1-D imaging of rotation-vibration non-equilibrium from pure rotational ultrafast coherent anti-Stokes Raman scattering

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We present one-dimensional (1-D) imaging of rotation-vibration non-equilibrium measured by two-beam pure rotational hybrid femtosecond/picosecond coherent anti-Stokes Raman scattering (fs/ps CARS). Simultaneous measurements of the spatial distribution of molecular rotation-vibration non-equilibrium are critical for understanding molecular energy transfer in low temperature plasmas and hypersonic flows. However, non-equilibrium CARS thermometry until now was limited to point measurements. The red shift of rotational energy levels by vibrational excitation was used to determine the rotational and vibrational temperatures from 1-D images of the pure rotational spectrum. Vibrational temperatures up to 5500 K were detected in a CH₄/N₂ nanosecond-pulsed pin-to-pin plasma within 2 mm near the cathode. This approach enables study of non-equilibrium systems with 40 μm spatial resolution.

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Rotation-vibration non-equilibrium and energy transfer is a key area of study for low temperature plasmas that enhances chemical reactivity in applications such as materials synthesis [1], plasma catalysis [2], plasma assisted combustion [3], CO₂ dissociation [4], and CH₄ reforming [5]. Furthermore, rotation-vibration non-equilibrium is critical for understanding the aerodynamics and thermal loading of space vehicles re-entering Earth's atmosphere at hypersonic velocities [6]. In these applications, there exist spatial gradients of temperature due to energy transport across the boundary layer in hypersonic vehicles [6] or the cathode sheath layer in low temperature plasmas [7]. Therefore, it is of great interest to be able to conduct spatially resolved measurements of rotation-vibration non-equilibrium in these systems. Hybrid femtosecond/picosecond coherent anti-Stokes Raman scattering (fs/ps CARS) is a powerful tool to measure species concentration and temperature on the picosecond time scale and tens of microns of spatial resolution, while avoiding non-resonant contributions to the CARS spectrum from four-wave mixing [8–16]. This is achieved by generating the Raman coherence through excitation by a spectrally broad femtosecond laser which provides both the pump and Stokes photons. A separate time-delayed spectrally narrow ps pulse scatters off this coherence and generates the CARS signal beam. Since the pump/Stokes and probe beam do not overlap in time, the non-resonant contributions are avoided but many Raman transitions can be probed at once through the broadband excitation.

Typically, simultaneous measurements of rotation-vibration non-equilibrium of molecules like N₂ measured by CARS uses the vibrational Q-branch and its rotational structure to determine the rotational and vibrational temperatures [17]. However, the spectral resolution for the fs/ps CARS approach is determined by the spectral width of the ps probe pulse. To resolve the rotational structure, a spectral resolution less than 1 cm⁻¹ is required [18,19], which is a major challenge for many fs/ps CARS systems since they use the fs laser to generate ps probe pulses with ~2 to 3 cm⁻¹ resolution [13–15]. As a result of the challenges associated with simultaneous measurements, rotation-vibration non-equilibrium can instead be determined by measuring the rotational S-branch and vibrational Q-branch separately as was done in [20]. However, it is desirable to have simultaneous measurements of rotation-vibration non-equilibrium to understand energy transfer between the rotational and vibrational modes. To overcome the spectral resolution difficulty, a dual-pump CARS scheme was applied for simultaneous measurement of the rotational S-branch and the vibrational Q-branch of N₂ and determination of both rotational and vibrational temperatures in a N₂ plasma [15]. However, this approach requires four beams, which adds complexity to setup and alignment. More importantly, measurements of the vibrational Q-branch typically require an optical parametric amplifier (OPA) to produce the correct Raman shift in the Stokes beam. This limits the feasible amount of energy in the Stokes beam and makes it difficult to extend the measurement to one-dimensional (1-D) imaging. Therefore, it would be desirable to measure rotation-vibration non-equilibrium using the rotational CARS spectrum instead, where
many photon pairs are available to generate the Raman coherence. Consequently, 1-D imaging is feasible with a 2-beam phase matching scheme [11,12,16]. To our knowledge, only translated point CARS measurements of rotation-vibration non-equilibrium have been done [15,17] and 1-D imaging has not yet been reported.

It has been shown previously that vibrationally excited states of diatomic molecules exhibit a frequency shift in their rotational spectra due to rotation-vibration coupling and can be used as a measure of both rotational and vibrational temperature [21,22]. In fact, vibrational excitation shifts of the rotational levels have been considered for modelling fs/ps rotational CARS measurements of flames and plasmas [15,16]. However, in these fs/ps CARS studies either the rotational modes were assumed to be in equilibrium with the vibrational modes [16] or the probe pulse had insufficient spectral resolution to resolve the vibrational shifts [15]. Furthermore, interfering beat patterns in the time-domain were observed in [15] due to simultaneous sampling of transitions spaced 1 cm⁻¹ apart by a probe pulse with ~3 cm⁻¹ transform-limited spectral full-width at half maximum (FWHM). As a result, it was argued that dual-pump CARS was needed to determine the rotational temperature in non-equilibrium systems.

