Design and Performance of an Acoustic Levitator System Coupled with a Tunable Monochromatic Light Source and a Raman Spectrometer for In Situ Reaction Monitoring

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ABSTRACT: The design and performance of a custom-built reaction chamber combined with an acoustic levitator, a tunable monochromatic light source, and a Raman spectrometer are reported. The pressure-compatible reaction chamber was vacuum-tested and coupled with the acoustic levitator that allows contactless sample handling, free of contingent sample requirements such as charge and refractive index. The calibration and performance of the Raman spectrometer was studied utilizing gated detection and three different gratings that can be interchanged within seconds for a desired resolution and photon collection range. A wide range of 186–5000 cm⁻¹ Raman shift, with a small uncertainty of ±2 cm⁻¹, can be recorded covering a complete vibrational range in chemical reaction monitoring. The gating of the detector allowed operation under the room light and filtration of unwanted sample fluorescence. The in situ reaction perturbation and monitoring of physical and chemical changes of samples by the Raman system were demonstrated by degradation of polystyrene by monochromatic UV light and photobleaching of a potato slice by visible light. This instrument provides a versatile platform for in situ investigation of surface reactions, without external support structures and under controlled pressure and radiation conditions, relevant to various disciplines such as materials science, astrochemistry, and molecular biology.

1. INTRODUCTION

In recent years, acoustic levitation⁴ has been utilized in a range of scientific disciplines including but not limited to analytical chemistry, astrochemistry, materials science, and pharmaceuticals.⁵−⁷ Acoustic levitation has also been used in contactless mixing of two droplets to initiate a chemical reaction⁹,¹⁰ and to transport protein crystal samples in crystallography experiments.¹¹ In acoustic levitation, a liquid droplet, a solid particle, or a suspension is levitated slightly below the pressure nodes of the ultrasonic standing wave. The major advantages of such a levitation technique lie in the removal of often complicating effects of a contacting surface on the sample. The absence of a contacting surface or a wall also increases the sensitivity of the spectroscopic detection techniques such as infrared spectroscopy, Raman spectroscopy, mass spectrometry, and X-ray diffraction utilized to study the chemical and physical modifications of the levitated droplet/particle. Furthermore, a levitator can be used to probe the differences between the physical and chemical processes on the bulk samples and the surfaces of sub-millimeter size levitated particles.⁶,¹² Such tiny particles floating in the given gas medium provide ideal laboratory conditions simulating the dust particles or droplets in the atmospheres of planets and exoplanets.⁵ Moreover, the levitated particle can respond quickly to temperature and radiation perturbations due to their small total heat and radiation dose capacity. Hence, a levitation device provides a versatile platform to study the physical and chemical processes without support structures as well as creates less-contingent sample trapping requirements, such as electric charge and refractive index.

In combination with a controlled gas and radiation environment, acoustic levitation provides a strong and relatively unexplored avenue for studying how a sample might behave in a variety of atmospheric conditions. For example, acoustic levitation has been gaining traction in the space science community for its potential in low-gravity simulation experiments.¹³,¹⁴ The ability to suspend micrometer-sized particles or droplets in a given gas medium provides ideal laboratory conditions for simulating interplanetary dust particles and micrometeoroids that can be found in the atmospheres of planets and exoplanets. In such laboratory experiments, gas–surface, heterogeneous reactions are of particular interest to investigate cloud and haze dynamics.⁵,¹⁵ In a gas–surface reaction investigation, reactions of unsaturated hydrocarbon gases on the surface of silicon single particle had been performed,⁵ where selected hydrocarbon gases are adsorbed on the surface and polymerized to refractory...
carbonaceous compounds whose spectroscopic signatures matched with the meteoritic sample providing insights into the possible formation conditions of the carbonaceous compounds in the meteoritic sample. A model mechanistic computational study showed that 1,3-butadiene (H₂CHCHCH₂) can adsorb exothermically on the silicon surface propagated by a second H₂CHCHCH₂ molecule adsorption with further release of energy. Free radical formation and further chain propagation likely lead to the removal of hydrogen atoms with increasing carbon content to form the complex product mixture. This finding might explain the lower than expected methane to carbon monoxide (CH₄/CO) ratio in the hot Neptune GJ 436b, where the abundant methane photochemically converts to higher order hydrocarbons and ultimately to refractory graphitic carbon in the presence of catalytic silicon surface. Furthermore, the liquid droplets are often used as a reaction medium for varieties of physical and chemical reaction processes, such as nucleation and crystallization in a container processing, but recently, such reactions have been the subject of investigation under levitation conditions. However, the time scales of such reactions may be longer than the evaporation time of these small liquid droplets. Hence, it is of prime interest to prolong the evaporation lifetime of liquid droplets. Recently, experiments have been performed on the evaporation times and trends of microliter volumes of common solvent methanol and water under normal container processing and under the acoustic levitation conditions. Results showed several fold increase on lifetime of the droplet in the later conditions. Evaporation of solutions resulted in recovery of the solute material but with trapped solvent inside the residue under ambient conditions.

