CHEMICAL PRODUCTION OF EXCITED NF

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The a1Δ and b1Σ+ states of NF, NCl and NBr can be efficiently generated by reactions of F+HN3 followed by halogen atom reactions with the azide radical, N3. Scalability of these systems is sensitive to the buildup of N3 concentrations since N3+N3 reactions undermine the overall efficiency. Large concentrations of N3 are encountered when the halogen atom reaction with N3 is significantly slower than its formation step, F+HN3. Hence scaling of NF(a1Δ) is not feasible but that of NCl(b1Σ+) and NBr(b1Σ+) shows promise. A diagnostic tool sensitive...
to optical gains or absorptions of $<2 \times 10^{-6} \text{cm}^{-1}$ was developed. In the NCl flame, the medium was transparent everywhere to the diagnostic probe, indicating fast removal of NCl($b^1\Sigma^+$) by reactions with Cl and/or F atoms which are present in large excess. Strong absorptions were observed in the NBr flame in accord with an expected short radiative lifetime of the $b^1\Sigma^+$ state. Energy transfer from NF($a^1\Delta$) to atomic Bi generating the $6^2D_{3/2}$ metastable state proceeds at gas kinetic rates. Using a single frequency CW dye laser, the first high resolution spectrum Bi $5^4S_{3/2}-6^2D_{3/2}$ was measured, and the radiative lifetime of the $6^2D_{3/2}$ state determined. Although an NF-Bi gain experiment was not performed, the appropriate diagnostic system was demonstrated.
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I. INTRODUCTION

Since the initial demonstration by Clyne and White\(^1\) that electronically excited NF could be produced by a fast bimolecular reaction, nitrogen halide diatomics have drawn considerable interest as candidates for chemically pumped lasers. This interest stems in part from the electronic structure of these molecules, shown in Fig. 1, in the form of a potential energy diagram for NF. The nitrogen halides are isoelectronic to the group VIA diatomics, and hence have low lying \(a^1\Delta\) and \(b^1\Sigma^+\) excited electronic states. Transitions from these states to the \(X^3\Sigma^-\) ground state are nominally forbidden, and the radiative lifetimes of the excited states may be quite long, particularly for the lighter members of the group. Malins and Setser\(^2\) have reported the lifetime of NF\((a^1\Delta)\) to be \(\tau_{\text{rad}} = 5.6\) s. Hence, excited nitrogen halides may be considered as energy storage agents capable of transferring their energy to a desired lasing species, or, for the shorter lived states, as lasing molecules themselves. The \(a^1\Delta\) and \(b^1\Sigma^+\) states exhibit little or no Franck-Condon shift with respect to the ground state, and consequently their emission spectra, in the near IR and visible regions, respectively, are dominated by strong \(\Delta v = 0\) sequences with very narrow bandwidths.\(^3,4\)

Figure 1 also serves to demonstrate the major drawbacks to be overcome by any potential nitrogen-halide-based laser. Since there is no Franck-Condon shift and the major transitions terminate on levels which are thermally populated at room temperature, the pumping reaction must be
Figure 1. Potential energy diagram for the \( \chi^3\Sigma^- \), \( a^1\Delta \), and \( b^1\Sigma^+ \) states of NF molecule.
dynamically constrained to produce the excited states in order to generate a population inversion. Also, the $b^1\Sigma^+$ and $a^1\Delta$ excited states lie only $\sim 20$ and $\sim 40$ kcal/mole below the dissociation limit of the ground state (for NF), respectively, and therefore nitrogen halide molecules in these states are susceptible to reactive quenching processes in which the N-X ($X = \text{halogen}$) bond is broken. Any effective nitrogen halide laser system must overcome these two problems: it must be dynamically constrained to produce the excited states, and the gas mixture must not contain appreciable concentrations of reactive quenchers.

Herbelin and co-workers at the Aerospace Corporation have shown the chemical system studied by Clyne and White accommodated at least the first of these requirements. These authors have reported that the reaction of hydrogen on deuterium atoms with NF$_2$ radicals produces NF($a^1\Delta$) in very large proportion:

$$H + \text{NF}_2 \rightarrow \text{HF} + \text{NF}(a^1\Delta) \quad (1)$$

Herbelin's work has suggested that the branching ratio for formulation of NF($a^1\Delta$) in this process is essentially unity, and this conclusion has been recently corroborated by the experiments of Cheah and Clyne. The dynamic constraint operative in the system is thought to be spin conservation. It has been argued that the hydrogen atom attacks the nitrogen in NF$_2$ to form an excited singlet difluoramine intermediate, which must decompose to singlet HF and singlet NF. The Aerospace group has also shown that the $b^1\Sigma^+$ state of NF
can be produced by near-resonant energy pooling between NF(a¹Δ) and a variety of metastable energy transfer agents.⁵,⁷ These metastables include vibrationally excited HF, O₂(a¹Δ), and I*(5²P₁/₂). Sloan and co-workers⁸ have presented data which suggest that energy pooling with excited HF, a process involving Δν = 2 transitions in the HF, occurs at a rate comparable to rotational relaxation in this molecule (i.e., with nearly gas kinetic efficiency).

The principal drawback to the NF₂ system is that hydrogen atoms, necessary reagents for the production of NF(a¹Δ), are also efficient reactive quenchers of both the a¹Δ and b¹Σ⁺ states:

\[ \text{H} + \text{NF}(a¹Δ, b¹Σ⁺) \rightarrow \text{HF} + \text{N} \]  

Reaction (2) is believed to produce electronically excited N(2D) atoms.⁵,⁶ Since reaction (2) is quite rapid \( k_2 \approx 10^{12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \), it could severely limit the performance of any laser based on the NF₂ system.

To date, reactions analogous to process (1) have not been demonstrated for the production of excited NCl or NBr. Several years ago, however, Clark and Clyne⁹ reported the generation of NCl(1Σ⁺) and NBr(1Σ⁺) by the reactions of chlorine or bromine atoms with ClN₃. Emissions were observed from only the 1Σ⁺ states of the excited molecules. These authors postulated that the excited species were produced by reactions of Cl or Br atoms with gas phase N₃ radicals, themselves generated by abstraction of the chlorine atom from ClN₃.
Clyne's work formed the basis of the preliminary experiments performed at the Rockwell Science Center in 1978, in which the reaction of fluorine atoms with gas phase HN\textsubscript{3} was studied.\textsuperscript{10} These experiments were performed using a discharge-flow apparatus in which the reagents were mixed in a linear flow tube. Fluorine atoms were produced by microwave discharge of F\textsubscript{2}/Ar or CF\textsubscript{4}/Ar mixtures. Gas phase HN\textsubscript{3} was produced by metering a 75% aqueous solution of H\textsubscript{2}SO\textsubscript{4} onto a bed of NaN\textsubscript{3}; the gaseous HN\textsubscript{3} product was swept from the reactor by a stream of argon. A small portion of the HN\textsubscript{3}/Ar stream was bled into the vacuum system and admitted to the fluorine atom flow through a movable injector. The spectrum of the reaction flame was dominated by emissions from NF(b\textsuperscript{1}Σ\textsuperscript{+}) near 529 nm and NF(a\textsuperscript{1}Δ) near 874 nm (the Δν = 0 sequences of transitions to the ground state), and included weak HF vibrational overtone emission (from levels up to ν = 5) and nitrogen first positive (B\textsuperscript{3}Π\textsubscript{g} - A\textsuperscript{3}Σ\textsuperscript{+}\textsubscript{u}) emission. The NF(b\textsuperscript{1}Σ\textsuperscript{+}) emission was observed to scale quadratically with the NF(a\textsuperscript{1}Δ) intensity, which at low densities varied roughly linearly with the HN\textsubscript{3} flow rate. The experiments were performed with a large excess of fluorine atoms.

On the basis of these results, the following mechanism was postulated:

\begin{align*}
F + HN_3 &\rightarrow HF^* + N_3 \quad (3) \\
F + N_3 &\rightarrow NF(a^{1}\Delta) + N_2 \quad (4)
\end{align*}
\[ \text{NF}(a^1\Delta) + \text{HF}^*(v) \rightarrow \text{NF}(b^1\Sigma^+) + \text{HF}^*(v - 2) \] (5)

Addition of quenchers of vibrationally excited HF to the flow (e.g., CO₂), was found to dramatically reduce the NF(b^1\Sigma^+) intensity, while increasing the NF(a^1\Delta) intensity, as expected from the proposed mechanism. The limiting rate in the system was determined by monitoring the temporal profile of the NF(a^1\Delta) emission. For kinetics pseudo-first order in fluorine atoms, this rate constant was found to be \( k_{\text{lim}} = 2 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \). It was not known whether this rate constant applied to reaction (3) or reaction (4).

As noted above, reaction (4) must be strongly constrained to produce the a^1\Delta state of NF for the F + HN₃ chemical system to be a viable laser candidate. In fact, strong constraints are expected on the basis of spin conservation. To illustrate this point, Fig. 2 shows a diagram indicating the relative energy of initial and final states for the analogous Cl + N₃ reaction. The reaction is assumed to proceed through a real or virtual ClN₃-like intermediate (the Cl + N₃ case was chosen for this illustration since the heat of formation of ClN₃ has been measured). The Cl (2P) and N₃(2Π) reagents may combine on either singlet or triplet reaction coordinates. The singlet path lies lowest in energy; the energy of the intermediate has been shown in the figure as the ground state (1A') of the ClN₃ molecule. The singlet intermediate should correlate directly to NC₃(a^1\Delta) or NC₃(b^1\Sigma^+). Judging from the onset of the absorption spectrum of ClN₃, corresponding to transitions to the lowest lying singlet state, the lowest triplet states of this molecule should lie at energies greater than that of the Cl + N₃ reagents. Hence, an
Figure 2. Energetics of the Cl + N\(_3\) reaction. (The reaction should proceed via a ClN\(_3\)-like intermediate lying at an energy higher than that of the free ClN\(_3\) molecule, which is shown in the figure).
energy barrier should exist in the adiabatic triplet coordinate, effectively preventing direct formation of ground state \( ^3\Sigma^- \) NCl. Production of the ground state might still occur via crossing of the singlet and triplet coordinates in the product channel, however. It is expected that such processes will be improbable for the lighter NX molecules (smaller spin-orbit coupling), but might be significant for heavier species such as NBr or NI.

A preliminary test of this hypothesis was made by measuring the photon yield for emission from NF(a\( ^1\Delta \)). These experiments were performed by calibrating the absolute photon collection efficiency of the detection system of the apparatus, using the \( O + NO \) flame as a chemical actinometer. An estimate of the peak density of NF(a\( ^1\Delta \)) was determined from \( N = \alpha I/A \), where \( I \) is the measured intensity, \( A \) is the NF radiative lifetime, and \( \alpha \) is the calibration factor. These experiments suggested that the yield, defined as the NF(a\( ^1\Delta \)) density relative to the initial \( HN_3 \) density, was in excess of 40%. Although these results were only semi-quantitative, they supported our hypothesis concerning the role of spin constraints in the system.

Subsequent to these preliminary experiments, a contract for further research on nitrogen halide systems was awarded to the Rockwell Science Center by the Air Force Weapons Laboratory. This report describes the work performed and results obtained in this effort, which spanned the period from December 1978 through July 1981. The overall objectives of the program were as follows:
1. To determine the behavior of the F + HN₃ system upon scaling to densities commensurate with a small scale gain demonstration.

2. To assess similar chemical systems producing electronically excited NCl and NBr.

3. To demonstrate optical gain on those nitrogen halide systems which could be scaled to the appropriate densities.

The first segment of the program addressed scaling of the NF system. In view of the radiative lifetimes of the b¹Σ⁺ and a¹Δ states of this molecule, scaling to densities required for a gain of 10⁻⁴ cm⁻¹ required the development of completely new systems for the generation of fluorine atoms and HN₃. The second phase of the program involved small-scale measurements on azide flames producing excited NCl and NBr, using techniques similar to those employed in the preliminary experiments described above.¹⁰ The third phase of the program, gain experiments, involved the development of an apparatus with which gains or absorption as small as 10⁻⁴ per round trip transit through the medium could be measured. This system was based on measurement of the phase shift of a modulated probe beam transmitted through a resonant cavity surrounding the medium, and was similar to a system described by Herbelin and co-workers for the measurement of mirror reflectivities.¹⁴ In the final phase of the program, this gain diagnostic was used in preliminary measurements on the NF(a¹Δ) - Bi(6⁴S₃/₂) energy transfer system.
During the period of time spanned by this program, a related effort directed toward measurement of the fundamental properties of the nitrogen halide molecules and the kinetics and mechanisms of their formation was underway in our laboratory. Data obtained in the program were used extensively in interpreting the results described in this report.
II. CHEMICAL GENERATION OF EXCITED NF

As noted above, the primary objective of this first part of the program was to assess the ability of the $F + HN_3$ chemical system to scale to densities at which an optical gain of $10^{-4}$ cm$^{-1}$ would be produced on the $b^1\Sigma^+ + \chi^3\Sigma^-$ transition of NF. The first issue to consider is the gain levels to which our preliminary experiments$^{10}$ corresponded. The optical gain (or absorption) coefficient at the center of a single rotational line in the 0,0 band of the $b^1\Sigma^+ - \chi^3\Sigma^-$ transition is given by the following expression:

$$G = \frac{2}{\Delta \nu} \frac{\ln 2}{\pi} \cdot \frac{\lambda^2}{8\pi} \cdot A_{ij} \left( N_i - \frac{g_i}{g_j} N_j \right)$$

(6)

where $\Delta \nu$ is the linewidth of the transition, $\lambda$ is its wavelength, $A_{ij}$ is the rate of spontaneous emission from the upper (i) to the lower (j) state, $g_i$ and $g_j$ are the degeneracies, and $N_i$ and $N_j$ are number densities, respectively. For the conditions of our experiment, $\Delta \nu$ should be close to the Doppler linewidth for NF near 529 nm, 1.1 GHz at 350 K. Since each rotational level in the $b^1\Sigma^+$ state can radiate to five different levels in the ground $\chi^3\Sigma^-$ state (corresponding to the $^5\Sigma$, $^3\Pi$, $^1\Pi$, $^3\Sigma$, and $^1\Delta$ branches of this transition$^3$), the measured spontaneous emission rate for the $b^1\Sigma^+$ state,$^{15}$ $A = 44$ s$^{-1}$, must be divided by five to obtain the radiative rate for an individual transition. Since we presume the spin constraints favoring production of
excited state to be strong, we will assume the ground state density, \( N_j \), to be zero. Finally, assuming an equilibrium distribution among the rotational levels of the \( b^1\Sigma^+ \) state corresponding to \( T = 350K \), and furthermore that only the \( v' = 0 \) vibrational level is populated (nearly 90% of the emission found in the experiments was from \( v' = 0 \)), \( N_i = 10^{11} \text{ cm}^{-3} \). These conditions result in a gain given by \( G = 1 \times 10^{-7} \text{ cm}^{-1} \). Hence, a scale up of approximately three orders of magnitude is required for a gain of \( 10^{-4} \text{ cm}^{-1} \), corresponding to \( N_i = 10^{14} \text{ cm}^{-3} \), or a total \( NF(b^1\Sigma^+, v' = 0) \) density of \( \approx 10^{15} \text{ cm}^{-3} \). In order to approach these densities in a flow system, it was necessary to consider completely different sources of fluorine atoms and HN\(_3\) capable of producing torr-level partial pressures of these species in the flow.

1. **Generation of Gaseous HN\(_3\)**

   The principal constraint taken into account in the design of the HN\(_3\) generator was that at no time was the vapor to condense, nor was the HN\(_3\) partial pressure to exceed 100 torr. This constraint was applied strictly as a safety precaution, since condensed HN\(_3\) is known to be unpredictably explosive. The 100-torr limit placed on the pressure was to insure that condensation could occur nowhere in the system at room temperature. The vapor pressure of liquid HN\(_3\) is not well known; safe operation with gaseous HN\(_3\) pressures as high as 200 torr have been reported, however.*

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*A.P. Baranavski, private communication, Naval Research Laboratories, 1981.
Either of two basic concepts may be applied to the generation of large scale flows of HN₃. First, a large ballast tank might be filled with HN₃ to a pressure of ~ 100 torr; the tank would then be evacuated through the reactor during a run. Assuming that in the reactor the gases passed through a cross-sectional area of 5 cm² at a linear velocity of 5000 cm s⁻¹, a partial pressure of 1 torr of HN₃ would correspond to a flow rate of ~ 1 mole s⁻¹. At this flow rate, a ten-minute run would evacuate a tank with a volume of ~ 100 L. Clearly, this approach is intractable from the standpoint of both safety and ease of operation. A more feasible idea is the use of a device which continuously generates HN₃ for consumption in the reactor. Such a generator would simply be a scaled-up version of the H₂SO₄/NaN₃ reactor used in the preliminary experiments. A diagram of this smaller-scale generator is shown in Fig. 3. Solid NaN₃ is contained in a three-necked flask, as shown. A 75% aqueous solution of H₂SO₄ is added dropwise to the NaN₃ via a long section of 1-mm stainless steel tubing attached to a syringe. The tubing enters the flask through a rubber septum. The syringe is driven at a metered rate by a syringe pump (Sage Instruments). The HN₃ vapor produced by the reaction is swept from the reactor by a stream of argon. The reactor effluent is passed through a trap containing anhydrous CaSO₄ ("Drierite") to remove unwanted water vapor from the stream prior to being vented to a fume hood (the entire apparatus was positioned behind a blast shield inside the hood). For a typical chemiluminescence experiment, the H₂SO₄ flowed at ~ 10 drops per minute, resulting in an HN₃:Ar proportion in the effluent stream of ~ 1:10. Since the generator was operated at atmospheric pressure, the partial pressure
Figure 3. Small-scale HN₃ generation of gas phase HN₃.
of HN₃ in the vessel was ~ 80 torr. As an additional precaution against the formation of condensed HN₃, the reaction vessel was partially immersed in an oil bath heated to ~ 325K. The HN₃/Ar mixture was bled into the vacuum system through a metering valve for use in the chemiluminescence experiments.