In this Letter, we present 1-D two-beam fs/ps CARS imaging of spatial gradients in rotational and vibrational temperatures of N₂ in a CH₄/N₂ pin-to-pin nanosecond pulsed discharge near the cathode.

We show that with a spectrally narrow picosecond probe pulse, vibrational contributions to the pure rotational CARS spectrum can be directly measured. The required number of beams reduces to two, which simplifies the setup for 1-D CARS imaging.

The theory of fs/ps CARS modelling has been described previously for rotation-vibration equilibrium [10,13] and non-equilibrium environments [15]. Only a brief explanation of the CARS model and illustrative examples of the non-equilibrium spectra will be presented. CARS is a third-order optical non-linear process that utilizes the difference frequency of the pump and Stokes photons to match a resonant molecular rotational or vibrational transition. The time-delayed probe pulse scatters off the coherently excited molecules which generates the CARS signal at the anti-Stokes Raman frequency. In the time-domain, the intensity of the CARS signal can be expressed as follows [10]:

\[
I_{\text{CARS}}(t) \propto \left| P^3(t) \right|^2 = \left( \frac{c}{\hbar} \right)^3 \int_0^\infty dt_2 \left( R_{\text{CARS}}(t_2) \times E_{\text{Stokes}}^* (t + \tau_2 - t_2) \times E_{\text{pump}} (t + \tau_2 + \tau_1 - t_2) \right)^2
\]  

where \( P^3(t) \) is the third-order polarization, \( E_{\text{pump}} \) and \( E_{\text{Stokes}} \) are the electric fields of the pump and probe beams, \( E_{\text{Stokes}}^* \) is the conjugate electric field of the Stokes beam, \( \tau_1 \) is the time delay between the pump and Stokes beams, \( \tau_2 \) is the time delay between the Stokes and probe beams, and \( t \) is an integration variable representing coherence time scale between the pump/Stokes and the probe beams. The molecular response, \( R_{\text{CARS}} \), can be written as:

\[
R_{\text{CARS}} = \sum_v \sum_f I_{v,f-\nu,f+2} \times \exp \left( \frac{\hbar}{\lambda} \left( i \Delta E_{v,f-\nu,f+2} - \frac{1}{2} I_{v,f-\nu,f+2} \right) \right)
\]

where \( I_{v,f-\nu,f+2} \) is the Boltzmann-weighted Raman transition intensity modified by the Herman Wallis Factor [23]. Placzek-Teller coefficients, and the polarization anisotropy due to vibrational anharmonicity [24], \( \Delta E_{v,f-\nu,f+2} \) is the energy level difference of the rotational S-branch transition, \( I_{v,f-\nu,f+2} \) is the Raman linewidth calculated by a Modified Exponential Gap model. Vibrational levels \( v = 0 \) to \( v = 5 \) were considered and the rotational linewidths were assumed to be the same across all vibrational levels. Rotation-vibration non-equilibrium was considered using a two-temperature Boltzmann distribution similar to [15].

The N₂ rotational energy levels were calculated as a function of the vibrational energy level as follows:

\[
F(v, J) = B_v (J(J+1)) - D_v (J(J+1)^2)
\]

where \( F \) is the rotational energy of an N₂ molecule in a vibration-rotation state \( (v,J) \) while \( B_v \) and \( D_v \) are the rotational energy level-dependent rotational and centrifugal constants. \( B_v \) and \( D_v \) cause red shifts in the rotational energy levels with increasing vibrational energy through rotation-vibration coupling. For molecules in rotation-vibration non-equilibrium, the vibrationally-excited levels cannot be ignored and will appear at these shifted frequencies in the rotational CARS spectrum [21,22]. This enables the detection of rotation-vibration non-equilibrium from pure rotational CARS.

Fig. 1 shows a demonstration of the advantage of using a long ps probe for detection of rotation-vibration non-equilibrium. A time delay scan of the probe pulse with respect to the pump/Stokes beams was simulated for 60 Torr N₂ at a rotational temperature, \( T_{\text{rot}} \) of 400 K and a vibrational temperature, \( T_{\text{vib}} \) of 3500 K. In Fig. 1, only the duration of the probe pulse was changed from 65 ps (left) to 6 ps (right). With the 65 ps probe, the ~0.23 cm⁻¹ spectral width was narrower than the 1 cm⁻¹ spacing between adjacent vibrational levels. Each transition was probed separately without any spectral overlap. With the 6 ps probe, heating was observed due to simultaneous sampling of neighboring vibrational levels, cited in [15] to be detrimental for accurate rotational CARS thermometry. The 65 ps probe avoids these effects on the rotational spectrum.

Fig. 2 provides an illustration of the shifted Raman transitions for N₂ at a rotational temperature of 500 K and vibrational temperature of 4000 K. The underlying peaks are the vibrationally-shifted rotational Raman transitions simulated with a long probe pulse. The overall profile shown is the total convolution of these peaks with the spectrometer slit function, experimentally fit from room temperature rotational spectra of N₂. As seen in Fig. 2, even if the vibrationally-shifted peaks cannot be individually resolved by the spectrometer, their contribution can be determined from the one-sided broadening of the rotational transition. Vibrational broadening can be detected starting from a vibrational temperature of 1500 K and becomes prominent above 2000 K.

