Localized time accurate sampling of nonequilibrium and unsteady hypersonic flows: methods and horizons

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Abstract
Modern “non-intrusive” optical methods are providing revolutionary capabilities for diagnostics of hypersonic flow fields. They generate accurate information on the performance of ground test facilities and provide local time accurate measurements of near-wall and off-body flow fields surrounding hypersonic test articles. They can follow the true molecular motion of the flow and detect nonequilibrium states and gas mixtures. They can be used to capture a wide range of turbulent scales and can produce highly accurate velocity, temperature and density measurements as well as time-frozen images that provide intuitive understanding of flow phenomena. Recent review articles address many of these methods and their applications. The methods highlighted in this review are those that have been enabled or greatly improved by new, versatile laser systems, particularly including kHz rate femtosecond lasers and MHz rate pulse burst lasers. Although these methods can be applied to combusting environments, the focus of this review is on external high Mach number flows surrounding test articles and wind tunnel core flow properties. The high repetition rates enable rapid time evolving flows to be analyzed and enable the collection of large data sets necessary for statistical analysis. Future capabilities based on the use of atomic vapor filters and on frequency tunable, injection locked MHz rate lasers are promising.

1 Introduction
As maneuverable air platforms move up in speed from subsonic through supersonic to hypersonic, satisfactory safety, reliability and performance become increasingly difficult to achieve. The trade-off between performance and reliability grows ever more critical, with the loss of a few percent in performance potentially leading to a lost mission capability. The platform design task is made even more challenging by the high cost of flight tests and the constraints of ground testing capabilities. In addition, flight tests diagnostics are sparse and maneuvers are severely restricted and often cannot fully simulate mission requirements. To overcome these limitations, high fidelity and validated computational models are needed. These models must incorporate the true physics of the air flow and must be able to predict performance, including laminar to turbulent transition, nonequilibrium phenomena, dissociation, ionization, unsteadiness, shock interactions and separation as well as perturbations caused by ablation, surface degradation and structural coupling. Air breathing platforms add another level of complexity for modeling, including air fuel mixing, flame holding, ignition, combustion efficiency, unstart, sensitivity to maneuvers and thrust generation.

The challenge to the diagnostic community is twofold (1) to capture the relevant physics with sufficient fidelity to inform the modeling community and (2) to provide methods for validating model predictions in relevant environments. In both cases, the data need to be captured with temporal and spatial resolutions that enable detailed processes to be followed and understood. Other diagnostic challenges include the development of sensors to provide feedback control for flight operations and for safety monitoring. For these phenomena to be properly measured, sampling must occur at rates ranging from kiloHertz to megaHertz, often with resolution better than one millimeter.

Laser-based methods for the measurement of hypersonic flows date back more than 40 years (Zimmermann and Miles...
These advances have been enabled by advances in laser technology, including the invention of the tunable dye laser, advancements in high pulse energy solid-state laser technologies, efficient harmonic conversion, injection and narrow linewidth control, rapid multiple pulse capabilities, and femtosecond and picosecond laser technologies. These have been coupled with advances in digital camera and intensifier capabilities. Many of the earlier approaches are now being leveraged by the development of MHz rate pulse burst lasers, kHz rate femtosecond lasers and high-speed, intensified cameras that use rapid on-chip storage capabilities.

For this review, we will focus on methods that are enabled or enhanced by high repetition femtosecond lasers and by narrow linewidth MHz rate pulse burst lasers and produce time accurate measurements of local nonequilibrium and unsteady flow properties, where “time accurate” refers to sampling that is not time or multiple pulse averaged. These laser systems, together with the methods they are enabling and methods that are on the near horizon, have the potential to revolutionize our ability to acquire critical data for model development and code validation in high enthalpy hypersonic flows. Some also produce instantaneous images of flow properties that enhance intuitive understanding of complex interaction.

Commercially available femtosecond lasers typically utilize titanium sapphire as a gain medium since it has a very broad gain bandwidth, enabling the generation and amplification of sub-picosecond pulses. Typical pulse widths are 50 to 100 femtoseconds centered around 800 nm in the near infrared. Pulse outputs of the systems considered here are in the 7 mJ to 14 mJ range and pulse repetition rates are 1 kHz to 10 kHz, with the pulse energy dropping at the higher repetition rates. Femtosecond ytterbium-based fiber lasers operating at 1030 nm are also available, with commercial amplified systems operating at pulse repetition rates from 100 kHz to 1 MHz, but with lower pulse energies (< 1 mJ) and longer pulse lengths (~300 fs). It may be useful to recall that for Gaussian “transform limited” pulses, the full width and half maximum uncertainty product of time and frequency is $\Delta t \cdot \Delta f = 0.44$. Thus, 50 femtosecond pulses have a frequency width of $8.8 \times 10^{12}$ Hz or, at 800 nm, about 20 nm. A 1 mJ, 50 fs pulse corresponds to 20 Giga-Watts, so the femtosecond lasers are useful for nonlinear interactions due to their high peak power, but they are not useful for high-resolution spectroscopy due to the wide spectral bandwidth. The relationship between frequency width and pulse time indicates that if the frequency spectrum is constrained, for example, by a spectral filter, then the pulse length is increased. This provides flexibility in the pulse length and spectral performance of femtosecond lasers—their pulse lengths can be controlled by spectral filtering methods based either on dichroic spectral filters or diffraction gratings. Femtosecond lasers can be frequency up-converted with good efficiency due to the high intensity, and operation at second, third and fourth harmonics is easily achieved. They are also efficient at driving optical parametric oscillators and amplifiers, which enable broad tunability over the UV, visible and infrared. For hypersonics, femtosecond lasers have enabled femtosecond laser electronic excitation tagging (FLEET) (Michael et al. 2011; Miles et al. 2018), femtosecond/picosecond hybrid coherent anti-Stokes Raman scattering (hybrid CARS) (Prince et al. 2006) and greatly enhanced two-photon absorption laser-induced fluorescence (TALIF) (Goehlich et al. 1998).

