Infrared electric field sampled frequency comb spectroscopy

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Probing matter with light in the mid-infrared provides unique insight into molecular composition, structure, and function with high sensitivity. However, laser spectroscopy in this spectral region lacks the broadband or tunable light sources and efficient detectors available in the visible or near-infrared. We overcome these challenges with an approach that unites a compact source of phase-stable, single-cycle, mid-infrared pulses with room temperature electric field–resolved detection at video rates. The ultrashort pulses correspond to laser frequency combs that span 3 to 27 μm (370 to 3333 cm⁻¹), and are measured with dynamic range of >10⁶ and spectral resolution as high as 0.003 cm⁻¹. We highlight the brightness and coherence of our apparatus with gas-, liquid-, and solid-phase spectroscopy that extends over spectral bandwidths comparable to thermal or infrared synchrotron sources. This unique combination enables powerful avenues for rapid detection of biological, chemical, and physical properties of matter with molecular specificity.

INTRODUCTION

The resonant interaction of infrared light with matter provides rich structural and functional information in complex biological (1, 2), chemical (3–5), and physical systems (6, 7). Notable examples include studying conformational changes in proteins (8, 9), tracking chemical and ultrafast dynamics (10–13), measuring vibrational spectra with high resolution (14, 15), and exploring the origins of chirality (16, 17). Light sources with full mid-infrared (MIR; 3 to 25 μm) coverage that simultaneously address multiple molecular vibrational modes with user-specified spectral and temporal resolution are beneficial for all these applications, as well as the growing field of infrared spectro-imaging (18–20). Owing to the disparate and demanding criteria (e.g., brightness, bandwidth, and resolution), existing technologies such as thermal sources, synchrotrons, tunable lasers [quantum cascade and semiconductor lasers (21, 22)], and parametric oscillators (23, 24) often lack the desired properties or become increasingly complex. Adding to this complexity, infrared photodetection requires cryogenic cooling for low-noise operation and exhibits poor quantum efficiencies above 12 μm. In this work, we introduce a simple yet general framework for the generation of coherent MIR light spanning 3 to 27 μm and direct electric field detection of the corresponding near–single-cycle pulses. Complete infrared electric field readout at video rates using room temperature near-infrared (NIR) photodetectors resolves millions of phase-locked frequency comb modes that comprise the infrared spectrum. We highlight these advantages in precision spectroscopy of ammonia (25, 26) and carbon dioxide (27) with simultaneous spectral coverage and resolution not achieved with other laser sources. Moreover, with brightness and scan-free acquisition rates 100 times greater than found in conventional Fourier transform spectrometers, our approach provides increased sensitivity and speed in recording the full infrared spectra of biological and molecular compounds in the condensed phase.

EXPERIMENTAL SYSTEM

In our experimental system, intra-pulse difference-frequency generation (DFG) in quadratic nonlinear media provides a simple and robust source of ultrashort infrared pulses with an intrinsically stable carrier-envelope phase (CEP) (28, 29). The corresponding multi-octave instantaneous bandwidth allows us to probe multiple ro-vibrational excitations from 500 to 2500 cm⁻¹. Leveraging the CEP stability, we implement dual frequency comb electro-optic sampling (EOS) (Fig. 1A), which directly measures the MIR electric field and removes the need for cryogenically cooled photodetectors. In dual-comb EOS, the electric field of the MIR pulse train (with repetition rate fᵢ + Δfᵢ) is sampled by sum-frequency (SF) mixing with an NIR few-cycle pulse train (with repetition rate fᵢ). The nonlinear interaction results in an infrared field–dependent polarization rotation of the sampling pulse that is resolved via ellipsometry using room temperature InGaAs photodetectors (Fig. 1A) (30, 31). Because of the repetition rate offset, the sampling pulse scans across the MIR electric field automatically and a complete electric field measurement is made every Δfᵢ⁻¹ ≈ 20 ms.

Dual-comb sampling of radiation (also known as asynchronous optical sampling) has been used in the terahertz domain (optical period, τᵢTHz ~ 1 ps) using >50-fs-long optical pulses (32, 33). However, its extension to the MIR has been limited by the required subcycle timing jitter and few-cycle sampling pulses (<10 fs). We overcome these challenges and measure a mutual timing jitter of 80 as between the two frequency combs, corresponding to <20-mrad residual phase noise for a 10-μm field. This exacting criterion is seen intuitively in the frequency domain, where the nonlinear sampling results in a new SF comb, with the entire MIR spectrum folded into each free spectral range (Fig. 1B) (34, 35). Thus, coherent multiheterodyne beating occurs in the NIR, with the sampling pulse serving as a local oscillator comb.

Applications of EOS in the MIR, including sensitive spectroscopy, have been shown previously (30, 31, 36–38). However, previous measurements in the MIR used a mechanical translation stage to vary the time delay, restricting the resolution and acquisition time to...

