Evolution of Combustion-Generated Particles at Tropospheric Conditions

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ABSTRACT

This paper describes particle evolution measurements taken in the Particulate Aerosol Laboratory (PAL). The PAL consists of a burner capable of burning jet fuel that exhausts into an altitude chamber that can simulate temperature and pressure conditions up to 13,700 m. After presenting results from initial temperature distributions inside the chamber, particle count data measured in the altitude chamber are shown. Initial particle count data show that the sampling system can have a significant effect on the measured particle distribution: both the value of particle number concentration and the shape of the radial distribution of the particle number concentration depend on whether the measurement probe is heated or unheated.

Nomenclature

\( B \) Buoyancy flux
\( d_0 \) Nozzle diameter
\( d^* \) Equivalent source diameter, \( d^* = d_0 \sqrt{\frac{\rho_0}{\rho_\infty}} \)
\( g \) Gravitational constant
\( J \) Momentum flux
\( l_c \) Coflow length scale
\( l_M \) Morton length scale
\( m \) Mass flux
\( Q \) Volume flux
\( R \) Richardson number
\( r \) Radial coordinate
\( Re \) Reynolds number
\( T \) Temperature
\( u \) Velocity
\( x \) Axial distance from the jet exit plane
\( x_E \) Axial distance from the virtual origin to the jet exit plane
\( \rho \) Density
\( \theta \) Normalized temperature difference, \( \frac{T-T_\infty}{T_0-T_\infty} \)
\( \zeta \) Particle number concentration

Subscripts

\( 0 \) Nozzle exit
\( CL \) Centerline
\( \infty \) Chamber
1 Introduction

Although particulate emissions from aircraft engines were initially of concern because of the visible smoke [1, 2], recent research has focused on the health and environmental effects of ultrafine particulate emissions. Due to the negative health effects of ultrafine particles, the regulations on particles smaller than 2.5 microns have become increasingly stringent [3]; most particulates emitted by aircraft are smaller than 2.5 microns [1, 4].

A major environmental effect of aircraft particulate emissions is their effect on the the formation and properties of cirrus clouds in the troposphere; however, this effect is not well understood [5]. Several areas require further research, including the microphysical and chemical processes governing the evolution of aviation particulates. Previous work has sought to characterize aircraft engine emissions using combustion rigs, on-ground engine tests [4, 6–8], and measurements taken in the aircraft engine plumes at altitude [6]. The work described in this paper complements these efforts.

This paper describes measurements taken in the Particulate Aerosol Laboratory (PAL) at NASA Glenn Research Center. In this unique facility, a combustor capable of burning jet fuel exhausts into an altitude chamber capable of simulating temperature and pressure conditions up to 45,000 ft, allowing the particle evolution to be studied at altitude conditions. A scanning mobility particle sizer (SMPS) measures the particulate profile at the combustor exit and in the altitude chamber. The probe used to sample the exhaust plume in the altitude chamber can be placed at discrete axial positions that range from the exhaust nozzle exit to the top of the altitude chamber. This paper will present initial measurements of the particle size distribution taken from the combustor and measurements of the particle size distribution taken in the altitude chamber.

2 Facilities and Instrumentation

2.1 Facility Description

The Particulate Aerosol Laboratory (PAL) consists of a burner connected to an altitude chamber, as shown in Figure 1. The burner is capable of burning conventional jet fuels and alternative fuels such as Fischer-Tropsch. A 1.6-m long transition pipe...
The chamber diameter is 0.597-m, and the height of its cylindrical test section in 1.83-m. Four window ports, each 1.52-

m tall by 0.102-m wide, are located 90° apart from each other. The window ports can be fitted with windows, instrumentation plates, or blanks. When a window is installed, two panes of glass are used, and the space in between the panes is kept at vacuum; this prevents condensation on the windows and also provides good insulation. (When a blank or instrumentation plate is used, it is a single thickness of aluminum and does not provide good insulation.) For the measurements reported here, three ports contained windows and one port contained an instrumentation plate; Figure 1a shows the instrumentation plate that was used during these measurements.