The other new laser capability is the burst mode or pulse burst laser (Wu et al. 2000; Wu 2000). This laser is capable of emitting high energy pulses at up to or even exceeding 1 MHz rates, enabling rapid imaging of dynamic phenomena when paired with a high framing rate camera. Commercial pulse burst lasers are now available and their operational capabilities are being upgraded at a notable pace (Slipchenko et al. 2021). Typical individual pulses have a temporal width of ~10 ns and recent progress has extended the burst duration to many milliseconds and pulse energies to hundreds of millijoules (Felver et al. 2020), enabling quasi-continuous imaging of hypersonic flows and combusting environments. Shortening the pulse to sub-picoseconds (Roy et al. 2014) extends the pulse burst laser to achieve more effective nonlinear optical diagnostics. The laser is based on Nd:YAG technology and operates at 1.064 microns, but it is efficiently frequency doubled to 532 nm and can be tripled to 355 nm or quadrupled to 266 nm. It operates with a continuous wave source laser which is sliced and amplified through a series of amplifier stages. Since the source laser has a naturally very narrow linewidth, the output of the pulse burst laser has a single longitudinal mode and a very narrow linewidth. This differs significantly from standard Nd:YAG lasers that naturally operate with multiple longitudinal modes and must be injection locked to constrain operation to a single mode. The consequence is that the pulse burst laser is ideal for applications that require narrow linewidth such as filtered Rayleigh scattering (FRS) (Forkey et al. 1996) and holography. It can also serve as a pump for frequency tunable
optical parametric oscillators (Jiang et al., 2011), extending the burst capability to virtually any wavelength from ultraviolet to the far infrared. Recent work on operating with much shorter pulse lengths, on the order of 100 picoseconds, has enabled high repetition rate nonlinear processes including the development of picosecond laser electronic excitation tagging (PLEET). (Jiang et al. 2017a, b).

2 Diagnostic approaches based on femtosecond lasers

2.1 Femtosecond laser electronic excitation tagging (FLEET)

Femtosecond laser electronic excitation tagging (FLEET) takes advantage of the high intensity and short pulse length of the femtosecond laser to drive a high-order multi-photon interaction that leads to the dissociation of nitrogen in air, in pure nitrogen or in nitrogen containing gas mixtures. When the nitrogen subsequently recombines, the two nitrogen atoms initially form an excited electronic state of molecular nitrogen which fluoresces in the red to near infrared spectral region, corresponding to the “first positive” emission band of molecular nitrogen. Due to the time required for recombination and the rather long lifetime of the upper excited state, this fluorescence continues for many tens of microseconds. The region of excitation is well localized by the focused femtosecond laser, creating a well-defined line extending through the focal, or Rayleigh region of the laser. Depending on the focusing, the tagged line can be many centimeters long. That line then moves with the flow and subsequent imaging by a fast gated intensifier captures the line displacement as a function of the delay time between tagging and the intensifier “on” time. In this manner, the velocity of the flow can be precisely measured and any velocity fluctuations due to turbulence or shear become immediately evident due to the distortion of the line. The intensifier can be gated “on” multiple times to follow the same tagged line as it evolves with time, enabling imaging at MHz rates even with the tagging only occurring at the laser pulse repetition rate. Thus, “Lagrangian” measurements of flow properties are produced. In this manner, FLEET differs from vibrationally excited nitric oxide monitoring (VENOM) (Sanchez-Gonzalez et al. 2012), Raman excitation plus laser-induced fluorescence (RELIEF) (Noullez et al. 1997) and other approaches that require separate tagging and interrogation lasers.

Recent experiments performed at the AEDC Hypervelocity Wind Tunnel 9 have provided accurate measurements of flow velocity in freestream and boundary layer nitrogen flow at Mach 10, 14 and 18, at very low pressure (down to 0.35 Torr) and density (3 × 10⁻³ kg/m³) (Dogariu et al. 2019a, 2021). These experiments have been enabled by the development of transportable instrumentation for FLEET velocimetry capable of withstanding the operating conditions of large-scale wind tunnel facilities.

