using the output of a 1 m laser as the pump for single pass, single cell spectral observations. To perform these experiments, two different e-beam laser devices were used. The first experiments, which were carried out in FY '79, involved the (10 x 10 x 100) cm³ device. It was run with conventional optics and provided ~5 J in 600 ns for the few experiments we tried. Under these conditions, the output was collected by a 6-inch Dynasil UV-1000 quartz plano-convex lens (50 cm f1) and focused into the center of our high-pressure H₂ cell (~10 atm). We observed output spectra corresponding to S₁, S₂ and AS₁ (see Figure 22). From color Polaroid open-shutter photographs, it appeared that the output consisted of principally parametric four-wave conversion processes as was evidenced by the observation of annular rings (see Ref. 6). Also, the intensity
LONG PULSE LENGTH XeF* SRS EXPERIMENT
(10 ATM H₂)

Figure 22  Spectra Giving Qualitative Indication of Relative
Intensities of XeF Pumped H₂ (AS₁, P, S₁, S₂)
correlations in the Stokes radiation \((AS_1 \succ S_2)\) supported this contention.

More recently, we have been performing stimulated Raman scattering experiments using another 1 m long, e-beam initiated XeF laser as a pump. The e-beam was a high energy (400 kV) Marx bank driven device (Systems, Science and Software, model 450) operated at current densities near 18 A/cm\(^2\). The electrons entered the laser cavity through a 100 x 4 cm, 1 mil Kapton foil, with two external magnets providing a guide field to confine the beam and give uniform deposition into the gas. The laser mix consisted of 0.1% NF\(_3\), 0.5% Xe and the balance Ne, at a total pressure of 60 psi. The active laser cavity was fitted with Dynasil UV-1000 quartz windows set at the Brewster angle. Several cavities were tried. Initial experiments, similar to those performed using the 10 cm aperture device, used a conventional flat-flat resonator; the resulting output was focused into the center of a small H\(_2\) Raman cell and significant conversion to the blue-green was observed (see Figure 23). Later experiments utilized a positive branch confocal unstable resonator with a magnification of 2.35. The two aluminum coated front surface cavity mirrors, with a nominal reflectivity of 92\% at 353 nm, were spaced 267 cm apart. The output coupler was an aluminum scraper mirror set at 45\°, which produced a square exit beam with a 4 x 4 cm outer...
Figure 23 Blue-Green Optical Conversion of XeF\(^*\) (1 Meter Device)
dimension and a centered obscuration (see Figure 24). Typical output energy was ~1 J in 0.4 µsec. Measurements of energy through a pinhole indicated focused (50 cm focal length plano-convex lens) intensities > 0.7 GW/cm². This output beam was of sufficient quality that it could be collected by various focal length lens and focused into the center of hydrogen Raman cells of varying overall lengths (the cell lengths were varied to reduce the fluence incident on the input windows). In this way, significant generation of a variety of Stokes shifted radiation could be produced (see Figure 25).

In these initial experiments, it was our objective to convert the output of this long pulse length XeF laser efficiently to stimulated Raman output. To perform these experiments and provide insight into the underlying physics operable under our conditions, the following experimental parameters were varied to explore the Raman process: focal length, pressure, and pump intensity (see Figure 26).

A change in the focal length, since intensity X length is a constant, is expected to provide a changing distribution of angles which could influence the phase matching conditions for four-wave processes. With regard to the desired Raman process, provided the depth of field at the focus is small compared to the cell length, there should be no impact.
Figure 24  Photograph of 1-m Xenon Fluoride Laser
Figure 26  Experimental Parameters Used to Explore Raman Process
Similarly, since at our operating pressures, the Raman gain is constant and independent of pressure, variations in pressure are expected to affect dispersion, although very insensitively due to the nonresonant wavelength regions involved. Again, the expectation is to vary the ability of various parametric processes to phase match and, therefore, alter the output observed.

Lastly, and most significantly, are variations in XeF pump intensity to observe threshold behavior of the various Stokes shifted radiation:

\[ I_s(L) = I_s(0) e^{gsIp^L} \]

The Raman gain is given by:

\[ gs = \frac{\lambda_s^2}{hv_s} \frac{N}{\pi \Delta v_R} \frac{\partial \sigma}{\partial \Omega} \]

where \( \lambda_s \) is the Stokes wavelength in the medium, \( N \) is the number density of molecules, \( \frac{\partial \sigma}{\partial \Omega} \) is the spontaneous Raman scattering cross section and \( \Delta v_R \) is the full width at half maximum of the Raman linewidth in \( H_2 \). A plot of first Stokes gain versus wavelength is provided in Figure 27.