This small-scale generator produced flows of HN₃ (in the reactor) on the order of 0.01 mmol s⁻¹. Given the extremely slow flow rates of H₂SO₄ and Ar used, it seems likely that it would be capable of generating perhaps four times this flow rate at its performance limit. Hence, an increase by a factor of ~ 25 was required for the scaling experiments. For stoichiometric reactions between H₂SO₄ and NaN₃, the consumption rates of these two species would be 1.6 cm³ min⁻¹ of concentrated H₂SO₄ (96%) and 3.9 g min⁻¹ of NaN₃. Tests with the small scale generator showed that H₂O dilution of the H₂SO₄ was necessary to produce an appreciable reaction rate with the solid NaN₃; either the concentrated H₂SO₄ was too viscous to adequately wet the solid, or some dissolution of the azide salt is required. Hence, the total H₂SO₄:H₂O = 3:1 flow rate in the scaled generator would be ~ 2.1 cm³ min⁻¹.

In scaling the small reactor, several additional constraints must be accounted for. First, the rate may be limited by wetting of the NaN₃, i.e., diffusion of the acid solution through the solid. Also, it would be undesirable to use a carrier gas in an atmospheric pressure generator for the large system, since the small proportion of HN₃ in the carrier would require a substantial pressure (~ 10 torr for 1 torr of HN₃) in the flow reactor. The apparatus constructed to accommodate these constraints and produce the large HN₃ flow rates required is shown in Fig. 4. The reaction chamber consisted of
Figure 4. Apparatus for the generation of large flows of gaseous scale HN₃.
a large Pyrex flask (≈ 2 l) which was rotated by a common rotary evaporator (Buchler Instruments). Rotation served to tumble the NaN₃, in order to produce a more uniform wetting pattern. The aqueous acid solution was contained in a reservoir shown to the left of the figure, and was metered into the reaction vessel via a linear array of six droppers extending from a common feed line. The acid flow was regulated by a small metering valve. The tumbler-dropper design was the result of small-scale tests using water and NaCl. These tests showed the wetting pattern of the tumbled salt to be good, and indicated that for vacuum conditions, the dropper design with the liquid flow regulated by a valve was necessary for adequate flow control.

The reactor effluent passed through a large trap containing "Drierite" (anhydrous CaSO₄) prior to entering the flow system. The pressure in the reactor was continuously monitored with a Validyne inductance manometer, as shown in the figure. In addition to the normal path to the flow tube, a secondary vacuum system was attached to the generator for regulation of the reactor pressure and disposal of the HN₃ vapor in the reactor after a run. This system consisted of a large toggle valve (operated remotely by a simple chain), a trap containing soda lime (to reactively remove the HN₃ from the flow), and a small pump vented up the hood. Initially, a 100-torr check valve was installed in parallel with the toggle valve to insure that the pressure did not exceed this value; the check valve was eventually removed, however, since it continually malfunctioned. In practice, the pressure was very easily regulated with the toggle valve.

A second reagent reservoir (see Fig. 4) contained an aqueous basic solution, typically NaOH. Addition of this solution to the system was used to
rapidly quench the reaction. The basic solution was admitted upstream of the metering valve such that large amounts might be added rapidly. In practice, this system was used only to stop the reaction at the end of a run. Both the acid and base solutions could be drained from the reservoir via a system which consisted simply of a three-way valve (which acted as an exit port) and a filter flask connected to the house vacuum system. The entire HN₃ reactor was constructed in the fume hood, and a Lucite blast shield erected in front of it. A photograph of the apparatus is shown in Fig. 5.

This generator design was quite successful in that it was found to generate copious quantities of gaseous HN₃. A reactor pressure of 100 torr was readily achieved, and could be easily regulated by variation of the acid flow and use of the toggle valve described above. Two systems were installed to monitor the flow rate of HN₃ produced by the reactor. The reactor effluent (from the Drierite trap) was admitted to the flow system via a large regulating valve. The monitoring systems were installed between the generator and this valve. First, a UV absorption system was used to measure the purity of the HN₃. The absorption was measured through a 1.0-cm-long cell, using the 253.7 nm line of a mercury lamp as a light source. The HN₃ extinction coefficient is 17.9 L mole⁻¹ cm⁻¹ at this wavelength, yielding an absorption of ~20% for a pressure of 100 torr. Absorptions corresponding to essentially pure HN₃ were observed for experiments at higher HN₃ flow rates. Experiments at lower flow rates (at the beginning of each run) indicated smaller HN₃ percentages, suggesting that some air was still being purged just after start-up. The second monitoring system measured the flow rate of HN₃ admitted to the flow reactor. Standard mass flowmeters could not be used in this case,
Figure 5. Photograph of the large-scale HN$_3$ generator.
since the back pressure was only 100 torr (close to the value of the limiting pressure drop across such a meter). Hence, HN₃ flow rates were monitored by measuring the pressure drop down a 5.0-cm length of 0.6-mm tubing using an MKS Baratron differential capacitance manometer. The pressure drop was calibrated against known flow rates of N₂ for a back pressure of 1.0 atm. The N₂ flow rates were calibrated against HN₃ (back pressure 100 torr) by measuring the time required to fill a fixed volume. Large-scale tests of the HN₃ generator showed that HN₃ flow rates well in excess of 0.3 mmol s⁻¹ could be achieved, yielding partial pressures in the flow tube greater than 300 mtorr at ~ 5000 cm s⁻¹. The experiments indicated that substantially larger flow rates could be achieved by increasing the H₂SO₄ flow.

2. Large-Scale Discharge-Flow Reactor

Fluorine atom flows of the magnitude required for the experiments might be generated by a precombustor, a kilowatt microwave discharge, or by a high current DC discharge through parent gases. Of these three, the DC discharge is the most readily applicable to a laboratory scale apparatus. The discharge unit used in the F + HN₃ experiments was originally part of a small CW HF laser based on a design by Hinchen¹⁷ and subsequent refinements by Spencer.² The device consisted of a Pyrex discharge tube surrounded by a water-cooled jacket (Lucite). Eight nickel rods arranged in an annulus about the centerline of the tube formed the cathode of the discharge. The anode was simply the other stainless steel endplate of the tube. The power supply used

²D.J. Spencer, private communication, Aerospace Corp., 1975.
in the experiments was a Universal Voltronics 20-kV, 600-mA power unit. In
previous experiments with this system, fluorine atoms were produced by passing
the discharge through SF₆-He mixtures. The device produced an optimum concen-
tration of ~2 torr of atoms for flow rates of SF₆ = 4 mmol s⁻¹ and He = 7.5
mmol s⁻¹ at total pressure of 6.5 torr and a linear velocity of 6000 cm
s⁻¹.¹⁹

The DC discharge tube was connected to the flow reactor via a water
coupled section of copper tubing, 2.5 cm in diameter and 20 cm long. This
section acted as the anode of the discharge. It was positioned so as to
laterally offset the discharge tube from the flow reactor by ~9 cm. This
offset was necessary to isolate light generated in the discharge from the flow
tube. The entire discharge tube was mounted on an optical rail inside an
opaque Lucite box.

A simple linear design was adopted for the initial flow reactor; it
was a 22-mm-i.d. section of quartz tubing 1.0 m in length. The tube was
cooled by passing water through an outer jacket, also of quartz. The inner
surface was coated with halocarbon wax to protect against etching by the
fluorine atom stream. The tube was attached to both the discharge unit and
the vacuum system by Ultra-Torr "O"-ring fittings. Figure 6 shows a detail of
the connection to the anode of the discharge. For the initial experiments,
HN₃ was injected into the stream of fluorine atoms via a stainless steel ring
fabricated from 1.6-mm tubing. The ring was perforated with eight 0.5-mm
holes, positioned to inject the HN₃ upstream into the atom flow. The
regulating valve for control of the flow is shown just above the injector.
The injector assembly was electronically isolated from the discharge anode by
Figure 6. Detail of the linear flow tube showing the anode of the dc discharge, the ring injector for admission of HN3 to the flow, and the water-cooled flow tube.
a 6.0-mm-thick Teflon disc. This configuration effectively prevented arcing of the discharge to the injector.

The flow reactor was continuously evacuated by a Kinney 500-ft³/min pump located under the laboratory building. A commercial scrubber capable of removing fluorine from the gas stream was installed on the exit port of the pump to prevent venting of the raw reactor effluent to the atmosphere. The gas pressure in the flow tube was measured with an MKS Baratron capacitance manometer. The flow rates of reagent gases other than HN₃ (SF₆ or F₂, Cl₂, He, N₂, and NO) were measured using a Tylan mass flowmeter. These flowmeters were calibrated for N₂ flows. Actual flow rates for the gases used were obtained using conversion factors supplied by the manufacturers. Measurements of the pressure rise caused by known flow rates of N₂ showed that the linear velocity in the 22-mm-i.d. flow tube was nearly 8000 cm s⁻¹ for a pressure of 10 torr. Minor throttling of the pump reduced the velocity to 5000 cm s⁻¹. A photograph of the DC discharge and flow tube assembly is shown in Fig. 7.

Initial tests of the discharge unit were made using SF₆/He mixtures. The discharge was very unstable for these mixtures, and rather small concentrations of fluorine atoms were produced, as evidenced by the addition of Cl₂ to the stream via the injector. In this case, F atoms react rapidly with Cl₂ to produce Cl atoms, and recombination of the chlorine atoms results in a characteristic red emission:

\[ F + Cl₂ \rightarrow ClF + Cl \]  

(7)
Figure 7. Photograph of the discharge-flow apparatus.
Ganguli and Kaufman\textsuperscript{18} have shown these reactions to be an effective chemiluminescent titration for fluorine atoms, and this method was used in our preliminary experiments on the F + HN\textsubscript{3} system\textsuperscript{10}. Molecular fluorine was found to be far superior to SF\textsubscript{6} as a parent species for fluorine atoms. Addition of Cl\textsubscript{2} to a stream of discharged He/F\textsubscript{2} resulted in a very bright Cl\textsubscript{2} afterglow whose intensity could be easily monitored with a small monochromator (Oriel 0.16 m) and an uncooled RCA 1P28, photomultiplier tube. A typical titration curve (Cl\textsubscript{2} flow rate vs intensity) is shown in Fig. 8. The data shown yield an endpoint corresponding to a fluorine atom flow rate of 0.40 mmoles s\textsuperscript{-1}, or a partial pressure of about 450 mtorr for the conditions of the run. Comparison to the F\textsubscript{2} flow rate indicated that \textasciitilde~35\% of the F\textsubscript{2} molecules were dissociated. Similar tests were performed for other F\textsubscript{2} flow rates and the results are shown in Fig. 9. The data indicated that the fractional dissociation remained approximately constant as the F\textsubscript{2} flow rate was increased. An F\textsubscript{2} flow rate of \textasciitilde~1 mmole s\textsuperscript{-1} produced nearly 1 torr of fluorine atoms. Further titrations were performed just after the apparatus had been changed to incorporate a larger injector (see Fig. 9). These resulted in a smaller fractional dissociation (\textasciitilde~25\%, curve B in the figure), presumably due to heterogeneous recombination of the atoms on the metal surface.
Figure 8. Results of a typical Cl₂ titration for the determination of the fluorine atom flow rate. (At the endpoint, the flow rate of Cl₂ is equal to the flow rate of fluorine atoms.)
Figure 9. Variation of the fluorine atom flow rate with the $F_2$ flow rate in the discharge-flow apparatus. (Curve A represents data taken with the small ring injector in place; curve B shows data obtained with the larger tube injector in the apparatus.)
Large-Scale Tests of the $F + HN_3$ and $F + DN_3$ Reactions

The first large scale tests were performed by adding $HN_3$ to the fluorine atom stream via the small ring injector described above. Additions of $HN_3$ to give a pressure rise of up to 250 mtorr above the ~ 5.0 torr pressure of the $F/F_2/He$ stream resulted in a brilliant yellow-green flame about 10 cm in length. About 10 seconds after the introduction of the $HN_3$, the emission rapidly collapsed to a short orange flame. We believe this behavior to be caused by thermal decomposition of the $HN_3$ inside the injector prior to reaction. The ring injector was itself engulfed in the flame and acted as the principal pressure drop in the system. It seems likely that the decomposition was caused by the relatively high pressure inside the injector, coupled with the temperature rise produced by the flame. This problem was completely eliminated by replacing the ring injector with an open-ended 5-mm-diameter tube pointed down the flow direction. Under these conditions, the regulating valve acted as the principal pressure drop and no decomposition flame was produced. Preliminary measurements of the $NF(b^1\Sigma^+)$ and $NF(a^1\Delta)$ emissions in the flame, near 529 nm and 874 nm, respectively, were made with uncooled RCA 1P28 and RCA 4832 (GaAs) photomultiplier tubes, respectively. Narrow band interference filters (bandwidth ~ 5 nm) were used to isolate the NF emissions in each case. For F atom partial pressures on the order of 0.5 torr, the data in the low $HN_3$ flow rate regime ($< 0.1$ mole s$^{-1}$) reproduced the small-scale flow tube results, i.e., the $b^1\Sigma^+$ emission rose to a faster than linear rate with increasing $HN_3$, and the $a^1\Delta$ emission rose roughly linearly, falling off at the higher flow rates. We note, however, that because of the
finite bandwidth of the interference filter, the near IR signal recorded may include emission from excited $N_2$ ($B^3\Pi_g$) (the $v', v'' = 2,1$ band of the first positive emission) in addition to NF(a$^1\Delta$). Neither the visible nor near-IR signals appeared to peak and decay before all the F atoms were consumed (as indicated by titration with Cl$_2$ downstream).

Further experiments with the discharge-flow system were performed by using DN$_3$ instead of HN$_3$, so that quenching of NF(a$^1\Delta$) by vibrationally excited HF (to produce NF(b$^1\Sigma^+$)) might be avoided. DN$_3$ was produced, using D$_2$SO$_4$/D$_2$O mixtures in the azide generator described above. Prior to the actual generation of DN$_3$, the generator and flow lines were passivated with D$_2$O for a period of ~24 hours. Infrared analysis of the generator effluent under operating conditions showed the azide product consisted of > 90% DN$_3$.

NF(a$^1\Delta$) emission produced by the F + DN$_3$ reaction was monitored with the 9-pin GaAs photomultiplier tube noted above. A narrow band interference filter centered at 875.9 nm was positioned in front of the PMT, and an aperture of 2.5 mm limited the light collection. Spectra of the F + DN$_3$ emissions were recorded both with and without the filter, using a 0.5-m monochromator (McKee-Pederson) and a cooled GaAs PMT. The spectra showed that the filter in fact transmitted only the NF(a$^1\Delta$) emission near 874.2 nm and effectively blocked the nearby $N_2$ first positive emission. The light collection efficiency of the detection apparatus (uncooled GaAs PMT) was calibrated by using the O + NO reaction as a chemical actinometer. In this technique, a known concentration of O atoms is produced by adding NO (through the injector used for HN$_3$) to a stream of N atoms (produced by passing $N_2$ through the
discharge) until an endpoint is reached at the extinction of the afterglow. At the endpoint, the O atom flow rate is equal to the NO flow rate. Further addition of NO to the system produced NO\(_2\) emission from the O + NO reaction, for which the absolute photon yield has been measured.\(^{13}\) Measurements using this technique yield a parameter \(\alpha\) which, when multiplied by the measured intensity, gives the flow rate of photons produced by the reaction. Since the NF(a\(^1\Delta\)) intensity was measured at a given point in the flow tube, the photon flow rate can be equated to NA, where \(N\) is the NF(a\(^1\Delta\)) number density in the emitting volume and \(A\) is the spontaneous emission rate (0.18 s\(^{-1}\)).\(^2\) Hence,

\[
[NF(a^1\Delta)] = \alpha I / A \tag{10}
\]

The value of \(\alpha\) determined in the calibrations was quite reproducible, in the sense that calibrations were made at several points down the length of the linear flow tube, and each measurement gave the same value of \(\alpha\) within the accuracy of the input parameters.

