Dalton’s and Amagat’s laws fail in gas mixtures with shock propagation

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A shock propagating through a gas mixture leads to pressure, temperature, and density increases across the shock front. Rankine-Hugoniot relations correlating pre- and post-shock quantities describe a calorically perfect gas but deliver a good approximation for real gases, provided the pre-shock conditions are well characterized with a thermodynamic mixing model. Two classic thermodynamic models of gas mixtures are Dalton’s law of partial pressures and Amagat’s law of partial volumes. We measure post-shock temperature and pressure in experiments with nonreacting binary mixtures of sulfur hexafluoride and helium (two dramatically disparate gases) and show that neither model can accurately predict the observed values, on time scales much longer than that of the shock front passage, due to the models’ implicit assumptions about mixture behavior on the molecular level. However, kinetic molecular theory can help account for the discrepancy. Our results provide starting points for future theoretical work, experiments, and code validation.

INTRODUCTION

In 1802, John Dalton’s publication in *Memoirs of the Literary and Philosophical Society of Manchester* formulated the law of additive (or partial) pressures (1), stating that the total pressure in a non-reactive gas mixture—at constant temperature and volume—is equal to the sum of the partial pressures of the component gases.

In 1880, French physicist Emile Hilaire Amagat published his findings while researching the compressibility of different gases (2). Amagat’s law of partial volumes states that the total volume of a gas mixture is equal to the sum of the partial volumes each gas would occupy if it existed alone at the temperature and pressure of the mixture (3). While some advancements have been made in experiments of shock interaction with a single-component gas (4, 5), much less is known about multicomponent gas mixtures.

Shock interactions with gas mixtures are relevant to many engineering problems, including gas-cooled reactor power plants (6, 7), mixing processes in supersonic and hypersonic combustion (8–11), and astrophysical phenomena (12, 13). Our experiment was originally designed to determine which thermodynamic law (Dalton or Amagat) is more suitable for predicting properties of gas mixtures interacting with a planar shock wave. Post-shock properties are obtained using the Rankine-Hugoniot equations (14–18), which calculate post-shock values, such as pressure, temperature, and density, based on incident shock Mach number and upstream (preshock) conditions.

For a proper comparison of theoretical and experimental values of post-shock properties, pressure and temperature must be measured immediately before (downstream of) and immediately after (upstream of) the shock front. Pressure measurements are not difficult: High-frequency response pressure transducers (PTs) are readily available. Temperature measurements are more challenging. Thermocouples are intrusive and lack the necessary response time [O(10^{-6}s)]. Infra-red (IR) detectors, on the other hand, have ultrafast response times [O(10^{-8}s)] and are inherently nonintrusive. Here, we present temperature measurements using an InfraRed Associates mercury-cadmium-telluride (MCT), liquid nitrogen-cooled IR detector operating at 77 K, with a response time of 60 ns (6 × 10^{-8}s). Coupled with a Thorlabs stabilized broadband IR light source, with a color temperature of 1500 K, the MCT detector provides line-of-sight bulk temperature measurements both before and after the shock.

For our experiment, we selected two highly disparate gases forming a binary gas mixture: sulfur hexafluoride (SF₆) and helium (He). SF₆ and He are relatively inexpensive and nontoxic and have highly contrasting properties (molecular weight, viscosity, and specific heat), presenting an extreme (and hopefully easy to interpret) case of a mixture with easily distinguishable components. Two molar concentrations of each gas were chosen: 50%/50% (50/50) and 25%/75% (25/75) SF₆ to He, respectively.