Figure 1 shows FLEET measurements taken at Mach 18 in AEDC Tunnel 9. The composite image in Fig. 1a shows FLEET tagged lines captured at five successive time steps after a single-shot laser pulse tagging: 2, 7, 12, 17 and 22 microseconds. The thin tagged laser line is broadened by diffusion in the low pressure environment, but the center of the displaced line can be found to good accuracy by fitting the broadened profile with a Gaussian curve, as shown by the red dots in Fig. 1a. The repetition rate of the laser pulse tagging was 1 kHz, and measurements were taken over the ~6 s
run time, which includes the ~3.5 s fully developed Mach 18 flow and the transient flow. The almost perfect linear displacement with time fit shown in Fig. 1b is a demonstration of a steady freestream flow velocity over the 22 μs observation interval. The time evolution of FLEET velocity measurements shown in Fig. 1c indicates that the velocity during the run changes due plenum gas heating. Figure 1c also shows that the precision of velocity measurements was better than 0.2% during the ~3.5 s run, and the FLEET velocity estimates agreed very well with the predicted velocity from plenum temperature measurements.

An example of boundary layer velocity measurements using FLEET is shown in Fig. 2. Two sets of measurements (I) and (II) were taken during two different Mach 18 runs performed at the same operating parameters in order to capture the flow patterns inside (II) and outside (I) the boundary layer of the 1.5 m diameter test section of the tunnel. The two image sets (I) and (II) consist of 500 time-averaged single shots collected in fully developed Mach 18 nitrogen flow, and converted to velocity measurements. The velocity uncertainties in the outside region (I) are smaller than the data point in the figure. Uncertainties in the inner region (II) are indicative of the turbulent boundary layer. In Fig. 2, the mean streamwise velocity ($U$) is normalized to the local edge velocity obtained from the first set of measurements (I). The measurement configuration has allowed recovering the boundary layer velocity profile as close as to ~2 cm from the tunnel surface, and the measured boundary layer thickness of ~32 cm is in good agreement with the theoretical prediction (Dogariu et al. 2021).

FLEET velocimetry has been also demonstrated in other complex flow environments, such as in combustion (DeLuca et al. 2017), in the NASA Langley high-pressure cryogenic tunnel (Burns et al. 2018a, 2018b), in bladeless wind turbines (Fisher et al. 2020a), in the Quiet Mach 6 tunnel at Purdue (Fisher et al. 2021), in a high-enthalpy arc-jet tunnel (UTA 2020) and in the wake of a cone at Mach 8 at Sandia (Zhang et al. 2019). Multiple line FLEET has been applied at Sandia (Zhang et al. 2020) for measurements in an over-expanded unsteady supersonic jet.

Several refinements of this technique have addressed the concerns related to the energy deposition. Depending on the laser pulse energy, that heating associated with the FLEET tagging step can be on the order of 200 K within the tagged line (Edwards et al. 2015). This increase in temperature raises concern about possible perturbations to the flow measurement. Operating with the Ti:sapphire laser frequency doubled to 400 nm improves the tagging by reducing the pulse energy required and narrowing the width of the tagged line (Dogariu et al. 2019a). Selective two-photon absorptive resonance femtosecond laser electronic excitation tagging velocimetry (STARFLEET) (Jiang et al. 2016) has been developed to minimize the heating. STARFLEET moves the tagging wavelength into the ultraviolet at 202.25 nm, a wavelength which overlaps a two-photon transition to a pre-dissociative state of molecular nitrogen. This resonant approach reduces the energy required for dissociation by a factor of 30 compared to FLEET at 800 nm and significantly reduces heating. Recent experiments at NASA have applied STARFLEET for measurements in cryogenic tunnels (Reese et al. 2020).

2.2 Hybrid femtosecond/picosecond coherent anti-Stokes Raman scattering (Hybrid CARS)

The four-wave mixing of coherent anti-Stokes Raman scattering (CARS) leaves molecules in their original states, so there is no intrinsic heating, as compared to FLEET, which relies on dissociation. Two pump beams that are separated in wavelength by the Raman shift coherently drive the selected molecules and a probe beam interacts with those molecules producing the CARS output beam. The CARS beam emerging from the sample region is highly directional due to phase matching, so background luminosity is easily rejected. Localization is achieved by crossing the pump beams in the sample region (Eckbreth 1996). Earlier work on CARS spectroscopy applications for temperature and species concentration used nanosecond laser systems in combustion environments (Eckbreth 1980) and more recently in high enthalpy arc facilities (Gülhan et al. 2018). For these applications, the operation has been at repetition rates of 10 to 30 Hz due to the pulse energies required to achieve intensities high enough to provide good signal strength. CARS suffers from
a non-resonant background which involves a similar non-linear interaction and thus satisfies the same phase matching and wavelength relationships. Due to coherent interference, this background is particularly troublesome for the measurement of other than major species and for the measurement of closely spaced spectral features. Suppression of this background in nanosecond applications is achieved by selective polarization, at the cost of reduced signal power.