The intensity of the NF(a\(^1\Delta\)) emission produced by the reaction rose rapidly with the initial addition of DN\(_3\) to the stream of fluorine atoms, gradually rolling off to a peak value for DN\(_3\) flow rates in excess of the F atom flow rate. The time decay of the excited NF in the direction of the flow was not rapid, the signal falling an order of magnitude over a distance of \(\sim 15\) cm (\(\tau_d \sim 2\) ms). The yields (relative to the DN\(_3\) flow rate) measured at maximum intensity were extremely small, on the order of 5\%. A number of observations suggested that this result was caused by slow mixing in the flow
reactor. First, visual observation of the flame (i.e., the green \( \text{NF}(b^1\Sigma^+) \) emission produced by trace \( \text{HN}_3 \)) indicated a sheath of high intensity surrounded by a low intensity core, presumably containing unreacted \( \text{DN}_3 \). Increasing the \( \text{DN}_3 \) flow rate lengthened the flame, but did not affect the peak \( \text{NF}(a^1\Delta) \) intensity. Also, observation of the \( \text{N} + \text{NO} \) titration reaction \((k = 10^{-11} \text{ cm}^3\text{ molecule}^{-1}\text{ s}^{-1})\) at the high flow rates used for the \( \text{F} + \text{DN}_3 \) system yielded endpoints which were measurable only at distances greater than 25 cm downstream of the injector. Use of a perforated sphere mixing element on the \( \text{DN}_3 \) injector (the open-ended 5-mm tube noted above) did not improve the results, and eventually it became very hot, leading to \( \text{DN}_3 \) decomposition as noted with the ring injector.

Hence, mixing of the reagents appeared to be a fundamental problem, in that mechanical mixing (as by the perforated sphere) required a pressure drop across the mixing element, resulting in catastrophic \( \text{DN}_3 \) decomposition as the uncooled element was heated by the reaction. Since the pressure in the system was necessarily on the order of 5 torr (the production of large concentrations of \( \text{F} \) atoms by the discharge required a large diluent flow), some form of mechanical mixing was necessary. Hence, at this point it was decided to replace the linear flow tube by a Hinchen-Spencer type water-cooled mixing head.\(^{17}\) This head is basically a transverse flow device in which the fluorine atom/diluent flow from the discharge is passed through a cooled channel 10 m wide and 3 mm high. Reagents were injected into the stream by a

\(^{17}\)D.J. Spencer, private communication, Aerospace Corp., 1975.
linear array of holes, 30 on each of the top and bottom surfaces of the channel. For our experiments, the ends of the channel were closed by CaF$_2$ windows 5.0 cm in length.

A diagram of the transverse flow arrangement is shown in Fig. 10. The apparatus is quite similar to that described previously, except that the flow tube was replaced by the mixing head. Titrations of the fluorine atom flow in the reaction zone were performed by addition of Cl$_2$ as in previous experiments. These tests indicated that the discharge through F$_2$/He mixtures produced a maximum of 250 mtorr of fluorine atoms; discharges through SF$_6$/He produced a maximum of 100 mtorr of atoms. One advantage of the transverse flow arrangement was that the concentration of DN$_3$ in the reaction zone could be determined directly by UV absorption measurements near 205 nm. The light source used was a D$_2$ lamp whose output was folded through the reactor (see Fig. 10) to give an absorption pathlength of 20.3 cm. DN$_3$ present at 100 mtorr yielded an absorption of $-10\%$ for those conditions. These measurements resulted in a much improved measure of the true DN$_3$ flow rate. The DN$_3$ absorption was observed to be completely removed by the stream of F atoms from the discharge, indicating good mixing in the region probed by the experiments.

NF(a$^1\Delta$) emissions produced in the reactor were observed by using the uncooled GaAs PMT as before. The light collection efficiency of the system was again calibrated, using the O + NO reaction, and the initial N + NO titration yielded a good endpoint in the observation zone, a further indication of good mixing. The time decay of the NF(a$^1\Delta$) produced in the reactor was monitored by movement of the detector down the 5.0-cm observation window in the direction of flow. The experiments again indicated a relatively
Figure 10. Schematic diagram of the 10-cm transverse flow apparatus, showing the optical train for direct detection of DN3.
slow time decay, although faster than in the linear flow tube. For peak F and DN3 concentrations, the excited NF(a1Δ) decayed in ~ 4 cm. Absolute concentrations of NF(a1Δ) were measured vs the DN3 flow rate by recording the calibrated PMT response and the calibrated Baratron flowmeter reading with an x-y plotter. Typical results are shown in Fig. 11. The data shown were measured, using SF6 as a source of fluorine atoms, such that the partial pressure of F atoms was ~ 100 mtorr. The temperature in the water cooled system was measured, using a Pt/10% Rh-Pt thermocouple, and found to be only a few degrees above ambient. The data show that the NF(a1Δ) concentration rises very rapidly with the addition of DN3 to the system, reaching a flat peak at ~ 3.9 x 10^{14} \text{ cm}^{-3} (~12 \text{ mtorr partial pressure}) for a DN3 concentration several times that of the fluorine atoms. Clearly, the NF(a1Δ) yield relative to the DN3 flow rate declines rapidly with increasing DN3 density. We note, however, that over most of the density regime indicated in the figure, fluorine atoms were the limiting reagent. Since, according to our postulated mechanism (reactions (3) and (4)), two F atoms are required to produce one NF(a1Δ) molecule, the limiting density in this and similar experiments was about 1.6 x 10^{15} \text{ cm}^{-3} (~50 \text{ mtorr partial pressure}). Relative to this value, the peak NF(a1Δ) yield indicated by Fig. 11 is ~ 24%. The fact that the yield at the F atom-DN3 stoichiometry point was only one-third this value indicates that something was awry in the experiment, however, or that the simple model of Eqs. 3 and 4 does not hold at high reagent densities. If in fact the fluorine atoms were acting as a limiting reagent, the NF(a1Δ) density would level off or decline for DN3 densities beyond the stoichiometry point.
Figure 11. The absolute density of NF\(^{(a^1\Sigma)}\) produced by the F+DN\(_3\) reaction in the transverse flow apparatus vs the DN\(_3\) flow rate. (The initial partial pressure of fluorine atoms was 100 mtorr.)
The spectrum of the emission produced in the transverse flow head in the vicinity of 874 nm was recorded with a 0.5-m monochromator and a cooled GaAs PMT. The PMT response was measured with a PAR photon counting apparatus, and the results are shown in Fig. 12. The $\text{NF}(a^1\Delta) + X^3\Sigma^-$ transition is still the dominant feature, but $N_2$ first positive emissions have become quite strong. The spectrum exhibits no evidence of $\text{HF} \; v = 3 + v = 0$ overtone emission in this region.

In order to more thoroughly investigate the discrepancy between the high density data and the simple model supported by our preliminary low density experiments, it was decided that experiments directed toward measuring the $\text{NF}(a^1\Delta)$ yield vs $D_N$ flow rate would be performed for low reagent densities. For this purpose, our original flow tube apparatus\textsuperscript{10} was reassembled.

4. Low Density Yield Experiments

Low density yield experiments on the $F + \text{HN}_3$ reaction were performed with a cylindrical Teflon flow tube which has been thoroughly described in the literature.\textsuperscript{10} In brief, the tube has an internal diameter of 3.1 cm. Atoms are created by passage of parent gas/diluent mixtures (in this case $\text{F}_2/\text{Ar}$) through a 100 W microwave discharge positioned ~50 cm upstream of the fixed observation port. $\text{HN}_3$ entered the system through a movable slide tube internal to the Teflon flow tube; the outer surface of this tube was Teflon coated to prevent excessive atom recombination. Movement of the slide allowed time resolution of the emissions observed at the view port.
Figure 12. Spectrum of emission in the vicinity of 870 nm produced by the the F+DN₃ reaction in the transverse flow apparatus. (The spectrum shows contributions from both excited N₂(Γ₉g) and NF(a¹A).)
The fluorine atom densities produced by the microwave discharge were again calibrated using the \( \text{Cl}_2 \) titration. These tests indicated that nearly 100 mtorr of F atoms could be produced for large \( \text{F}_2 \) flows. In the first experiment in which the apparatus was used, HN\(_3\) was obtained from the large-scale generator, with the goal of probing the reaction at initial HN\(_3\) pressures of 10 to 50 mtorr and total pressures on the order of 1.0 torr. After the very first such experiment, however, an explosion occurred in the laboratory during clean-up of the HN\(_3\) generator. The explosion occurred approximately one hour after the run, and was apparently caused by condensed HN\(_3\) in the reaction vessel. At this point, experiments on the F + HN\(_3\) system ceased, pending a satisfactory re-design of the generator to insure complete removal of the HN\(_3\). Several weeks were spent on designing and testing a removal system based on reaction of HN\(_3\) with aqueous HNO\(_2\). The products of this reaction are \( \text{H}_2\text{O} \) and the innocuous gases \( \text{N}_2 \) and \( \text{N}_2\text{O} \). Although preliminary tests indicated that such a system was feasible, it was apparent that assembly of a generator incorporating this system would be a sizable engineering effort well outside the scope of the program. Hence, all further experiments were performed with the small-scale atmospheric pressure generator described above and shown in Fig. 3. This generator was capable of producing HN\(_3\) pressures in the flow tube as large as 20 mtorr.

Emissions produced within the flow tube were observed through the fixed observation port, which was a 2.5-cm-diameter \( \text{CaF}_2 \) window sealed to the tube via a Viton "O" ring. The emission was dispersed by a 0.5-m monochromator and detected by the cooled GaAs PMT and photon counting system. The light
collection efficiency of the system was calibrated, using the O + NO reaction as described above. The calibration procedure yielded a value of $\alpha$ (see Eq. (10) above) in fair agreement with previous experiments using this apparatus. Photon yields for the production of NF($a^1\Delta$) were measured as a function of several experimental variables. A number of techniques were used in these experiments which should make the data more reliable than that obtained in our preliminary measurements. First, a mass flowmeter was used to measure the Ar/HN$_3$ flow rate, resulting in more precise knowledge of the HN$_3$ concentration in the system. Perhaps a more significant change was the measurement of initial NF($a^1\Delta$) densities by extrapolation of the time decay of the emission signal measured over many ms to $t = 0$. Since the emissions from excited HF and N$_2$ decay rapidly in time, this technique effectively avoids the problem of spectral overlap from these species. Furthermore, the effect of additives on the NF($a^1\Delta$) intensity could be observed with greater clarity.

Figure 13 shows the exponential time decay of the NF($a^1\Delta$) emission observed for several experimental conditions. The lowest intensity data (closed circles) indicate an NF($a^1\Delta$) decay rate $\lambda_d = 243$ s$^{-1}$. The next higher curve (open circles) represents data taken with $\sim 223$ mtorr of CO$_2$ added to the system. It is clear that the addition of CO$_2$ significantly enhances the NF($a^1\Delta$) intensity (by more than a factor of 2 at $t = 0$), while not appreciably affecting the decay rate. Addition of still more CO$_2$ had little effect.
Figure 13. Time decay of $\nu_\text{F}(a-\Delta)$ emission produced in a low density flame.
Increasing the HN₃ flow rate while maintaining a moderate CO₂ pressure increased the NF(a¹Δ) intensity, but had only a slight effect on the decay rate (closed squares). Based on data such as this, it was originally thought that CO₂ enhanced the NF(a¹Δ) intensity by quenching vibrationally hot HF, which is itself a quencher of NF(a¹Δ). Were this hypothesis correct, the dramatic effect observed would indicate that in the absence of CO₂ a significant fraction of the NF(a¹Δ) is converted to the b¹Σ⁺ state, as per Eq. 5 above. A similar effect on the NF(a¹Δ) intensity was observed, however, when SF₆ was added to the mixture. Hence, it seems more likely that the role of these additives is to stabilize an intermediate (N₃) and/or vibrationally relax the excited NF.

Photon yields for the production of NF(a¹Δ) by the F + HN₃ reaction were measured by extrapolating intensities measured downstream to t = 0, as discussed above, for flows including a moderate amount of CO₂. The data were obtained with a fixed monochromator slit width of 750 mm (spectral bandpass ~ 2.5 nm). Measurements were made for several HN₃ densities at two different fluorine atom flow rates. The results of these experiments are shown in Fig. 14. The data indicate a very rapid initial decline of the yields with increasing HN₃ density, leveling off somewhat for initial HN₃ pressures over 5 mtorr. The yield at t = 0 would appear to be near unity. We note that this result may be somewhat fortuitous, however, owing to the uncertainties of the input parameters used for the calibration. The yield for a given HN₃ density was significantly improved by increasing the fluorine atom pressure, as shown in the figure.
Figure 14. Photon yields for NF(\(a^1\Sigma\)) emission produced in a low density flame, for two initial fluorine atom pressures. (The solid lines represent results of the calculations described in the text.)
In order to investigate the chemical processes contributing to the yield data shown in Fig. 14, a very simple kinetic code describing the production of NF(a$^1\Delta$) in the system was assembled. The model incorporated the following processes:

\begin{align*}
F + HN_3 & \xrightarrow{k_3} HF + N_3 \quad (3) \\
F + N_3 & \xrightarrow{k_4} NF(a^1\Delta) + N_2 \quad (4) \\
N_3 + HN_3 & \xrightarrow{k_{11}} \text{Products} \quad (11) \\
N_3 + N_3 & \xrightarrow{k_{12}} \text{Products} \quad (12)
\end{align*}

The rate constants used in the model were $k_3 = 1.5 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, $k_4 = 2 \times 10^{-12}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, $k_{11} = 2 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, and $k_{12} = 8 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. The coupled differential rate equations corresponding to processes (3), (4), (11) and (12) were integrated to yield time dependent values of the densities of F atoms, HN$_3$, N$_3$ and NF($a^1\Delta$) using a fourth order Runge-Kutta routine. The calculations were performed with a DEC 11V03 laboratory computer. Yield data obtained from the calculation (yield = peak NF($a^1\Delta$) concentration + initial HN$_3$ concentration) are shown as the solid lines in Fig. 14. As is apparent from the figure, the
agreement between experiment and calculation is quite good, suggesting that processes (11) and (12) may account for the falloff in the yield with increasing density.

In both the small-scale experiments and the calculations, the initial density of fluorine atoms was many times greater than the density of HN₃. It is apparent that, since k₃ is much greater than k₄, a substantial concentration of N₃ builds up as the HN₃ density increases, allowing the side reactions (11) and (12) to occur. These side reactions are the probable sources of the N₂ first positive emission in the flame, which was observed to increase with increasing HN₃ density. Piper and co-workers have in fact reported the observation of N₂ B³Σg⁻A³Σ⁺ emission from reaction (12). Hence, the yield of NF(a¹Δ) falls off with increasing HN₃ density, but it seems likely that the NF population remains heavily inverted. Increasing the fluorine atom density enhances the rate of reaction (4) relative to reactions (11) and (12), and hence increases the yield, as shown in Fig. 14.

This simple model cannot predict the behavior at higher densities shown in Fig. 11. The calculations indicate that the only way for the yield to be high at this regime is for the rates of reactions (3) and (4) to become comparable, either by some dependence of the rate constants on the experimental condition (e.g., temperature), or by some artificial limit on both the rates, as in slow mixing. Under no circumstances does the model predict the strange stoichiometry apparent in Fig. 11.
The behavior of NF(b$^1\Sigma^+$) observed in the experiments was in good agreement with previous measurements. The intensity of the b$^1\Sigma^+$ + X$^3\Sigma^-$ emission increased quadratically with increasing HN$_3$ density (indicative of the formation of the b$^1\Sigma^+$ state by the pooling reaction, process (5)), reaching a peak at which an abrupt leveling off took place. This behavior is reflected in the photon yields measured for emission from NF(b$^1\Sigma^+$), shown in Fig. 15. The peak yield measured was ~ 5 x 10$^{-4}$. Two factors evident in the time decay of the emission may contribute to this low value. First, there appeared to be a mixing time of about 1 ms. Hence, extrapolation of the decay of the emission to t = 0 would yield an intensity substantially greater than the peak value observed. Second, the emission exhibited a very rapid decay, dropping nearly two orders of magnitude in only 4 ms. This result indicates a severe quenching problem. The most obvious potential quenchers are F atoms and F$_2$, which are present at much larger concentrations than other species in the mixture. For a fixed F$_2$ flow rate, variation of the F atom flow by changing the microwave power had little or no effect on the b$^1\Sigma^+$ intensity, indicating that quenching by fluorine atoms is negligible. The b$^1\Sigma^+$ intensity was found to be quite sensitive to the F$_2$ flow rate, however, indicating that molecular fluorine may be an efficient quencher of excited NF.

It is evident from the data shown in Fig. 14 that the efficient production of excited NF by the azide process should not scale to the high densities required for a high-energy chemical laser system. Although the experiments at higher densities (Fig. 11) indicated a substantial "yield" (relative to fluorine atoms), it is apparent that the chemistry responsible
Figure 15. Photon yields for NF(5/2+) emission produced in a low density flame.
for this result is quite complex. Hence, a second portion of this program was begun, directed toward assessing the utility of similar chemistry for the production of excited NCl, NBr, or NI.
III. CHEMICAL PRODUCTION OF EXCITED NCl, NBr, and NI

The results presented in Section I above indicated that, although the behavior of the F + HN\textsubscript{3} system at low densities is well understood, it does not logically extend to the high density regime. At high densities, the system appears to be more complex, and it seems likely that achieving a complete understanding of the mechanisms operative in this regime would require a considerable effort. Hence, before undertaking such a task, it is reasonable to ask if there are similar systems of greater promise which should be considered first. The most obvious such systems are azide mechanisms for the production of excited singlet NCl, NBr, or NI. These heavier nitrogen halides have potential advantages over NF in that their larger transition moments should yield higher gains for a given density of excited species. For example, comparison of the gain of an individual line in the $b^1\Sigma^+ + \chi^3\Pi^-$ transition of NCl to the analogous transition in NF, using Eq. (6) above, yields the following relationship:

$$\frac{G(\text{NCl})}{G(\text{NF})} = 2.15 \frac{\text{ANC}1}{\text{ANF}}$$

(13)

where rotational equilibrium at 350 K has been assumed and $\text{ANC}1$ and $\text{ANF}$ are the respective spontaneous emission rates. Since $\text{ANC}1/\text{ANF} = 35,2,12$ the gain of the NCl transition is about 75 times the gain in NF for the same inversion density. Presumably a transition in NBr would have still higher gain, but for
such a heavy molecule, spin-forbidden production of ground state NBr \( (X^3\Sigma^-) \) by the azide reaction (see Fig. 2) might be a serious consideration.