Even though the experiments described above approximately simulate the temperature and pressure conditions in selected exoplanet atmospheres, such as hot Jupiters and Neptunes, and provide a glimpse of the initial reaction and products, systematic improvements in laboratory simulation conditions such as host star radiation in the atmospheres, as well as detection techniques, are highly desirable. By utilizing monochromatic light from UV to IR range, it is possible to obtain information on the wavelength-dependent reactions on the surface of levitated particles. Such experiments might also yield valuable information on the underlying, hitherto largely unknown, reaction mechanism. Here, we describe the design and performance of a pressure-compatible reaction chamber equipped with an acoustic levitator and a tunable monochromatic light source (240–2400 nm) as the radiation source and demonstrate its applicability in inducing the chemical and physical changes on polystyrene microparticles. Photobleaching of a sliced potato sample with visible light was demonstrated for the instrument’s potential applications in chemical monitoring of agricultural products and environmental samples. To record the reaction changes in situ, a custom-built Raman spectroscopic system was interfaced with the reaction chamber and acoustic levitator. While the reaction chamber and levitator enclosure designs are similar to that reported by Brotton and Kaiser, addition of the tunable light source and the three-grating Raman spectrometer system adds into the instrumental capability. While such capability is enhanced in our instrument, Brotton and Kaiser have additional probes of IR and UV−vis spectroscopies. Calibration, performance, and application in monitoring chemical signatures during sample physical and chemical transformation are described in detail. Such Raman spectral recordings over a time period also allow kinetic studies in formation, destruction, or transformation reactions on surfaces as well as identification of reactants and products based on their vibrational signatures.

2. RESULTS AND DISCUSSION

2.1. Reaction Chamber and Levitation Device. The design of the reaction chamber is similar to that reported by Brotton and Kaiser with fewer access ports. The experimental
Figure 2. Schematic of the Raman spectroscopic system (not drawn to the scale). Details of parts and timing sequence are given in the text.
that reported by Bennett et al.25 and Brotton and Kaiser.4 In addition to photon collection optics, the major difference in our system from the above-mentioned systems lies in three available gratings that can be interchanged within seconds to achieve desired spectral resolution and range as well as smaller camera pixel size of 13 μm compared to 26 μm.

