High-power broadly tunable picosecond IR laser system for use in nonlinear spectroscopic applications

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We developed a high-power tunable picosecond IR laser system suitable for nonlinear spectroscopic investigations. We employ a Ti:sapphire regenerative amplifier that produces 1.9-ps, 17-cm⁻¹, 800-nm pulses at 1 kHz as a pump source. White-light generation in ethylene glycol and optical parametric amplification in potassium titanyl phosphate are used to produce the IR pulses. The tuning range extends from 2.4 to 3.8 μm in the idler and 1.0 to 1.2 μm in the signal. A total efficiency (signal plus idler) as high as 20% was achieved. The spatial, spectral, and temporal characteristics of the IR beam are presented along with vibrational sum-frequency spectra from a glass/octadecyltrichlorosilane/air interface. © 1996 Optical Society of America.

1. INTRODUCTION

The study of interfaces buried between two condensed phases has grown rapidly in the past decade. Nonlinear-optical methods such as second-harmonic generation and sum-frequency generation (SFG) are techniques that have made significant contributions in this area. For the case of IR–visible SFG, one can achieve both interfacial and molecular selectivity and thus study an interface on a molecular level.1–6 Crucial to the development of these nonlinear optical methods has been the development of nanosecond tunable IR laser systems necessary to perform SFG experiments. The development of picosecond higher-peak-power tunable IR laser systems has been somewhat slower. One reason is that the IR pulses must have both high peak power and a narrow spectral bandwidth so that different vibrational modes can be resolved. Systems producing nanosecond pulses have narrow bandwidths but low peak power, whereas systems producing femtosecond pulses have high peak power but large bandwidths. We developed a high-peak-power tunable IR laser system that is optimized with respect to both of these limitations. It is based on a Ti:sapphire regenerative amplifier that produces 1.9-ps, 17-cm⁻¹, 800-nm pulses at a repetition rate of 1 kHz. The conversion of the 800-nm pulses into tunable high-power IR pulses is achieved through optical parametric amplification of a white-light continuum.

Parametric processes such as optical parametric generation and optical parametric amplification (OPA) in nonlinear crystals have been used extensively to generate tunable IR pulses.7,8 In each of these processes a photon pair is created at the expense of a pump photon with the wavelengths governed by the phase-matching conditions in the nonlinear crystal and the wavelength of the pump photon. By convention, the shorter-wavelength component is termed the signal and the longer-wavelength component termed the idler. The majority of these systems are based on Nd:YAG pump lasers and produce pulses with durations from tens of picoseconds to nanoseconds and bandwidths from several wave numbers to a few tenths of a wave number.9–12 More recently, regeneratively amplified Ti:sapphire lasers have been used to produce subpicosecond tunable IR pulses with bandwidths of the order of 100 wave numbers.13–15 In fact there are some commercially available systems that produce tunable femtosecond IR pulses. These systems, however, are not particularly well suited to nonlinear spectroscopic applications owing to the large bandwidths and the low pulse energies produced. Further, the majority of these systems use β-barium borate or lithium triborate for the nonlinear crystal, which limits the tuning range to 2.4 μm. We employ a Ti:sapphire regenerative amplifier that produces 1.9-ps pulses at 800 nm and a bandwidth of 17 cm⁻¹ as our pump source. The IR generation is based on white-light-continuum generation in ethylene glycol and OPA in two independently pumped potassium titanyl phosphate (KTP) crystals. This technique has been developed in the femtosecond regime11,16–18 but has not been exploited in the smaller-bandwidth picosecond regime. White-light generation was chosen to provide an intense seed covering the entire signal wavelength region, 1–1.6 μm. Our first attempt at IR generation involved seeding the first KTP crystal with the entire IR portion of the continuum from 1 μm to 1.6 μm. This, however, produced signal and idler bandwidths in excess of 100 cm⁻¹, with the large bandwidth being a result of the large acceptance bandwidth of KTP in the 2.6–3.8 μm region. The bandwidth can be narrowed by supplying the KTP crystal with a seed that is narrower than the acceptance
bandwidth of the crystal. We accomplished this by using a grating to disperse the continuum spatially before seeding the first KTP crystal. With the use of the grating the portion of the IR continuum that actually seeds the first crystal spans approximately 7 cm\(^{-1}\). Narrowing the output pulses by narrowing the seed pulses before amplification has also been used in the femtosecond regime.\(^{11,10}\) KTP was selected as the nonlinear crystal because of the desired tuning range, 1–4 μm, its figure of merit, which is proportional to the conversion efficiency, and its availability. Further, type II (ordinary idler, extraordinary signal, and ordinary pump) phase matching in the \(xz\) plane of the KTP crystals was chosen because of a higher \(d_{eff}\) and larger tuning range for an 800-nm pump. We use two amplification stages mainly because higher energies can be achieved but also because a second crystal angle tuned in the opposite direction has the effect of compensating for the beam displacement from the first crystal.