For the case of stimulated Raman scattering building from the initial spontaneous noise, a total gain of \(-30\) is required. This suggests that sequential Raman processes, i.e., \( I_p \) providing \( S_1 \); \( I_{s_1} \) producing \( S_2 \), etc., involves a very large total gain length. Parametric generation of higher order Stokes radiation, not being a gain
Figure 27  First Stokes Gain for Forward SRS in $H_2$
process, however, can be quite easily initiated, e.g., see Figure 28. From the data in this figure, it is readily seen that at threshold levels for the generation of $S_1$ at 414 nm, $S_2$ at 500 nm and $S_3$ at 624 nm are simultaneously present. This is very strong evidence for the presence (at high conversion efficiency) of parametric processes. These data were taken at 26 atm and used a hard focal length experimental setup (1.35 m). The interpretation of these parametric processes and their continued interplay and influence on the observed stimulated Raman output were a central thrust of our experimental program.

To perform these experiments, several techniques to vary the focal length were used. One (depicted in Figure 29) was to utilize a focusing mirror; the other technique involved varying the focus by the relative position of two lenses, then turning the corner with a flat 100% reflector. Both techniques gave essentially the same Raman results, although the lens/flat arrangement minimized astigmatism. By passing the output of the Raman cell through a LiF prism, the beam was dispersed into spatially resolved individual beams so that each could be sampled (in separate experiments) by an energy meter. The results for a typical set of experiments are summarized in Table 2. Here data are presented for two pressures: 7 and 70 atm. The distribution of measured radiation as energy percent as well
Figure 28  SRS Threshold Data (1.35 m focus: 26 atm H₂)
Figure 29 Energy Conversion Measurement Experimental Setup
TABLE 2
CONVERSION EFFICIENCIES MEASURED FOR XeF* PUMPED H₂
(1-m Device Experiments)

<table>
<thead>
<tr>
<th></th>
<th>ENERGY (PHOTON) %</th>
<th></th>
<th>ENERGY (PHOTON) %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7 AMG</td>
<td></td>
<td>70 AMG</td>
</tr>
<tr>
<td>PUMP</td>
<td>51 (51)</td>
<td></td>
<td>54 (54)</td>
</tr>
<tr>
<td>S₁</td>
<td>26 (30)</td>
<td></td>
<td>31 (36)</td>
</tr>
<tr>
<td>S₂</td>
<td>14 (20)</td>
<td></td>
<td>6 (8)</td>
</tr>
<tr>
<td>S₃</td>
<td>~1 (1.6)</td>
<td></td>
<td>~1 (1.3)</td>
</tr>
<tr>
<td>AS₁</td>
<td>~2 (1.8)</td>
<td></td>
<td>~1 (1)</td>
</tr>
<tr>
<td>TOTAL</td>
<td>94% (~104%)</td>
<td></td>
<td>93% (~100%)</td>
</tr>
</tbody>
</table>
as photon conversion is shown. This type of energy bookkeeping allowed us to calibrate the optical multichannel analyzer (Tracor Northern) and use it as a secondary standard allowing for a much more rapid data collection and reduction rate.

Typical data for various focal lengths as a function of pressure are shown in Figures 30 through 33. The data from the 1.35 m focus and the 3.6 m focus show evidence for significant effects from parametric processes (e.g., $S_3$ magnitude). A check on the 3.6 m interpretation is found in Figure 34 where simultaneous thresholds are observed as before.

At the softer focus (4.6 m and 5.1 m), suppression and virtual elimination of $S_3$ was observed as well as enhanced conversion efficiency (see Figures 35 and 36). These data were recorded using spectrally filtered photodiodes which monitored the XeF input and output as well as $S_1$ and $S_2$ radiation. High depletion of the XeF pump (> 70%) was observed both by integrating the area under these traces as well as by using the OMA. This increase in pump depletion is consistent with a decrease in parametric processes which may, for example, regenerate the pump (e.g., $S_1 + S_1 + S_2 + P$).

