



へととういう

1. N. P.

Start Press

MICROCOPY RESOLUTION TEST CHART NATIONAL BUREAU OF STANDARDS-1963-A



C\_EILE COPY



UILU-ENG-84-2548

FINAL REPORT ON

ONR CONTRACT

N00014-82-K-0209

"VISIBLE AND NEAR-INFRARED DISSOCIATION LASERS"

Prepared for

Dr. M. B. White Office of Naval Research 495 Summer St. Boston, MA 02210

Prepared by

J. G. Eden K. P. Killeen Department of Electrical and Computer Engineering University of Illinois 1406 W. Green St. Urbana, Illinois 61801

July 1984

84

Approved for public release Distribution Unlimited

007

13

07

DISTRIBUTION STATEMENT A

LECTE

JUL 1 9 1984

B

REPORT DOCUMENTATION PAGE	READ INSTRUCTIONS
I. REPORT NUMBER	BEFORE COMPLETING FORM TACCESSION NO. 3. RECIPIENT'S CATALOG NUMBER
FD-1	4143 DIGI-ENC-84-2548
. TITLE (and Subtilie)	S. TYPE OF REPORT & PERIDD/ddveker
VISTRIE AND NEAR-INFRARED DISSOCIATION	Final Report
VISIBLE AND MEAN INTRAKED DISSOURTION	21 Jan. 1982 to 20 Apr. 19
	UILU-ENG-84-2548
· AUTHOR(+)	CONTRACT OR GRANT NUMBERS
J. G. Eden	N00014-82-K-0209
PERFORMING ORGANIZATION NAME AND ADDRESS	10 PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS
Dept. of Electrical and Computer Engin	Project /10
1406 W. Green St., Urbana, II, 61801	rioject 410
1. CONTROLLING OFFICE NAME AND ADDRESS	12 REPORT DATE
Office of Naval Research	July 1984
800 N. Quincy St.	13. NUMBER OF PAGES
Arlington, VA 22217	ontrolling Office) 15 SECURITY CLASS. (of this report)
	UNCLASSIFIED
	154. DECLASSIFICATION DOWNGRADING SCHEDULE
6 DISTRIBUTION STATEMENT (of this Report)	PORTONI CTATEMENT & )
( -	DISTRIBUTION STATEMENT A
DISTRIBUTION UNLIMITED	Approved for public released
7. DISTRIBUTION STATEMENT (of the abatract entered in Block	k 20, 11 dillerent from Report)
7. DISTRIBUTION STATEMENT (of the abairact entered in Block	k 20, Il dillerent from Report)
7. DISTRIBUTION STATEMENT (of the obstract entered in Block 18. SUPPLEMENTARY NOTES	a 20, 11 dillereni Iram Report)
7. DISTRIBUTION STATEMENT (of the obsidect entered in Block 18. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse side if necessary and identif	k 20, 11 dillereni Iram Report) 17 by block number)
<ul> <li>7. DISTRIBUTION STATEMENT (of the observed entered in Block</li> <li>18. SUPPLEMENTARY NOTES</li> <li>9 KEY WORDS (Continue on reverse side if necessary and identif</li> <li>Visible, laser, tunable, excimer, inje</li> <li>slave amplifier, blue-green, red, mole</li> </ul>	<pre>x 20, 11 dillerent from Report) // by block number/ ection locking, master oscillator, cule, iodine</pre>
<ul> <li>7. DISTRIBUTION STATEMENT (of the abstract entered in Block</li> <li>10. SUPPLEMENTARY NOTES</li> <li>9 KEY WORDS (Continue on reverse side of necessary and identify</li> <li>Visible, laser, tunable, excimer, inje</li> <li>slave amplifier, blue-green, red, mole</li> </ul>	<pre>// 20, 11 dillerent from Report) // by block number/ ection locking, master oscillator, cule, iodine</pre>
<ul> <li>DISTRIBUTION STATEMENT (of the observed in Block</li> <li>SUPPLEMENTARY NOTES</li> <li>KEY WORDS (Continue on reverse side if necessary and identify</li> <li>Visible, laser, tunable, excimer, inje</li> <li>slave amplifier, blue-green, red, mole</li> <li>Experiments have been conducted i</li> <li>cadmium monoiodide (CdI) laser has bee</li> <li>region centered at 657 nm, complete lo</li> <li>for injected laser intensities of 5 W</li> <li>green excimer amplifier at 506 nm h</li> <li>the UV and green bands of I<sub>2</sub> has been</li> </ul>	<pre>tr 20, if different from Report)  tr br block number)  cction locking, master oscillator, cule, iodine</pre>
<ul> <li>DISTRIBUTION STATEMENT (of the obside of entered in Block</li> <li>SUPPLEMENTARY NOTES</li> <li>KEY WORDS (Continue on reverse side if necessary and identify</li> <li>Visible, laser, tunable, excimer, inje slave amplifier, blue-green, red, mole</li> <li>ABSTRACT (Continue on reverse side If necessary and identify</li> <li>Experiments have been conducted i cadmium monoiodide (CdI) laser has bee region centered at 657 nm, complete lo for injected laser intensities of 5 W I2 green excimer amplifier at 506 nm h the UV and green bands of I2 has been signal gain coefficient at 506 nm is</li> <li>CONN 1473 EDITION OF 'NOV 65 IS OBSOLETE</li> </ul>	<pre>// 20, 11 dillerent from Report) // by block number) // by block number // by</pre>
<ul> <li>DISTRIBUTION STATEMENT (of the abstract entered in Block</li> <li>SUPPLEMENTARY NOTES</li> <li>KEY WORDS (Continue on reverse side if necessary and identify</li> <li>Visible, laser, tunable, excimer, inje slave amplifier, blue-green, red, mole</li> <li>ABSTRACT (Continue on reverse side if necessary and identify</li> <li>Experiments have been conducted i cadmium monoiodide (CdI) laser has bee region centered at 657 nm, complete lo for injected laser intensities of 5 W I2 green excimer amplifier at 506 nm h the UV and green bands of I2 has been signal gain coefficient at 506 nm is</li> <li>P 1 JAN 73 1473 EDITION OF 1 NOV 65 IS OBSOLETE</li> </ul>	Ty by block number) Section locking, master oscillator, cule, iodine ()) ()) ()) ()) ()) ()) ()) ()
17. DISTRIBUTION STATEMENT (of the absilact entered in Block 18. SUPPLEMENTARY NOTES 19. KEY WORDS (Continue on reverse side if necessary and identify Visible, laser, tunable, excimer, inje slave amplifier, blue-green, red, mole 20. ABSTRACT (Continue on reverse side If necessary and identify Experiments have been conducted i cadmium monoiodide (CdI) laser has bee region centered at 657 nm, complete lo for injected laser intensities of 5 W I <sub>2</sub> green excimer amplifier at 506 nm h the UV and green bands of I <sub>2</sub> has been signal gain coefficient at 506 nm is D <sup>ronm</sup> 1473 EDITION OF I NOV 65 IS OBSOLETE	<pre>// 20, if different from Report) // by block number) // by block number //</pre>