The shock tube consists of two sections: driver and driven. The driver section is pressurized (with nitrogen) to a value depending on the desired strength of the shock wave. The driven section is pressurized with the test gas mixture, up to one of three different initial pressures: 39.3, 78.6, or 118 kPa (the average local atmospheric pressure in the laboratory is approximately 78.6 kPa). A thin-film polyester diaphragm separates the two sections. Once both sections have been evacuated using a vacuum pump, the driver is then filled with the driver gas, and the driven section is filled with our test gas mixture. When the driver and driven sections are at the desired pressure, a pneumatically driven stainless steel rod, tipped with a broad arrowhead, ruptures the diaphragm, sending a planar shock into the driven section. Four PTs, located on the top of the driven section, record the pressure pulse from the shock wave as it passes. The MCT detector and IR source are located coincident with the fourth downstream PT, providing nearly instantaneous temperature measurements immediately before and immediately after the shock. For details of experimental methods and theoretical evaluation of post-shock properties, see Materials and Methods.

Each gas mixture was tested at three driver pressures (1006, 1145, and 1282 kPa), and each of these was applied to three initial pressures (39.3, 78.6, and 118 kPa) in the driven section, providing experimental datasets at nine distinct pressure ratios (overpressures, Pᵣ), where
pressure ratio is defined as the ratio of driver pressure to the driven pressure ($P_r = P_{\text{driver}}/P_{\text{driven}}$). Figure 1A is an experimental signal trace at a pressure ratio of $P_r = 10.9$, for a 50%/50% mixture of SF$_6$/He.

The black line shows the signal (V) from the fourth downstream PT, and the magenta line shows the signal (V) from the MCT detector. Average values of the signals from each PT are determined in a 2-ms window after shock impact (dashed blue line). This average value, combined with a calibration curve for each PT (provided by the manufacturer), provides us with the post-shock pressure, $P_2$ (kPa). To obtain the incident shock velocity, $u_1$ (m/s), the distance between two successive PTs is divided by the time it takes the shock wave to travel between them. Measurements of temperature for the MCT are obtained by taking the maximum value of the signal within the same 2-ms window (Fig. 1A). We conducted a rigorous calibration experiment with the MCT detector and IR light source to determine post-shock temperature, $T_2$ (K).

**RESULTS**

We conducted multiple experiments for each gas mixture, with a minimum of six experiments for each of the nine pressure ratios. $P_r$ ranges from 8.54 to 32.6 for the 50/50 SF$_6$/He mixture and from 8.56 to 33.0 for the 25/75 SF$_6$/He mixture. Experimental results in Figs. 1 to 4 were produced by statistical analysis (19) of the dataset of measurements described above and correspond to mean values for each pressure ratio. Theoretical predictions are calculated as follows: The equations of state (EOS) are used to characterize the component gases. An ideal gas EOS is used for He, while a virial expansion—up to the fourth virial coefficient—is used for SF$_6$. Inputs for the EOS are the initial pressure ($P_1$) and temperature ($T_1$) in the driven section of the shock tube. Once the components have been characterized (the process includes calculations of density, specific heat ratio, thermal expansion coefficient, isothermal compressibility, and speed of sound for both He and SF$_6$), we use Dalton’s and Amagat’s laws to determine thermodynamic coefficients for the gas mixture. Specifically, we are interested in the specific

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Fig. 1. Time history of recorded signals and post-shock pressure dependence on initial pressure ratio. (A) Sample PT and MCT detector signal traces, showing locations of incident and reflected shocks, as well as the 2-ms averaging window (dashed blue lines) used in data analysis. The black line is the signal trace from the fourth downstream PT, and the magenta line is the MCT signal trace. (B to D) Post-shock pressure, $P_2$, versus pressure ratio, $P_r$, for a 50/50 (by mole) binary mixture of SF$_6$ and helium, respectively. (E to G) Post-shock pressure versus pressure ratio for a 25/75 binary mixture of SF$_6$ and helium. In all panels (B to G), black symbols correspond to experimental values, blue symbols represent Dalton’s law predictions, and red symbols denote Amagat’s law predictions. Vertical error bars correspond to total uncertainty in post-shock pressure $P_2$. Horizontal error bars are omitted as they do not extend past the physical size of the symbols. The driven pressure associated with each dataset is displayed above the corresponding panel.
heat ratio ($\gamma$) and the speed of sound ($a$). These values, combined with the incident shock speed ($u_i$), are used as inputs to the Rankine-Hugoniot equations, which are then used to determine post-shock temperature and pressure. Predictions displayed in Figs. 1 to 4 correspond to mean values for each pressure ratio. This process, and the corresponding statistical analysis, is explained in detail in Materials and Methods.