2.2 Instrumentation

A TSI scanning mobility particle sizer (SMPS) system model 3936NL76 — consisting of a TSI model 3776 condensation particle counter (CPC), a TSI model 3080 electrostatic classifier, and a TSI model 3085 nano differential mobility analyzer — was used to measure the particle size distribution. The CPC was also used alone to measure the particle count.

A blunt-tipped sampling probe with a 0.32-cm inner diameter was inserted into the transition pipe facing normal to the flow. This probe was used to sample the particle distribution at the burner exit. To minimize agglomeration and condensation in the sampling system, the combustion products were diluted with dry nitrogen slightly downstream of the probe tip.

A blunt-tipped sampling probe was also used to measure the particle count in the altitude chamber. The probe face was parallel to the primary flow direction except for the measurements at 0.18 m downstream of the nozzle exit at 12,190 m standard day conditions; in this single case, the probe face was approximately normal to the primary flow direction. This probe had a 0.4-cm inner diameter and could be covered with heat tape to prevent ice formation inside the probe. It was connected to a translation stage that allowed it to be moved radially.

2.3 Test Conditions

Four burner conditions (see Table 1) and four altitude chamber conditions were studied. The altitude chamber conditions corresponded to standard day temperatures and pressures at 6,100 m; 9,140 m; and 12,190 m [9] as well hot day temperature and pressure conditions at 9,140 m [10] (see Table 2). The chamber Reynolds number indicates that the flow was turbulent for all cases. As Figure 2 shows, the baseline temperature in the altitude chamber (when no combustion products are added) corresponded to standard day temperatures and pressures at 13,700 m.
TABLE 1: Burner conditions

<table>
<thead>
<tr>
<th>Condition Number</th>
<th>Fuel/Air Flow Rate</th>
<th>Total Mass</th>
<th>$l_M$ at</th>
<th>$x_E$ at</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>kg/min m m m cm cm cm cm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.045 0.166 2.16 2.82 3.02 4.55 3.0 3.1 2.9 2.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>0.032 0.230 2.98 3.88 4.18 6.28 3.0 3.1 2.9 2.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>0.017 0.342 4.50 5.88 6.28 9.43 3.1 3.2 3.0 2.9</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>4</td>
<td>0.007 0.336 4.99 6.65 6.75 9.95 3.5 3.6 3.4 3.3</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

FIGURE 3: Combustion product temperature as combustion products exit the nozzle into the altitude chamber at x=0. For all cases the temperature rapidly returned to the baseline temperature as the radial distance $r$ increased, so only the region near the nozzle exit is shown.

3 Facility Temperature Distribution

Because the particle evolution depends strongly on temperature, the temperature distribution in the altitude chamber was characterized; the temperature distribution depends on the conditions at the nozzle exit, the conditions in the altitude chamber, and the type of flow field that is established. From several nozzle diameters downstream of the nozzle exit to the axial location where the effects of the altitude chamber walls become important, the flow field in the PAL should act as either a buoyant jet or a momentum-dominated jet. The relative effects of jet exit momentum and buoyancy are usually expressed in terms of the Richardson number, $R$, or the ratio of the axial distance from the nozzle exit, $x$, to the Morton length scale, $l_M$ [11, 12]. These are defined in terms of the jet volume flux, $Q$, the jet momentum flux, $J$, and the jet buoyancy flux, $B$, as:

$$R = \frac{Q(B/\rho_\infty)^{\frac{1}{2}}}{(J/\rho_\infty)^{\frac{1}{4}}}$$

$$l_M = \frac{(J_0/\rho_\infty)^{\frac{1}{2}}}{(B_0/\rho_\infty)^{\frac{1}{2}}}$$