The reactions of Cl and Br atoms with \( \text{N}_3 \) hold further potential advantage over the NF case in their energetics. These reactions liberate \(-66\) kcal/mol and \(-53\) kcal/mol, respectively, sufficient for the direct population of the \( b^1\Sigma^+ \) states of NCl and NBr. In contrast, the exothermicity of reaction (4) is marginal for the production of NF \( (b^1\Sigma^+) \), and in fact our experiments indicated that this state is generated predominantly by the resonant E-V process Eq. (5).

Azide mechanisms for the production of excited NCl and NBr were studied some years ago by Clark and Clyne.\(^9\) These authors observed that the reactions of Cl and Br atoms with ClN\(_3\) produced emissions from the \( b^1\Sigma^+-X^3\Sigma^- \) transitions of NCl and NBr, respectively, with no evidence of the \( a^1\Delta \) states. Strong \( N_2 \) first positive emissions were also observed, and free \( N_3 \) radicals were detected in absorption. From the presence of the \( N_3 \) radicals, the authors concluded that the Cl + \( N_3 \) reaction was much slower than Cl + ClN\(_3\). Combourieu and co-workers\(^{23}\) and Jourdain, et al.\(^{21}\) obtained somewhat contradictory results in studies of the Cl + ClN\(_3\) system using mass spectrometric techniques. No evidence of \( N_3 \) buildup was found in these experiments, and a computer fit to the data suggested a very fast rate for the Cl + \( N_3 \) reaction.

In small-scale discharge-flow experiments performed in our laboratory prior to the present program, chlorine atoms were added to the F + HN\(_3\) system
as a logical extension of the titration of fluorine atoms with molecular chlorine. The introduction of chlorine to the system produced a brilliant red flame concentrated near the HN$_3$ injector. Scans of the emission spectrum revealed many bands of the $b^1\Sigma^+ - X^3\Sigma^-$ transition in NCl.\textsuperscript{24} In addition, emission bands in the vicinity of 1.1 \mu m were identified as the $a^1\Delta-X^3\Sigma^-$ transition of NCI.\textsuperscript{24} In contrast to the NF experiments, the intensities of both the NCl $a(1\Delta)$ and $b(1\Sigma^+)$ emission varied linearly with the HN$_3$ flow rate, suggesting that both states may be direct products of the pumping reaction. An analysis of the time dependence of the emissions and their behavior as the reagent concentrations were varied suggested the following mechanism for the production of excited NCl:\textsuperscript{19}

\begin{align*}
F + HN_3 & \xrightleftharpoons[k_3]{k_3} HF + N_3 \tag{3} \\
Cl + HN_3 & \xrightleftharpoons[k_{14}]{k_{14}} HCl + N_3 \tag{14} \\
F + N_3 & \xrightleftharpoons[k_4]{k_4} NF(a^1\Delta) + N_2 \tag{4} \\
Cl + N_3 & \xrightleftharpoons[k_{15}]{k_{15}} NCl(a^1\Delta,b^1\Sigma^+) + N_2 \tag{15}
\end{align*}

Rate constants were deduced as follows: $k_3 > 1 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}, k_{14} > 1 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}, k_4 = 2 \times 10^{-12}$ cm$^2$ molecule$^{-1}$ s$^{-1}.$
and $k_{15} = 1 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ (from Jourdain, et al.$^{21}$). It is expected that the “hybrid” system reactions (3) and (15) should yield a superior performance relative to the analogous NF system (reactions (3) and (4)), since the more closely matched rates of these processes should reduce the steady-state concentration of N$_3$, and hence the effect of side reactions involving this species. Indeed, it was observed that addition of Cl$_2$ to the F+ HN$_3$ flame effectively quenched the N$_2$ emission while producing the red NCl flame.

In a similar manner, the addition of Br$_2$ to the F+ HN$_3$ system was found to produce emissions from both the b$^1\Sigma^+$ and a$^1\Delta$ excited states of NBr. In this case, however, the N$_2$ first positive emissions were not quenched, suggesting that the Br + N$_3$ reaction may be slower than F + HN$_3$.

In view of these results, the objective of this portion of the program was simply to select the optimum halogen atom-azide molecule chemical system. Both the spectroscopic characteristics of the emitters and the kinetics of the pumping reactions were to be considered. Within this context, the following issues were addressed:

1. The production of N$_3$ radicals, and the possible utility of reagents other than HN$_3$ for this purpose.

2. The possible production of excited NI molecules by azide reactions.
3. The relative proportions of the $b^1 \Sigma^+$ and $a^1 \Delta$ states produced by the NCl and NBr systems.

4. The absolute yield of excited singlet molecules produced in each of the nitrogen halide systems.

With the exception of Task 1, the experiments were performed with the small-scale discharge-flow system described previously.\textsuperscript{10}

1. The Production of N\textsubscript{3} Radicals

To date, the only tested technique for observation of the N\textsubscript{3} radical in the gas phase involves optical absorption on the bands of the $2 \pi_g + 2 \pi_u^+$ transition near 270 nm.\textsuperscript{25} This method was used by Clark and Clyne in their early experiments on ClN\textsubscript{3} systems.\textsuperscript{9} Since the concentration of N\textsubscript{3} in our system is likely to be extremely small relative to the initial densities of the reagents, such an absorption experiment required a long absorption path-length. Hence an apparatus was constructed to generate a long pathlength F + HN\textsubscript{3} flame, through which a beam of UV light might be passed two or more times. This apparatus, shown in Fig. 16, is essentially a transverse flow system in which HN\textsubscript{3} is admitted to a stream of halogen atoms via a 15.0-cm-long Teflon tube perforated with an array of holes. The flow reactor is a rectangular box with internal dimensions 15.0 cm x 2.0 cm. In the experiments, fluorine atoms were created by a passage of an F\textsubscript{2}/Ar mixture through two parallel microwave discharge cavities as shown in the figure. Chlorine or
Figure 1.6. Schematic diagram of the 15-cm transverse flow apparatus, showing the optical arrangement used for detection of Hg radicals. (T1 and T2 represent the positions of thermocouples, and P1, P2, and P3 are pressure taps.)
bromine atoms could be produced by addition of the molecular halogens to the fluorine atom stream just downstream of each of the discharge zones. For these experiments, the position of the HN₃ injector was fixed just upstream of two 5.0-cm-long windows on each side of the flow reactor, as indicated in Fig. 16. The windows were CaF₂ rectangles cemented to the reactor wall. The temperature and pressure of the gas stream could be measured at either of two points in the reactor, as shown in the figure. Pressures were measured with an MKS Baratron capacitance manometer. The gas temperature was measured by using Pt/10% Rh:Pt thermocouples referenced to 273 K.

The light source used in the absorption experiments was an Oriel deuterium lamp. The source beam was collimated by an iris and two lenses prior to being reflected through the reactor twice by a concave reflector. The collimation of the beam was necessary so that both legs of the optical path might pass through the rather short flames generated in this apparatus (see below). The total absorption length was very nearly 30.0 cm. The light was dispersed by a Jarrell-Ash 0.25-m scanning monochromator and detected with an RCA 1P28 photomultiplier tube. In order to detect small absorptions, a null detector was incorporated into the apparatus so that its offset capabilities might be used.

For a total pressure of 500 mtorr, the linear velocity in the flow reactor was measured to be ~1200 cm s⁻¹. The tandem microwave discharge worked well in the sense that it was quite stable. Addition of Cl₂ to the discharged F₂/Ar stream yielded a visible Cl₂ afterflow which appeared to be uniform over the reactor volume.
Gaseous HN₃ was produced by using the small-scale generator discussed above and shown in Fig. 3. Nitrogen was used as a carrier gas for these experiments, and the generator was found to easily produce flows of 5%-10% HN₃ in N₂. ClN₃ was generated by passage of a dilute Cl₂/He mixture (3% Cl₂) over H₂O moistened NaN₃ suspended on glass wool in an U-tube. Figure 17 shows a diagram of the apparatus constructed for this purpose. The effluent from the Cl₂-NaN₃ reactor was passed through a second U-tube containing Drierite to remove unwanted H₂O from the stream. As with the small HN₃ generator, the ClN₃ generator was operated at atmospheric pressure, a small amount being bled into the flow reactor via a metering valve. The entire apparatus was assembled in a fume hood behind a Lexan blast shield. This apparatus was found to efficiently convert Cl₂ to ClN₃, and conversion efficiencies (determined by UV absorption by the ClN₃) were measured to be 80% to 100% for some experiments.

The addition of HN₃ to the stream of fluorine atoms resulted in the familiar bright green flame dominated by NF b¹Σ⁺ → X³Σ⁻ transitions. The flame appeared to extend across the full width of the reactor, although its length was very sensitive to the reagent flow rates. The flame was typically shorter than the 5.0-cm length of the window. Addition of Cl₂ to the system produced the bright red NCl flame, with characteristics similar to those observed for NF.

Measurements of the absorption spectrum of the F + HN₃ flame in the 270-nm region indicated several features characteristic of N₃. These features were visible only for very low F atom flow rates with HN₃ in excess. The
Figure 17. Small scale apparatus for the generation ClN₃.
spectrum is shown in Fig. 18. Definitive absorptions by four $N_3$ features are apparent in the figure, with the 000-000 band of the $2^3\sigma_g + 2^3\pi_u^+$ transition having the greatest strength. The maximum absorption observed for this transition was approximately 4% over the 30 cm pathlength. Substitution of Cl$N_3$ for HN$_3$ in the system resulted in a similar spectrum of comparable strength. Although a quantitative comparison between the $N_3$ yields in the HN$_3$ and Cl$N_3$ systems could not be made, since the F atom flow rate was too low to be accurately measured (and hence reproduced), it would appear from these results that these yields are not least comparable.

As expected, the $N_3$ absorption diminished upon addition of Cl$_2$ to the F + HN$_3$ system (since Cl + $N_3$ is a faster reaction than F + $N_3$). This phenomenon had been previously inferred from the observation that the addition of Cl$N_3$ quenched the $N_2$ first positive emission present in the flame, thought to be caused by reactions involving $N_3$.

In the present experiments, the measured $N_3$ absorption was correlated with the $N_2$ emission by monitoring the latter, using an uncooled GaAs PMT and a broadband filter centered at 750 nm. For this purpose, a 6.0-mm orifice previously used to pass a thermocouple into the center of the flame was sealed with a quartz window and used as a view port. As expected, the $N_2$ emission increased at a faster than linear rate as the $N_3$ concentration was increased by the addition of HN$_3$ to the system. Since, however, changes in the $N_3$ concentration could be caused only by changing the HN$_3$ flow rate, these results suggest only that the production of excited $N_2$ is second order in HN$_3$ products; i.e., it may be caused by $N + N_3$, $N_3 + N_3$, or $N + N + M$ reactions.
Figure 18. Absorption spectrum of N₃ radicals observed in a low density F+HN₃ flame.
In previous experiments, it was noted that addition of CO$_2$ or SF$_6$ to the F + HN$_3$ or F/Cl + HN$_3$ systems resulted in a large increase (by as much as a factor of two) in the intensity of emissions from excited NCI or NF.$^{10,19}$ Emissions from both the $a^1A$ and $b^1E^+$ states exhibited this effect. One possible explanation was that CO$_2$ or SF$_6$ collisionally relaxed vibrationally excited N$_3$ molecules (produced by the F + HN$_3$ reaction) which might otherwise dissociate. Since the absorption experiments described above monitor the ground vibrational level (000) of the N$_3$(2$^2_g$) state, this hypothesis was tested by monitoring the effect of added SF$_6$ on the N$_3$ absorption. Little or no effect on the N$_3$ (000) concentration was found, placing the relaxation theory in doubt. It is possible, of course, that collisional relaxation might populate levels below the predissociation zone but above the ground state; it would seem likely that some effect on the 000 population would be observed, however. Hence, no definitive explanation is at hand for the SF$_6$/CO$_2$ phenomenon at the present time.

Possible Sources of Excited NI Molecules

At this point in the program, the small discharge-flow apparatus$^{10}$ was reassembled for study of the kinetics of chemiluminescent reactions producing excited NCI, NBr, or NI. A number of experiments were performed in which NI chemiluminescence was sought from systems analogous to those producing excited NCI and NBr. Molecular iodides (HI and I$_2$) were added to the F + HN$_3$ system, such that the following sequence of reactions might occur:
Very little information is available concerning the NI molecule. Phillips\textsuperscript{26} deduced an NI bond strength of approximately 40 kcal/mol from \( \text{N}_2^+ - \text{I}_2 \) chemiluminescence experiments. If this value is correct, the \( \text{I} + \text{N}_3 \) reaction would liberate \(-25\) kcal/mol, sufficient for exciting possible transitions only in the near IR. Miller and Andrews\textsuperscript{27} have suggested that the NI(\(^{1}\Sigma^+\)) state should be about 38 kcal/mol above the ground state.

Addition of HI to the flowing stream of fluorine atoms produced a very bright yellow-green flame. The spectrum of the flame exhibited many red degraded bands, which were identified as IF \( \tilde{B}^3\chi_o^+ + \chi^1\Sigma^+_g \) transitions. Addition of small quantities of HN\(_3\) to this flame (downstream of the HI inlet) had little effect; larger quantities of HN\(_3\) appeared to quench the flame. Similar results were obtained when HI was mixed with the HN\(_3\) stream prior to entering the flow reactor. The UV-visible emission spectrum of the flame, recorded with the cooled GaAs PMT and photon counter, exhibited no features attributable to NI.

A stream of I\(_2\) was produced by passing helium through a bed of solid iodine, yielding an I\(_2\)/He mixture containing about 0.3% I\(_2\). Addition of this mixture to the stream of F atoms in the flow reactor again produced a diffuse yellow-green flame, with a banded spectrum characteristic of IF. Addition of HN\(_3\) to this system downstream of the I\(_2\) inlet appeared to quench the flame.
When the I₂ and HN₃ were pre-mixed prior to entering the F atom stream, the presence of HN₃ was observed to greatly enhance the intensity of the IF flame, probably because of the heat release from the HN₃ reaction. As in the HI experiments noted above, no NI emission was evident in the UV-visible spectrum.

Assuming that the relationship between the energies of the a¹Δ and b¹Σ⁺ states of NI is roughly that found for the other nitrogen halides, near IR emission from the a¹Δ state would be obscured by HF overtone transitions when HN₃ is used as an N₃ source. Hence, infrared emission from NI was sought, using DN₃ and ClN₃ as reagents, with I₂ as the source of iodine atoms. The IR spectrum of the flames produced was measured with a 0.25-m Jarrell-Ash monochromator and a liquid-N₂-cooled intrinsic germanium detector. No NI emission was found in these experiments. Two possible explanations for these negative results seem plausible. First, the I + N₃ reaction may be too slow relative to the N₃ production rate, such that most of the N₃ radicals are lost through side reactions. Second, the spin rules (favoring production of excited singlet states) may be very weak in the I + N₃ system, owing to the large spin-orbit coupling in the iodine atom.

3. The Relative Yields of a¹Δ and b¹Σ⁺ States

Given the information obtained to date, it seems likely that X + N₃ reactions are strongly constrained by the spin rules to produce excited singlet NX molecules, at least for the lighter NF and NCl systems. There would not seem to be any constraint on the relative amounts of the b¹Σ⁺ and
a^1\Delta states produced, however, given that sufficient energy is available to populate both states. If these reactions primarily generate the lower lying a^1\Delta state, the most viable laser candidate might be the a^1\Delta + X^3\Sigma^- transition in NBr; on the other hand, if substantial proportions of the b^1\Sigma^+ states are produced, the best candidate might be the b^1\Sigma^+ + X^3\Sigma^- transition in NCl.

Experiments directed toward measuring the relative densities of the a^1\Delta and b^1\Sigma^+ states were performed by recording the emission spectra of flames generated in the small discharge-flow apparatus\textsuperscript{10} with a 0.25-m monochromator and an intrinsic germanium detector. This detector (North Coast Optics) is normally used for its sensitivity in the 1.0-\mu m region, but in previous experiments it was shown to also respond to the NCl b^1\Sigma^+ + X^3\Sigma^- transition near 665 nm. The relative peak intensities of the a^1\Delta and b^1\Sigma^+ emissions were measured by recording the time decays of these emissions and extrapolating back to t = 0. Figure 19 shows a log plot of typical decay curves recorded for the NCl flame.