Figure 1 shows plots of post-shock pressure ($P_2$) versus pressure ratio ($P_r$) for the 50/50 mix (Fig. 1, B to D) and the 25/75 mix (Fig. 1, E to G). The corresponding initial pressures (in the driven section) are displayed above each panel (Fig. 1, B to G). Vertical error bars correspond to total uncertainty in post-shock pressure. Horizontal error bars corresponding to uncertainty in pressure ratio ($P_r$) do not extend past the physical size of the symbols and are therefore omitted. Note the strong disagreement not only between experimental values and theoretical predictions but also between the two mixture concentrations. Not only do these results disagree with both Dalton’s law and Amagat’s law well beyond experimental uncertainty, the disagreement varies with experimental value, leaving no clear answer: Which law describes the experiment better overall?

Figure 2 shows the relationship between post-shock pressure ($P_2$) and incident shock speed ($u_i$) for the 50/50 SF$_6$/He mixture (Fig. 2A) and the 25/75 SF$_6$/He mixture (Fig. 2B). In both plots, black symbols denote experimentally measured values, blue symbols correspond to Dalton’s law predictions, and red symbols represent Amagat’s law predictions. Vertical error bars in both plots denote total uncertainty in post-shock pressure. Horizontal error bars on experimental values correspond to total uncertainty in incident shock speed. Note that horizontal error bars for both Dalton’s and Amagat’s laws are withheld, as the uncertainty in incident shock speed for both thermodynamic laws is identical to those for experimental measurements.

Black arrows in Fig. 2 (A and B, upper left corner) show the direction of increasing driver pressure, from 1006 to 1282 kPa. While it seems that experimental values are closer to Dalton’s law predictions for the 78.6- and 118-kPa datasets, data for 39.3 kPa is inconclusive. Again, note the discrepancies between respective initial pressures and between the two mixtures. The next step is to evaluate post-shock temperature and compare it with theoretical predictions.

Figure 3A is a plot of post-shock temperature ($T_2$) versus pressure ratio for the 50/50 SF$_6$/He mixture, and Fig. 3B is the same plot for the 25/75 SF$_6$/He mixture. Vertical error bars for experimental measurements and theoretical predictions correspond to total uncertainty in post-shock temperature. Note that the inputs used to calculate predictions for both Dalton’s and Amagat’s laws are the initial pressure ($P_1$) and temperature ($T_1$) in the driven section of the shock tube, which are measured values. Therefore, these predictions will not fit smoothly on a curve; the curve fits shown in Fig. 2 (A and B) serve merely as visual aids.

Again, there is strong disagreement between experimentally obtained values of post-shock temperature and theoretical predictions. For the 50/50 mixture, it seems that experimental values are closer to Dalton’s law predictions, and for the 25/75 mixture, experimental values are closer to Amagat’s law predictions; these discrepancies are not random. What could be responsible for this systematic disagreement?