The fs/ps CARS setup is shown in Fig. 3 and is similar to the setup.
beams were routed through the imaging setup in Fig. 3. The CARS beam was sent to a beam dump, while the probe and CARS signal produced and detected

At the hollow fiber. The spectrally broadened pulse was compressed by a high-voltage switch at 20 Hz (PVX 4130, DEI) triggered by a delay generator (DG645, SRS). The gas mixture used was 40% CH₄ and 60% N₂ at 60 Torr, and gas was flowed to flush the discharge region between voltage pulses. To measure close to the cathode surface, both the probe and pump/Stokes beams were partially blocked by a knife edge and imaged to the probed region. This created a sharp edge on the incoming beams and reduced light scattering from the cathode surface. The line-spread function was determined by the method in [12] and was ~40 μm.

All measured spectra were divided by the excitation profile determined from the non-resonant background in pure CH₄. The corrected spectra were then fit through nonlinear least-squares fitting based on interpolation of a library of pre-calculated spectra, similar to past fs/ps CARS studies [16]. The fitting parameters were the rotational and vibrational temperatures, and offsets in the baseline intensity and the frequency axis. Bi-linear interpolation was used for the spectral library look-up of the rotational and vibrational temperatures. Shown in Fig. 4 is an example of a fit of a non-equilibrium rotational CARS spectrum with \( T_{rot} = 340 \text{ K} \) and \( T_{vib} = 3600 \text{ K} \). There is a visible "bump" on the low-frequency side of each rotational peak from the higher vibrational levels of N₂. The squared residuals were plotted for an equilibrium fit \( T_{rot} ^{0}, T_{vib} ^{0} \) versus a non-equilibrium \( T_{rot} \neq T_{vib} ^{0} \) fit. At the locations of the expected contributions from the vibrationally excited N₂, the residuals show a sharp increase for the equilibrium fit. However, these peaks in the residuals disappear once rotation-vibration non-equilibrium is considered.

It is known from past CARS studies that the vibrational distribution function may not be Boltzmann [15,17]. From Fig 2, even at vibrational temperatures of 4000 K, vibrational levels higher than \( v=1 \) make minor contributions to the spectral profile. Therefore, the reported temperature is the \( T_{rot} \) temperature where the ground and first excited vibrational state are used to define the temperature for a Boltzmann rotational distribution. Since the CARS signal was generated with a spectrally narrow probe pulse, detection of non-Boltzmann vibrational distributions would only require a spectrometer with high enough resolution to differentiate the individual vibrationally-shifted peaks. At best, vibrational levels
up to $v = 2$ or $3$ could be detected with such a spectrometer, while $v = 5$ was detected with dual-pump CARS [15]. This is because the vibrational Raman cross-section scales with $v+1$, whereas the rotational Raman cross-section increases by a factor between 1 to 1.5 for $v < 10$ in $N_2$ [24]. The trade-off for higher sensitivity is increased complexity and lower Stokes energy if an OPA is used.

As a demonstration of this method, a 1-D measurement of rotation-vibration non-equilibrium distribution is presented in Fig. 5. The image was taken 20 $\mu$s after the voltage pulse and three pixel rows were binned post-processing, resulting in $\sim 42 $ $\mu$m per point. Error bars were determined based on the 95% confidence intervals of the fits. There was significant rotation-vibration non-equilibrium and spatial gradients in both temperatures, where the peak rotational and vibrational temperatures along the laser sheet were 380 K and 5500 K, respectively. The vibrational temperature exhibited a strong spatial dependence within the first mm of the cathode with a peak at 860 $\mu$m from the cathode. After the peak, rotation-vibration non-equilibrium remained, and the vibrational temperature was consistently above 2500 K.

In summary, simultaneous measurements of rotation-vibration non-equilibrium in 1-D from pure rotational fs/ps CARS were demonstrated for the first time in a CH/N$_2$ nanosecond-pulsed pin-to-pin discharge. The rotation-vibration non-equilibrium was detected using the red shift of the rotational constant with increasing vibrational energy level. This approach enabled the use of two-beam phase matching for straightforward implementation of 1-D imaging with a spatial resolution of 40 $\mu$m. Molecules in non-equilibrium other than N$_2$ can be imaged as well, since only rotation-vibration coupling in the rotational energy levels is required. Measurements in a nanosecond-pulsed plasma showed strong spatial gradients in the rotational and vibrational temperatures starting within 150 $\mu$m of the cathode surface. The time evolution of the 1-D rotation-vibration non-equilibrium can be obtained by adjusting the electronic delay to the high voltage switch. Such measurements will be the subject of future work. Finally, 2-D measurements of rotation-vibration non-equilibrium may be possible like in equilibrium rotational fs/ps CARS [25] through hyperspectral imaging [26]. This would allow validation of 2-D simulations of non-equilibrium systems critical for development of future plasma technologies and hypersonic vehicles.

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