The ratio of the peak (t = 0) concentrations of the b^1\Sigma^+ and a^1\Delta states is given by the following expression:

\[
\left( \frac{[b^1\Sigma^+]}{[a^1\Delta]} \right)_{t=0} = \left( \frac{I(b^1\Sigma^+)}{I(a^1\Delta)} \right)_{t=0} \cdot \frac{A(a^1\Delta)}{A(b^1\Sigma^+)} \cdot \frac{R(a^1\Delta)}{R(b^1\Sigma^+)}
\]

where the I's are measured intensities, the A's are spontaneous emission rates, and the R factors represent the spectral response of the monochromator-
Figure 19. Relative intensities of NF(a^1Δ) and NF(b^1Σ^+) emissions produced in a low density F+NH₃ flame. (The numbers in parenthesis refer to NF(a^1Δ) intensities. The data shown were obtained with a detector which had an anomalously low sensitivity to the a^1Δ emission.)
detector assembly. If it is assumed that \( A(a^1\Delta)/A(b^1\Sigma^+) = 4 \times 10^{-3} \) (from measurements of the radiative lifetimes of these states in \( \text{NF}_2 \)), the concentration ratio can be calculated from the data if the R factors are known. To this end, the response of the detection system was calibrated with a blackbody source at known temperatures. The value of \( R(a^1\Delta)/R(b^1\Sigma^+) \) was found to be \( \sim 32 \).

Relative peak densities of the \( a(1\Delta) \) and \( b(1\Sigma^+) \) states of NCI and NBr were measured for several experiments in which either HN\(_3\) or ClN\(_3\) was used as the azide source. For the NBr experiments, bromine atoms were produced by the addition of a \( \text{Br}_2/\text{Ar} \) mixture to the flow reactor just upstream of the discharge zone. The mixture, containing 17% \( \text{Br}_2 \), was prepared in a passivated \( 35 \) \( \text{L} \) tank prior to the experiments. Data obtained from these experiments are shown in Table I. The data for the NCI and NBr systems are quite consistent, suggesting that a similar process is responsible for the generation of excited species in each case, as expected. For the HN\(_3\) reactions, about 14% of the excited singlets are in the \( b^1\Sigma^+ \) states near \( t = 0 \), such that the \( b^1\Sigma^+ + X^3\Sigma^- \) transitions have much higher "gain" than the \( a^1\Delta + X^3\Sigma^- \) transitions. Using Eq. (6) above, we find that the HN\(_3\) data shown in Table I indicate that

\[
\text{Gain (} b^1\Sigma^+ + X^3\Sigma^-\text{)}:\text{Gain (} a^1\Delta + X^3\Sigma^-\text{)} = 56,
\]

if it is assumed that the relative inversion densities are given by the measured density ratios. In sharp contrast, the ClN\(_3\) reactions produce 1% or less of the excited singlets in the \( b^1\Sigma^+ \) states, in agreement with inferences noted from the preliminary spectroscopic experiments described above. If in fact the \( b^1\Sigma^+ \) and \( a^1\Delta \) states are both produced by the reactions of chlorine or bromine atoms with \( \text{N}_3 \).
radicals, this difference must be caused by quenching of the \( b^1\Sigma^+ \) states by \( \text{CIN}_3 \) or reaction products. In subsequent experiments, in which rate constants for the collisional quenching of \( \text{NCI}(b^1\Sigma^+) \) were measured, it was found that quenching by \( \text{CIN}_3 \) is in fact very rapid. The rate constant for this process was measured to be \( k(\text{CIN}_3) = 1.5 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \).\textsuperscript{28}

<table>
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<tr>
<th>Molecule</th>
<th>Azide Source</th>
<th>( I(b^1\Sigma^+)/I(a^1\Delta) )\textsuperscript{a}</th>
<th>( [b^1\Sigma^+]/[a^1\Delta] )\textsuperscript{b}</th>
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<tr>
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<td>CaSO(_4)</td>
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<td>0.005</td>
</tr>
</tbody>
</table>

\( \textsuperscript{a} \)Corrected for the spectral response of the detection system.

\( \textsuperscript{b} \)Calculated assuming \( A(a^1\Delta)/A(b^1\Sigma^+) = 4 \times 10^{-3} \).

It should be noted that the measured relative densities do not necessarily correspond to the intrinsic branching ratios for \( b^1\Sigma^+/a^1\Delta \) production by the \( \text{N}_3 \) reactions, since the time dependences of the emissions have not been taken into account. In view of the rapid radiative rates of the \( b^1\Sigma^+ \) states,\textsuperscript{12} it is likely that their measured time decay represents a formation rate, whereas the decay of the \( a^1\Delta \) emissions is likely to be
dominated by collisional quenching processes. Hence, the intrinsic $b^1\Sigma^+/a^1\Delta$ branching ratios are likely to be somewhat higher than the measured density ratios. We note in this regard that a statistical distribution between the two states would yield $[b^1\Sigma^+]:[a^1\Delta] = 1:2$. In any case, it is clear from the data that the $b^1\Sigma^+ + X^3\Sigma^-$ transitions in flames produced by the HN$_3$ reactions hold the greatest promise for lasing. In order to completely assess this promise, however, the actual yields of these states relative to the reagent flows must be measured. These experiments are addressed in the following section.

4. NCl and NBr $b^1\Sigma^+ + X^3\Sigma^-$ Photon Yields

Measurements of the photon yields of the $b^1\Sigma^+ + X^3\Sigma^-$ transitions were complicated by two factors. First, the radiative lifetimes of the $b^1\Sigma^+$ states of NCl and NBr were unknown at the time of the experiments. Hence, a direct calculation of the peak excited state density, as had been done for measurements of the NF$(a^1\Delta)$ yield, was not possible. Second, the short time length of the flames (for the typical reagent densities used in the experiments) would make comparison to the $0 + NO$ reaction difficult, since calibration by this method assumes a uniform intensity distribution over the volume probed by the detector. In order to solve both of these problems, it was decided to stretch the flames out in time by using very low reagent concentrations. This condition yields a diffuse flame over the full reactor volume. Furthermore, since collisional quenching is not likely to be significant at such low densities, the time integrated intensity profile will
approach the photon yield calculated using the radiative lifetime (i.e., the local excited state density at the peak intensity for higher concentration conditions). For the low density conditions, the photon yield is given by

$$Yield = \frac{\alpha}{\int_0^t I(t) \, dt}$$

where $\alpha$ is the calibration factor obtained from the $O + NO$ procedure and $HN_3$ is assumed to be the limiting reagent.

In the experiments performed, the concentrations of the atoms and $HN_3$ were reduced until the time decay of the $b^1\Sigma^+$ emissions stretched several cm down the length of the flow tube. Care was taken to ensure that the systems were still pseudo first-order in atoms ($F$, $Cl$, or $Br$ atoms), i.e., that $HN_3$ was in fact the limiting reagent. Typically, full-time resolution of the flames required fluorine atom pressures less than 1.0 mtorr (as determined by $Cl_2$ titration) with roughly equivalent $Br$ or $Cl$ atom flows. The $HN_3$ flow rate was normally about 10% of the total atom flow rate. The role of $HN_3$ as the limiting reagent was verified by observing the sensitivity of the emission signal to the $HN_3$ flow rate.

Under such conditions, the $NCI \ b^1\Sigma^+ + \chi^3\Sigma^-$ emissions generated by the $F/Cl + HN_3$ reaction exhibited a fully resolved rise and fall, as shown in Fig. 20. The data were taken with a 0.25-m Jarrell-Ash monochromator with
Figure 20. Time-resolved intensity of the dependence of NCl (b^1Σ^+) → X^3Σ^− emission produced in a low density flame. (□), P(HN) = 0.034 mtorr; (●) P(HN3) = 0.019 mtorr.
1.0-mm slits, fixed at the wavelength of maximum intensity for the $\Delta \nu = 0$
sequence of the NCI $b^1\Sigma^+ + X^3\Sigma^-$ transition near 665 nm. Analysis of the data
shown indicates a risetime $\tau_{\text{rise}} = 3$ ms and a decay time $\tau_{\text{decay}} = 3.5$ ms. In
view of the short radiative lifetime of NCI($b^1\Sigma^+$), $\tau_{\text{rad}} = 630$ ps, it seems
likely that the measured time dependence tracks the $N_3$ concentration in the
system. Hence the observed rise and fall should correspond to the rates of
the $F + HN_3$ and $Cl + N_3$ processes, although which rate dominates the rise is
uncertain. Given pseudo first-order kinetics and an $F$ or $Cl$ atom density of
approximately $1.6 \times 10^{13}$ cm$^{-3}$, the rate constant for the rise would be
$k_{\text{rise}} = 2 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ and the rate constant for the fall would
be $k_{\text{fall}} = 1.7 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. These values are in good agreement
with previous measurements of rate constants for the $F + HN_3$ and $Cl + N_3$ reac-
tions, and add credence to our hypothesis that the steady state $N_3$
density is minimized by the closely matched rates of these two processes.

Similar experiments performed for the NBr flame showed considerable
interference in the NBr $b^1\Sigma^+ + X^3\Sigma^-$ spectrum by $\Delta \nu = 3$ bands of the $N_2$
first positive transition produced at very low atom concentrations. Hence, the NBr
measurements were made at somewhat larger atom densities than those used for
the NCI experiments, and spectroscopic scans were made at several times during
the course of the experiment to make certain that the intensity being measured
was in fact due to $b^1\Sigma^+ + X^3\Sigma^-$ transitions in NBr. The data obtained in these
experiments are shown in Fig. 21. Measurements were made for three $HN_3$ flow
rates, and the peak intensities scaled in a roughly linear fashion with the
$HN_3$. The decay rates calculated from the data are nearly equal for the three
Figure 21. Time-resolved intensity of the NBr($b^1\Sigma^+$) – $X^3\Sigma^+$ emission produced in a low density flame. (A, $P(\text{HN}_3)=0.030$ mtorr, $\lambda_d=634$ s$^{-1}$; B, $P(\text{HN}_3)=0.014$ mtorr, $\lambda_d=659$ s$^{-1}$; C, $P(\text{HN}_3)=0.0058$ mtorr, $\lambda_d=598$ s$^{-1}$.)
experiments and are about twice the magnitude of the decay rates found for NCl (Fig. 20). This faster decay could be caused by the larger atom flow rates used in the NBr experiments. The NBr data shown in the figure exhibit an unresolved rise, clearly much faster than in the NCl case. We interpret these data to indicate that the rise of the emission is characteristic of the F + HN₃ rate and the decay characteristic of Br + N₃. For an estimated Br atom density of 8.2 × 10¹³ cm⁻³, the decay corresponds to a rate constant k = 7.2 × 10⁻¹² cm³ molecule⁻¹ s⁻¹ for the Br + N₃ process. As noted earlier, the discrepancy between the F + HN₃ and Br + N₃ rates could account for the formation of excited N₂ (from N₃ side reactions) in the system.

Conversion of these data to photon yields requires calibration of the light collection efficiency of the detection apparatus. This calibration was performed, using the O + NO reaction as a chemical actinometer, as described above. The calibration factor α was measured at the wavelength of the 0,0 bands of the b¹Σ⁺ → X³Σ⁻ transitions in NCl and NBr, yielding the values α(NCl) = 5.0 × 10⁸ cm⁻³ and α(NBr) = 5.5 × 10⁸ cm⁻³. The uncertainties in determining these values are actually much greater than the agreement would indicate. The major source of error occurs in the N + NO titration producing O atoms. The endpoint NO flow rates in the titration were found to vary by as much as 25%. Other errors (e.g., in the measurements of the total pressure, the gas flow rates, and the emission intensities) should have been much smaller, probably less than 10%. Hence we estimate the overall random error in the measurement of α to be about 50%. Since the published value for the absolute rate constant for photon production by the O + NO reaction has

75
approximately the same uncertainty, the measured photon yields are uncertain within a factor of 2.

NCl and NBr \( b^1\Sigma^+ + \chi^3\Sigma^- \) photon yields were calculated by time integrating the measured intensity profiles, correcting by \( a \), and dividing by the initial HN\(_3\) density as in Eq. 19 above. The results obtained from this procedure are given in Table 2. Given the overall uncertainties in the measurements as discussed above, the yields for both the NCl and NBr transitions are about 10\%. No trend with increasing HN\(_3\) flow rate was found, suggesting that these values should be close to a low pressure limit.

Table 2

<table>
<thead>
<tr>
<th>Molecule</th>
<th>( P_0(\text{HN}_3) ), mtorr</th>
<th>Yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCl</td>
<td>0.019</td>
<td>0.07</td>
</tr>
<tr>
<td>NCl</td>
<td>0.034</td>
<td>0.15</td>
</tr>
<tr>
<td>NCl</td>
<td>0.049</td>
<td>0.16</td>
</tr>
<tr>
<td>NBr</td>
<td>0.030</td>
<td>0.07</td>
</tr>
<tr>
<td>NBr</td>
<td>0.014</td>
<td>0.04</td>
</tr>
<tr>
<td>NBr</td>
<td>0.006</td>
<td>0.07</td>
</tr>
</tbody>
</table>

The values for the \( b^1\Sigma^+ + \chi^3\Sigma^- \) yields shown in Table 2 are close to those found for the ratios of the peak densities of the \( b^1\Sigma^+ \) and \( a^1\Delta \) states, as shown in Table 1. Hence, it is tempting to speculate that all of the HN\(_3\)
added to the system is converted to NCI or NBr in either the a^1\Delta or b^1\Sigma^+ states. This inference is of course subject to the error bounds of both measurements, and as noted above, it is possible that the nascent b^1\Sigma^+ : a^1\Delta branching ratios are actually greater than the measured density ratios. We note as well that the density ratios were measured for much larger reagent concentrations than were the b^1\Sigma^+ photon yields. Nonetheless, it seems likely that, at low densities, a large fraction of the initial HN_3 is converted to excited singlet nitrenes.

5. Kinetic Model for the Production of NCI b^1\Sigma^+

The experiments described above yielded information regarding the rates of processes producing excited NCI, the relative proportions of the b^1\Sigma^+ and a^1\Delta states of NCI present in the flame, and the absolute yield of the b^1\Sigma^+ state at low reagent densities. In parallel experiments, the radiative lifetime of the NCI (b^1\Sigma^+) state and rate constants for its quenching by collisions with a variety of species were determined. Hence, it is reasonable to consider all of these results along with additional data from the literature in order to estimate the gain levels to be expected in the NCI flame. A similar body of information is not available for the NBr system at the present time.

Table 3 shows the kinetic model assembled for the NCI system. The model assumes that the Cl + N_3 reaction produces only excited singlet NCI, with the b^1\Sigma^+:a^1\Delta branching ratio taken from the yield data and peak density ratios presented above. Rate constants for the reactions of N_3 radicals with
HN\textsubscript{3} and N\textsubscript{3} were taken from the work of Jourdain and co-workers.\textsuperscript{21} Although these authors report a range of values for these rates, we have chosen the largest rate constants for the model since these were consistent with the measured drop in the NF(a\textsuperscript{1}A) yield with increasing HN\textsubscript{3} flow (see Section 2.0 above). In some of the calculations described below, an additional reaction was added to the model, in which ground state NCl(X\textsuperscript{3}Σ\textsuperscript{−}) was removed from the system by ClF. This process was assigned the rate constant \( k = 1 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \), in accord with the rate of quenching of NCl(b\textsuperscript{1}Σ\textsuperscript{+}) by ClF.
Table 3
Kinetic Model for the NCl Flame

<table>
<thead>
<tr>
<th>Process</th>
<th>Rate Constant</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pumping Reactions:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$F + HN_3 \rightarrow HF + N_3$</td>
<td>$1.5 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(21)$^a$</td>
</tr>
<tr>
<td>$Cl + HN_3 \rightarrow HCl + N_3$</td>
<td>$1.0 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(21)</td>
</tr>
<tr>
<td>$Cl + N_3 + NCl(b^{1\Sigma^+}) + N_2$</td>
<td>$2.2 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(22)$^a$</td>
</tr>
<tr>
<td>$\quad + NCl(a^1\Delta) + N_2$</td>
<td>$1.3 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(22)$^a$</td>
</tr>
<tr>
<td><strong>Side Reactions:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$F + N_3 \rightarrow NF(a^1\Delta) + N_2$</td>
<td>$2.0 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(10)(21)</td>
</tr>
<tr>
<td>$N_3 + HN_3 \rightarrow H + 3N_2$</td>
<td>$2.0 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(23)</td>
</tr>
<tr>
<td>$N_3 + N_3 \rightarrow 2N_2$</td>
<td>$8.0 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(22)</td>
</tr>
<tr>
<td><strong>Quenching Processes:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$NCl(b^{1\Sigma^+}) + ClF \rightarrow NCl(X^{3\Sigma^-}) + ClF$</td>
<td>$1.2 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(30)</td>
</tr>
<tr>
<td>$\quad + F_2 \rightarrow NCl(X^{3\Sigma^-}) + F_2$</td>
<td>$2.2 \times 10^{-13} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(30)</td>
</tr>
<tr>
<td>$\quad + Cl_2 \rightarrow NCl(X^{3\Sigma^-}) + Cl_2$</td>
<td>$1.7 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$</td>
<td>(30)</td>
</tr>
<tr>
<td><strong>Radiation:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$NCl(b^{1\Sigma^+}) \rightarrow NCl(X^{3\Sigma^-} + h\nu$</td>
<td>$1590 \text{ s}^{-1}$</td>
<td>(12)</td>
</tr>
</tbody>
</table>

$^a$From data presented in this report.