2. EXPERIMENTAL LAYOUT

The Ti:sapphire regenerative amplifier laser system consists of a Ti:sapphire oscillator (Coherent Mira-900 basic) pumped by an Ar\(^+\) ion laser (Coherent Innova-310), a stretcher–compressor (Quantronix 4822), and a Ti:sapphire regenerative amplifier (Quantronix 4810) pumped by the second harmonic of a Nd:YLF laser (Quantronix 527 DP-H). The output of the laser system (720 μJ, 800 nm) is first split with a partially reflecting mirror (M1) that sends 460 μJ to the optics for nonlinear conversion and leaves 260 μJ for the SFG experiments. Figure 1 depicts the optical setup used to convert the 800-nm photons into IR photons. This 460 μJ is split into three pump lines with two 50% reflecting mirrors, M2 and M3. The three lines are used for pumping the white-light generator WL and the two KTP (5 mm × 5 mm × 5 mm) crystals OPA1 and OPA2. For each line a polarizer/half-wave-plate combination is used to select the polarization and power in that line. The white-light pump passes through lens L4, which focuses 110 μJ of the 800-nm light into the ethylene glycol. The ethylene glycol flows through a 10 mm × 10 mm × 50 mm quartz cell (WL) that limits the effect of bubble formation on the white light generated by boiling of the ethylene glycol. The bubble formation is further minimized by cooling the ethylene glycol to 16 °C. The white light generated then passes through filter F1 (RG-850), which transmits only the IR portion of the continuum and blocks the visible and the remaining 800-nm pump. The IR portion of the continuum is brought to focus at the position of OPA1 by lens L3. The bandwidth of the seed-supplied OPA1 is narrowed by grating G1 (600 g/mm, blazed at 750 nm). The white-light seed, approximately 7-cm\(^{-1}\) bandwidth, is then amplified in OPA1 with 120 μJ of the 800-nm light telescoped to a beam diameter, 2\(\omega_0\), of 355 μm (64 GW/cm\(^2\)). The IR seed and the 800-nm pump are combined with dichroic mirror M4, which reflects 800 nm and transmits 1–1.6-μm light. The temporal overlap of the seed and the pump in OPA1 is controlled by delay line D1, which incorporates a hollow retroreflector. After this first stage of amplification, signal energies as high as 10 μJ and idler energies as high as 2.5 μJ are obtained. Since higher energies are desired, another amplification stage (OPA2) located 560 mm from OPA1 is employed. Filter F2 (80% transmission from 2.2–4.1 μm) is used to block the amplified signal and the 800-nm pump and to transmit the idler for amplification in OPA2. The idler is used for the seed in OPA2 because higher energies and better spatial beam profiles are attained in this configuration (see the following discussion). The idler from OPA1 is focused to a beam diameter, 2\(\omega_0\), of 760 μm at the position of OPA2 by a CaF\(_2\) lens, L2. The idler generated in OPA1 is amplified in OPA2 with 160 μJ of the 800-nm light, which is telescoped to a beam diameter, 2\(\omega_0\), of 598 μm (30 GW/cm\(^2\)). The idler and the 800-nm pump are combined with the use of dichroic mirror M5 that reflects...
800 nm and transmits 1.5–4.0 µm. The temporal overlap of the seed and the pump in OPA2 is controlled by a second delay line D2. One of two filters is used in the position of F3: the first for the measurement of the idler energy (80% transmission from 2.2–4.1 µm) and the other for the measurement of the signal energy (RG-1000). The IR output from OPA2 is then collimated with lens L1 for the SFG experiments. Except for the CaF2 lenses, L1 and L2, all other lenses are broadband antireflection coated BK7, and all mirrors not mentioned are either broadband near-IR high reflectors for the 800 nm or protected gold for the IR.