The use of CARS for measurement of species mole fractions, temperature and nonequilibrium states has been revolutionized by the development of the hybrid femtosecond/picosecond CARS approach (Prince et al. 2006; Pestov et al. 2007; Dogariu 2015; Miller et al. 2010; Dedic et al. 2017; Dogariu and Pidwerbetsky 2012). The hybrid fs/ps approach can achieve CARS with a single femtosecond laser operating at kilohertz rates and suppresses the non-resonant background by temporal delay of the probe. The low energy (~ mJ) of the femtosecond pulse is offset by its short pulse length, leading to exceptionally high intensities and thus very efficient nonlinear pumping. By their nature, the femtosecond pulses are broadband, so they can be used to simultaneously drive many modes and even many molecular species into coherent oscillation. The proper Raman shifted wavelengths required for pumping are generated from the initial laser pulse by optical parametric mixing or by continuum light generation in hollow waveguides. Since the non-resonant interaction has essentially no lifetime and the pumped states all have coherent lifetimes much longer than femtoseconds, separating the timing of the pump and probe pulses eliminates the non-resonant background. Varying the time delay of the probe pulse allows sampling the coherent evolution of the excited states, and that temporal evolution can be converted into the Raman spectrum (Lucht et al. 2006). Alternatively, probing with a single narrow linewidth laser pulse and observing the spectrum with a spectrometer produces a single-shot measurement of a wide spectral region (Prince et al. 2006; Pestov et al. 2007). In this case, minimizing the time separation interval between the femtosecond pump beams and the probe beam is important to avoid coherent mode beating in the observed spectrum arising from the multiple time-evolving molecular states. Thus, the probe delay needs to be long enough to avoid non-resonant background and short enough to avoid significant mode beating. The delayed narrow linewidth probe pulse produces a broadband width, phase matched coherent output beam which is sent into a spectrometer and provides a spectral readout of the populations of all the coherently driven molecular states. The spectral resolution is limited by the linewidth of the probe pulse. In many cases, sufficient resolution can be achieved with a picosecond probe pulse that has a much narrower spectral width than the femtosecond pump pulses. The time delayed picosecond pulse can be created from a femtosecond pulse by spectral filtering (a narrower spectrum produces a longer pulse), so the same femtosecond laser can be used for all the required pulses (Dogariu 2015). Each laser pulse captures a broad spectrum, so the multiple molecular state populations associated with local species mole fractions as well as rotational and vibrational temperatures can be sampled at the laser pulse repetition rate.

Recent experiments at AEDC Tunnel 9 have applied this method to the measurement of the rotational and vibrational temperatures at a one kHz rate throughout the entire run times for multiple runs at Mach 10, 14 and 18, leading to the determination that the vibrational mode is far from equilibrium (Dogariu et al. 2019b, 2021). The left panels of Fig. 3 show examples of the measured single-shot nitrogen rotational and vibrational spectra during a Mach 18 run. The vibrational spectrum shown in the top left panel of Fig. 3 contains rotational lines, so both the vibrational and the rotational temperatures can be extracted. The bottom left panel shows the pure rotational spectrum. A modified CARSFT (Palmer 1989) code was used to generate the best fits for the spectral shapes of the synthetic spectra and is shown in red in the left two panels of Fig. 3. Since the CARSFT routine does not account for the time delay between pump and probe, it cannot be used reliably to determine the temperature for hybrid fs/ps CARS, particularly for the rotational temperature due the fast decoherence of the rotational modes and strong dependence on the pulse delays. In order to use this code to obtain the temperatures for the experiments conducted at Tunnel 9, a relative calibration was performed by comparing the CARSFT simulations with spectra recorded in a heated oven with temperatures ranging from ambient to 1,400 K. Best fits were achieved by adjusting the parameters included in the code, such as pulse bandwidths, saturation and pressure. The long coherence time of the vibrational modes led to good fits as shown in Fig. 3. At the very low rotational temperatures in the hypersonic flows in Tunnel 9, there are very few rotational modes active, so the decoherence is relatively slow. In this case, the calibration procedure used to determine the rotational temperature found that the CARSFT routine gives values within the experimental errors.

The time evolution of the vibrational temperature derived from the observed N2 Q-branch ($\Delta v = 0$, $\Delta \neq \pm 1$) shown on the right side of Fig. 3 was obtained during a Mach 18 run. The combined experimental and fitting uncertainties are shown in the figure. The results found vibrational temperatures greater than 1100 K and rotational temperatures of ~ 35 K. The vibrational temperature increase seen during the three second run is due to an increase in plenum temperature, $T_p$, shown by the red line. This lack of vibrational equilibration is a characteristic of rapid expansions that are typical of hypersonic ground test facilities and needs to be taken into account in the modeling for accurate agreement.
Current work at Sandia is expanding the hybrid CARS to the measurement along a line, opening the door to spatially resolved measurements of rotational and vibrational temperatures as well as molecular species distributions (Chen et al., 2020).