The Isoplane spectrograph and CCD camera were connected to a computer interface and controlled by LightField software (Princeton Instruments), which also allowed data recording and processing. Synchronization of the excitation laser, spectrograph, and CCD camera was done as follows. The spectrometer and CCD were controlled by LightField via the USB and GigE interface. A function generator (Quantum Composers, PG9514) outputs a TTL pulse via BNC connection that triggered the CCD camera at the desired repetition rate (typically 1 Hz to 1 kHz). The output from the camera circuit board then triggered the external input of the excitation laser. The output from the laser further triggered the spectrometer and camera with adjustable gated delay and width for scattered photon collection. A typical gate delay of 300 ns from laser excitation time and a gate width of 250 ns were used in the data described in this paper. The Isoplane spectrograph was equipped with a turret containing three different dispersion gratings of size 40 mm × 40 mm with 300, 600, and 1200 grooves/mm with 500 nm blaze wave length, which provides about 70% efficiency in the collection range. For 100 μm entrance slit and cyclohexane sample, the full width at half peak maximum were 38, 18, and 8 cm⁻¹ for 300, 600, and 1200 grooves/mm gratings, respectively. The observable Raman shift range for these gratings was 5000, 2500, and 1000 cm⁻¹. The LightField software allowed selection of a particular grating and center wavelength within few seconds. The gating of the detector allowed filtration of background light as well as an unwanted fluorescence signal from the sample. Such gating ability not only allows operation of the data collection at normal room light but also increases the Raman signal-to-noise ratio. Note that our system employs sample at a long distance from collection optics and spectrometer as opposed to commercial Raman system where sample-scattered photons are collected at a close distance to the spectrometer as opposed to commercial Raman system where sample-scattered photons are collected at a close distance to the spectrometer.

2.3. Spectrometer Calibration and In Situ Reaction Monitoring. The Isoplane spectrograph provides the scan range of 0–1400 nm, wave length accuracy of 0.2 nm, and sub-pixel repeatability utilizing an AccuDrive grating scan system controlled by the LightField command. The dispersed light from the spectrograph was recorded by PI-MAX4 iCCD in its 1024 × 1024 pixels. The initial calibration and wavelength accuracy check of the camera pixels was performed by utilizing a neon lamp source in the visible range of electromagnetic spectrum by placing it outside the entrance slit of the Isoplane spectograph. Spectra were recorded using neon emission lines for each of the three gratings using 650 nm center wavelength in the camera pixel. First, the bandwidths were examined for different slits of 10, 20, 30, 40, 50, and 100 μm size. Below 20 μm, the bandwidths did not change. Hence, we analyzed bandwidths and range of the spectrometer at a 20 μm slit width by calibrating neon emission lines. Linear fits correlating camera pixel number to atomic emission wavelength were used. For 300 grooves/mm grating, the spectrometer range was 200 nm with a band full width at half-maximum (fwhm) of 1 nm (20 cm⁻¹). For 600 grooves/mm grating, the spectrometer range was 100 nm with a band fwhm of 0.52 nm (10 cm⁻¹). For 1200 grooves/mm grating, the spectrometer range was about 50 nm with a band fwhm of 0.19 nm (5 cm⁻¹). Hence, in our spectrometer, the best possible band fwhm is 5 cm⁻¹. For our fixed laser excitation at 532 nm, the center wave length of the detector was adjusted to a desired value to capture the Raman scattered photons. The current system utilizes optics to collect the Stokes shift but anti-Stokes shift can be recorded by adjusting the center wavelength within a few seconds utilizing the LightField control. Stokes shift range of 0–4500 cm⁻¹ covers most of the vibrational region, which can be achieved in a current set up with 300 grooves/mm. The 600 grooves/mm grating provides better resolution and still can cover the desired range in most experiments. In the case of higher desired resolution, 1200 grooves/mm grating can be used to monitor the Raman peaks within a short range, but at least two different recordings are required to cover the long range due to its 1000 cm⁻¹ range for one recording. Figure 4 shows the Raman shift calibration using cyclohexane for three different gratings mentioned above with typical photon collection times of 50–500 μs. The 50–500 μs collection time equals to 200 ns of gate width, 50 on-CCD collections, and 5–50 of such cycles. Dispersive slit images were recorded and converted to cross sections for graphing. Cross sections were generated by binning each CCD pixel column summing the intensities from each pixel column to produce a single data point for each pixel column. For each grating, a graph of spectroscopic peak maximum pixel number versus Raman shift was created, where the Raman shift values were taken from the NIST database.26 While linear equation fittings yielded satisfactory results, quadratic fits to the data points gave best results with errors of ±5, ±2, and ±2 cm⁻¹ for 300, 600, and 1200 grooves/mm gratings, respectively. The errors represent the standard deviation of Raman shift values from the NIST data. Data were taken at 100 μm slit width, which provided sufficient resolution to monitor the cyclohexane peaks even at lowest resolution grating. The peak at zero Raman shift arises due to