The threshold behavior (Figure 37) at the softest focus investigated (5.1 m - essentially limited by room dimensions) shows distinct, different thresholds for
Figure 30  Significant $S_3$ Production at Hard Focus (1.35 m)
Plot of $S_n/S_1$ vs $H_2$ Pressure
Figure 31 Reduced $S_3$ Production at Softer Focus (3.6 m) Plot of $S_n/S_1$ vs $H_2$ Pressure
Figure 32 $S_1$ Dominates at Soft Focus (4.6 m) Plot of $S_n/S_1$ vs $H_2$ Pressure
Figure 33 Pressure Insensitive $S_1$ and $S_2$ Output at Softest Focus Studied (5.1 m), Plot of $S_n/S_1$ vs $H_2$ Pressure
Figure 34  SRS Threshold Behavior (3.6 m focus)
CONVERSION EFFICIENCY ~ 70%

Figure 35  Raman Temporal Signals Observed at Soft Focus (4.6 m) Showing Pump Depletion of 70%
Figure 36  Raman Temporal Signals Observed at Softest Focus Studied (5.1 m) Showing Pump Depletion of 72%
Figure 37  Different Thresholds Observed for $S_1$ and $S_2$ at 5.1 m Focal Arrangements
S1 and S2 and an indication that S1 might be decreasing at the highest pump energies studied (~1.3 J). To properly interpret these results, a thorough knowledge of the spatial properties of the pump beam is required; since the nature of the spatial beam profile impacts the Raman result (see Figure 38). Most modeling has considered only collimated beams with either "flat top" or Gaussian spatial profiles in the near field. Our unstable resonator is expected to produce a "near flat top" profile (depending on its beam quality) and focus to a pattern similar to that shown at the bottom of Figure 38. With such a beam, multiple Stokes output is expected due to sequential conversion in those portions of the beam at highest intensities, just threshold S1 production for the less intense portions, and no conversion at all for the smallest lobes.

To improve the conversion depicted by the data presented in Figure 37, we improved our cavity optics (including use of dielectric coated 100% reflectors) and alignment procedures; this allowed us to achieve higher W/cm² due to improved beam quality as manifested by a reduction in energy threshold and alteration in the S2 to S1 ratio (see Table 3). After laser beam quality improvements, increased S2 production was observed at similar overall conversion efficiency. These data suggest near complete conversion of the depleted XeF radiation to Raman output; further improvements in conversion efficiency is limited by the
Figure 38 Spatial Beam Profile Influence on Raman Output
TABLE 3
LASER BEAM QUALITY IMPROVEMENTS HAVE RESULTED IN ENHANCED $S_2$ PRODUCTION

<table>
<thead>
<tr>
<th>OLD</th>
<th>NEW</th>
</tr>
</thead>
<tbody>
<tr>
<td>~1.2 J</td>
<td>~1.2 J</td>
</tr>
<tr>
<td>~6 XDL</td>
<td>~2 XDL</td>
</tr>
<tr>
<td>≥70%</td>
<td>≥ 70%</td>
</tr>
<tr>
<td>~0.7</td>
<td>~1.35</td>
</tr>
<tr>
<td>$\frac{S_2}{S_1}$ RATIO</td>
<td></td>
</tr>
<tr>
<td>~25%</td>
<td>* 100</td>
</tr>
</tbody>
</table>

BLUE–GREEN OUT
$\frac{XeF \text{ IN}}{XeF \text{ OUT}}$ 35%

$\Rightarrow$ CORRESPONDS TO
0.4 J IN 0.4 μsec
beam quality of the device, and the fact that these experiments utilize a focused beam.

In summary, experiments on a 10 mJ, 6 ns device showed significant conversion (66%) to $S_1$ could be achieved using XeF as a pump. Scaling these results to a 1-2 J, 400 ns, 1 m scale size device was reasonably straightforward and showed the importance of eliminating parametric processes for Raman control which was achieved by going to very soft focal beam experiments (see Figure 39). Also, over 70% (energy) pump depletion was routinely observed. This degree of conversion is consistent with the nature of the pump spatial beam quality and the qualitative far field intensity distribution discussed earlier.
Figure 39  Spectrographic Evidence for Focal Length Influence on Minimizing Parametric Processes
SECTION II
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