The time dependent behavior of the concentrations of the various species present in the flame was determined by integrating the rate equations corresponding to the processes in the model using a fourth order Runge-Kutta
routine. Finite mixing was taken into account for every bimolecular process by using the following form for the rate equations:

\[
\frac{d[X]}{dt} = \pm \frac{1}{xy} \frac{1}{M} \left( \frac{1}{k[X][Y]} \right)^{-1}
\]  

(20)

where \( k \) is the bimolecular rate constant and \( M \) is the fixed mixing rate.

Figure 22 shows the results of the calculation for the initial partial pressures \( P^0(\text{Cl}) = P^0(\text{F}) = 50 \) mtorr and \( P^0(\text{HN}_3) = 10 \) mtorr (densities were calculated assuming a temperature of 350 K). The mixing rate for the data shown was \( M = 50 \text{ torr s}^{-1} \) (or \( 1.4 \times 10^{18} \text{ cm}^{-3} \text{ s}^{-1} \)). This value corresponds to a mixing rate about one-fifth the value of the \( \text{F} + \text{HN}_3 \) reaction rate. The calculation shows that for these conditions the peak \( \text{NCI}(b^1\Sigma^+) \) concentration (and hence the peak emission intensity) should occur about 270 \( \mu \text{s} \) after the initial contact of the reagents. The \( \text{N}_3 \) density exhibits a broad peak occurring much earlier, at about 100 \( \mu \text{s} \). The ground state density rises slowly, reaching a peak at \( t > 500 \) \( \mu \text{s} \). The calculations corresponding to Fig. 22 included the \( \text{NCI}(X^3\Sigma^-) + \text{CIF} \) process noted above; exclusion of this process had only a minor effect on the results. The inversion density, \( \Delta N \), corresponding to these results was calculated as follows:

\[
\Delta N = N(b^1\Sigma^+) - \frac{1}{3} N(X^3\Sigma^-)
\]  

(21)
Figure 22 Calculated time evolution of selected species in the NCl flame for $P(\text{HN}_3)=10$ mtorr, $P(\text{Cl})=P(\text{F})=50$ mtorr.
The peak inversion for these conditions is predicted to be \( \Delta N = 0.62 \text{ mtorr} \) or about \( 1.7 \times 10^{13} \text{ cm}^{-3} \) at 350 K. Assuming all of the excited \( \text{NCl}(b^1\Sigma^+) \) molecules to be in the lowest vibrational level, and given rotational equilibrium at 350 K, this inversion corresponds to a gain \( G = 5.8 \times 10^{-5} \text{ cm}^{-1} \) for a single transition from the most populated rotational level \((J = 13)\). We have assumed that the four branches of the NCl transition \((Q_p, Q_R, Q_p, \text{ and } S_p)\) have roughly equal radiative rates given by one fourth the measured value.\(^{12}\)

The effect of mixing on the system was investigated by performing a series of calculations in which the peak inversion density and the time elapsed from the initial contact of the reagents to the peak inversion was determined for various values of \( M \). The results of these calculations are shown in Fig. 23. The initial pressures used were \( P^0(\text{HN}_3) = 10 \text{ mtorr} \) and \( P^0(\text{Cl}) = P^0(\text{F}) = 50 \text{ mtorr} \), again at 350 K. The peak inversion density exhibits a steep rise as the mixing rate increases from 0 to \( 1 \times 10^{18} \text{ cm}^{-3} \text{ s}^{-1} \), reaching a plateau at about \( \Delta N = 1.7 \times 10^{13} \text{ cm}^{-3} \) for faster mixing rates. Values of \( M \) as high as \( 3 \times 10^{20} \text{ cm}^{-3} \text{ s}^{-1} \) resulted in relatively small changes from this value. The time to peak inversion exhibits a behavior nearly inverse to that found for the magnitude of the peak inversion. It drops very rapidly as \( M \) increases from 0 to \( 1 \times 10^{18} \text{ cm}^{-3} \text{ s}^{-1} \), then levels off at about 200 us.

Calculations were also performed for somewhat higher reagent densities to probe the initial scaling of the system. For an initial \( \text{HN}_3 \) pressure of 20 mtorr and 100 mtorr each of chlorine and fluorine atoms, the results indicated an inversion density \( \Delta N = 3.8 \times 10^{13} \text{ cm}^{-3} \) at 350 K,
Figure 23 Dependence of the calculated inversion density (•) and time to peak inversion (■) on the mixing rate.
corresponding to a gain of $1.3 \times 10^{-4}$ cm$^{-1}$ for a single line in the 
NCl $b^1\Sigma^+ \rightarrow X^3\Sigma^-$ transition. The mixing rate used was $M = 2.8 \times 10^{20}$
cm$^{-3}$ s$^{-1}$. Clearly, if the model were to accurately reflect the kinetics of 
the real system, the demonstration of a laser at densities near these would 
seem feasible.
IV. GAIN MEASUREMENTS

As noted previously, the kinetic model for the NCl system predicts a single line gain on the order of $5 \times 10^{-5}$ cm$^{-1}$ for reagent densities typical of the experiments performed in our laboratory. For a transverse flow reactor with an active length of 10 cm, the predicted round trip gain would be $1 \times 10^{-3}$. Measurement of an optical gain of this magnitude by using standard single-pass techniques would be extremely difficult. Recently, however, Herbelin and co-workers$^{14}$ demonstrated a multiple-pass method capable of measuring absorptions or gains of $1 \times 10^{-4}$ or less. Although these authors used this technique for the measurement of mirror reflectivities, it can be easily extended to measurements of attenuation or amplification in optical media. The experiment (the details of which have been thoroughly described in the literature)$^{14}$ is based on measurement of the lifetime of photons injected into a very low loss resonant optical cavity surrounding the medium to be probed. This lifetime, $\tau$, is related to the average number of round trip cavity transits, $n$, by $\tau = 2nL/c$, where $L$ is the cavity length and $c$ is the speed of light. The round trip transmission is given by $T = \exp(-1/n)$. This factor defines the mirror reflectivities, $T = R_1 R_2$, for a cavity surrounding a vacuum or medium with negligible losses. For a medium with optical gain, $T_{total} = T_{mirrors} T_{medium} = T_{mirrors} (1 + G)$.

In principle, the cavity transmission might be measured directly by monitoring the attenuation of the transmitted beam. In this case, the experiment is simply a multiple-pass absorption measurement. Deconvolution of
the data would be difficult, however, since the average number of cavity transits would not be determined. A much superior method involves measurement of the optical time delay, $\tau$, caused by the extended path length. This can be accomplished by amplitude modulation of the CW probe beam and measurement of the phase shift, $\delta$, between the incident and transmitted light. The photon lifetime in the cavity, $\tau$, is simply related to the tangent of the phase shift by the following expression:

$$\tau = \frac{2nL}{c} = \tan \frac{\delta}{2\pi f}$$

(22)

where $f$ is the modulation frequency. Hence, the round trip gain of the medium is given by

$$G = \frac{4\pi fL}{c} \left( \frac{1}{\tan \delta_0} - \frac{1}{\tan \delta} \right)$$

(23)

where $\delta_0$ refers to the phase shift determined by the finite mirror reflectivities.

In practice, the sensitivity of the experiment is optimized by choosing a modulation frequency such that $\tan \delta$ varies around 1.0 (i.e., $\delta = 45^\circ$), and by using a large cavity length, $L$, such that the time delay is large for a given number of cavity transits. For typical high reflectivity mirrors ($R = 0.998$), the round trip transmission with no medium would be $T = 0.996,$
coring to \( n = 250 \). Given the relationships between \( \tau \), \( n \), and \( \tan \delta \) given in Eq. (22), a modulation frequency \( f = 95 \) kHz would yield \( \tan \delta = 1.0 \) for \( L = 100 \) cm and this value of \( n \). This frequency is well within the range of acousto-optic or electro-optic modulators. An optical gain in this cavity of \( 1 \times 10^{-3} \) would result in an increase of \( \tan \delta \) from 1.0 to 1.33 (or a change in the phase angle \( \delta \) from 45° to 53°).

1. **The Apparatus for Gain Measurements**

   A schematic of the optical train and electronics of the gain apparatus is shown in Fig. 24. The light source used was a Coherent Radiation model 699-21 ring dye laser pumped by a model CR-3000K krypton ion laser. The ion laser could be operated on many different lines from the UV to the deep red portion of the spectrum. Broadband operation of this laser using optics for the yellow-green output (appropriate for pumping the Rhodamine 101 dye used in the NCl/NBr gain experiments) produced nearly 4.0 W of output. Tuning of the ring dye laser is accomplished with a birefringent filter and two etalons internal to the laser cavity, producing an output linewidth of a few MHz (limited by the frequency jitter of the device). The dye laser output is actively stabilized by continuous comparison to a temperature stabilized reference cavity. For optimum conditions, the "single frequency" output of the dye laser was \( \sim 150 \) mW for \( \sim 3.5 \) W of input power with Rhodamine 101 dye, with the laser tuned near the wavelength of the NCl \( b^1\Sigma^+ \leftrightarrow x^3\Sigma^- \) transition. The output of the dye laser was analyzed with a 1.0-m monochromator (Interactive Technology) and a spectrum analyzer (Tropel model 240). The dye laser system...
Figure 24. Schematic diagram of the optical train and electronics used for gain measurements.
was mounted on an air-suspended optical table; it was found that this was necessary to avoid mode-hopping during a laser scan. The laser could be automatically scanned over frequency ranges as large as 30 GHz by movement of the intracavity etalons.

The actual gain experiment was located in a separate laboratory adjacent to that in which the laser was mounted. The laser beam passed through a small hole in the wall separating the rooms. Amplitude modulation of the beam was accomplished with a Coherent Associates model 304 acousto-optic modulator. This device operates from DC to 3 MHz, with a transmission greater than 80%. The modulator was driven by a Wavetek model 148 function generator. The laser beam exhibited considerable divergence over the path from the laser to the gain experiment, and it was necessary to reduce its diameter with a small telescope prior to entering the modulator. The modulated beam was injected into the optical cavity surrounding the flow tube via the finite transmission of the mirrors. Light transmitted by the cavity was detected with a cooled GaAs photomultiplier tube (RCA C31034).

The phase shift caused by the presence of the resonant cavity in the beam was measured with a PAR 5204 lock-in analyzer. This device monitors both quadrature and in-phase components of the input signal with a phase stability of \( \pm 0.2^\circ \). The tangent of the phase shift was measured by the following procedure. First, the cavity was de-tuned and the phase of the lock-in amplifier (referenced to the modulator) adjusted such that the quadrature signal was zero. The cavity was then re-tuned to maximize the signal observed by the PMT (this corresponded to a tightly focussed spot transmitted by the
cavity, indicating that stable transverse cavity modes had set up). At this point, the phase shift caused by the resonator was determined by the ratio of quadrature and in-phase components of the transmitted light, as follows:

\[
\tan \delta = \frac{\sin \delta}{\cos \delta} = \frac{I_{\text{quadrature}}}{I_{\text{in-phase}}} \tag{24}
\]

The value of \( \tan \delta \) was monitored continuously by using the ratiometer capability of the lock-in analyzer. Data were recorded, using an X-Y plotter, with the X coordinate driven by the scan control of the laser and the Y coordinate by the output of the lock-in.

The flow reactor used in the experiments was a modified version of the transverse flow device discussed above and shown in Fig. 16. Two basic modifications were made for the gain experiments. First, a new \( \text{HN}_3 \) injector was used, such that its position with respect to the optical axis of the cavity might be varied. The new injector was the full 15-cm length of the flow reactor with 45 \( 1.0 \)-mm-diameter holes for admission of \( \text{HN}_3 \) into the flow stream. The position of the injector could be moved over a distance of \( \sim 5 \) cm. The second change involved replacement of the windows previously cemented to the sides of the flow reactor by \( 1.91 \)-cm-diameter "Ultra-Torr" fittings. These fittings were connected to the mirror mounts (Oriel) by \( 1.91 \)-cm-diameter flexible stainless steel tubing, to give a total cavity
length on the order of 120 cm. A diagram of the modified low reactor and cavity is shown in Fig. 25.

A variety of mirrors with high reflectivities in the red visible region (appropriate to the NCl and NBr transitions) were tested in the apparatus. The mirrors ultimately used in the experiment were obtained as a loan from Dr. J.M. Herbelin of the Aerospace Corporation. Phase shift measurements at 70 kHz indicated a round trip transmission of 0.9968 for a cavity formed from these mirrors, corresponding to a reflectivity of 0.9984 for each mirror. This result was in excellent agreement with the value of 0.9989 quoted by Dr. Herbelin. This measurement also showed the phase stability of the experiment to be excellent. Measurements of tan \( \gamma \) made over a period of one hour indicated that tan \( \delta \) varied by only \( \pm 0.01 \), corresponding to a baseline sensitivity in the cavity transmission of \( \pm 0.00003 \). In practice, the phase stability was found to vary considerably over many experiments, such that a typical sensitivity was on the order of \( \pm 0.00005 \).

Following assembly of the apparatus, tests were performed to check the sensitivity of the measurement to the presence of absorbing species in the flow reactor. The first such test involved passage of known amounts of \( \text{I}_2 \) through the cavity. Tellinghuisen\textsuperscript{29} has reported absorption coefficient for \( \text{I}_2 \) continuous absorption in the region 600 nm - 800 nm. For these experiments, a stream of \( \text{I}_2 \) vapor was obtained by passing \( \text{N}_2 \) through a saturator containing solid \( \text{I}_2 \) at room temperature. The \( \text{N}_2 - \text{I}_2 \) flow was metered into the cavity, where it was mixed with a flow of argon used to purge the cavity optics. The experiments were performed by using the broadband output of the dye laser.
Figure 25. Diagram of the transverse flow reactor and optical cavity used for gain measurements.
(i.e., tuned only by the birefringent filter, with the etalons out of the cavity) at 673.0 nm. No appreciable changes in tan δ were observed for flows of N₂ and Ar with no I₂ present, relative to values obtained for the cavity under vacuum. The addition of I₂ resulted in a marked change in tan δ, however. Following Eq. 23 above, the I₂ concentration in the cavity is related to the measured phase shift by the following expression:

\[
[I₂] = \frac{1}{εₐ} \cdot \frac{2πfL}{c} \cdot \frac{1}{\tan δ₀} - \frac{1}{\tan δ}
\]  

(25)

where ε is the I₂ absorption coefficient at the wavelength of the probe laser and t is the absorption length of the medium, 15 cm. Hence a plot of the I₂ concentration, obtained from the measured flow rate of the mixture, vs the measured values of \((1/\tan δ₀) - (1/\tan δ)\) should be linear with a slope characteristic of ε. Figure 26 shows such a plot for the data obtained in these experiments. The slope of the line drawn through the data yields an absorption coefficient ε = 24 & mole⁻¹ cm⁻¹, in fair agreement with the value ε = 44 & mole⁻¹ cm⁻¹ reported by Tellinghuisen.²⁹ As is apparent in the figure, there is considerable scatter in the data, which suggests a sensitivity to a minimum of ~ 500 p/m in absorption. It seems likely that the source of both the scatter and the discrepancy with Tellinghuisen's value of ε lies in the determination of the I₂ concentration, which was simply obtained from the measured I₂/N₂ flow rate and the I₂ vapor pressure in the
Figure 26. Plot of the $I_2$ partial pressure in the optical cavity vs the change in $(\tan \delta)^{-1}$ measured near 673 nm. (The slope of the line yields an absorption coefficient $\delta = 24 \text{ l mole}^{-1} \text{ cm}^{-1}$.)
saturator. If appreciable quantities of I₂ absorbed on the walls of the tubing or reactor vessel downstream of the saturator, the concentration of I₂ would be overestimated, and hence the value of c underestimated.

A second test of the system was made by investigating the production of NH₂ by the reaction F + NH₃ + NH₂ + HF. NH₂ is known to have a strong absorption on its X²B₁ → A²A₁ transition near 630 nm.¹ For this test, fluorine atoms were generated by passage of an F₂/He mixture through the microwave discharges, and NH₃ was added to the flow via the movable injector bar. The dye laser was operated in the "single frequency" mode, scanning through 30-GHz segments. These experiments showed that single absorption features in NH₂ are very easily observed with the phase shift apparatus. A typical absorption spectrum recorded near 630.1 nm is shown in Fig. 27. Four absorption features are evident in the spectrum, labelled A, B, C, and D. The center frequency of the scan is not well known, since it lies outside the wavelength zone over which our monochromator was calibrated (calibrations were performed, with neon lines as wavelength markers). If in fact this wavelength is close to 630.1 nm, the most intense features in the spectrum (A and C) correlate well with the 414-404 and 515-505 transitions if the 080-000 band in NH₂.² This identification would require the center wavelength to be 630.18 nm in air. The features labelled B and D do not correlate with tabulated lines of the NH₂ spectrum. We note, however, that due to the nonlinear relationship between optical density and tan δ, these features are in fact quite weak relative to A and C, and hence may not have been seen before. The true absorption strengths of features A, B, C, and D have the relative values
Figure 27. Absorption spectrum of the $X^2B_1 \rightarrow \tilde{A}^2A_1$ transition of NH$_2$ near 630.1 nm, measured with the phase shift apparatus.
11.0 : 0.7 : 4.0 : 0.1, respectively. NH₂ is apparently a long-lived species in the F + NH₃ flame, since movement of the NH₃ injector over a distance of a few cm had little or no effect on the spectrum. Hence, although these data suggest that the frequency selection and scanning mechanism of the system work well, they do not contain information regarding matching of the flame geometry to the region probed by the laser.

