LETTER

Biofuel blending reduces particle emissions from aircraft engines at cruise conditions

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Aviation-related aerosol emissions contribute to the formation of contrail cirrus clouds that can alter upper tropospheric radiation and water budgets, and therefore climate¹. The magnitude of airtraffic-related aerosol-cloud interactions and the ways in which these interactions might change in the future remain uncertain¹. Modelling studies of the present and future effects of aviation on climate require detailed information about the number of aerosol particles emitted per kilogram of fuel burned and the microphysical properties of those aerosols that are relevant for cloud formation². However, previous observational data at cruise altitudes are sparse for engines burning conventional fuels^{2,3}, and no data have previously been reported for biofuel use in-flight. Here we report observations from research aircraft that sampled the exhaust of engines onboard a NASA DC-8 aircraft as they burned conventional Jet A fuel and a 50:50 (by volume) blend of Jet A fuel and a biofuel derived from Camelina oil. We show that, compared to using conventional fuels, biofuel blending reduces particle number and mass emissions immediately behind the aircraft by 50 to 70 per cent. Our observations quantify the impact of biofuel blending on aerosol emissions at cruise conditions and provide key microphysical parameters, which will be useful to assess the potential of biofuel use in aviation as a viable strategy to mitigate climate change.

The global aviation sector contributes approximately 5% of the current anthropogenic radiative forcing, owing to direct emissions of fossil-fuel CO₂ (28 mW m⁻²) and the formation and evolution of contrails and contrail-induced cirrus clouds (50 mW m⁻²)^{1,3-5}. Of these effects, the largest uncertainties are associated with aviation-induced cloudiness, both directly from contrail-induced cirrus clouds and indirectly from the contribution of black carbon, organic and sulfate aerosols that may act as cloud condensation nuclei and ice nuclei^{1,6-8}. With emissions of CO₂ from fuel expected to more than double by 2050, aviation-related contributions to radiative forcing may increase to 3–4 times the year 2000 levels⁵. Consequently, some governments are exploring ways to curb these emissions, and the International Air Transport Association (IATA) has targeted carbon-neutral growth by 2020 and a 50% reduction in carbon emissions by 2050 (ref. 9).

Sustainable biojet fuels are a promising route for mitigating greenhouse gas emissions. However, many challenges remain before aviation biofuels can be widely adopted, particularly with regard to cost and sustainability. Jet fuels are more highly refined than the biofuels used for surface transportation, with the latter perhaps presenting a "better biomass opportunity cost"¹⁰. However, unlike for aviation, there are many alternative energy solutions for surface transportation, other than liquid hydrocarbon-based fuels, that are realizable in the near future^{10,11}. Biojet fuels consist of a mixture of C₉–C₁₆ hydrocarbons that are typically formed via transesterification and subsequent hydroprocessing of plant and animal oils to produce a hydrotreated esters and fatty acids (HEFA) fuel that has many of the properties of petroleumderived jet fuels^{12,13}. Promising plant-based feed stocks for future aviation biofuels include Jatropha, Camelina and algae¹².

Biojet fuels have potential as a future aviation fuel source that is not dependent on fossilized carbon and that contains near-zero levels of sulfur and aromatic species, which are commonly present in petroleumbased jet fuels at levels of several hundred parts per million by mass (p.p.m.m.) sulfur and around 20% aromatics by volume. Previous laboratory and ground test experiments using bio-based fuels or synthetic Fischer–Tropsch fuels produced from natural gas and coal feed stocks show that the absence of sulfur and aromatic species within the fuel substantially reduces the sulfate and black carbon particle emissions from aircraft engines^{14–16}. These results are important for reducing the impact of aviation on local air quality near airports and suggest that similar reductions are likely to be observed at high-altitude cruise conditions; however, the engine operating conditions on the ground (for example, temperature, pressure, fuel flow rates, fuel/air ratio and maximum thrust) are very different from those in flight.