The pump laser system and the IR laser system were characterized in terms of power, wavelength, spatial bandwidth, temporal pulse width, and spatial beam quality. Wavelengths were measured with a 0.3-m crossed Czerny–Turner monochromator with a 300 grooves/mm grating blazed for 3 µm and an indium antimonide (InSb) detector. Bandwidths (full width at half-maximum intensity [FWHM]) were measured with the monochromator or with a grating (1200 or 1800 grooves/mm) in combination with a one-dimensional CCD array (UniData BP2048). The monochromator was used in different orders of diffraction depending on the wavelength of light being examined. Specifically, the pump-beam, the signal-beam, and the idler-beam spectra were obtained in fourth-, third-, and first-order diffraction, respectively. Further, the resolution of the monochromator was approximately 3 cm⁻¹ at the pump and the signal wavelengths and approximately 1 cm⁻¹ at the idler wavelengths. The resolution of the grating/CCD combination was approximately 4 cm⁻¹ and was mainly limited by the beam diameter. Powers were measured with a pyroelectric joulemeter (Molecron J3-09), and temporal pulse profiles were characterized by autocorrelations (Inrad 5-14B) and/or cross correlations (FWHM).

Spatial beam quality was characterized by knife-edge measurements across the transverse profile of the beams. Assuming an approximately Gaussian transverse spatial profile, the distance between knife-edge positions passing 16% and 84% of the total beam energy was used to approximate the Gaussian beam radius ω, which has the standard definition of the half-width at 1/e of maximum for the field amplitude, or the half-width at 1/e² of maximum for the intensity. This is in turn related to the FWHM for the intensity by FWHM = (2 ln 2)¹/²ω = 1.177ω. The measured pump- and idler-beam sizes were then used to estimate the M² parameter (i.e., times diffraction limited) for the pump and the idler beams.

3. Ti:SAPPHIRE LASER CHARACTERIZATION

A titanium:sapphire (Ti:sapphire) regenerative amplifier system based on the chirped pulse amplification technique is used as the pump source and thus is a primary determinant of the spectral, the spatial, and the temporal characteristics of the output of the tunable IR system. It is therefore quite important that those same spectral, spatial, and temporal characteristics of the regenerative amplifier pump source be well understood.

The grating-based stretcher employed in the chirped pulse amplification technique is a particularly convenient location for manipulation of the spectral, and therefore temporal, properties of the amplifier seed pulses. At the midpoint of the stretcher the various wavelength components are spatially dispersed, allowing the possibility of spectral filtering. Other researchers have used sophisticated amplitude and phase masks specifically to tailor the amplified pulses in an attempt to control coherently the optically driven processes. For the present application, all that is required is a simple reduction of the bandwidth. This is accomplished by the insertion of an aperture at the midpoint of the stretcher, limiting the seed bandwidth passed to the amplifier by the stretcher. This technique must be used with caution, however. The stretching process relies on the temporal chirping of a wide spectral bandwidth. If this bandwidth is narrowed, the corresponding temporal width of the chirped pulse also becomes narrower, leading to higher peak powers in the amplifier. (In contrast to the transform-limited case, in which pulse width and bandwidth are inversely related, pulse width and bandwidth are linearly related in the highly chirped case for a given amount of dispersion.) The grating pulse stretcher must be modified to provide greater dispersion in order to stretch these spectrally narrower pulses to a safely amplified pulse width. This can be most easily accomplished by insertion of a different grating with a higher groove density and therefore greater pulse stretching per unit bandwidth.