where

$$Q = \int_0^\infty \bar{u}(x,r) 2\pi r dr,$$

$$B = \int_0^\infty g \Delta \rho \bar{u}(x,r) 2\pi r dr,$$

$$J = \int_0^\infty \rho \bar{u}^2(x,r) 2\pi r dr.$$

$u$ is the axial velocity, $\rho$ is the density, and the subscripts 0 and $\infty$ refer to conditions at the jet exit and in the altitude chamber, respectively. In the jet-like limit, $R$ and $x/l_M$ are small; in the plume-like limit, $R$ approaches 1 and $x/l_M$ is large. Papanicolaou and List [12] have shown that $R \propto x$ in the jet-like limit and $R$ is constant in the plume-like limit. They have also shown that flows with $x/l_M < 1$ exhibit jet-like scaling and flows with $x/l_M > 5$...
For the measurements reported here, the axial locations of interest range from several nozzle diameters to 71 cm downstream of the nozzle exit. In this region, $R$ is small and $x/l_{M} \ll 1$ (see Table 1), so the effects of buoyancy should not be significant. A similar length scale analysis of the effects of the cold nitrogen coflow [13] shows that the coflow effects should also be small. Therefore, the flow field is momentum-dominated and should follow nonbuoyant jet scaling laws.

For momentum-dominated jets, it can be shown [14,15] that a conserved scalar, such as the normalized temperature difference, $\theta = \frac{T_{0} - T_{\infty}}{T_{0} - T_{\infty}}$, is inversely proportional to the normalized distance from the virtual origin, $(x + x_{E})/d^{*}$, where $x$ is the distance from the jet exit, $x_{E}$ is the distance from the virtual origin to the jet exit, and $d^{*} = d_{0} \sqrt{\frac{\beta_{0}}{\rho_{\infty}}}$. Diez & Dahm [11] \(^1\) show that for a momentum-dominated or buoyant jet the distance $x_{E}$ that the virtual origin is upstream of the nozzle exit is given by:

$$ x_{E} = l_{M} \left\{ \left( \frac{l_{2}}{l_{1}} \right)^{2} \frac{r^{2}}{c_{\delta} c_{u,j}} \right\}^{-\frac{1}{2}}, $$

where

$\begin{align*}
    l_{1} &= 0.262, \\
    l_{2} &= 0.131, \\
    c_{\delta} &= 0.36, \\
    c_{u,j} &= 7.2
\end{align*}$

In other words, $\theta = c_{\theta} (\frac{x + x_{E}}{d^{*}})^{-1}$, where $c_{\theta}$ is approximately 5.4 \(^{1}\)The formula given here does not match the one given by equations 42 and 47 in [11]. There seem to be several typographical errors in [11], with the missing exponent of -1/2 in equation 47 being the most important. The missing exponent can be found by following the derivation described in the text of [11] and substituting equation 46 into equation 44.

\(^{1}\)Furthermore, momentum-dominated jets are self-similar, so a conserved scalar normalized by its centerline value is a function of the radius normalized by the jet width, $\delta$; that is, $\theta/\theta_{CL} = f(r/\delta)$. For both momentum-dominated and buoyant jets, $\delta$ is proportional to axial distance from the virtual origin; therefore, $\theta/\theta_{CL} = f^{\prime} (\frac{r}{x + x_{E}})$. Figures 4a and 4b support this scaling: Figure 4a is consistent with $\theta/\theta_{0} = 5.4(\frac{x + x_{E}}{d^{*}})^{-1}$ scaling until the effect of the altitude chamber wall becomes important at $\frac{x + x_{E}}{d^{*}} \approx 130$. Figure 4b shows that $\theta/\theta_{CL}$ is a function of $r/(x + x_{E})$ for $r/(x + x_{E})$ values that are sufficiently far from the chamber walls. (When interpreting Figure 4b, note that the temperature increase due to the chamber walls is more important on the poorly-insulated instrument plate side of the chamber ($+r$) than on the relatively well-insulated window side ($-r$) and that as the axial distance increases the $r/(x + x_{E})$ value at which the wall effects become important decreases.) Also shown in Figure 4b are a curve fit to the data, the normalized concentration profile, $\xi/\xi_{CL}$, used by Diez & Dahm [11], and the normalized velocity profile, $u/\bar{u}_{CL}$, used by Hussein et al. [17]. The curve fit used to model the current data and the profiles used by the Diez & Dahm and Hussein et al are of the form $\xi = e^{-\alpha (\frac{r}{x + x_{E}})^{2}}$, where $\xi$ is $\theta/\theta_{CL}$, $\xi/\xi_{CL}$, or $u/\bar{u}_{CL}$. The width parameter $\alpha$ is the value of $\theta/\xi_{CL}$ is between the values for $\xi/\xi_{CL}$ and $u/\bar{u}_{CL}$, further indicating that the jet scaling is appropriate for the normalized temperature difference $\theta$.