**DISCUSSION**

A compelling theoretical analysis of finite-strength shock propagation through a binary gas mixture was published by Sherman (20) for inert gas mixtures consisting of argon (Ar) and He (for a range of molar concentrations of each component). Using a continuum approach, Sherman focused on determining the structure of a shock wave of arbitrary strength, which includes ordinary diffusion, baro-diffusion, and thermal diffusion effects. He concluded that barodiffusion speeds up the heavier component and slows down the lighter
component relative to the mass velocity of the mixture (20). He also
determined that thermal diffusion will have the opposite effect, slow-
ing down the heavier component and speeding up the lighter one.
Therefore, thermal diffusion would (at least partially) counteract
barodiffusion within the shock wave. Sherman calculated his results
through numerical integration of the Navier-Stokes equations.
A primary assumption associated with this analysis was always im-
posing thermal equilibrium between species through the shock wave.
However, Sherman indicates that these assumptions (and corre-
spanding analysis) may not be valid for strong shock waves, or for
mixtures with large molecular mass ratios. He states that one might
intuitively expect that the maximum shock strength for which these
calculations give realistic predictions would be reduced (somewhat
intuitively) as the molecular mass ratio increases because of the
difficulty in maintaining thermal equilibrium between the gas com-
ponents (20).

In 1967, Bird (21) produced an interesting attempt to model shock
propagation through a binary mixture of 50/50 Ar-He, representing
the gas molecules as rigid elastic spheres—with the appropriate masses
and diameters—and compared his results with the analytical predic-
tions of Sherman. His model did not have the same temperature con-
straints as Sherman and predicted even greater differences in velocity,
temperature, and concentration profiles than the analytical profiles of
Sherman. Furthermore, Bird’s findings suggest that the temperature
non-equilibrium between species increases with low concentrations
of the heavy gas component and that this (non-equilibrium) can per-
sist for a considerable distance downstream of the shock (21). For
reference, the molecular mass ratio with respect to binary mixtures of
Ar-He is approximately 10, while the molecular mass ratio asso-
ciated with the current work (SF₆-He) is around 36.5.

The studies conducted by Sherman and Bird reasonably concluded
that shock propagation through gas mixtures with relatively massive
molecules would cause differences in their behavior, but the results
only pertained to gas molecule velocity, concentration, and tempera-
ture profiles—on time (and length) scales much smaller than these
considered in the current work. Perhaps the results shown in Figs. 1
to 3, taken in context with these early studies, suggest a different ex-
planation, beyond experimental uncertainty.

Is a kinetic molecular theory (KMT) (22, 23) explanation possible?
Dalton’s and Amagat’s laws predate KMT, but each law makes implicit
assumptions about reversibility. Both laws assume thermodynamic
equilibrium. However, while Dalton’s law assumes that the gases are
always perfectly mixed, Amagat’s law assumes that the gases will se-
parate over time. This assertion is interesting because that is exactly what
we see in experiments. The molecular mass of SF₆ is approximately
36.5 times greater than that of He. If the gases are not constantly mixed,
then they will begin to separate (under normal conditions, the gases
will separate within a few minutes). Before taking measurements de-
scribed here, our initial guess was that Amagat’s law might provide a
better prediction for these dissimilar gases. The time scale of that sepa-
ration, however, is much longer than that of the experiment associated
with the shock passage, thus leading to both Dalton’s law (infinite sepa-
ration time) and Amagat’s law (the gases in the mixture are in effect
always separated) failing to produce a match with experiment.

What relaxation time scale is relevant for the equilibrium status of
a shocked gas mixture? Relaxation time is defined as the time within
which a perturbed gas will reach statistical (thermodynamic) equi-
librium (24). In binary gas mixtures whose constituents have widely
different molecular masses ($M_{SF_6} \gg M_{He}$), disparate relaxation times
are manifest, governing the approach to equilibrium of the various
degrees of freedom (25). According to Mora and Fernández-Feria
(25), for such mixtures, the process of equilibration can be charac-
terized by three different relaxation times: two for self-equilibration
of the component gases and a third one associated with the slower pro-
cess of interspecies equilibration. For this analysis, these relaxation
times are the post-shock mean free time (or average time between mo-
lecular collisions) for each of the gas components ($\tau_{He}$ and $\tau_{SF_6}$) and the change in collision time between them ($\Delta \tau = \tau_{He} - \tau_{SF_6}$). What is