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electric field (with repetition rate such that it is contained within every Nyquist zone (sampled to Wollaston prism (WP), and balanced photodetectors (BPD) yields a signal, filtered with a bandpass filter (BPF). Ellipsometry using a quarter waveplate (QWP), (GaSe) crystal, with a time-domain signal-to-noise ratio (SNR) <10-fs NIR pulse, the MIR electric field is sampled in a gallium selenide spectral resolution (0.003 cm−1). The multi-octave instantaneous bandwidth we achieve is unparalleled when compared to other frequency comb sources (24, 42, 43) and enables simultaneous probing of multiple vibrational bands without the need for tuning. Figure 2B shows the absorption signatures of important compounds for atmospheric chemistry such as methane, ethane, carbon dioxide, and water vapor in both the functional group (1500 to 4000 cm−1) and molecular fingerprint regions (500 to 1500 cm−1). In particular, the vibrational spectra of the P, Q, and R branches of the O–C–O bending vibration of CO2 around 15 μm are measured using the OP-GaP MIR comb over a 2-m path length (Fig. 2C). Comparison with a model (44) shows that quantitative results are not limited by our dual-comb EOS detection. The intensity and phase of the absorption lines, which are pressure-broadened to 5 GHz, are resolved with individual comb teeth of the frequency comb (Fig. 2C). The comb teeth are stabilized to a 10-kHz level, which is not a fundamental limit, but represents the instrument line shape in our present spectrometer (45). With time-domain apodization of the electric field signal, 500-MHz resolution data are also shown, which exhibit higher SNR by a factor of √5 (Fig. 2D).

Ammonia is an important compound used in the agricultural and biopharmaceutical industries (26) and fundamental scientific studies including astrochemistry (7). In Fig. 3, we present our measurement of its ν3 vibration (A1, symmetric bend) across 770 to 1165 cm−1 (8.6 to 13 μm), where 118,000 comb teeth are individually resolved. Similar simultaneous spectral coverage has only previously been achieved with thermal sources (46) and scanning Fourier transform spectrometers, but here, we additionally leverage the intrinsic frequency accuracy and resolution that the frequency comb provides. Excellent agreement with the HITRAN model (44) is seen across the entire spectrum. We observe the splitting in the Q branch due to pyramidal inversion of the NH3 molecule, where the nitrogen atom undergoes room temperature quantum tunneling through the potential barrier formed by the H3 plane (47). The combination of the broadband high-resolution spectroscopy and room temperature photodetection renders our system powerful for studying quantum chemistry (48), interstellar composition (25), and mapping individual crystalline domains in nanoscale materials (49).

An advantage of our approach is that it provides full and simultaneous MIR spectral coverage at a user-defined resolution. Here, we use numerical apodization to yield resolution that is more appropriate for broadband condensed phase samples, but high-repetition rate frequency combs (50) or dynamic repetition rate switching (51) can be used to tailor the resolution and further increase data acquisition rates. We demonstrate these advantages with broadband liquid-phase absorption signatures of an R–(–)-carvone, a prototypical chiral molecule,
and the amide vibrations in a monoclonal antibody (NISTmAb), a reference for the biopharmaceutical industries (Fig. 4). In Fig. 4A, liquid-phase absorption spectra acquired over two octaves from 500 to 2500 cm$^{-1}$ are shown, with the $n_2$ (O–H–O bend) vibration at 1650 cm$^{-1}$ being saturated. In Fig. 4B, the absorption signature of a dried NISTmAb film is shown from the 1200- to 1800-cm$^{-1}$ region that identifies the amide I, II, and III bands. The amide bands in proteins are used to determine the folding, unfolding, and aggregating mechanisms (8, 52, 53). The center frequencies of the amide I (1636 cm$^{-1}$) and amide II bands (1549 cm$^{-1}$) indicate a β-sheet structure for the protein, agreeing with previous studies (8, 54).

CONCLUSION

In conclusion, we presented a comprehensive framework for MIR laser spectroscopy that unified the highly desired qualities of phase-stable single-cycle temporal waveforms and ultrabroadband spectral coverage with room temperature, video rate, electric field detection with bandwidth extending from 370 to 3333 cm$^{-1}$. Inheriting the robustness and simplicity from the NIR pulses derived from erbium fiber lasers, the compact source provides a benchtop (<0.5 m$^2$) footprint for infrared spectroscopy in the molecular fingerprint region and beyond. Quantitative high-resolution spectroscopy is demonstrated with electric field sampling in a spectral region beyond the reach of high-speed HgCdTe detectors. In addition to probing narrow linewidth transitions in gas phase, we probe wave number–scale transitions in liquid- and solid-state materials across 500 to 2500 cm$^{-1}$ and capture both functional group and molecular fingerprint regions. Such broadband spectroscopy is critical for strongly coupled interactions between infrared light and condensed phase matter that enable applications such as quantum computation within the internal degrees of freedom in a molecule (55). When combined with novel imaging techniques such as infrared atomic force microscopy and scanning near-field optical microscopy, the system described here would enable nanoscale spectro-imaging of samples that currently require the bandwidth and brightness of a synchrotron. Last, the near–single-cycle nature of our MIR light source can serve as a robust seed for parametric amplifiers in strong-field physics (56).