**2.3 Two-photon absorption laser-induced fluorescence (TALIF)**

Hybrid CARS and CARS in general rely on the presence of vibrational and rotation molecular states and are thus not applicable for measurements of atomic species; however, the presence of atomic species from dissociation is common in hypersonic flows and atomic species may be contaminants in high enthalpy ground test facilities. Standard laser-induced fluorescence of relevant atomic species is not possible in air since the wavelengths required are in the vacuum ultraviolet. This problem has been overcome by the combustion community through the development of two-photon absorption laser-induced fluorescence (Goehlich et al. 1998). In this case, the laser can operate at wavelengths that are not strongly absorbed by air, and through a two-photon interaction, can reach high lying atomic resonant states which subsequently fluoresce. Deriving quantitative values using this approach has been hampered by the nonlinear nature of the excitation, fluorescence quenching, ionization and stimulated emission, and, for gases with residual undissociated molecules, laser-induced dissociation. Moving to femtosecond pulse methods has alleviated the dissociation limitation by achieving the excitation of atomic species that are present in the test gas faster than the dissociation can occur. Contrary to expectations, relatively efficient excitation of states with linewidths far narrower that the linewidth of the femtosecond pulse occurs due to the multi-photon process, where the red and blue skirts of the spectrum offset each other and add to the required narrow transition wavelength (Kulatilaka et al. 2012). An advantage of the femtosecond approach is that the commercial femtosecond lasers operate at higher repetition rates than nanosecond lasers and have excellent stability from one pulse to the next, so fluctuations associated with the multi-photon nature of the excitations are minimized. Quenching can be measured by capturing the fluorescence at two or more times after excitation (Settersten et al. 2006), allowing extrapolation to unquenched densities. Stimulated emission and ionization are residual problems that limit laser fluence. Stimulated emission can be monitored by measuring the backward emission along the excitation laser path (Agrup and Alden 1994), and ionization can be monitored by microwave scattering from free electrons (Dogariu and Miles 2011). Possible methods of calibration include seeding a small but well-known concentration of a noble gas such as xenon and simultaneously exciting those atoms, providing an ability to ratio the TALIF from the noble gas with that of the atomic species (Niemi et al. 2001). Both xenon and oxygen have two-photon resonances that are at almost the same wavelength (225.5 and 224.3 nm for xenon and 225.6 nm for oxygen). Krypton and atomic nitrogen and atomic hydrogen are close as well (204.2 for krypton, 206.6 nm for nitrogen and 205.1 nm for hydrogen). The full width at half maximum of a Gaussian 50 fsec pulse

![Fig. 3 Left: Vibrational (top) and rotational (bottom) hybrid fs/ps CARS spectra (black) and best fit nonequilibrium calibrated CARSFT numerical simulations (red) for Mach 18 flow in AEDC Tunnel 9. Right: Comparison of 1 kHz rate hybrid fs/ps CARS vibrational temperature ($T_v$) measurements (black) in the test section and plenum temperature ($T_0$) (red) versus time for a three second run plus turn on and turn off transients](image-url)
at 266 nm is ~ 2 nm, so excitation of xenon and oxygen may be achieved simultaneously. Figure 4 shows the excitation and emission transitions for TALIF in atomic oxygen and nitrogen, together with the similar schemes for the atomic species used for TALIF calibration: Xe and Kr, respectively. Recent measurements in the arc-jet facility at the University of Texas at Arlington have demonstrated the capability of fs-TALIF to measure atomic species in high enthalpy hypersonic flows (UTA press release 2020). Figure 5 shows TALIF fluorescence from naturally occurring xenon in air (~ 80 parts per billion) following femtosecond two-photon excitation at 252.48 nm, demonstrating the high sensitivities that can be achieved.

3 Pulse burst laser enabled methods

The pulse burst laser enables numerous previously developed laser diagnostics to be implemented at high repetition rates and is particularly useful for acquiring data in short run time facilities and for the measurement of rapid interactions. A review of applications is provided by Thurow (2013) and more recently by Slipchenko et al. (2021)

3.1 Filtered Rayleigh scattering

The use of planar Rayleigh scattering for imaging supersonic and hypersonic flows provides instantaneous details of turbulent boundary layer structure, shock structure and shock wave/boundary layer interactions (Miles and Lempert 1990). Initial applications of MHz rate filtered Rayleigh scattering were for the imaging the temporal evolution of shock boundary layer interactions though Rayleigh scattering from carbon dioxide nano particles that formed in the core of low enthalpy supersonic and hypersonic flows. In this case, approximately 1% of CO₂ was introduced into the air upstream of the plenum (Erbland et al. 1997; Wu 2000; Poggie et al. 2004) and as the flow expanded through the nozzle, the CO₂ vapor condensed into a thin fog of CO₂ nanoparticles in the core region, but not in the boundary layer due to the higher temperature in that region. Thus, the CO₂ fog provided high contrast images showing the structure of the outer boundary layer and dynamic interactions with shocks and shock-induced separation. This approach takes advantage of the filtered Rayleigh capability by frequency doubling the pulse burst laser to 532 nm and placing an iodine vapor cell as a filter
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Iodine has numerous strong absorption lines that fall within the gain spectrum of the frequency doubled Nd:YAG laser. The narrow linewidth continuous wave seed laser is tuned so the second harmonic wavelength overlaps one of these strong iodine absorption lines. By illuminating the flow at an angle such that the light from the moving CO₂ particles is Doppler shifted out of the iodine absorption, the background from the stationary windows and walls that otherwise obscures the flow images is eliminated while the Doppler shifted light scattered from the flow passes through the filter and is imaged by the camera. Figure 6 shows sequential images taken with this approach.