### 4 Initial Particle Evolution Measurements

Particle measurements were taken both in the transition pipe (see Figure 1) and in the altitude chamber for burner condition 1 from Table 1 and at all chamber conditions listed in Table 2. For the particle measurements made in the transition pipe, the sample was diluted with nitrogen with a dilution ration of 1.91 to prevent condensation and agglomeration in the sampling line; the sample

<table>
<thead>
<tr>
<th>Altitude</th>
<th>Altitude</th>
<th>Day Type</th>
<th>Temperature</th>
<th>Pressure</th>
<th>Density</th>
<th>Chamber Reynolds Number</th>
<th>Nominal Chamber Coflow Velocity</th>
</tr>
</thead>
<tbody>
<tr>
<td>(m)</td>
<td>(ft)</td>
<td></td>
<td>(K)</td>
<td>(kPa)</td>
<td>(kg/m³)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6,100</td>
<td>20,000</td>
<td>Standard</td>
<td>249</td>
<td>46.601</td>
<td>0.653</td>
<td>17,900</td>
<td>0.7</td>
</tr>
<tr>
<td>7,620</td>
<td>25,000</td>
<td>Hot</td>
<td>259</td>
<td>37.650</td>
<td>0.490</td>
<td>18,700</td>
<td>1.0</td>
</tr>
<tr>
<td>9,140</td>
<td>30,000</td>
<td>Standard</td>
<td>228</td>
<td>30.149</td>
<td>0.459</td>
<td>25,800</td>
<td>1.4</td>
</tr>
<tr>
<td>12,190</td>
<td>40,000</td>
<td>Standard</td>
<td>217</td>
<td>18.823</td>
<td>0.303</td>
<td>32,100</td>
<td>2.5</td>
</tr>
</tbody>
</table>
FIGURE 4: (a) Normalized jet centerline temperature $\theta_{CL}$ as a function of normalized axial distance $(x + x_E)/d^*$ and (b) the normalized temperature distributions as a function of the normalized radial distance. The symbol color indicates the altitude conditions, the ‘outer’ symbol indicates the axial position, and the ‘inner’ symbol indicates the fuel-air ratio. Note that the effects of the chamber wall are important when $(x + x_E)/d^* > 130$.

was not diluted for measurements made in the altitude chamber.

4.1 Burner Measurements

A representative particle number distribution in the transition pipe is shown in Figure 5. Integrating the particle number distribution gives a total number concentration of $1.2 \times 10^7$ particles per standard cubic centimeter, where standard pressure and temperature are defined as 101.325 kPa and 298.15 K, respectively.

Note, however, that the particle measurements taken in the chamber suggest that the combustor particle output is not steady (see the next section). The variation of combustor particle output with time is being measured, and for future tests, two particle measurement systems will be used so that the particle distributions in the transition pipe and in the altitude chamber can be measured simultaneously.