| $P_e$ (50/50) | $n_{P_2}$ | $v_{P_2}$ | $t_{P_2}$ | $n_{He}$ | $v_{He}$ | $t_{He}$ | $n_{SF_6}$ | $v_{SF_6}$ | $t_{SF_6}$ |
|--------------|----------|----------|----------|---------|---------|---------|----------|---------|---------|
| 32.61        | 15 6     | 2.4469   | 6 5      | 2.7764  | 5 4      | 2.7764  |
| 10.88        | 20 10    | 2.2281   | 6 5      | 2.7764  | 5 4      | 2.7764  |
| 29.24        | 20 16    | 2.1199   | 5 4      | 2.7764  | 5 4      | 2.7764  |
| 14.60        | 21 11    | 2.2010   | 5 4      | 2.7764  | 5 4      | 2.7764  |
| 9.70         | 21 10    | 2.2281   | 5 4      | 2.7764  | 5 4      | 2.7764  |
| 12.83        | 22 15    | 2.1314   | 4 3      | 2.7764  | 6 5      | 2.5706  |
| 8.54         | 22 17    | 2.1098   | 6 5      | 2.5706  | 5 4      | 2.5706  |
remarkable about this method is that the mean free time is temperature and pressure dependent (see Materials and Methods).

Figure 4 is a plot of the change in collision time, $\Delta t$ (in picoseconds), versus incident shock velocity ($u_i$) for the 50/50 SF$_6$/He mixture (Fig. 4A) and the 25/75 SF$_6$/He mixture (Fig. 4B). Symbol type and color in Fig. 4 have been arranged similar to Fig. 2. Again, the black arrow in both plots points in the direction of increasing driver pressures. Note again the systematic discrepancies in these data, consistent with the spread of experimental data points. The negative values for $\Delta t$ are due to the fact that the average collision time for SF$_6$ is an order of magnitude greater than that of He. While Figs. 1 to 3 show discrepancies produced by Dalton’s and Amagat’s laws in predicting post-shock properties (pressure and temperature) on a macroscopic scale, Fig. 4 provides a context for these discrepancies, relating them to the simplest quantitative parameter describing the disparity between component gas behavior on a microscopic scale. Therefore, if the component gases behave differently, i.e., have a large difference in response time, on a macroscale, it is not reasonable to assume that these discrepancies manifest on a macroscale? This hypothesis is reinforced by the results obtained from Sherman (20) and Bird (21), especially considering the large molecular mass difference between the species.

The simple explanation that KMT can provide is that differences in the response time of the molecules account for the disagreement between theory and experiment. This explanation appears to agree with our data, at least qualitatively. These observations show that Dalton’s and Amagat’s laws fail to accurately describe the behavior of a gas mixture that underwent shock acceleration, with implications that the same failure can manifest in other non-equilibrium situations.

### MATERIALS AND METHODS

#### Experimental setup

Two gas mixtures were tested: 50%/50% and 25%/75% SF$_6$ to He, by mole, respectively. Experiments were conducted at the Shock Tube Facility in the Mechanical Engineering Department at the University of New Mexico. The shock tube itself is approximately 5.2 m long with a 2-m-long driver section and a 3.2-m-long driven section (7.62 cm inside square cross-section).