The challenge for the Rayleigh measurement of density and other parameters in unseeded hypersonic flows is to achieve sufficient pulse energies to provide good signal to noise from the low densities associated with hypersonic ground test facilities. The Rayleigh scattering cross section is greatly enhanced in the ultraviolet, so increases in pulse energy, extension to the ultraviolet and implementation of other filters to block out background scattering including interferometric filters (Cutler et al., 2020) and atomic vapors such as mercury at 254 nm and optically pumped barium at 355 nm (Hetlage and Limbach 2021) may enable this capability.

By tuning the laser across the absorption band of the vapor filter, filtered Rayleigh scattering can be used for imaging velocity, temperature and density by deconvolving the measured profile to give the lineshape and frequency shift of the Rayleigh scattering, (Forkey et al. 1998) This approach requires many image frames to be taken and pixel resolved deconvolution, so it is not useful for rapidly varying flows or short run time facilities. Single-shot imaging of specific flow properties can be achieved by tuning the laser relative to the vapor filter absorption spectrum, changing the collection angle or changing the vapor pressure (Miles et al., 2001). For example, when the Rayleigh scattering linewidth matches the filter linewidth, the scattering is independent of temperature (Feng et al. 2020) leading to a measurement of pressure, assuming an ideal gas. By tuning the laser to the edge of the vapor absorption profile, the scattering is sensitive to velocity (Thurow et al. 2005). That velocity sensitivity can be enhanced by changing the vapor absorption line profile with the addition of an inert gas that collision broadens the line (Elliott and Beutner 1999).

3.2 MHz rate laser-induced fluorescence (LIF)

The extension of the pulse burst laser to variable wavelength operation opens the door to its use for laser-induced fluorescence applications. The only candidate for LIF in pure air is oxygen, which for atmospheric temperatures requires a laser operating in the ultraviolet overlapping the Schumann Runge bands at wavelengths below 200 nm,. This has been achieved using ArF lasers operating at 193 nm (Laufer et al. 1990), but so far that wavelength region has not been reached with the pulse burst laser. If nitric oxide is naturally present from the high temperature in the plenum or from post-shock heating or it is seeded into the flow, it can be used for LIF with the frequency tunable MHz laser operating at 226 nm (Jiang et al., 2011). An example is the stereo imaging achieved in the 31-Inch Mach 10 Air Tunnel at NASA Langley (Medford et al., 2011). In this case, the air was seeded with nitric oxide from a spanwise slot on a wedge model and the evolution of the seeded air was followed after passing over a small cylindrical protuberance with stereo imaging at up to 500 kHz. The high repetition rate may also enable the vibrationally excited nitric oxide monitoring (VENOM) technique (Sanchez-Gonzalez et al. 2012).

3.3 Picosecond laser electronic excitation tagging (PLEET)

Significantly shortening the pulse to sub-nanoseconds while maintaining the pulse energy enables the MHz pulse burst system to achieve flow tagging through PLEET (Jiang et al. 2017a, b). This approach is similar to FLEET and was developed to overcome the lack of very high rep rate, high pulse energy (~ 1 mJ per pulse which is what is required for FLEET) femtosecond lasers. PLEET requires about 30 times more energy per pulse than FLEET, since the pulses are approximately 1000 times longer, and thus have much lower

Fig. 6 Mach 2.5 air flow over a 24° wedge taken at 500 000 frames per second with the pulse burst laser and an iodine “notch” filter. Images are sequenced in 2 μs time steps from left to right. Flow is from left to right, and light is scattered from CO₂ particles.
peak intensity. Significantly more heating of the tagged region occurs, and strong short-lived emission in the ultraviolet corresponding to the nitrogen “second positive” bands occurs over the first 500 nsec or so, followed by long lifetime “first positive” red and infrared emission characteristic of FLEET that is used for the displacement imaging. The tagged regions continue to fluoresce for longer times than FLEET, which may be associated with the higher temperature and higher energy deposition. Applications at 100 kHz in a Mach 6 Ludwieg tube at the Air Force Research Lab (Hsu et al. 2020) and 25 kHz measurements in transonic facilities at NASA have been conducted (Burns et al. 2018a, 2018b).