ŝ

UILU-ENG-84-2548

FINAL REPORT ON

ONR CONTRACT

N00014-82-K-0209

"VISIBLE AND NEAR-INFRARED DISSOCIATION LASERS"

Prepared for

Dr. M. B. White Office of Naval Research 495 Summer St. Boston, MA 02210

 $\langle \cdot \rangle$ 

Prepared by

J. G. Eden K. P. Killeen Department of Electrical and Computer Engineering University of Illinois 1406 W. Green St. Urbana, Illinois 61801

July 1984

## TABLE OF CONTENTS

\*\*\*\*\*

τ

K.

	Page
A. I <sub>2</sub> Green Amplifier	1
B. CdI Discharge-Pumped Laser	4
PUBLICATIONS UNDER ONR SUPPORT	11
DEGREES GRANTED UNDER ONR SUPPORT	11
PERSONNEL	11
APPENDIX	12



Our research in the study of tunable molecular lasers has been quite productive, specifically with regard to the work on the  $I_2$  green excimer laser and the cadmium-monohalide discharge laser.

# A. I<sub>2</sub> Green Amplifier

During this time, we have extensively investigated the optical properties of this bound-to-free transition in diatomic iodine under electron beam excitation of dilute mixtures of hydrogen iodide in high pressure rare gas buffers. Using a coaxial diode that was built to mate with a Febetron 706 electron beam generator as an excitation source, we have observed spontaneous emission and amplification on this transition centered at 505 nm with a bandwidth of ~ 150 Å. The details of this work have been reported in our attached paper: Gain on the green (504 nm) excimer band of  $I_2$ ; <u>Appl. Phys. Lett.</u> 43, 539 (1983). Figure 1 shows the gain and emission profiles for this green band in argon and neon gas diluents.