To begin an experiment, we separated the driver and driven sections of the shock tube with a thin-film polyester diaphragm. Both sections were then evacuated using an ITT Pneumatic vacuum pump to ~78.6 kPa. The driven (test) section was filled with the test mixture to a predetermined pressure (39.3, 78.6, or 118 kPa). The driver was filled with nitrogen to another predetermined pressure (1006, 1145, or 1282 kPa). The pressure in the driver section depends on the desired strength (or Mach number) of the shock wave (2). Once the pressure in both sections has stabilized, a pneumatically driven stainless steel rod tipped with a broad arrowhead punctures the diaphragm, sending a normal shock down the length of the driven section. Four PT’s (with a response time of ~1 µs), located on the top of the driven section (~0.8 m apart), record the pressure history of the shock wave as it passes. These data can then be used to determine the velocity of the shock wave, $u_i$, and the post-shock pressure, $P_m$. The MCT detector and IR source were placed on opposite sides of the shock tube, perpendicular to the optical axis and located coincident with the fourth downstream PT. Two zinc selenide (ZnSe) optical windows were placed on either side of the shock tube, mounted flush with the inside of the test section. These optical windows were used to create an airtight, unobstructed light path from the IR source, through the test gas in the driven section, to the sensor on the MCT detector. They were also used as broadband filters to decrease the wavelength range of the incoming light from the IR source to between 7 and 12 µm (the IR source outputs light between 0.5 µm and approximately 20 µm). A germanium notch filter, mounted on the MCT detector itself, further reduces the range of incoming light to between 7.5 and 9.0 µm. Note that the target range for these experiments was chosen to be between 7.5 and 8.5 µm, on the basis of the IR absorption spectrum of SF$_6$ (26).

Calibration curves that relate the signal (V) from the PTs to pressure (kPa) in the driven section were provided for each transducer by the manufacturer. For the MCT detector, a calibration experiment was conducted to determine the relationship between signal (V) and temperature (K). This experiment used an aluminum cylinder with ZnSe optical windows mounted on each side, with components and geometry identical to that of the shock tube setup described above. The cylinder chamber was evacuated with a vacuum pump and filled with the test gas mixture at a prescribed pressure [target pressures were determined from previous experiments (27)]. A helical coil resistance heater, placed on the inside of the calibration cylinder, was then activated to increase the temperature of the test gas to a predetermined value. A Vincent Associates Uniblitz LS6 laser shutter (6-mm aperture, 1.7-ms open time) was placed along the optical axis in front of the MCT detector, which effectively simulates an instantaneous increase in temperature, as would be seen by the sensor when the shock wave passes (2). Once the laser shutter was activated, the signal from the detector was recorded and used as a baseline for that pressure-temperature (P-T) combination. For each gas mixture, data were obtained at up to 25 P-T combinations, with a minimum of six measurements at each combination. This method provided ample data to determine a calibration curve for each gas mixture.

#### Thermodynamic models

The following theoretical analysis was used to determine post-shock pressure and temperature for both Dalton’s and Amagat’s laws. EOS

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| $P_T$ (25/75) | $n_p$ | $v_p$ | $t_p$ | $u_i$ | $t_m$ | $n_r$ | $v_r$ | $t_r$ |
|----------------|-------|-------|-------|-------|-------|-------|-------|-------|
| 33.03          | 20    | 12    | 2.1788| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
| 16.32          | 24    | 20    | 2.0860| 4     | 3     | 3.1824| 4     | 4     | 2.7764|
| 10.88          | 18    | 8     | 2.3060| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
| 29.16          | 28    | 21    | 2.0796| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
| 14.59          | 19    | 10    | 2.2281| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
| 9.73           | 21    | 12    | 2.1788| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
| 25.59          | 21    | 11    | 2.2010| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
| 12.79          | 20    | 13    | 2.1604| 4     | 3     | 3.1824| 6     | 5     | 2.5706|
| 8.56           | 18    | 10    | 2.2281| 6     | 5     | 2.5706| 6     | 5     | 2.5706|
are needed to characterize the gas components; the Ideal Gas (Eq. 1) EOS was used for He, and a virial expansion (Eq. 2) was used for SF₆

\[ P_v = RT \]  
(1)

\[ P = \frac{RT}{v} \left( 1 + \frac{B(T)}{v} + \frac{C(T)}{v^2} + \frac{D(T)}{v^3} + \frac{E(T)}{v^4} \right) \]  
(2)