4 Future capabilities

The continued development of laser technologies will significantly impact hypersonic diagnostics and potentially bring some of these methods to flight data acquisition systems. These advances will further the goal of capturing true time-frozen data at rates high enough to follow dynamic processes at locations that are relevant to vehicle design and to the determination of critical physical processes. For example, the development of 100 kHz to MHz rate, high pulse energy femtosecond laser systems will enable the application of FLEET, fs/ps CARS and TALIF at those repetition rates (Smyser et al. 2019). The extension of MHz rate lasers to this femtosecond regime unites both the femtosecond diagnostics with the MHz sampling capability, enabling rapid measurements in dynamic environments including both external hypersonic flows and internal flows for air breathing, SCRAM jet powered platforms. These approaches involve combinations of femtosecond seeding, amplification with broadband Nd:glass, and pulse shape control and represent yet another step in the laser technology that promises to significantly advance diagnostic capabilities. Preliminary demonstrations of this capability have been achieved both for FLEET (Fisher et al. 2020b) and hybrid CARS (Smyser et al. 2020).

An increase in the pulse energy of the pulse burst lasers will enable Rayleigh measurements of low density gases. Higher repetition rates may enable microsecond time-frozen volume imaging with spatially swept beams. Rapidly frequency swept pulsed lasers will enable other nonlinear processes such as frequency chirped coherent Rayleigh/Brillouin scattering to be implement for the measurement of velocity and temperature. (Gerakis et al., 2021). Multi-wavelength pulsed lasers may enable simultaneous calibration and single-pulse imaging of velocity fields and thermodynamic properties. Higher spectral purity lasers will further atomic and molecular filter capabilities.

The integration of atomic filters into the detection systems together with the development of higher pulse energy, narrow linewidth tunable pulse burst lasers provide new opportunities for hypersonic diagnostics. Two such approaches are currently being explored—slow light imaging spectroscopy (SLIS) (Miles and Dogariu 2020; Dogariu and Miles 2018) for imaging rotational Raman features and filtered low-angle Thomson scattering (FLATS) (Miles et al. 2020) for detecting low density ionization near leading edges and in boundary layers of hypersonic test articles. SLIS takes advantage of the delayed propagation of light through atomic vapor cells when the light wavelength falls close to an atomic resonance (Hau et al. 1999; Wang et al. 2000). The wavelengths of Rayleigh, Raman and Thomson scattering all follow the wavelength of the excitation laser source. Therefore, if the laser source wavelength is changed, their wavelengths are changed, so with a tunable laser, the Rayleigh, Raman or Thomson wavelengths can be tuned to fall close to an atomic vapor resonance. Other features of Rayleigh, Raman and Thomson scattering are that the scattering only occurs during illumination and that it does not suffer from quenching. In this way, Rayleigh, Raman and Thomson scattering differ substantially from laser-induced fluorescence which has an excited state lifetime and has its fluorescence significantly affected by quenching. The primary difficulty with Rayleigh, Raman and Thomson scattering is the very low scattering cross sections, so methods that remove background interference and enhance light collection may have significant impact on data acquisition capabilities for hypersonics. This is especially important for data acquisition in short run time shock and expansion tubes where long time averaging is not possible and for capturing dynamic phenomena and turbulent flow properties.

4.1 Slow light imaging spectroscopy (SLIS)

The two features of Rayleigh, Raman and Thomson scattering, frequency tunability and time synchronization, enable slow light imaging. Typical delays that can be achieved through atomic vapor cells without significant loss can be more than 20 ns, so if the illumination time is shorter than that, then the scattering light pulse delay will be greater than the pulse time and the delayed light can be easily distinguished by an intensifier time gated camera. The atomic vapor cell does not affect the propagation vectors of the delayed light, so images are preserved. This opportunity is particularly useful for rotational Raman imaging, since rotational Raman is significantly stronger than vibrational Raman, and Raman scattering is capable of providing measurements of temperature, species concentrations and nonequilibrium. The low scattering cross section, the proximity of strong Rayleigh and background laser scattering, the interleaved rotational spectral lines from other constituents
and the need for a spectrometer have previously limited the ability to image these spectral features. The acceptance bandwidth of the atomic filter is a few GHz, so assuming the laser is injection locked (narrow band), individual rotational Raman lines can be imaged. The product of collection solid angle and collection aperture (etendue) can be orders of magnitude greater than spectrometers. Candidate elements for atomic filters are cesium, potassium, rubidium, sodium and mercury. There is a fundamental trade-off between transmission and delay: longer delays result in lower transmission. The Raman scattering is greatly enhanced in the ultraviolet, so operation using atomic resonances in the ultraviolet is preferred, suggesting mercury at 245 nm or UV transitions in alkalis. Figure 7 shows the predicted delay for a 5 ns long transform limited pulse tuned to a transmission window between isotopic lines of mercury and passing through a 2” long atomic mercury vapor filter with the side arm temperature held at 100 C. A delay of up to 50 nsec is predicted with 60% transmission (Rad et al. 2020). Better performance may be achievable with isotopically purified mercury.