One interesting aspect of the work on  $I_2$  that was discussed only briefly in this article is the relationship between the upper molecular states of the green band and that for the 342 nm emission (the D' state). It was found that a strong inverse relationship exists between the gain measured on the 505 excimer band and the intensity of stimulated emission on the 342 nm band. Under the nominal conditions used to obtain ~ 1.2% cm<sup>-1</sup> amplification at 506 nm in  $I_2$ , the 342 nm D'  $\rightarrow$  A' transition superfluoresces with an intensity of 1.8 MW cm<sup>-2</sup>. When this is increased to 30 MW cm<sup>-2</sup> using a high Q cavity to enhance the circulating UV flux, the gain at 506 nm is completely extinguished. This is shown in Figure 2 which is a plot of gain on the  $I_2$  green band at 506 nm versus intracavity flux on the UV laser transition. Note the steep slope of the curve at 1.8 MW - cm<sup>-2</sup> which supports our earlier calculations that the 506 nm



ЧġҚ

Figure 1,



ANALY STREET STREET STREET

gain coefficient at this intensity is only ~ 20% of its small signal value. Though we do not as yet have the complete picture on the relationship between the upper states of these bands concerning energy defect and coupling, it is likely that suppressing the 342 nm UV superfluorescence will lead to a dramatic increase in the amplification measured in the green.

To obtain a measure of the ultimate gain possible from this visible excimer, more information is needed about this coupling as well as experiments designed to measure the extraction efficiency on this band. A high powered (> 10 MW -  $cm^2$ ) probe signal from an excimer pumped dye laser could be used to study the effect of visible amplification and UV (342 nm) emission versus probe signal intensity. These experiments could best be carried out using longer pulse e-beam pump devices which transfer greater energies to the excited manifold of states in I<sub>2</sub> than can be realized with our present source. This data should help provide a more complete idea of the viability of this system as a tunable green amplifier.

### B. CdI Discharge-Pumped Laser

Tunable, efficient sources of coherent radiation in the visible and near infrared are attractive for a variety of applications, including photochemistry communications and countermeasures. This requires gain media with a large degree of homogeneous broadening such that the energy stored in the upper laser level can be efficiently extracted at any wavelength in the gain spectrum.

As a demonstration of the tunability of new laser media in the visible, an injection locking experiment has been performed on a cadmium monoiodide UV-preionized, transverse discharge laser. This laser was recently developed here at the Gaseous Electronics Laboratory and to our knowledge is the only one of its kind in existence.

The injection locking experimental apparatus is shown in Fig. 3. The  $\sim 1$  µs long pulse from a Chromatix CMX-4 flashlamp pumped, tunable dye laser was injected into the CdI discharge cavity through the 1.35% transmitting mirror. The dye laser linewidth was narrowed to 0.3 cm<sup>-1</sup> with an internal etalon. Part of the dye laser pulse was split off to monitor its intensity with a calibrated vacuum photodiode. All intensities mentioned are those which pass through the input mirror, i.e., the actual intensity injected into the cavity. No attempt was made to mode match in this experiment. The timing circuit triggered the fast CdI discharge (120 ns FWHM current pulse) sometime during the longer dye laser pulse. The output spectrum through the 0.27% transmitting mirror was recorded with a PARC model 1450 optical multichannel analyzer using a model 1454 intensifying head coupled to a 0.6 m Hilger-Engis spectrometer. The spectro meter was operated in first order and the slits were adjusted to provide resolution of 1.4 Å.

100 A

Figure 4 shows the integrated output intensity dependence of the CdI laser on the injected wavelength for a constant injected intensity of ~ 5 W cm<sup>-2</sup>, which was the maximum intensity used in this experiment. The upper curve is the B + X fluorescence spectrum of CdI between 650 and 662 nm and the lower curve is a composite of the output spectra for nine different injected wavelengths. Each line is also the sum of ten shots to average out shot-to-shot variations. Between 655 and 660 nm, locking is complete at 5 W cm<sup>-2</sup> and tuning over this did not show any region where locking could not be achieved. The integrated output intensity varied by only a factor of 3 over this range. Outside this region, locking is always incomplete at 5 W cm<sup>-2</sup> although various amounts of energy could be extracted at the injected wavelength.

The effect of the injected dye laser intensity on the CdI laser output spectrum is seen in Fig. 5 where the dye laser wavelength is set at 653.1 nm.



للمتر في ال

. •

Figure 3.



sarah waxara

s - I scottil contrati manata antenza accessa accessa antenza

Figure 4.