The use of an aperture at the midpoint of the pulse stretcher, although simple to implement, requires care in analyzing pulse-width and bandwidth data of the amplified pulses. For our purposes, output pulses with bandwidths of 20 cm⁻¹ or less are desired from seed pulses that are ~100 cm⁻¹ FWHM. Over the central 20 cm⁻¹ of the seed bandwidth, the spectral intensity distribution is relatively flat, leading to an essentially rectangular spectrum for the amplifier output. Such a spectrum yields pulses with an intensity profile \( I(t) \propto (\sin^2 t)^2 \) = sinc² t. This is in turn related to the FWHM for the intensity by FWHM = (2 ln 2)¹/²ω = 1.177ω. The measured pump- and idler-beam sizes were then used to estimate the M² parameter (i.e., times diffraction limited) for the pump and the idler beams.
FIG. 2. Autocorrelation and spectrum of the pump (800-nm) beam: (a) autocorrelation and (b) the spectrum obtained with a monochromator (see text). The solid curves are a guide for the eye.

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The spatial quality of the beam was measured by knife-edge measurements. The compressed output of the regenerative amplifier was thus determined to be 1.2 times diffraction limited.

4. IR SYSTEM CHARACTERIZATION

Figure 3 shows the experimental and theoretical tuning curves for type II phase matching in KTP. Theoretical tuning curves were generated by imposing type II (o-ray pump, e-ray signal, and o-ray idler) collinear phase matching in the \(xz\) plane (\(\phi = 0^\circ\)) of the KTP crystal.26

The Sellmeier coefficients for KTP used are from Vanherzeele.27 The extent of the experimental tuning range is governed by several limitations. First is the transparency region of the KTP crystals, which is 350 nm to 4.5 \(\mu\)m. This is the ultimate limit in that light cannot be generated at wavelengths outside the transparency region. Next is the transmission region of the filters F2 and F3, which is 2.2 to 4.0 \(\mu\)m for each. With these filters, idler wavelengths above 4.0 \(\mu\)m or below 2.2 \(\mu\)m are not transmitted and thus are not available for experiments. The last limitation is the dichroic mirror M4, which reflects 800 nm and transmits 1.0 to 1.6 \(\mu\)m. This prevents us from generating light above 4.0 \(\mu\)m since wavelengths below 1.0 \(\mu\)m are necessary as the signal seed and are not transmitted. From this, one can see that, with the use of different filters and a different dichroic mirror, idler wavelengths to 4.5 \(\mu\)m can be achieved.

SFG experiments require high peak intensities for both the visible/near-IR pulse and the tunable IR pulse. In addition, good spatial and temporal overlap must be achieved. These requirements demand good spectral/temporal pulse characteristics as well as good spatial beam characteristics. The spectral, the temporal, and the spatial characteristics of pump, signal, and idler were all measured and analyzed with these considerations in mind.