Here we report airborne measurements of jet engine exhaust, sampled at cruise conditions, from engines burning both a blended biofuel and a conventional jet fuel. Research aircraft from NASA, the German Aerospace Center (DLR) and the National Research Council (NRC) Canada were equipped with state-of-the-art instrumentation and sampled the exhaust of the NASA DC-8 turbofan engines at atmospheric and engine conditions that are exclusively met in flight. The tests were conducted during 2013–2014 as part of the Alternative Fuel Effects on Contrails and Cruise Emissions Study (ACCESS) at NASA Armstrong Flight Research Center in Palmdale, California, USA.

The DC-8 source aircraft has four wing-mounted CFM56-2-C1 engines that can be fed fuel from any of four segregated fuel tanks within the wings. During the flight experiments, these tanks contained either a medium- or low-sulfur-content Jet A fuel, while a fuselage-mounted auxiliary tank contained an approximately 50:50 (by volume) blend of a low-sulfur-content Jet A fuel and a Camelina-based HEFA biojet fuel (see Methods).

The exhaust plumes from the left and right inboard DC-8 engines were sampled by research aircraft flying in a trailing formation at a distance of 30–150 m (plume age of about 0.15–0.75 s) behind the DC-8 (Fig. 1). This short distance assures that the plumes from specific engines did not mix. Three different fuels and three different engine thrust conditions were investigated, which bracket the range of realistic flight conditions on the DC-8 flight curve (Fig. 1d). Commercial aircraft typically fly at thrust conditions at or slightly above the 'maximum range' point, at which the quotient of drag and Mach

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Figure 1 | **Side and forward views of DC-8 contrails and the operational cruise curve. a**, Side view of the NASA HU-25 Falcon aircraft sampling the DC-8 contrail. **b**, **c**, Forward-looking views of the DC-8 contrails with the inboard engines throttled up to the high-thrust condition and the outboard engines throttled back to almost idle (b), and the reverse conditions (**c**). **d**, The operational flight curve for the DC-8 (red curve), assuming an average aircraft gross weight of 90,718 kg (200,000 lb) at 10,670 m (flight level, FL310); the red points highlight

number is minimized. Thrust can also be varied by the flight crew for schedule and fuel-burn considerations.

The use of the four-engine DC-8 is advantageous for in-flight engine emissions testing and overcomes sampling challenges associated with dual-engine flight tests for two reasons. First, the thrust settings of the two inboard engines (no. 2 and no. 3) can be varied over the range of typical flight operating conditions while adjusting the outboard engines (no. 1 and no. 4) to maintain a constant airspeed. Figure 1d shows the thrust settings for the DC-8 engines (blue points), which were set at an achievable cruise Mach number of 0.6 for the NASA and DLR chase planes. Second, we can sequentially probe emissions from both inboard engines burning both fuels during a single flight and under the same atmospheric conditions. This allows us to account for differences in engine performance that influence the engine-specific emissions indices. For example, the left inboard engine of the DC-8 emits 1.3–2 times more particles than does the right inboard engine, depending on the thrust setting.

The effect of thrust changes on the emissions of particles and, particularly, water vapour from the engines is visibly evident under contrail-forming conditions, under which the plume is supersaturated with respect to liquid water, satisfying the Schmidt–Appleman criterion¹⁷. A noticeable decrease in the concentration of ultrafine (less than 10 nm in diameter) particles was measured under contrail-forming conditions versus visually clear air conditions. This difference implies that ice particles collide with and uptake these small particles, thereby decreasing particle number. Consequently, we confine our analysis to the determination of engine emissions indices for clear-air (that is, non-contrail-forming) exhaust plumes only. This ensures that the reported emissions data are not affected by contrail processing in between the engine exhaust plane and the sampling inlet.