ÿ

Figure 5.

As the injected intensity increases, more energy is extracted at 653.1 nm and less on the naturally lasing line at 657.1 nm. For this case, locking is not quite complete at the maximum intensity of 5 W cm<sup>-2</sup> although the majority of the energy is emitted at 653.1 nm. This data is plotted graphically in Fig. 6, along with data at two other wavelengths: 655.1 nm and 657.1 nm. As shown in this graph, when injecting at 653.1 nm, energy begins to be extracted at the injected wavelength for intensities of only ~  $10^{-2}$  W - cm<sup>-2</sup> and the output increases exponentially up to 5 W cm<sup>-2</sup>. When injecting at 655.1 nm, energy begins to be extracted at intensities of  $10^{-3}$  W cm<sup>-2</sup> and the output increases until complete locking occurs at  $\leq 1$  W cm<sup>-2</sup>. Beyond that, a less than linear increase is seen. The increase is always less than linear at 657.1 nm but there is more than an order of magnitude improvement in the output over the uninjected laser by locking with just 5 W cm<sup>-2</sup>. The dye laser flux seeding the cavity allows threshold to be reached sooner and therefore more of the stored energy can be extracted.

Two important points should be noted. The first is that only weak intensities ( $\leq 5 \text{ W cm}^{-2}$ ) are needed to achieve complete locking over a 5 nm range between 655 and 660 nm. This is the result of the modest but significant peak gain coefficient of CdI, 2% cm<sup>-1</sup> at 657.1 nm, which drastically reduces competition from amplified spontaneous emission. The second point is that, as seen in Fig. 6, as the injected intensity increases, the difference in the output at different wavelengths decreases, indicating a high degree of homogeneous broadening. Figure 6 also indicates that with slightly higher injected intensities, complete locking should be achievable across the entire wavelength region of Fig. 4 between 650 and 662 nm. These results are now being prepared for publication.



Parkana, unnunn

ALL TAXABLE MANUAL ANALY ANALY ANALY ANALY ANALY ANALY ANALY

#### PUBLICATIONS UNDER ONR SUPPORT

- D. P. Greene and J. G. Eden, "Discharge pumped ZnI (599-606 nm) and CdI (653-662 nm) amplifiers," <u>Appl. Phys. Lett.</u>, vol. 42, p. 20, January 1983.
- D. P. Greene and J. G. Eden, "Lasing on the B-X band of cadmium monoiodide (CdI) and <sup>114</sup>CdI in a UV-preionized, transverse discharge," <u>Appl. Phys. Lett.</u>, vol. 43, p. 418, September 1983.
- D. P. Greene and J. G. Eden, "Transient absorption spectroscopy of a CdI discharge," (in preparation).
- 4. D. P. Greene and J. G. Eden, "On the origin of the 454 nm bound-free absorption band in a CdI discharge," (in preparation).
- D. P. Greene and J. G. Eden, "Injection locked, tunable output from a CdI transverse discharge laser," (in preparation).
- 6. K. P. Killeen and J. G. Eden, "Gain on the green (504 nm) excimer band of I<sub>2</sub>," <u>Appl. Phys. Lett.</u>, vol. 43, pp. 539-541, September 1983.
- 7. K. P. Killeen and J. G. Eden, "Coupling between the green and UV bands of  $I_2$ ," (in preparation).

### DEGREES GRANTED UNDER ONR SUPPORT

D. P. Greene, Ph.D. Degree in Electrical Engineering, August 1984.

#### PERSONNEL

J. G. Eden, D. P. Greene, K. P. Killeen

# Gain on the green (504 nm) excimer band of I<sub>2</sub>

APPENDIX

K. P. Killeen and J. G. Eden

Department of Electrical Engineering, University of Illinois, Urbana, Illinois 61801

(Received 15 June 1983; accepted for publication 7 July 1983)

Gain on the green excimer band  $(\lambda \sim 504 \text{ nm})$  of the iodine dimer has been observed in electronbeam-pumped mixtures of Ar (or Ne), and hydrogen iodide. A peak gain coefficient of  $\gamma > 1.1\%$  cm<sup>-1</sup> is measured at 506 nm with a tunable dye laser and the full width at halfmaximum (FWHM) of the gain spectrum in Ar diluent is 13 nm. Temporally resolved gain and fluorescence measurements show that the green emission originates from an I<sub>2</sub> ion pair state other than D'. However, strong collisional coupling between the upper states of the UV ( $\lambda = 342 \text{ nm}$ ) and green bands and superfluorescence on the UV band ( $I_{342} \sim 2 \text{ MW cm}^{-2}$ ) appear to limit the green gain coefficient to < 20% of its small-signal value. Consequently, suppression of superfluorescence on the UV D' $\rightarrow A$ ' bands of the homonuclear halogens should lead to a new family of excimer lasers with wavelengths extending from the green into the ultraviolet.