2. **Gain Experiments on the NCl and NBr Systems**

The NCl flame was generated in the transverse flow reactor in the manner described above. In brief, chlorine atoms were produced by adding Cl₂ to the F/F₂/He flow just downstream of the discharge zones. An HN₃/Ar mixture containing 5% to 10% HN₃ was added to the stream through the movable injector bar, generating the bright red NCl b¹Σ⁺ → X³Σ⁻ emission. Argon was also admitted to the stream via inlets which served to purge the flexible stainless steel tubing leading to the reflectors. For the initial tests, the total pressure in the reactor was ~800 mtorr, with the partial pressures of F and Cl atoms varying from 50 to 100 mtorr, and the partial pressure of HN₃ from 5 to 10 mtorr. The temperature in the reactor was measured by insertion of a chromel-alumel thermocouple in the flow. For the conditions noted above, the gas temperature was observed to rise very slowly, reaching only about 350 K after nearly an hour of operation.

NCl b¹Σ⁺ → X³Σ⁻ chemiluminescence was monitored by replacing the temperature tap directly over the optical axis (see Fig. 16) with a small CaF₂ window. NCl emission passing through this window was detected by using a
narrow band interference filter centered near 665 nm (to pass the $\Delta v = 0$ sequence of the $b^1\Sigma^+ - X^3\Sigma^-$ (transition) and a room temperature GaAs photomultiplier tube (RCA 4832). The NCl emission was maximized for roughly equivalent flows of fluorine and chlorine atoms, in agreement with our previous observations. For the conditions yielding optimum NCl chemiluminescence intensity, the emission was quite easily visible to the eye through the cavity end reflectors. The chemiluminescence appeared as a very well defined red spot on the mirrors, indicating that the losses in the medium were sufficiently small to allow stable transverse modes to set up. The shape of the spot was quite sensitive to the mirror positions, and a symmetric circular spot could be obtained with fine tuning. It was noted that the alignment for a circular spot was somewhat different from that corresponding to minimum cavity loss for the dye laser, indicating a slight spatial mismatch between the modes generated by the flame and those produced by injection of the dye beam into the cavity.

Laser scans in segments of 30 GHz were made in the region of the $Q^p$ head of the $0,0$ band of the NCl $b^1\Sigma^+ - X^3\Sigma^-$ transition. Although a number of scans were made over the region 664 nm - 666 nm, most of the effort was concentrated in the region near 664.4 nm, corresponding to transitions from rotational levels around $N = 13$, which should have the greatest population for an equilibrium distribution near 350 K. In this region, each 30-GHz scan should include two or possibly three NCl transitions in either the $Q^p$ or $Q^r$ branches. In all of the experiments, atomic species (F or Cl atoms) were in large excess over HN$_3$. No appreciable gain or loss was observed, although the
kinetic model discussed in Section III predicts a large gain (>$10^3$ p/m) for the conditions of these experiments. Moreover, absorption due to ground state NCl was not observed downstream of the flame, as predicted by the model. Although the experiment is less sensitive to absorption (because of the threefold degeneracy of the ground state), an absorption signal should have been observed with a strength approximately three to five times the intrinsic intensity of the noise (see Fig. 22).

In order to probe the effect of mixing on the system, the reagent flow rates (in particular the diluent flow) were adjusted in order to reduce the total pressure in the reactor to ~400 mtorr. Helium was used as a diluent in both the halogen and HN$_3$ streams. A simple diffusion calculation for these conditions suggests that the mixing time should be less than 50 $\mu$s. Experiments were again performed in which the partial pressures of F and Cl atoms were varied from 30 mtorr to 50 mtorr, and HN$_3$ partial pressures as high as 12 mtorr were achieved (for a gas temperature near 350 K). Other parameters varied included the total pressure (maintained below 500 mtorr), the identity of the diluent (He or Ar), the position of the HN$_3$ injector with respect to the optical axis of the cavity, and the probe laser wavelength. The laser wavelength was varied over several 30-GHz segments around the initial setting (664.4 nm), and finally over much wider zones. No evidence of gain or absorption was observed in any of the experiments. The limiting sensitivity of these tests can be determined from the magnitude and noise level of the baseline phase shift. For the conditions of the experiments, the phase shift was given by $\tan \delta = 1.08 \pm 0.03$, for a modulation frequency of
65 kHz. Using the relation \( \tan \delta = 4 \pi f n L / c \), the average number of cavity transits is found to be \( 317.4 \pm 9.4 \), corresponding to a round trip transmission given by \( T = 0.9968 \pm 0.0001 \). Hence, a signal to noise ratio \( S/N = 1 \) would correspond to a gain or absorption of about 200 p/m.

Further experiments were performed in which \( SF_6 \) was added to the reactive flow in the hope of increasing the intensity of the \( 0,0 \) band of the NCI emission. As noted above, experiments with the cylindrical flow tube indicated that addition of either \( SF_6 \) or \( CO_2 \) increased the \( 0,0 \) intensity by as much as a factor of 2, possibly due to vibrational relaxation within the \( b^1\Sigma^+ \) state or to the stabilization of a reaction intermediate such as \( N_3 \). Note in this regard that the kinetic model discussed in Section III above assumes population of only the \( v' = 0 \) level of the \( b^1\Sigma^+ \) state. In the transverse flow gain experiments, addition of \( SF_6 \) again resulted in a large increase in the \( 0,0 \) intensity. For a total pressure rise of 70 mtorr caused by \( SF_6 \) addition, the NCI emission (viewed through a narrow bandpass filter) doubled in intensity. Nevertheless, no gain or absorption was observed for these conditions. We note, however, that the mixing rate in the system may have been significantly reduced by the presence of large amounts of \( SF_6 \) in the flow.

One final series of these tests was performed on the NCI system in which an effort was made to reduce the intrinsic signal to noise level to the minimum possible value. This was accomplished by incorporating a signal averager (Tracor-Northern NS-570) into the data collection apparatus. This change resulted in an ultimate sensitivity to less than 50 p/m in gain or
absorption for 20 scans over the 30-GHz zone probed. As before, no variation in the baseline phase shift attributable to gain or loss was observed. Tests were made for several 30-GHz regions in both the 0,0 and 1,0 bands of the $b^1\Sigma^+ - \chi^3\Sigma^-$ transition. This result sets a limit on the maximum value of the inversion density present in the system, $\Delta N = [b^1\Sigma^+] - 1/3 [\chi^3\Sigma^-] < 1.6 \times 10^{12}$ cm$^{-3}$, for an initial HN$_3$ density of $\sim 2.7 \times 10^{14}$ cm$^{-3}$. The temperature in the system was measured to be 350 K just downstream of the optical axis, in agreement with previous results. Since no absorption was observed at any position downstream of the HN$_3$ injector (i.e., in time zones where most of the excited NCI had radiated), the maximum density of ground state NCI downstream was $[\text{NCI} (\chi^3\Sigma^-)] < 4.8 \times 10^{12}$ for similar conditions. Since precautions were taken to avoid any optical saturation effects, it would seem that the discrepancy must be due to reactive removal of excited NCI by a process not included in the model. This process cannot involve F$_2$, Cl$_2$, or ClF, in view of the known rate constants for NCI ($b^1\Sigma^+$) quenching by these species. The most likely reagent would appear to be chlorine or fluorine atoms. We note that the photon yield experiments for NCI and NBr (see Section III above) were performed with very low densities of these atoms, in an effort to maximize the time resolution of the flame. To test this hypothesis, the reaction

$F + \text{NCI} (b^1\Sigma^+) \rightarrow \text{products}$

was added to the kinetic model, with a rate constant $k = 1.5 \times 10^{-11}$ cm$^3$ mole$^{-1}$ s$^{-1}$ (about one-tenth the gas kinetic rate). The calculation predicted a maximum inversion density of $\sim 4.8 \times 10^{12}$ cm$^{-3}$ for the same initial conditions tested previously, which should be observable with a signal-to-noise ratio $S/N = 3$ for the optimum sensitivity. Hence, if such a
process is the source of the transparency of the NCl system, its rate must approach the gas kinetic value.

The NBr \( b^1\Sigma^+ - X^3\Sigma^- \) transition should be significantly more allowed than the analogous process in NCl, since the larger mass of the molecule should correlate to a much weaker spin constraint on the transition. In view of the measured radiative lifetime\(^{12} \) for NCl (\( b^1\Sigma^+ , v' = 0 \), \( \tau = 630 \mu s \)), it is expected that the lifetime for NBr (\( b^1\Sigma^+ \)) will be 100 \( \mu s \) or less. Hence, it was expected that gain experiments on the NBr system would yield strong absorption signals, since radiation from excited NBr should be much faster than its production by chemical reactions for the reagent densities typically used in the experiments. Moreover, NBr (\( b^1\Sigma^+ \)) radiation to the ground state should compete favorably with reactive processes removing the excited species, such as those inferred above. Because of the degeneracies of the \( b^1\Sigma^+ \) and \( X^3\Sigma^- \) states (1 and 3, respectively), the experiment is intrinsically less sensitive to absorption by a factor of 3. We note also that the 1:1 abundance ratio of the \(^7\)Br and \(^8\)Br isotopes will reduce the sensitivity still further. It was expected that the increased transition moment and the relatively fast buildup of NBr ground state molecules would compensate for these factors, however.

The NBr experiments began with assembly of a stable source of \( \text{Br}_2 \) vapor. A simple bubbler was set up for this purpose, with helium passed into liquid \( \text{Br}_2 \) via a coarse fritted disc. Since the helium pressure was regulated to about 900 torr in the bubbler, and the liquid \( \text{Br}_2 \) was held at ambient temperature, \( \text{Br}_2 \) vapor comprised about 20\% of the total \( \text{Br}_2/\text{He} \) flow. The NBr
flame was generated in the transverse flow apparatus by simply substituting the \( \text{Br}_2/\text{He} \) stream for \( \text{Cl}_2 \). The bright red flame was easily generated, and the flow of \( \text{Br}_2 \) was found to be quite stable and controllable. The flame intensity (observed with the uncooled GaAs PMT as described above) exhibited a sharp maximum with respect to the relative flow rates of \( \text{Br}_2 \) and \( \text{F}_2 \).

Gain/absorption measurements on the NBr system were performed in the manner noted above. The wavelength of the probe laser was set near \( \pm 3 \text{ nm} \), in the 0,0 band of the NBr \( b^1\Sigma^+ - x^3\Sigma^- \) transition. Since a high resolution study of this band has not, however, been reported, the distribution of individual features present within a given 30-GHz segment was unknown. In fact, NBr absorption features were easily found, with fourteen lines with strengths up to 2500 p/m being apparent within an 80 GHz-zone around 673.6 nm. This absorption spectrum is shown in Fig. 28. These data were obtained for an initial \( \text{HN}_3 \) pressure of 10 mtorr. It was observed that the absorption features were maximized by reducing the \( \text{F}_2 \) flow rate to a value well below that typically employed. This change resulted in Br atom densities well in excess over F atoms, and suggests that either the Br + \( \text{N}_3 \) rate is much slower than F + \( \text{N}_3 \), or that NBr \( (b^1\Sigma^+) \) is quenched by \( \text{F}_2 \) or fluorine atoms. Since, as noted above, results obtained using the cylindrical flow tube suggest that the Br + \( \text{N}_3 \) rate is actually faster than F + \( \text{N}_3 \), the quenching argument seems much more likely. Clearly, quenching by F atoms would agree with our inferences from the NCl gain experiments. Variation of the position of the \( \text{HN}_3 \) injector with respect to the optical axis indicated that the NBr absorption strength passed through a relatively flat peak about 1 ms.
Figure 28. Absorption spectrum of the $X^3\Sigma^+$ - $b^3\Pi$ transition of NBr produced by the F/Br + HN$_3$ reaction.
downstream of the injector. Some evidence of saturation of the $b^1\Sigma^+ - X^3\Sigma^-$ transition (i.e., an inverse dependence of the absorption strength on the incident laser energy) was found. This effect was completely eliminated by attenuating the probe beam (~100 mW) by a factor of $10^3$.

If it is assumed that the transitions observed correspond to rotational states near the peak of the equilibrium distribution for $T = 350$ K, and that the radiative lifetime of NBr ($b^1\Sigma^+$) is 100 µs, then the maximum absorption strengths observed in our experiments indicate a density of ground state NBr molecules of about $3 \times 10^{13}$ cm$^{-3}$. In view of its radiative lifetime, the concentration of excited NBr should be much less than that of the ground state. The calculated density corresponds to about 10% of the initial HN$_3$ density, in rough agreement with the measured photon yield for this system. Hence, it is feasible that the ground state NBr molecules are produced solely by radiation from the excited state.

3. The NF-Bi Energy Transfer System

Although the $a^1\Delta$ state of NF can be generated in high yield by the azide system discussed in Section 1, or by reactions of NF$_2$ radicals,$^5$ the very long radiative lifetime of this state, 5.6 seconds,$^2$ precludes lasing on the $a^1\Delta - X^3\Sigma^-$ transition. Hence it is reasonable to consider systems in which the energy stored in the excited NF is transferred to species more amenable to lasing, in a manner analogous to the $O_2^* - I^*$ system.$^{32}$ Capelle and co-workers$^{33}$ have shown that NF $a(^1\Delta)$ can participate in a very rapid energy transfer process with atomic bismuth, producing the $6^2D_{3/2}$ excited
state of the atom:

\[ \text{NF (a}^1\text{A}) + \text{Bi (6}^4\text{S}_{3/2}) + \text{NF (X}^3\text{Σ}^-) + \text{Bi (6}^2\text{D}_{3/2}) \]  \hspace{1cm} (26)

Since the energy defect for this process is only 16 cm\(^{-1}\) (the forward reaction being exothermic), it seems likely that under appropriate conditions an equilibrium would set up, much as in the \( \text{O}_2^* - \text{I}^* \) system. If it is assumed that the electronic, vibrational, and rotational partition functions of \( \text{NF (a}^1\text{A}) \) are equivalent to those in \( \text{NF (X}^3\text{Σ}^-) \), the equilibrium constant for Eq. 26 would be 1.62 at 300 K. Hence, addition of a small concentration of Bi to a stream of chemically generated \( \text{NF (a}^1\text{A}) \) could result in efficient excitation of the bismuth atoms. If the concentration of \( \text{NF (a}^1\text{A}) \) is much greater than that of bismuth, the nearly total inversion in NF would be reflected in the Bi atoms. For example, if 0.1 mtorr of Bi were added to a stream of gases containing 4 mtorr of \( \text{NF (a}^1\text{A}) \) and no \( \text{NF (X}^3\text{Σ}^-) \), establishment of the equilibrium would result in partial pressures as follows: 3.902 mtorr \( \text{NF (a}^1\text{A}) \), 0.098 mtorr \( \text{Bi (2}^3\text{D}_{3/2}) \), 0.002 mtorr \( \text{Bi (4}^4\text{S}_{3/2}) \) and 0.098 mtorr \( \text{NF (X}^3\text{Σ}^-) \). Energy might be drained out of the equilibrium pool by collisional quenching processes, spontaneous emission, or stimulated emission.

Clearly, one of the principal advantages of the NF-Bi system relative to \( \text{O}_2-\text{I} \) lies in the fact that nearly total population inversions can be produced chemically in NF, permitting maximum power extraction. Another advantage lies in the intrinsically higher gain of the Bi \( 6^2\text{D}_{3/2} - 6^4\text{S}_{3/2} \) transition, for which the spontaneous emission rate has been calculated to be
about four times greater than that for the $I \, ^{52}_{2}p_{1/2} - \, ^{52}_{3}p_{3/2}$ transition. In addition, potential systems advantages may be gained from the shorter wavelength of the bismuth transition, which occurs near 875.2 nm. A clear disadvantage lies with the difficulties in generating a flow of Bi atoms well mixed with a stream of excited NF. Moreover, a key question is the degree to which excited bismuth undergoes reactions with the fluorides present in the mixture (including NF) to produce BiF. The objective of this last portion of the program was to investigate these issues and, if possible, to demonstrate gain in the NF-Bi system by using the resonant cavity phase shift method.

The experiments began by testing sources of gas phase bismuth atoms. The diagnostic employed was absorption of the 306.8-nm resonance line emitted by a hollow cathode bismuth lamp. This line corresponds to the $7^{4}p_{1/2} - \, ^{6}D_{3/2}$ transition in atomic bismuth. In view of the large oscillator strength of this transition ($A = 1.9 \times 10^{8} \, s^{-1}$), a density of Bi atoms on the order of $10^{12} \, cm^{-3}$ would almost totally attenuate this emission over a pathlength of only 2.0 cm, assuming the linewidth of the lamp radiation to be approximately equal to the absorption linewidth of the Bi atoms in the flow. Curves of growth measured for the 306.8 nm line using known vapor pressures of bismuth generated in an oven suggested, however, that the lamp output was significantly broader than the absorption linewidth. Hence, this technique appeared to be an adequate diagnostic in the low density regime.