![Figure 3. Effect of spectrograph light entrance slit size on peak intensity and resolution.](https://doi.org/10.1021/acsomega.1c00921)
scattered excitation laser light from multiple surfaces entering through the entrance slit. Using the cyclohexane fingerprint region bands, the peak fwhm values were determined to be 63, 56, 61, and 61 cm$^{-1}$ for the four peaks recorded with 300 grooves/mm grating. For the 600 grooves/mm grating and same peaks, the fwhm values were 38, 35, 36, and 28 cm$^{-1}$, respectively. Such typical values of fwhm for a peak for different gratings were 60, 36, and 32 cm$^{-1}$, respectively, also illustrated by the 2930 cm$^{-1}$CH$_2$ asymmetric stretch peak. Pelletier reported liquid cyclohexane bands with a bandwidth of about 10 cm$^{-1}$ with a 3 s collection time and a 25 μm slit width utilizing a holoplex transmission grating. Our bandwidths are larger than those reported by Pelletier due to the larger slit width but needed much shorter collection time as a consequence. Note that these fwhm values are typical for this calibration and can be optimized for higher resolution by the larger slit width but needed much shorter collection time as a consequence. Note that these fwhm values are typical for this calibration and can be optimized for higher resolution by changing collection focus at the entrance slit and the slit width. The 2930 and 2850 cm$^{-1}$ bands arise due to a$\_g$−CH$_3$ asymmetric stretches. Assignments of these and complete vibrational bands of cyclohexane can be found elsewhere.26,28

2.4. Tunable Light Source for Photoreaction. A tunable monochromatic light source was interfaced via a view port to the reaction chamber as shown in Figure 1. The light source includes a 300 W/15 A ozone-free xenon arc lamp, lamp power supply and control, beam optics, and a Cornerstone 130 monochromator (Newport Corporation, TLS-300XU) with the output light range from 250 to 2400 nm. In the optimum output range of 300−1800 nm and a slit size of 280 μm, a spectral resolution of 0.7 nm and a bandwidth of 3.7 nm can be achieved with a typical beam output size of 20 mm and 600 lines/mm diffraction grating. A bandwidth of 8 nm was obtained with a 600 μm slit size. The wavelength accuracy was typically 1 nm with 5% beam uniformity. The output optical power was about 0.1 mW and the irradiance was 0.2 mW/cm$^2$ for a 0.5 cm$^2$ beam area between 350 and 600 nm, with as much as 3 times higher power in the 800−1000 nm range. The output beam was aligned using vertical and horizontal adjustment knobs on the lamp housing door, rear reflector adjustment knobs on the lamp housing side panel, and focused to the center of the chamber using a lens inside the chamber. The beam size can be adjusted with extra focusing lenses, as necessary. The TLS-300XU system was connected to a control computer using an USB interface and TracQ, Oriel’s instrument control, and data acquisition software. The TracQ software allowed quick library file load and control of output wavelength with an accuracy of about 1 nm.