Here, \( P \) is the pressure (N/m²), \( v \) is the specific volume (m³/mol), \( R = 8.314 \) J/mol·K is the universal gas constant, \( T \) is the temperature (K), \( B(T) \) is the second virial coefficient, \( C(T) \) is the third virial coefficient, and so forth (28). Algebraic expressions of the temperature provided by the hard-core square well model intermolecular potential are used to represent \( B(T) \) and \( C(T) \), while \( D(T) \) and \( E(T) \) are represented as polynomial functions of the inverse temperature (29)

\[ B(T) = b_0[1 - (\lambda^2 - 1)\Delta] \]  
(3)

\[ C(T) = \frac{1}{8} b_0^2 (5 - c_1 \Delta - c_2 \Delta^2 - c_3 \Delta^3) \]  
(4)

\[ D(T) = \sum_{n=0}^{3} d_n T^{-n} \]  
(5)

\[ E(T) = \sum_{n=0}^{3} e_n T^{-n} \]  
(6)

where \( \Delta = e^{\epsilon/k_b T} - 1 \) (\( k_b \) is Boltzmann’s constant) and the coefficients \( c_1, c_2, \) and \( c_3 \) are given by

\[ c_1 = \lambda^6 - 18 \lambda^4 + 32 \lambda^3 - 15 \]  
(7)

\[ c_2 = 2 \lambda^6 - 36 \lambda^4 + 32 \lambda^3 + 18 \lambda^2 - 16 \]  
(8)

\[ c_3 = 6 \lambda^6 - 18 \lambda^4 + 18 \lambda^2 - 6 \]  
(9)

Values for \( d_n \) (\( n = 1,2,3 \)), \( e_n \) (\( n = 1,2,3 \)), \( b_0, \lambda, \) and \( \epsilon/k_b \) are provided by Hurly et al. (29).

Inputs for the EOS are the initial pressure (\( P_1 \)) and temperature (\( T_1 \)) of the test gas in the driven section. Once the component gases have been characterized, we used Dalton’s law and Amagat’s law to determine thermodynamic coefficients for the mixture. We were specifically looking for the speed of sound, \( a \), and the specific heat ratio, \( \gamma \). These variables, coupled with the incident shock speed \( u_i \) (determined from experiment) were used as inputs for the Rankine-Hugoniot equations (Eqs. 10 to 13), which relate post-shock properties in terms of initial conditions (preshock) and incident shock Mach number \( M_1 \)

\[ M_1 = \frac{u_i}{a} \]  
(10)

\[ M_1^2 = \frac{M_1^2}{\gamma} + \frac{2(\gamma - 1)}{2\gamma(\gamma - 1)}M_1^2 - 1 \]  
(11)

\[ T_1 \left( 1 + \frac{\gamma - 1}{2} M_1^2 \right) = T_2 \left( 1 + \frac{\gamma - 1}{2} M_2^2 \right) \]  
(12)

\[ \frac{P_1}{P_2} = \frac{1 + \gamma M_2^2}{1 + \gamma M_1^2} \]  
(13)

The subscripts 1 and 2 correspond to conditions before and after the shock, respectively. Once the post-shock temperature (\( T_2 \)) and pressure (\( P_2 \)) for each thermodynamic law (Dalton and Amagat) have been calculated, we can directly compare the results with experimental values. All theoretical calculations and analysis of experimental data (including statistical analysis) were performed in MATLAB.

**Kinetic molecular theory**

Central to KMT are the following assumptions (30):

1. The size of the particle is negligibly small; i.e., the particles themselves occupy no volume, even though they have mass. At the maximum concentration of molecules in our experiments, the Van der Waals correction to pressure associated with molecular volume would not exceed 2.5%.

2. The average kinetic energy of a particle is proportional to the temperature (K).

3. Particle collisions are perfectly elastic; they may exchange energy, but there is no overall loss of energy.

The following equations were used for kinetic theory analysis. The mean free path (\( l \)) is given by

\[ l = \frac{k_b T}{\sqrt{2\pi d^2 P}} \]  
(14)

where \( k_b \approx 1.381 \times 10^{-23} \) m²·kg·s⁻²·K⁻¹ is the Boltzmann constant, \( T \) is the temperature (K), \( d \) is the kinetic diameter, which is 0.260×10⁻⁹ m for He and 0.550×10⁻⁹ m for SF₆, and \( P \) is the pressure (Pa).