The spectral and the temporal characteristics of the signal and the idler pulses produced by the optical para...
metric amplifiers were measured and compared with those of the pump laser system. Autocorrelations and spectra of both the pump at 800 nm and the signal at 1.09 μm are shown in Fig. 4. The autocorrelation of the pump beam has a FWHM of 2.5 ps, as stated previously, and exhibits the typical shape of an autocorrelation of a sinc² pulse, including slight sidelobes. The corresponding spectrum is flatter on the top than typical spectra of mode-locked lasers, and it is a result of the roughly rectangular spectrum that should result from the hard-aperture spectral filter. The spectrum and the autocorrelation of the signal pulses look somewhat different. The signal spectrum is slightly wider than the pump spectrum, which is to be expected, and it looks much more Gaussian in nature than the pump spectrum. A rough calculation of the bandwidth expected to fall within the spatial profile of the pump beam in the first KTP crystal (209-μm FWHM) yields

\[ \Delta \nu \approx a \nu^2 \cos \beta \left( \frac{d}{D} \right) \]

\[ = \left( \frac{9166 \text{ cm}^{-1}}{6000 \text{ cm}^{-1}} \right)^2 \cos 21^\circ \frac{0.209 \text{ mm}}{400 \text{ mm}} = 6.8 \text{ cm}^{-1}, \]

where \( a \) is the groove spacing of the grating, \( d \) is the width of the pump beam (FWHM), \( D \) is the distance from the grating to the first KTP crystal, and \( \beta \) is the angle of diffraction from the grating. The 17 cm⁻¹ pump bandwidth yields a lower limit for the signal, and the measured signal bandwidth (18.5 cm⁻¹) is slightly higher than this lower limit. The FWHM of the autocorrelation is significantly narrower than the pump autocorrelation (1.8 versus 2.5 ps), and the sidelobes are much reduced relative to the pump. This would suggest that the functional forms of the signal pulse and the spectrum are quite different from their pump counterparts. An autocorrelation of 1.8 ps implies a pulse width of approximately 1.3 ps (dividing by 1.44) and a time–bandwidth product of 0.72. This is less than the transform limit for sinc² pulses and approximately twice the transform limit for Gaussian or sech² pulses.

A similar analysis was performed for the idler. However, because of the wavelengths involved (2.5–3.8 μm), autocorrelations could not be done with the optics and the nonlinear crystals on hand. Instead, the idler pulses were cross correlated with the pump laser pulses in a KTP crystal by SFG, and a pulse width for the idler was extracted by treating the cross correlation as a convolution of two Gaussians (\( \Delta t_{\text{idler}}^2 = \Delta t_{\text{idler}}^2 + \Delta t_{\text{pump}}^2 \)). Idler spectra for three different idler wavelengths are shown in Fig. 5, and cross correlation measurements for the same three idler wavelengths are shown in Fig. 6. At 3.0 and 3.6 μm the spectra look quite Gaussian like, and they are nearly the same bandwidth as the pump pulse at 18 and 16 cm⁻¹, respectively. The cross correlations are also well approximated by Gaussians at these two wavelengths with FWHM of 2.3 ps at both wavelengths. Deconvolution of the pump pulse width of 1.9 ps yields idler pulse widths of 1.3 ps and time–bandwidth products of 0.70 and
Likewise, the cross correlations are similar except for a relative enhancement of the long decaying portion with respect to the central peak, which seems to occur on similar time scales. One possible explanation for this behavior is that a spectrally broad short pulse may coherently excite an ensemble of molecules. These molecules then undergo some sort of free-induction decay, giving rise to the decaying tails observed in the cross correlations and the structure observed in the spectra. As the idler-pulse bandwidth is decreased, the size of the excited ensemble diminishes, thereby decreasing the amplitude of the decaying portion of the cross correlation and the intensity of the spectral structure. This phenomena was not investigated in great detail.

The spatial characteristics of the idler beam were measured very carefully. The final idler beam was focused by a +125-mm focal-length CaF$_2$ lens, and a series of knife-edge measurements were performed. In this way a set of values for $\omega$ versus propagation distance was generated and fit to the equation $21$ that is shown along with the data in Fig. 8. The fit yields a value of $M^2$ of 3.45, implying that the idler beam is approximately 3.5 times diffraction limited.