To demonstrate the efficiency of the light source in chemical and physical transformations of a sample, experiments were performed by levitating a polystyrene sphere of diameter 0.99 mm (Cospheric, PSS-1.05) under room temperature and pressure conditions. Figure 5 shows the cross sections of dispersive slit images of polystyrene spheres at different UV irradiation times. Each trace is for a total of 25 μs of collection time. Three different traces of Raman signals at 0, 2, and 15 s of 250 nm photon irradiation times are shown from top to bottom graphs. The higher background counts at lower Raman shift values were due to fluorescent light resulting from polystyrene at lower wave lengths. These spectra contain at least three distinct regions with vibrational bands of polystyrene. Bands at lowest Raman shift represent −CH$_2$ rocking motion, at middle represent −CH$_3$ bending motion, and at higher Raman shift −CH$_3$ stretch and −CH stretch of phenyl group.29 Degradation of polymer structures, due to dehydrogenation reactions, became apparent after 2 s of irradiation time while complete loss of structures was observed after 15 s. Such transformation was also visible physically, with white to black color change before and after irradiation, as shown in the right panel of Figure 5.

Figure 6 shows the effect of photobleaching by 532 nm visible light on a freshly sliced potato sample. Even with optimized gating conditions of a detector, the Raman spectrum is dominated by fluorescence light from the sample, masking the Raman peaks as shown in the top trace in Figure 6. Up to 1 h of photobleaching by the tunable light source at two wavelengths, 450 and 532 nm did not change the spectrum significantly, likely due to insufficient number of photons. Spectra were then recorded for photobleaching utilizing a 532 nm laser with a 60 mW power at 2, 4, 10, 15, and 30 min of excitations. Significant change in the spectra were observed on first 2 min where the fluorescence reduced and two Raman peaks became clearly visible. To prevent the photobleaching...
during recording of the spectra, very short excitation times (6 μs) were used. A small reduction in the background and an improvement in the spectra were observed after that, as shown in the 30 min spectra in Figure 6a. Two broad Raman bands near 2900 and 3400 cm\(^{-1}\) were attributed to C–H stretching and O–H stretching vibrations from carbohydrate and water since the typical carbohydrate and water contents in potatoes are 19 and 77%, respectively.\(^{30}\) Figure 6b shows the baseline-corrected, deconvoluted spectra for 30 min irradiation in the high-frequency region. Amorphous starch has low-frequency bands in the fingerprint region but too weak to be detected in our setup. Gaussian fits to the data with an \(R^2\) value of 0.982 were obtained. Experimental spectrum is shown in solid black curve while the cumulative overall fit is shown in dashed gray curve. The C–H stretching region can be fit with a major peak at 2923 cm\(^{-1}\) (red) and a minor peak at 3009 cm\(^{-1}\) (green). Similarly, the O–H stretching region can be fit with two peaks at 3224 cm\(^{-1}\) (blue) and 3398 cm\(^{-1}\) (cyan) with a relative magnitude of roughly 40:60%, respectively. These values match fairly well with previously reported spectra for potato.\(^{31}\) These results demonstrate that the 532 nm laser can be used as an effective photobleaching source as well as Raman excitation source.

3. CONCLUSIONS

The design and fabrication of a process chamber with controlled gas pressure was presented. While a levitator was utilized to hold a liquid droplet or a solid particle sample inside the chamber, a tunable wavelength light source was used as a reaction perturbation source. The physical and chemical changes at the gas–surface interface were monitored by a custom-built Raman spectrometer system. Three different gratings can be interchanged within seconds for a desired spectroscopic range and resolution. The gating capability of the detector allows operation at room light and selective detection of Raman photons from fluorescence. The performance of the device is demonstrated by degradation of polystyrene by UV light and photobleaching of potato by visible light. This versatile platform for \textit{in situ} reaction monitoring can be applied in varieties of fields, such as in astrochemistry-simulating heterogeneous atmospheric chemical reactions, in molecular biology in detection and monitoring of organic compounds, and in materials science to visualize surface-catalyzed reactions. The \textit{in situ} spectroscopic monitoring over time also allows investigation of kinetics of chemical reactions.

[FIGURE 6. (a) Raman spectra of a freshly sliced potato as a function of 532 nm photon irradiation times. Photobleaching of the sample allows reduction in fluorescence background and enhanced Raman signatures. (b) Baseline-corrected and deconvoluted spectra for 30 min of irradiation time in the high-frequency region.]
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