FIGURE 5: Particle number distribution in the transition pipe for burner condition 1 ($f/a = 0.045$)

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4.2 Initial Measurements in the Altitude Chamber

Initially, attempts were made to take particle evolution measurements in the altitude chamber with standard day pressure and temperature conditions corresponding to 9,140 m. Unfortunately, at these conditions, the water vapor in the jet condenses, and in some cases freezes, in the sampling probe. Therefore, heat tape was wrapped around the sampling probe and the altitude chamber conditions were changed to the considerably warmer hot day conditions at 7,620 m. For these chamber conditions and burner condition 1, Figure 6 shows the particle number concentration as a function of radial distance at axial distance 0.30 and 0.51 m downstream of the jet exit. (Note that in Figure 6 and in all other figures showing the particle number concentration, the number concentration is given in particles per actual cubic centimeter at the particle counter, not in particles per standard cubic centimeter. At the particle counter, the temperature is approximately standard temperature but the pressure is the altitude chamber pressure.)

After successfully measuring the particle number concentration with a heat tape-wrapped probe at hot day conditions corresponding to 7,620 m, initial particle number concentrations were measured at standard day conditions corresponding to 6,100 m, 9,140 m, and 12,190 m, as shown in Figures 7 and 8. In all cases, it was possible to measure the particle number concentration with the heat tape turned on. However, when the heat tape was turned off, it was not possible to measure the number concentration near the jet centerline at 6,100 m and 9,140 m because the number concentration was above the maximum concentration that the CPC could measure.

(Note that in several of the plots, there appear to be two separate particle curves. This is most apparent in the measurements taken with the heat tape on at 6,100 m conditions in Figure 7a. In all cases, the measurements were taken at different times, and the difference between the measurements appear to be a real change in the particle concentration, not a lack of precision in the CPC. For example, the “upper” curve in 7a was taken earlier in the day than the “lower” curve. The change in particle concentration could be caused by either a change in the chamber conditions or a change in the particle output from the combustor. Because the chamber conditions were constantly monitored to ensure the temperature and pressure remained as constant as possible, it seems to be more likely that a change in particle output from the combustor was responsible for the change in the chamber particle concentration. In the future, the combustor particle size distribution will be monitored continuously; for the measurements reported here, the same CPC was used for the combustor and the chamber particle measurements and these measurements could not be taken simultaneously.)

In addition, icing continued to be a problem. It was possible to prevent ice from forming in the sampling probe and at the sampling probe tip by avoiding sudden changes in the radial location of the probe and by briefly turning on the heat tape to melt the ice if it began to form. However, it is unclear how many of the particles measured when the heat tape was turned off were present in the plume and how many were formed in the probe.

Unfortunately, it is also unclear that measurements taken when the heat tape was turned on accurately reflect the number of particles in the jet plume. For the heated-probe measurements taken at 12,190 m conditions, the peak particle concentration occurs off the centerline, which is an unexpected result. However, for the heated-probe measurements taken at all other chamber conditions, the peak particle concentration occurs on the centerline, as expected. Research groups measuring the particulate emissions of jet engines on the ground have also noticed unexpected and as yet unexplained effects when using heated probes\(^2\), so it appears that using a heated probe is not an ideal solution.

Figure 9 shows the normalized particle number concentration profiles, \(\phi_{\text{max}} / \phi_{\text{max}} \text{ vs. } r/(x_{\text{E}})\). The profiles at chamber conditions corresponding to altitudes of 6,100 m and 9,140 m measured with a heated probe were consistent with the normalized temperature profiles, \(\theta/(\theta_{\text{L}})\). Both profiles at a chamber condition corresponding to 7,620 m, also taken with a heated probe, collapsed onto each other but the profiles were narrower than the normalized temperature profile. The profiles taken with an unheated probe at a chamber condition corresponding to 12,190 m are wider than the normalized temperature profile, and the profiles taken with a heated probe at the same chamber conditions have a maximum value off the centerline. Both of the heated- and unheated-probe 12,190 m profiles are very different in shape than the normalized temperature profiles.

Figure 10 shows the maximum concentration at each axial location and chamber condition as a function of the normalized downstream distance, \(x/d^*\). If the particle number concentration is a conserved scalar, the maximum concentration will decrease with axial distance from the nozzle exit. However, as the temperature in the jet cools, volatile particles will form; if a large enough number of volatile particles is formed, the particle number concentration will increase with axial distance. In addition, the heated probe may revaporize some or all of the volatile particles formed in the jet. (Because all particle measurements are done at room temperature, some particles may also revaporize in the sampling line between the altitude chamber and the condensation particle counter for both the heated and unheated probe cases.)