Figure 9(a) shows the experimental power curves for

Fig. 5. Spectra of the idler beam at various wavelengths: (a) 2.6 $\mu$m, (b) 3.0 $\mu$m, and (c) 3.6 $\mu$m. Solid curves are a guide for the eye.

0.62 at 3.0 and 3.6 $\mu$m, respectively. As with the signal pulses, these are smaller than for $\text{sinc}^2$ pulses but are approximately twice transform limited for Gaussian or $\text{sech}^2$ pulses. Note also that there are no readily apparent sidelobes in the cross correlations.

Some rather interesting behavior occurs in the region near 2.6 $\mu$m. Water vapor is known to absorb in this region and causes significant disruption of the spatial and the temporal characteristics of the idler beam as it propagates through several feet of room air before being measured. The spectrum shown in the top panel of Fig. 5 shows a relatively narrow (ca. 10-cm$^{-1}$ FWHM) central peak riding on an irregular background extending over 80 cm$^{-1}$. The corresponding cross correlation is shown in the top panel of Fig. 6. Note the asymmetry and the leading peak (ca. 5-ps FWHM) followed by a relatively long decay (tens of picoseconds). Interestingly, these temporal and spectral profiles are actually a considerable improvement over the behavior observed before the pump bandwidth was narrowed from 21 cm$^{-1}$ to 17 cm$^{-1}$ (idler bandwidths decreased from 25–30 cm$^{-1}$ to 16–18 cm$^{-1}$). Two corresponding sets of measurements are shown in Fig. 7. The spectra exhibit similarly placed features, but the side peaks and background are larger relative to the main peak in the case of the spectrally broader pump.
at 2.7 μm, which is caused by absorption of H₂O vapor in air, as discussed above (note that there is no corresponding dip in the signal energy). The second feature is a peak in both the idler and the signal energies at 3.6 and 1.04 μm, respectively, which is caused by three interacting effects. First, the intensity of the seed from the white light is increasing dramatically as the signal approaches 800 nm. Second, the transmission of the dichroic mirror M4 is decreasing as the signal wavelength approaches 1.0 μm. Finally, the value of $d_{\text{eff}}$ for type II phase matching in the $xz$ plane decreases as one goes to smaller angles according to $d_{\text{eff}} = d_{24} \sin \theta$, and thus the conversion efficiency decreases. The competing effect is the creation of a peak in both the idler and the signal energies. Figure 9(b) shows the photon conversion efficiency for both the signal and the idler corresponding to the energies in Fig. 9(a). The photon conversion efficiency was calculated with the energies from Fig. 9(a) and the total 800-nm pump energy (white light plus OPA1 plus OPA2 equal to 390 μJ) for each wavelength. Thus the reported efficiencies are an overall efficiency of the entire system and not an individual efficiency for the OPA process in a KTP crystal. From this graph one can see that a total efficiency (signal plus idler) as high as 20% is achieved.

5. SUM-FREQUENCY-GENERATION SPECTRA

The glass/octadecyltrichlorosilane (OTS)/air interface has been extensively studied with SFG,28–30 and we use it here as a comparative test for our laser system. Guyot-Sionnest et al. have reported three peaks in the C–H spectral region, all of which can be assigned to C–H stretches of the terminal CH₃ group. The first peak lies at 2878 cm⁻¹ with a bandwidth of 7 cm⁻¹ and has been assigned to the symmetric (s) stretch. A second peak at 2942 cm⁻¹ with a bandwidth of 9 cm⁻¹ has been attributed to the Fermi resonance between the s stretch and the overtone of the C–H bending mode. A third peak at 2964 cm⁻¹ has been assigned to the two nearly degenerate ($d_s$ at 2969 cm⁻¹ and $d_d$ at 2958 cm⁻¹) asymmetric (d) stretches of the terminal CH₃ group with bandwidths of 9 cm⁻¹ each. Further, under $P_{d_s} P_{\text{vis}} P_{\text{ir}}$ polarization conditions the asymmetric (d) stretch and the symmetric (s) stretch were present with the Fermi resonance obscured by the asymmetric (d) stretch. Under $S_{d_s} S_{\text{vis}} P_{\text{ir}}$ polarization conditions, only the symmetric (s) and Fermi resonance were present.