MATERIALS AND METHODS

Phase locking

We used two self-referenced Er:fiber lasers with nominal repetition rate, $f_r = 100$ MHz, to generate the MIR light and sampling pulse for dual-comb EOS. The lasers were made mutually coherent by optical phase locking each one to a cavity-stabilized continuous-wave laser at 1550 nm. The absolute linewidths of the frequency combs were tied to...
the 10-kHz linewidth of the continuous-wave laser. The line-position accuracy of the comb modes is $10^{-12}$, which is given by the precision with which we measured the lasers’ repetition rates. While appropriate for the spectroscopy shown here, these are not fundamental limits, and sub-hertz linewidths and uncertainties below $10^{-15}$ can be implemented if required. Moreover, while single-measurement resolution of 100 MHz ($0.003 \text{ cm}^{-1}$) is shown here, by stepping the repetition rates of the combs, the modes of the combs could be continuously swept across the full 100-MHz free-spectral range, thereby providing resolution down to the comb-tooth linewidth (57). Locking was achieved via feedback to the laser that was provided using a combination of an intra-cavity electro-optic modulator at a 1-MHz bandwidth and a piezo-electric transducer at a 10-Hz bandwidth. The integrated residual optical phase noise is $\sim 100$ mrad, corresponding
to a relative timing jitter of 83 as. In this work, $\Delta f_c$ is in the range of 40 to 50 Hz. However, we verified coherent averaging at $\Delta f_c = 15$ Hz, corresponding to a 1.5-fs timing resolution.

**Few-cycle NIR pulse synthesis**

A nonlinear amplification scheme in erbium-doped fiber, followed by spectral broadening in normal-dispersion highly nonlinear fiber, provides the bandwidth for a 10-fs NIR pulse that is compressed in bulk fused silica. Residual third-order dispersion was compensated using a pair of chirped mirrors. Frequency-resolved optical gating (FROG) was used to measure the pulse duration.

**MIR generation**

The few-cycle pulse was focused into a quadratic nonlinear crystal using an off-axis parabolic mirror ($f = 25$ mm) to drive intra-pulse DFG and produce MIR light. We generated light from 3- to 5.5-µm light in a 1-mm-thick PPLN with power ranging from 500 µW to 1.2 mW. In OP-GaP, a 1-mm-thick fanout crystal with grating periods in the range of 50 to 65 µm was used to generate broadband long-wave infrared light. The spectrum spanning 4 to 20 µm was generated in a 63-µm grating period and corresponded to 100 to 200 µW. The center frequency of the spectrum was determined by calculating its center of mass after deconvolving the spectrum using the response function. In a 500-µm-thick GaSe, type II phase matching was used, and the broadband spectrum provided approximately 2 µW across the entire band.

**EOS detection**

The SF generation (SFG) from the electro-optic crystal was filtered using a 25- or 50-nm bandpass filter around 1300 nm and analyzed using a broadband quarter waveplate followed by a high-extinction Wollaston prism. The light was focused onto a commercial InGaAs balanced photodetector (BPD). The differential photocurrent, $\Delta I / I \propto \Delta n \propto (d_{\text{eff}} n_0) E_{\text{MIR}}$, is proportional to the SFG-induced refractive index change (or, equivalently, nonlinear phase shift) (58). $d_{\text{eff}}$ is the frequency-dependent second-order susceptibility of the electro-optic crystal, $n_0$ is the refractive index of the NIR carrier frequency in GaSe, and $E_{\text{MIR}}$ is the MIR electric field. For the near-single-cycle OP-GaP MIR output, based on a focused beam waist of 100 µm² and a pulse duration of 30 fs with an average power of 100 µW, we estimated that the peak electric field in the GaSe crystal is $E_{\text{MIR}} = 15$ kV/cm. To trigger the dual-comb readout at the 50-Hz refresh rate, we used a nonlinear cross-correlation signal acquired via SFG between the two few-cycle NIR pulses in a 1-mm-thick barium borate crystal that was detected with a Si avalanche photodiode.

**EOS response function**

Although the temporal increment in the dual-comb EOS configuration is ≈5 fs ($\Delta f_c = 50$ Hz), the finite pulse duration of the sampling pulse (10 fs) imposes a signal response that must be removed by deconvolution to attain the true electric field waveform of the infrared radiation. As this measurement can be considered a convolution in time domain, the “true” electric field may be acquired by deconvolving the response function of EOS (59, 60). The electric field of the SFG is proportional to the envelope of the NIR sampling pulse, i.e., $E_{\text{SFG}} \propto d_{\text{eff}} \exp(i\Delta k z_{\text{TM}}) \times E_{\text{NIR}} E_{\text{MIR}}, \Delta k = \frac{1}{c} \left( n(\omega_{\text{MIR}}) \omega_{\text{MIR}} - n(\omega_{\text{NIR}}) \omega_{\text{NIR}} \right)$ is the phase mismatch in the SFG process, and $E_{\text{NIR}}$ is the sampling pulse envelope. By computing the phase mismatch and acquiring $E_{\text{NIR}}$ via FROG, a response function can be constructed (fig. S1). De-

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**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/6/eaaw8794/DC1

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**REFERENCES AND NOTES**

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