PACS numbers: 42.55.Hq, 33.20.Kf

The well-known molecular iodine (I2) ultraviolet laser at 342 nm arises from the D' (1432,  ${}^{3}\pi_{2g}) \rightarrow A'(2431, {}^{3}\pi_{2u})$ transition of the molecule.<sup>1-3</sup> First demonstrated in 1975, this laser has now been pumped by both electron beam<sup>4-7</sup> and optical excitation.<sup>8-11</sup> The collisional processes responsible for I<sub>2</sub> excited state production in the former case have been extensively studied<sup>12</sup> and the predominant mechanism for formation of the lowest lying  $I_2$  ion pair states  $(I^+I^-)$  is collisional deactivation of the I\* (\*P) metastables by iodinecontaining molecules such as HI, CF<sub>3</sub>I, or I<sub>2</sub>. Optical pumping of the laser is possible because the I<sub>2</sub> absorption band centered near 190 nm results in the production of excited  $D^{1}\Sigma_{+}^{+}$  molecules.<sup>13</sup> In the presence of a buffer gas, these molecules collisionally relax to the lowest lying ion-pair state,  $D'^{3}\pi_{2e}$ , which is the upper energy level for the UV laser. Both the spontaneous emission and laser spectra for the  $D' \rightarrow A'$  band exhibit well-developed vibrational structure since the transition is bound-bound<sup>2</sup>. Output energies up to 13 J have been reported for an  $I_2$  laser optically pumped by the radiation from an open discharge.<sup>10</sup>

In recent work in this laboratory on the iodine-monofluoride laser, a strong green emission band peaked near 504 nm was observed when mixtures of Ar and hydrogen iodide (HI) were irradiated by a relativistic electron beam. Substituting CF<sub>3</sub>I for the HI, the green continuum is again present (though considerably weaker) and underlies the long-wavelength portion of the IF 490-nm band. Hemmati and Collins,<sup>14</sup> Guy et al.,<sup>3</sup> Baboshin et al.,<sup>13</sup> and apparently Shaw et al. [see Fig. 1(b) of Ref. 11] have previously observed this I<sub>2</sub> band for rf discharge or optical (ArF laser or atomic iodine emission) excitation, but this is. to our knowledge, the first observation of the emission for electron beam pumping. References 3, 13, and 14 attribute this green continuum to the  $D'^{3}\pi_{2a} \rightarrow 2332, {}^{3}A_{2u}$  transition of I<sub>2</sub> which is bound  $\rightarrow$  free (cf. Fig. 1). However, preliminary time-resolved fluorescence measurements indicate that the green and UV bands of  $I_2$  do not share the same ion pair upper state (D') but both levels lie close to one another in energy. The experimental evidence which led to this conclusion will be described in detail elsewhere. The transition responsible for the green  $I_2$  band is, nonetheless, bound-free and so a laser on this transition

would be continuously tunable and could correctly be called an excimer laser.

This letter describes the first measurement of net gain on the green excimer band of  $I_2$  in *e*-beam excited mixtures of Ar (or Ne) and HI. The peak gain coefficient ( $\gamma$ ) of 1.1% cm<sup>-1</sup> is observed at 506 nm and the full width of the gain spectrum is ~13 nm.

A schematic diagram of the experimental apparatus is shown in Fig. 2. Excitation of the HI/rare gas mixtures is provided by a Febetron 706 *e*-beam generator (10-J stored energy) and a coaxial diode of 23-cm active length. The Febetron delivers 3-ns, 600-kV pulses to a cathode consisting of strips of 25- $\mu$ m-thick titanium foil situated longitudinally along the inner wall of a 5.5-cm o.d. aluminum tube. Aluminum anodes were fabricated from 7.6-mm i.d. (1-mm wall thickness) tubing by machining the wall (over the central 23cm region of the tube) down to 250  $\mu$ m. The desired wall thickness (~125 $\mu$ m) was then obtained by etching the tubes



FIG. 1. Partial energy-level diagram for molecular iodine showing the wellknown  $D' \rightarrow A'$  UV band and the green excimer transition (after Refs. 1 and 14). The position of the argon molecular excited states is also indicated. The upper level for the green band appears to be an ion pair state lying above but close to D'.