The results from Figure 10 are inconclusive. At the warmest chamber temperature — which occurs at the 7,620 m hot day conditions — the maximum particle concentration decreases with axial distance when the probe is heated. At the coldest chamber temperature, — which occurs at the 12,190 m standard day conditions — the particle concentration increases with axial distance when the probe is heated. So far, because more volatile particles will form at the cold 12,190 m standard day conditions...
FIGURE 6: Particle count at 7,620 m hot day conditions and at (a) $x=0.30$ m and (b) $x=0.51$ m downstream of the nozzle exit. Heat tape is wrapped around the particle probe for both cases.

FIGURE 7: Particle number as a function of radial location at (a) 6,100 m conditions and $x=0.30$ m and (b) 9,140 m conditions and $x=0.71$ m. Red symbols indicate that the heat tape wrapped around the particle probe was turned on, and blue symbols indicate that it was turned off.

than at the much warmer 7,620 m hot day conditions, this could be consistent with volatile particles forming in the jet and then revaporizing in the heated probe. However, unheated probe measurements at the 12,190 m standard day conditions show particle number concentration decreasing with increasing axial distance; although this could be consistent with particles revaporizing in the sampling line, it is inconsistent with the heated probe measurements.

When particulate emissions measurements are taken on the ground or in a test cell, the particulate sample is diluted at the probe tip to prevent condensation and to freeze the particulate chemistry. In this study, the altitude chamber particulate measurements were not diluted because the mixing in the jet dilutes the combustion products by a factor of between five and ten to one, depending on axial location. However, because the ambient temperature in the altitude chamber is much colder than the temperature in the jet — especially on the jet centerline and at the lower axial locations — more dilution may be necessary. For
future measurements, diluting the particulate sample so that the vapor pressure of water in the sample is below the saturation vapor pressure at the ambient chamber conditions could prevent condensation in the probe.

Before further particle measurements are made in the altitude chamber, it would be best to do a modeling study of the particle sampling system. This study could help determine the degree of condensation in the unheated particle probe and the effects of heating the particle probe. It could also provide guidance for choosing a sampling flow rate and help determine if a larger diameter sampling probe or tip dilution could reduce the particle probe effects.

**FIGURE 8**: Particle concentration with and without heat tape at 12,190 m standard day conditions at (a) 0.18 m and (b) 0.71 m downstream of the nozzle exit. Red symbols indicate that the heat tape wrapped around the probe was turned on, and blue symbols indicate that it was turned off.

**FIGURE 9**: Normalized particle number distribution plotted with the curve fit for $\theta/\theta_{CL}$. Unfilled symbols are used for measurements taken when the heat tape was turned on, and filled symbols are used for measurements taken when the heat tape was turned off.
5 Conclusions

Initial temperature and particle number concentration profiles were measured in the Particulate Aerosol Laboratory. This unique facility simulates engine exhaust at altitude by exhausting hot combustor exhaust into an altitude chamber. The small nitrogen coflow in the altitude chamber is kept at pressure and temperature conditions corresponding to aircraft cruising altitudes. Results from the normalized temperature distribution, $\theta$, show that the combustor exhaust flow field follows momentum-dominated jet scaling. Initial particle number concentration measurements were taken with a heated probe at four altitude chamber conditions and with an unheated probe at one altitude condition. Results show that whether the probe is heated or unheated affects the particulate measurements. Particle number concentrations measured with an unheated probe can be more than five times as great as those measured with a heated probe. In addition, the shape of the radial particle number concentration profile can be different when using an unheated probe than when using a heated probe. A modeling study of the particle sampling system should be done before further testing to determine the effects of probe heating and of condensation in the particle probe.

REFERENCES


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Combustion; Particulates; Emissions; Particles; Aerosol