Figure 10 shows the SFG spectra of a saturated monolayer of OTS on glass in air under different polarization conditions obtained with our laser system. Each spectrum was generated by averaging 800 laser shots at each point in 0.002-μm increments. With the 1-kHz repetition rate of our system this was easily achieved in scans that lasted approximately 40 min. Figure 10(a) shows the SFG spectrum under $P_{d_s} P_{\text{vis}} P_{\text{ir}}$ polarization conditions from our laser system. The asymmetric (d) stretch is the dominant peak present and occurs at 2964 cm⁻¹ with a width of 21 cm⁻¹. One can also see a small peak at 2879 cm⁻¹ that is due to the symmetric (s) stretch and a shoulder at 2943 cm⁻¹ that is due to the Fermi reso-
Fig. 9. Pulse energy and photon conversion efficiency for signal and idler beams: (a) output energy and (b) photon conversion efficiency. Solid curves are a guide to the eye.

nance. This is what is expected since the $P_{slg}P_{vis}P_{ir}$ polarization condition contains all the $\chi$ elements. Figure 10(b) shows the SFG spectrum under $S_{slg}S_{vis}P_{ir}$ polarization conditions. For this case the symmetric (s) stretch occurring at 2879 cm$^{-1}$ with a width of 19 cm$^{-1}$ and the Fermi resonance occurring at 2943 cm$^{-1}$ with a width of 23 cm$^{-1}$ are the dominant peaks present. The bandwidths of the individual peaks can be found by deconvolving the SFG widths with the 17-cm$^{-1}$ Gaussian laser bandwidth by assuming a Lorentzian vibration width. From this we get bandwidths of 4 cm$^{-1}$, 10 cm$^{-1}$, and 7 cm$^{-1}$ for the symmetric (s), Fermi resonance, and asymmetric (d) modes, respectively.

Fig. 10. Sum-frequency spectra of a monolayer of OTS on glass in air under different polarization conditions: (a) $P_{slg}P_{vis}P_{ir}$ and (b) $S_{slg}S_{vis}P_{ir}$. Solid curves are a guide to the eye.

6. CONCLUSION
We have demonstrated that intense broadly tunable IR pulses are produced by this system. Further, the spatial, the spectral, and the temporal qualities of the IR beam make it an attractive source for nonlinear spectroscopic applications. The moderately narrow bandwidths and the broad tuning range will allow for the study of a large number of condensed phase interfacial molecules by means of their vibrational modes. The 16–18 cm$^{-1}$ bandwidths reported here are well suited for studies of this type in which the bandwidths of the molecular vibrations are relatively large. Our tuning range spans and IR region in which molecules containing OH, NH, and CH bonds can be studied. We have demonstrated the ability to obtain SFG spectra in the CH stretching region with studies on the glass/OTS/air interface. We have also conducted preliminary SFG experiments on alcohol/air and n-alkane/air interfaces with great success in the CH stretch region. Water absorption of the IR beam as it propagates through air interfered with our ability to obtain spectra in the OH region from the glass/OTS/air and the alcohol/air interfaces. We are in the process of eliminating this limitation by removing the water in the beam path and further reducing the IR bandwidth. The practical limit of the bandwidths attainable by modifying the regenerative amplifier system is 5–10 cm$^{-1}$, which would improve the resolution somewhat and virtually eliminate the water absorption effects. We are also in the process of extending the tuning range beyond 4.0 $\mu$m so that we
can study a wider variety of molecules. We will employ other nonlinear crystals such as AgGaS$_2$ and KNbO$_3$ to achieve tunability beyond 4.0 $\mu$m.

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