Appl. Phys. Lett. 43 (6), 15 September 1983 0003-6951/83/180539-03\$01.00



FIG. 2. Schematic diagram of the experimental apparatus. Either an Ar<sup>+</sup> or tunable dye laser provided the probe beam and the grating served to suppress the background spontaneous emission. The turning mirror was removed to obtain the emission spectra.

in a column of 5 M NaOH solution for  $\sim 8$  min. The active volume of this device is 12 cm<sup>3</sup>. A ribbed Lexan insulator sleeve was installed on the cathode stem to prevent flashover to ground. Aluminum Brewster's angle window mounts also served as the means to introduce the gases to the interior of the anode and the diode pressure was maintained at  $\sim 10^{-6}$  Torr with an oil diffusion pump system.

Research grade rare gases and electronic grade HI were used as supplied by the manufacturers and the gases were premixed in a steel cylinder. Partial pressures below 100 Torr were measured with a capacitance manometer. Passivation of the system was accomplished by allowing 100 Torr of HI to stand in the vacuum manifold, mixing cylinder, and anode for 24 h.

Fluorescence spectra were recorded by viewing the *e*beam cell at one end with a Hilger-Engis 0.6-m spectrograph and a PAR optical multichannel analyzer (OMA). The first order dispersion of the spectrograph is  $\sim 1.4$  nm/mm, giving an overall detection system resolution of  $\sim 0.2$  nm.

To probe the excited medium for gain, the beam from an Ar ion laser or a flashlamp-pumped, tunable dye laser (Chromatix CMX-4;  $\Delta\lambda \sim 0.08$  nm) was directed along the axis of the anode. The steps taken to minimize interference from background fluorescence included placing irises and a diffraction grating in the optical path and locating the vacuum photodiode (S-20 surface) 5 m from the end of the e-beam diode. The transmitted probe laser intensity was subsequently displayed on a storage oscilloscope. Low probe intensities  $(< 10 \text{ kW cm}^{-2})$  were obtained with neutral density filters, thus ensuring that the probe flux did not approach the estimated saturation intensity of the green band. Also, the temporal history of the I, UV and green fluorescence was recorded with the same photodiode and interference filters (UV:  $T_{\text{max}} = 55\%$  at  $\lambda_0 = 350$  nm,  $\Delta \lambda = 55$  nm; green:  $T_{\text{max}}$ = 77% at  $\lambda_0$  = 500 nm,  $\Delta\lambda$  = 38 nm).

Spontaneous emission and gain waveforms from a 0.2% concentration of HI in 5930 Torr of Ar mixture are illustrated in Fig. 3. Actually, the 342-nm transition is super-

Appl. Phys. Lett., Vol. 43, No. 6, 15 September 1983

fluorescing, as evidenced by the spectral and temporal narrowing of the emission. (Note that the intensity scales for the green and UV bands shown at the top of the figure are fixed relative to one another.) For a probe laser wavelength of 506 nm, the oscillogram shown in Fig. 3(b) was observed. The strong absorption that is evident throughout the pump pulse is characteristic of electron-beam-excited rare gases<sup>15</sup> and this time period is immediately followed by a gain pulse of 8– 10 ns full width at half-maximum (FWHM). This feature is due entirely to gain in the active medium—background spontaneous emission was not observable on this intensity scale.

Probing the excited gases with various  $Ar^+$  and dye laser wavelengths between 495 and 512 nm has resulted in the I<sub>2</sub> green band gain spectra presented in Fig. 4. Each of the gain coefficient data points in the figure is the average of at least three experimental trials. Part (a) shows that, for Ar/HI mixtures, maximum gain is observed at 506 nm ( $\gamma = 1.1\%$  cm<sup>-1</sup>) and the FWHM of the entire profile is ~13 nm. Most of the absorption features that are so prominent in this spectrum [and (b) as well] are too broad to be attributed to excited atomic states and, as noted by Zamir *et al.*,<sup>15</sup> appear to be due to a transition of  $Ar_2({}^{3}\Sigma)$  molecules to a Rydberg state. None of the absorptive transitions in (a) or (b) appear to involve the I<sub>2</sub> molecule itself.