4.2 Filtered low-angle Thomson scattering (FLATS)

Filtered low-angle Thomson scattering provides a method to suppress background interference and extend the sensitivity range of Thomson scattering. It takes advantage of the “coherent” nature of Rayleigh/Brillouin and Thomson scattering compared to the “incoherent” nature of Raman scattering. This difference arises due to the fact that Raman scattering leads to a change in the population of molecular energy states, and thus, there is no coherence associated with the scattering process. Raman scattered photons have random phase and the frequency shift from the illumination laser is set by the energy levels involved. Rayleigh/Brillouin and Thomson scattering do not lead to a change in energy states, so the scattered photons retain the phase of the incident photons. Frequency shifts are associated with the observed net phase modulation from the laser source to the detector and arise from the motion of the scattering entities (atom, molecule, ion or electron) leading to broadening. In the forward direction, the observed net phase modulations of all scattered photons are the same and no broadening occurs from the thermal motion. When collected at 90 degrees to the illumination direction, the conventional collection configuration, the electron Thomson scattering linewidth extends beyond the rotational Raman shifts due to the very low mass and, consequently, the high thermal velocity of the electrons (Fig. 8). Measurement of electron density and temperature requires extracting the magnitude and lineshape of the electron Thomson scattering. Typical electron densities surrounding vehicles flying at Mach 13 are below 10^{13}/cc (Surzhikov 2018), making the measurement extremely difficult due to the overlapping Raman lines from the much higher densities of neutral molecular species.

The narrowing of the scattering in the forward direction enables the selective narrowing of the Thomson scattering relative to the Raman lines and thus the separation of the

**Fig. 7** Predicted delay for a 5 nsec pulse passing through a 2” long mercury vapor cell. The left panel shows the variation in delay as the laser is tuned across the mercury vapor absorption features and the black highlight shows the delay when the laser frequency falls between isotopic absorption lines as shown by the black line on the right hand panel.
Thomson from the Raman. The Rayleigh/Brillouin scattering and ion Thomson scattering are also narrowed by the same process, so inserting an atomic filter and tuning the illumination laser to the atomic filter absorption feature eliminates the Rayleigh/Brillouin and ion Thomson scattering and passes the Raman and electron Thomson scattering (Zaidi et al. 2002; Lee and Lempert 2002). The spectrum can then be analyzed by passing that light into a spectrometer. Since the Thomson scattering is well separated from the Raman, overlap is avoided and the Thomson scattering spectrum and amplitude can be directly recorded. Background plasma luminosity is broadband, and narrowing of the Thomson scattering improves the rejection of that broadband light by spectral filtering. The spectrometer can be eliminated by configuring the atomic cell as a prism to disperse the Thomson scattering and enable measurement of the Thomson scattering spectrum (Finkelstein et al. 1998). With the laser tuned to the atomic line center, the background laser scattering, Mie scattering, Rayleigh and ion Thomson scattering are blocked and only the electron Thomson and Raman scattering pass. The dispersion is strongest near the atomic resonance, so the electron Thomson scattering is strongly dispersed, whereas the Raman scattering only weakly dispersed. Figure 9 shows the configuration for the capture of forward scattering and the subsequent spectral filtering using an atomic vapor prism.

In order to enhance this effect, a multi-prism cell can be employed. The Thomson scattering cross section is \( \sigma = 7.94 \times 10^{-26} \text{ cm}^2/\text{sr} \), independent of wavelength, so operation in the near infrared is preferred in order to minimize the Rayleigh and Raman/Brillouin background. Candidate atomic vapors are rubidium at 780.0 nm and 794.7 nm, potassium at 766.5 nm and 769.9 nm, and cesium at 852.1 nm and 894.3 nm.

5 Summary

New approaches are revolutionizing the diagnostic capabilities for hypersonic ground test facilities, driven primarily by the availability of high-performance femtosecond and pulse burst laser capabilities. Molecular tagging has enabled the tracking of flow for measurements of velocity and acceleration as well as for the capture of turbulent structure and boundary layer profiles. Single-shot broad spectral
width femtosecond/picosecond CARS is providing real-time measurements of species mole fractions as well as localized rotational and vibrational temperatures. Atomic species are being tracked by TALIF, which is particularly useful at the low static pressures encountered in hypersonic facilities, since quenching rates are minimized. Seeding the flow with a small concentration of a noble gas provides a method for calibration. The extension of the pulse burst laser to frequency tunable narrow linewidth operation through the implementation of seeded optical parametric oscillators has opened the door to rapid LIF and potentially to the use of atomic filters for measurements of Raman and Thomson scattering at MHz rates. Extending the tunability range farther into the ultraviolet may enable rapid imaging of temperature and oxygen density through Schumann Runge LIF methods. Achieving picosecond and femtosecond pulses at MHz rates opens up the option for high rate nonlinear optical diagnostics, including flow molecular flow tagging. The continued development of laser capabilities along with detection methods shows great promise for more near-term advances based on rotational Raman and Thomson scattering.

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