Mean molecular speed (\( \mu_m \)) is obtained via

\[ \mu_m = \sqrt{\frac{8 TR_s}{\pi}} \]  
(15)

where \( R_s \) is specific gas constant: for He, \( R_s = 2.0773 \times 10^{-5} \) J/kg·K, and for SF₆, \( R_s = 56.9269 \) J/kg·K.

Average collision time (\( \tau \)) is calculated using

\[ \tau = \frac{l}{\mu_m} \]  
(16)

The change in average collision time, \( \Delta \tau \), is simply the average collision time of He, \( \tau_{He} \), minus the average collision time of SF₆, \( \tau_{SF_6} \)

\[ \Delta \tau = \tau_{He} - \tau_{SF_6} \]  
(17)

**Statistical analysis**

A comprehensive statistical analysis was performed on all experimental data, according to steps outlined by Wheeler and Ganji (19). Statistical analysis on all measurements (pressure, temperature, and velocity) begins with outlier rejection using the Modified Thompson
and the largest value is selected. This value is compared with the
product of \( t \) (tabulated with respect to \( n \)) times the SD \( S \). If the value of \( \delta \) exceeds \( tS \), then this value can be rejected as an outlier (only one value is eliminated for each iteration). The mean and SD of the
remaining values are then recomputed, and the process is repeated
until no more outliers exist. Note that \( n \) decreases with each outlier
rejection.

Pressure measurements were obtained using multiple devices.
Therefore, an estimation of the combined degrees of freedom according
to the Welch-Satterthwaite formula (Eq. 18) is necessary (19).

\[
\nu_x = \frac{\left( \sum \frac{S_i^2}{n_i} \right)^2}{\frac{1}{n} \sum \frac{1}{v_i}}
\]  

(19)

where \( v_i \) is the degrees of freedom for the measuring device and \( \nu_x \) is the
value of the combined degrees of freedom for variable \( x \). Degrees of freedom
for temperature and velocity measurements are simply \( \nu_x = n - 1 \).
When \( \nu_x \) has been determined, Student’s \( t \) distribution value (1) is found
on the basis of a 95% level of confidence. The total random uncertainty
in the mean value \( P_x \) is computed by

\[
P_x = \frac{S_x}{\sqrt{n}}
\]  

(20)

where \( S_x/\sqrt{n} \) is the estimate of the SD of the mean.

Systematic uncertainty for each variable is determined using the
mean value for the measurement and manufacturer-supplied
information, such as linearity, hysteresis, and uncertainty in the
measuring device. Sources of systematic error are PTs, pressure
gauges, oscilloscopes, and the MCT detector. Total systematic error
is given by

\[
B_x = \left( \sum \frac{B_i^2}{n_i} \right)^{1/2}
\]  

(21)

where \( B_i \) is the systematic error for measuring device \( i \). Once all
sources of random and systematic error have been determined, the
total uncertainty in the mean \( (W_x) \) is given by

\[
W_x = \left( B_x^2 + P_x^2 \right)^{1/2}
\]  

(22)

The mean value for a given set of measurements is used in all plots,
and error bars represent total uncertainty (\( \pm \)) in the mean value.

Table 1 details the number of measurements \( n \), degrees of freedom
\( \nu \) (combined or otherwise), and the \( t \) distribution value for
post-shock pressure \( (P_s) \), incident shock speed \( (u_i) \), and post-shock
temperature \( (T_s) \) for the 50/50 binary mixture of \( \text{SF}_6 \) and He. \( P \)_s
denotes the pressure ratio according to mean values of driver and driven
pressures for a given set of experiments. Table 2 details statistics for
a 25/75 mixture of \( \text{SF}_6 \) and He. All statistical analysis was performed
in MATLAB.

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