Figure 4(b) confirms the assignment of the dominant absorption lines in (a) to the Ar<sub>2</sub> excimer. In this case, Ne is the diluent and the dip in the Ar/HI gain spectrum at  $\lambda \simeq 504.5$  nm has vanished. Also, if one corrects for the smaller stopping power of neon as compared to argon, the peak  $\gamma$  in (b) is only slightly less than that for the Ar buffer. Only the central portion of the Ne/HI spectrum was examined in order to confirm the expected disappearance of the 504-nm absorption line. The top curves in (a) and (b) are the fluorescence spectra which have been corrected for the response of the spectrograph and OMA. Aside from the obvious absorption features, these bands are structureless continua, as would be expected.

The dependence of the small-signal gain coefficient at 506 nm on the argon pressure and the gas mixture composition is shown in Fig. 5. The best compromise between optimizing  $I^{*}({}^{4}P)$  formation<sup>12</sup> and minimizing quenching of the upper state occurs for  $P_{Ar} \simeq 4000$  Torr and an HI concentration of at least 0.5%.

We note from Fig. 3 that the peak 506-nm gain is measured following the e-beam pumping pulse and just after the



FIG. 3. (a) Fluorescence and (b) gain waveforms for e-beam excited  $I_2$ . The 342-nm superfluorescing pulse has a full width ~ 1.1 ns. In (b) the gain observed at  $\lambda = 500$  nm for a 0.2% HI in Ar ( $p_{101A1} \simeq 5930$  Torr) mixture is shown. The peak signal in the oscillogram corresponds to a gain coefficient of 1% cm<sup>-1</sup>. Gain coefficients as high as 1.2% cm<sup>-1</sup> have been observed.

K. P. Killeen and J. G. Eden



FIG. 4. Spontaneous emission and gain spectra for electron beam excited mixtures of HI and (a) argon or (b) neon. The minimum detectable gain coefficient is  $\sim 0.1\%$  cm<sup>-1</sup> and the maximum concentrations of Ar in Ne and Ne in Ar are both less than 1 ppm. The top traces in (a) and (b) are the flourescence spectra as recorded with an OMA.

peak in the 342-nm band intensity. The large optical flux at 342 nm that arises from superfluorescence on the  $D' \rightarrow A'$  band strongly depletes the D' population before gain is measured on the green transition. Consequently, since the upper states for the UV and 504-nm bands are collisionally coupled, the gain coefficients in Fig. 4 are "saturated." Therefore, under these experimental conditions, the measured gain coefficients are likely well below their small-signal values.

To estimate the green (506 nm) small-signal gain coefficient, let the frequency full width and the radiative lifetime  $\tau$ , for the  $D' \rightarrow A'$  band be  $\sim 10^{13} \text{ s}^{-1}$  and 7 ns, respectively. <sup>11,16</sup> Consequently, a conservative estimate<sup>12</sup> for the stimulated emission cross section for this UV band is  $\sigma_{SE} \simeq 6 \times 10^{-16} \text{ cm}^2$  and the saturation intensity is  $I_{sat} = hv(\sigma_{SE}\tau_r)^{-1} \simeq 0.14 \text{ MW cm}^{-2}$ . Using a Gen-Tec energy detector, 1 mJ of energy at 342 nm was found to be contained in the 1.1-ns FWHM pulse (so  $I_{peak} \sim 1.8 \text{ MW cm}^{-2}$ ). Then, if the  $D' \rightarrow A'$  transition is homogeneously broadened, the population inversion and  $\gamma$  at peak intensity are both only 8% of their small-signal values. Assuming that the populations of the lower states of the green and UV bands (one dissociative perhaps,  ${}^{3}\Delta_{2u}$ , and the other A', respectively) are negligible, then the inversion on the 504-nm I<sub>2</sub>



FIG. 5. Dependence of the green gain coefficient on the composition and total pressure of the Ar/HI mixture. The dye laser probe wavelength is 506 nm. band is also  $\sim 1/13$  that present before stimulated emission occurs on the 342-nm transition. Consequently, the presence of a distributed absorber at 342 nm in the gain medium may allow the peak gain coefficient at 506 nm to rise beyond 5% cm<sup>-1</sup>. Viewed from another perspective, saturating the green transition with a dye laser (or HgBr oscillator) pulse should yield reasonable extraction efficiencies.