Two sources of bismuth atoms were tested. The first of these was thermal decomposition of trimethylbismuth (TMB), a technique used by Herbelin.
and co-workers at the Aerospace Corporation. In our experiments, a stream of gaseous TMB was obtained by either premixing the vapor with argon in a storage vessel (TMB:Ar = 1:24) or by bubbling a rare gas directly through the liquid. The second method tested involved reactions of BiCl₃, a solid with a room temperature vapor pressure of ~2 torr. In this case, a stream containing BiCl₃ vapor was generated by passing argon through a fritted disc which supported a sample of BiCl₃ powder.

Several tests of bismuth atom generation from decomposition of TMB were performed with the small cylindrical flow tube. In the first experiments, the TMB/Ar mixture was admitted to a stream of atomic and molecular fluorine (prepared by microwave discharge) diluted in argon. A blue-white flame appeared at the point where the TMB entered the flow, but the intensity of the 306.8-nm resonance line was not attenuated. Similarly, no absorption was found at any point downstream of the TMB injector. In a second group of experiments, molecular hydrogen was added to the flow in excess of the total fluorine flow (atoms and molecules) about 20 cm upstream of the TMB injector, producing a highly diffuse blue-white flame. Addition of TMB to this mixture produced an absorption of nearly 30%. It was noted in these experiments that the flow tube became very hot. In subsequent experiments, TMB absorptions were found only when the flow tube was hot, suggesting that Bi atoms are produced by thermal decomposition of the TMB.

Further experiments on the generation of gas phase Bi atoms were performed by using the 15-cm transverse flow reactor, since this all-metal system was more amenable to the high temperatures apparently required for TMB.
dissociation. For this purpose, the mirror and purge line assemblies used for gain experiments were replaced by quartz end windows, and the output of the bismuth hollow cathode lamp was directed down the axis of the reactor. Fluorine atoms were produced by passage of an F$_2$/He mixture through the two parallel microwave discharges, and H$_2$ was admitted just upstream of the main chamber of the reactor (i.e., through the ports used for Cl$_2$ or Br$_2$ in the NCl and NBr experiments described above). In the initial experiments performed with this apparatus, TMB or BiCl$_3$ was admitted via the movable injector normally used for HN$_3$. Dilute mixtures of these reagents in rare gases were obtained as described above. Strong absorptions (as high as 80%) were obtained when TMB was added to the F$_2$-H$_2$ flame, and transmission of the 306.8-nm resonance line was restored when the flow of either TMB, H$_2$, or F$_2$ was turned off. The H$_2$ was always in excess over F$_2$, such that all of the fluorine atoms were reacted away prior to contact with the TMB. Absorptions were also found in experiments in which BiCl$_3$ was used as a reagent. In this case, bismuth atoms are probably formed by sequential stripping of chlorine atoms from the molecule by hydrogen atoms present in the flow. The absorption signals produced by BiCl$_3$ were quite erratic, however, and could not be reliably generated for given flow conditions.

Further experiments were performed for the TMB system in an effort to quantify the yield of bismuth atoms. For these experiments, the configuration of the apparatus was slightly modified, such that the TMB was premixed with the H$_2$ stream prior to entry of the gases into the reactor. In this case, the optical axis (the point in the flow where Bi atoms were detected) was about
12 cm downstream of the TMB/H₂ inlet. The temperature of the flowing gases was measured 12 cm downstream of the optical axis with a chromel-alumel thermocouple referenced to room temperature. For partial pressures of TMB in the flow varying from $1.4 \times 10^{-2}$ mtorr to $1.0 \times 10^{-1}$ mtorr, the density of Bi atoms present downstream was determined from the transmission of the 306.8-nm resonance line and the known oscillator strength of this $4p - 4S$ transitions. Since the lineshape of the emission from the lamp was unknown, it was assumed that $\alpha = \Delta \nu_{\text{lamp}} / \Delta \nu_{\text{gas}} = 1.5$, and that the lamp output was not self-reversed. Although these arbitrary assumptions result in a very large uncertainty in the calculation of the Bi densities, it seems unlikely that $\alpha$ would be less than 1.5, and hence the values obtained may be considered to be lower limits. The data obtained in these experiments are shown in Table 4. Bismuth densities as high as $10^{10}$ to $10^{11}$ atoms cm⁻³ were produced, and there is a clear correlation between increasing temperature and increasing density. Comparison of the calculated densities with the TMB flow rates indicates that a significant proportion of the parent molecules were dissociated at the higher temperatures.

A series of similar experiments were performed in which D₂ was substituted for H₂ as a fuel in the reactive flow. The results of these tests are also shown in Table 4. The data indicate Bi densities and yields quite similar to those found for the H₂ experiments under comparable conditions. This result supports the hypothesis that the TMB is dissociated by thermal excitation, rather than by an energy transfer process involving vibrationally excited HF or DF.
Table 4

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Temp (K)</th>
<th>( \phi_{\text{Bi}} ) (atoms cm(^{-3}))</th>
<th>Yield*</th>
</tr>
</thead>
<tbody>
<tr>
<td>H(_2)</td>
<td>366</td>
<td>(1.1 \times 10^{10})</td>
<td>0.02</td>
</tr>
<tr>
<td>H(_2)</td>
<td>355</td>
<td>(1.3 \times 10^{10})</td>
<td>0.03</td>
</tr>
<tr>
<td>H(_2)</td>
<td>355</td>
<td>(2.9 \times 10^{10})</td>
<td>0.01</td>
</tr>
<tr>
<td>H(_2)</td>
<td>373</td>
<td>(1.5 \times 10^{10})</td>
<td>0.02</td>
</tr>
<tr>
<td>H(_2)</td>
<td>423</td>
<td>(1.2 \times 10^{11})</td>
<td>0.17</td>
</tr>
<tr>
<td>D(_2)</td>
<td>366</td>
<td>(8.3 \times 10^{9})</td>
<td>0.01</td>
</tr>
<tr>
<td>D(_2)</td>
<td>452</td>
<td>(1.3 \times 10^{11})</td>
<td>0.17</td>
</tr>
<tr>
<td>D(_2)</td>
<td>482</td>
<td>(9.4 \times 10^{10})</td>
<td>0.13</td>
</tr>
</tbody>
</table>

*Yield = \( \phi_{\text{Bi}} \times \rho_{\text{TMB}} \)

No evidence indicating a falloff in the Bi yield with increasing TMB flow rate was found, within the range of densities which could be probed by the resonance absorption method. In order to monitor the generation of greater densities of atoms at higher TMB flow rates, absorption on the \(4S_{3/2} - 2D_{3/2}\) transition was directly measured by the resonant cavity phase shift method. These experiments required that the CW dye laser be operated at the wavelength of this transition, 875.2 nm. Lasing at this wavelength was obtained by pumping HITC dye dissolved in a mixture of propylene carbonate and ethylene glycol with the red lines of the krypton ion laser.\(^{36}\) For an input power of \(\sim 5\) W, a stable, single frequency output of \(\sim 150\) mW was readily
obtained near 875 nm. Steady performance at this level over a period of hours was achieved by passing the dye solution through a cooler which chilled it to \( \sim 9^\circ C \).

By using the output of the dye laser as a probe, a search was made for absorption by the \( ^4S_{3/2} \rightarrow ^2D_{3/2} \) transition in bismuth vapor generated by heating solid bismuth. Two bismuth sources were used for these tests. The first was a quartz heat pipe, with a wick \( \sim 10 \) cm in length. The bismuth transition was observed with this apparatus, but the poor signal to noise ratio of the spectrum obtained and the large density gradient undoubtedly present in the tube precluded any quantitative analysis. Much better results were obtained by using an apparatus in which a sealed quartz cell containing solid bismuth was suspended inside a commercial tube furnace (Marshall Product Co.). The cell, which was 14.7 cm in length, was positioned in the center of the 32-cm-long oven, the ends of which were sealed with insulating material save for a 5-mm-diameter aperture for transmission of the laser beam. The cell temperature was measured with a shielded chromel-alumel thermocouple, data from which indicated no measurable temperature gradient along the length of the cell. Temperature measurements made with an optical pyrometer agreed with the thermocouple readings to within \( \pm 10 \) K. The incident laser intensity \( (I_0) \) was monitored with a photodiode integral to the dye laser system. The transmitted intensity \( (I) \) was detected by a room temperature GaAs PMT (RCA 4832), whose response was monitored by a Keithley 610C electrometer. The optical density of the bismuth vapor \( (\text{OD} = \log \frac{I_0}{I}) \) was measured directly by using the ratiometer capability of a PAR 5204 lock-in analyzer. Spectra
were recorded by an X-Y plotter, with the X channel driven by the ring laser scan control and the Y channel by the output of the ratiometer. Typical scans were 30 GHz wide, as calibrated by an etalon with a 7.5-GHz free spectral range.

A typical spectrum of the Bi $6^4S_{3/2} - 6^2D_{3/2}$ transition recorded in this manner is shown in Fig. 29. The spectrum exhibits nine of the ten hyperfine lines expected for a magnetic dipole transition between these two states ($\Delta F = 0, \pm 1$), and the relative intensities of the lines are in excellent agreement with calculated values for such a transition. Only the 4,4 transition is missing, since its intensity is calculated to be only 0.3% of that of the most intense feature, the 6,6 line. The hyperfine splittings evident in the spectrum are in good agreement with previously reported values obtained from the spectra of allowed transitions terminating on the $6^4S_{3/2}$ and $6^2D_{3/2}$ states. No evidence of electric quadrapole transitions ($\Delta F = 0, \pm 1, \pm 2$) is apparent in the spectrum.

Since the optical pathlength was fixed for the closed cell experiment, measurements of the total integrated absorption coefficient of the transition were possible. For small laser powers, the integrated absorption coefficient is related to the spontaneous emission rate, A, by the following expression:

$$\int k_v \, dv = \frac{\lambda_0^2}{8\pi} \frac{g_2}{g_1} A N L$$

(27)
Figure 29. High resolution spectrum of the $6^2S_{3/2} - 6^2D_{3/2}$ transition in atomic bismuth, showing nine $F', F''$ hyperfine lines.
where $g_2$ and $g_1$ are the degeneracies of the $6^2D_{3/2}$ and $6^4S_{3/2}$ states, respectively, $N$ is the number densities of bismuth atoms, and $k_v \lambda = 2.303 \ OD(v)$. A plot of $2.303 \left(8\pi/A_0^2 \lambda\right) \int OD(v)dv$ vs $N$ is therefore linear with a slope equal to $A$. Such a plot for the data obtained in several experiments is shown in Fig. 30. The total intensities of the nine lines were integrated with a planimeter, and data recorded for various oven temperatures. The densities indicated in Fig. 30 were obtained from the vapor pressure data tabulated by Hultgren and co-workers. The slope of the plot yields a spontaneous emission rate $A = 22.5 \pm 1.4 \ s^{-1}$. This value is in reasonable agreement with that calculated by Garstang, given possible systematic errors in the vapor pressure data.

This information was used in experiments directed toward detecting free bismuth atoms produced by thermal dissociation of TMB in the transverse flow reactor. A portion of the CW probe laser beam was passed through the bismuth so that the laser might be locked on to the frequency of the $6,6$ hyperfine line. The beam was amplitude modulated with an acousto-optic device and injected into a resonant cavity surrounding the flow reactor, as in previous experiments. The cavity reflectors were loaned to us by Dr. J.M. Herbelin of the Aerospace Corporation. A value of tan $\theta$ near 1.0 was obtained for a modulation frequency of 21 kHz with no reagent flow, implying an average photon lifetime corresponding to 1015 round trip passes, or individual mirror reflectivities of 0.9995. This result is in good agreement with data from Dr. Herbelin's laboratory. As in the experiments described above, TMB diluted in helium was added to a flow in which fluorine atoms reacted with excess $H_2$. 
Figure 30. Plot of the integrated absorption coefficient of the Bi $6^{4}S_{3/2}$ - $6^{2}D_{3/2}$ transition vs the bismuth density. (The slope of the plot yields a spontaneous emission rate $A=22.5\pm1.4$ s$^{-1}$ for the $6^{2}D_{3/2}$ state.)
generating a temperature in excess of 100°C. The TMB entered the system through the movable injector bar. Addition of large flows of TMB to the system resulted in a drop in tan δ from 1.0 to 0.7, corresponding to a round trip absorption of about 400 p/m. The absorption signal disappeared when the laser frequency was moved off the 6,6 line and reappeared when the laser was tuned to the nearby 5,4 line. The magnitude of the absorption was very unsteady, however. In a subsequent experiment, a steady absorption was obtained by using large flows of undiluted TMB. The flow rate used corresponded to a TMB density in the reactor of \( \sim 6.5 \times 10^{14} \text{ cm}^{-3} \), to be compared to a bismuth atom density of \( \sim 10^{12} \text{ cm}^{-3} \) calculated from the magnitude of the absorption signal. Clearly, the yield of Bi atoms relative to the TMB flow is only on the order of 0.1%.

These data clearly indicate that a density of bismuth atoms sufficient for the observation of absorption, and hence gain, can be produced in the flow reactor using TMB. The dissociation yield is very low, however. Since undissociated TMB quenches Bi \( (6^2D_{3/2}) \) at a near gas kinetic rate,\(^{41}\) it is unlikely that optical gain on the \( 6^4S_{3/2} - 6^2D_{3/2} \) transition could be demonstrated under these conditions.
V. CONCLUSIONS

The data presented above support the hypothesis that a similar mechanism is operative in all of the halogen atom-azide molecule reactions investigated. This mechanism involves the generation of \( N_3 \) free radicals by the \( F + HN_3 \) reaction, followed by reaction of the \( N_3 \) with \( F, Cl, \) or \( Br \) atoms. Spin constraints on the \( N_3 \) reactions lead to the preferential population of accessible singlet states of the products, in particular the \( a^1\Delta \) and \( b^1\Sigma^+ \) states of the nitrenes (NX). In none of the experiments was any evidence suggesting direct population of ground state nitrenes \( (X^3\Sigma^{-}) \) obtained.

The scalability of these systems, i.e., the rate at which the density of excited products increases with increasing reagent flow, is a sensitive function of the relative rates of formation and decay of \( N_3 \) radicals in each system. In the NF case, the yield of excited singlets falls off rapidly because the rate of the \( F + HN_3 \) reaction is nearly an order of magnitude greater than the rate of \( F + N_3 \). This situation causes a buildup of the steady-state concentration of \( N_3 \) radicals, which can undergo very rapid second-order side reactions producing molecular nitrogen. Hence, the \( F + HN_3 \) system will not scale to the densities required for an efficient NF or NF-Bi chemical laser.

Effective scaling would appear to be much more probable for the NCl system, since the rate constants of reactions producing and removing \( N_3 \) are nearly equivalent in this case. The sum of all the information currently
available concerning the kinetics of processes producing excited NCl, side reactions of N₂, collisional decay processes, and NCl radiative decay, leads to the prediction of easily measurable gain on the b¹Σ⁺ → X³Σ⁻ transition of NCl. It seems very likely that the optical transparency indicated by the gain experiments described above results from removal of excited NCl (b¹Σ⁺) prior to radiation, probably by reaction with halogen atoms present in excess in the flow. The rates of these reactions are unknown. Even if their rate constants approached the gas kinetic value, however, it is possible that gain on the NCl b¹Σ⁺ → X³Σ⁻ transition might be demonstrated by employing stoichiometric flows of HN₃ and halogen atoms, i.e., to remove the excess atoms. The flow rates of HN₃ required for stoichiometric reaction, while maintaining the absolute rates necessary to produce excited NCl on a time frame competitive with its radiative relaxation, cannot be produced with the apparatus used in these experiments, however.

The absorption spectrum obtained for the NBr system demonstrates the viability of the resonant cavity phase shift technique for the measurement of small gains or absorptions. Although very little is known concerning the decay kinetics of NBr (b¹Σ⁺), the appearance of the absorption spectrum is in accord with the expected short radiative lifetime of this species. This shorter lifetime would make NBr a higher gain system than NCl (and perhaps more easily demonstrated at low densities), but kinetics information from the flow tube experiments suggests that NBr may not scale as well as NCl. The rate constant for Br + N₃ appears to be intermediate between Cl + N₃ and F + N₃, and about one-half as large as that for F + HN₃. This inference is
supported by the appearance of $N_2$ first positive emission in the NBr flame. More detailed information concerning the kinetics of NBr production and decay is necessary to fully assess the potential of this system.

The data presented above on the bismuth system represent the first high resolution measurement of the spectrum of the $6^4S_{3/2} - 6^2D_{3/2}$ transition, and the first direct measurement of the radiative lifetime of the $6^2D_{3/2}$ state. Although an NF-Bi gain experiment was not performed, the issues to be addressed by such an experiment were defined and an appropriate diagnostic system was demonstrated. The major hurdle to be overcome before a gain measurement can be performed is the demonstration of an adequate source of Bi atoms. Thermal dissociation of trimethylbismuth or reactions of BiCl$_3$ would not appear to be viable sources. For a small-scale gain measurement, simple evaporation of elemental bismuth may be the most appropriate method. An efficient thermal or chemical source must be developed for a large-scale laser device, however.
REFERENCES