Although saturated, the gain coefficients measured here are comparable to those for the XeF ( $C \rightarrow A$ ) band. With the same apparatus, the gain coefficient at  $\lambda = 488$  nm was found to be 1.5% cm<sup>-1</sup> for a 1:2:370 mixture of NF<sub>3</sub>:Xe:Ar (total gas pressure again  $\simeq 6000$  Torr). (In this case, the UV  $B \rightarrow X$  band was not superfluorescing.)

In summary, gain on the green excimer band of  $I_2$  has been observed in *e*-beam pumped rare gas/HI mixtures. The room-temperature operation of this system, its large homogeneously broadened gain profile, and attractive spectral position, coupled with the demonstrated energy storage capacity of the  $I_2$  ion-pair excited state manifold, make the band appear promising. Perhaps more importantly, the green band of  $I_2$  is only representative of a family of molecular transitions. Since few excimer gas laser systems exist (and particularly in the visible), one is encouraged to search for similar continua in the other homonuclear halogens  $Br_2$ ,  $Cl_2$ , and  $F_2$ .

The authors wish to thank S. B. Hutchison for constructing portions of the *e*-beam diode and developing the anode fabrication technique and M. L. Dlabal for initial spontaneous emission measurements on the green I<sub>2</sub> band. Also, the excellent technical assistance of Y. Moroz, A. B. Wilson, K. Kuehl, F. Ore, C. Henderson, L. McWhorter, D. Watterson, and K. Flessner is appreciated. The support of this work by the Office of Naval Research (M. White) under contract N00014-82-K-0209 is gratefully acknowledged.

- <sup>1</sup>R. S. Mulliken, J. Chem. Phys. 55, 288 (1971).
- <sup>2</sup>J. Tellinghuisen, Chem. Phys. Lett. 49, 485 (1977).
- <sup>1</sup>A. L. Guy, K. S. Viswanathan, A. Sur, and J. Tellinghuisen, Chem. Phys. Lett. **73**, 582 (1980).
- <sup>4</sup>R. S. Bradford, Jr., E. R. Ault, and M.L. Bhaumik, Appl. Phys. Lett. 27, 546 (1975).
- <sup>1</sup>J. J. Ewing and C. A. Brau, Appl. Phys. Lett. 27, 557 (1975).
- <sup>6</sup>A. K. Hays, J. M. Hoffman, and G. C. Tisone, Chem. Phys. Lett. **39**, 353 (1976).
- <sup>7</sup>D. J. Eckstrom, B. E. Perry, and K. Y. Tang, J. Appl. Phys. **50**, 3068 (1979).
- <sup>8</sup>N. G. Basov, J. S. Datskevich, V. S. Zuev, L. D. Mikheev, A. V. Startsev, and A. P. Skirokikh, Sov. J. Quantum Electron. 7, 352 (1977) [Kvant. Electron. (Moscow) 4, 638 (1977)].
- <sup>9</sup>L. D. Mikheev, A. P. Shirokikh, A. V. Startsev, and V. S. Zuev, Opt. Commun. 26, 237 (1978).
- <sup>10</sup>N. G. Basov, V. S. Zuev, L. D. Mikheev, and Yu. Yu. Stoilov, Izv. Akad. Nauk SSSR Ser. Fiz. 44, 1516 (1980).
- <sup>11</sup>M. J. Shaw, C. B. Edwards, F. O'Neill, C. Fotakis, and R. J. Donovan, Appl. Phys. Lett. 37, 346 (1980).
- <sup>12</sup>M. V. McCusker, in *Excimer Lasers*, edited by C. K. Rhodes (Springer, Berlin, 1979), pp. 67–86.
- <sup>11</sup>V. N. Baboshin, L. D. Mikheev, A. B. Pavlov, V. P. Fokanov, M. A. Khodarkovskii, and A. P. Shirokikh, Sov. J. Quantum Electron. 11, 683 (1981) [Kvant. Electron. (Moscow) 8, 1138 (1981)].
- <sup>14</sup>H. Hemmati and G. J. Collins, Chem. Phys. Lett. 67, 5 (1979), and Refs. 8 and 9 cited therein; Chem. Phys. Lett. 75, 488 (1980).
- <sup>15</sup>E. Zamir, D. L. Huestis, H. H. Nakano, R. M. Hill, and D. C. Lorents, IEEE J. Quantum Electron. QE-15, 281 (1979).
- <sup>16</sup>M. C. Sauer, Jr., W. A. Mulac, R. Cooper, and F. Grieser, J. Chem. Phys. 64, 4587 (1976).

Appl. Phys. Lett., Vol. 43, No. 6, 15 September 1983

K. P. Killeen and J. G. Eden