Particle number and mass concentrations, as well as trace-gas concentrations of carbon dioxide (CO_2) , carbon monoxide (CO) and

the maximum continuous thrust, range and endurance points. The blue points correspond to the ACCESS engine thrust settings, with the arrows included to guide the eye to the corresponding thrust condition on the flight curve; the blue points denote the mean fuel flow rate ± 1 arithmetic standard deviation (a.s.d.) during plume sampling of all fuels at altitudes of 9,140–10,970 m (30,000–36,000 ft). The numbers correspond to the values given in Extended Data Table 2. Note that all emissions indices reported here are for clear air (that is, under non-contrail-forming conditions).

nitrogen oxides (NO_x), were sampled by the chase aircraft using sample inlets mounted on the crown of the DLR and NASA Falcon aircraft and below-wing-mounted instruments on the NRC T-33. The amount of a given species (number of particles or integrated mass) emitted per kilogram of fuel burned is the species emissions index, which is computed assuming that the carbon content of the fuel is constant and completely converted to CO₂ on a mass basis (see Methods). This assumption implies a CO₂ emissions index of 3,160 g kg⁻¹ (note that here 'kg⁻¹' denotes 'per kilogram of fuel').

Table 1 shows the geometric mean (multiplied or divided by one geometric standard deviation (*1 g.s.d.)) particle and trace-gas emissions indices for multiple plume intercepts of the no. 3 engine exhaust plume while it was operating at medium cruise thrust (corresponding to 'maximum range' in Fig. 1d). Summary statistics for the additional thrust conditions and the no. 2 engine are provided in Extended Data Tables 3-7. Particle number emissions indices for the conventional petroleum-based jet fuels are of the order of 10¹⁴-10¹⁵ kg⁻¹, with the non-volatile number emissions index closer to 10¹⁴kg⁻¹ and the black-carbon-equivalent mass emissions index near 15 mg kg⁻¹. These emissions indices, and particularly those for the non-volatile particles, fall towards the lower end of reported emissions indices from previous flight test experiments conducted during the 1990s, which is attributable to efficiency gains implemented in the high-bypass CFM56 series of turbofan engines relative to the lower-bypass ATTAS Rolls/Royce/ Snecma M45H Mk501 engines, the P&W JT3D and JT9D engines, and the GE CF6 series engines¹⁸⁻²². Sulfur doping of the low-sulfur-content Jet A from 22 ± 13 p.p.m.m. to 416 ± 37 p.p.m.m. had no discernible effect on the engine particle emissions for particle diameters exceeding 5-10 nm (the lowest detectable sizes in this study); however, it is known that changes in the sulfur content of fuel affect the near-field aerosol number emissions index by markedly increasing the number of particles with diameters of less than 5 nm (ref. 23). In addition, most

Table 1 | Emission indices for no. 3 engine under medium thrust at cruise conditions

	Medium-sulfur-content Jet A fuel	Low-sulfur-content Jet A fuel		50:50 HEFA:low-sulfur-content Jet A fuel blend	
Emissions index (per kg of fuel)	Emissions index	Emissions index	Ratio	Emissions index	Ratio
Total particle ($D_p > 5$ nm) number	-	$1.94 imes 10^{15} st 1.53$	-	$1.58 imes 10^{15} st 1.64$	-
Ultrafine particle (5 nm $< D_p < 10$ nm) number	-	$1.40 imes 10^{15} \pm 1.62$	-	$1.24 \times 10^{15} \pm 1.74$	-
Fine particle ($D_p > 10$ nm) number	$6.51 \times 10^{14} {\rm *} 1.14$	$7.01 \times 10^{14} {\rm *} 1.55$	1.08	$3.36 \times 10^{14} {\rm \ast} 1.33$	0.52****
Volatile fine particle number	$3.86 \times 10^{14} \pm 1.08$	$3.52 \times 10^{14} \pm 1.74$	0.91	$1.86 imes 10^{14} st 1.38$	0.48****
Non-volatile fine particle number	$2.63 \times 10^{14} \pm 1.26$	$3.43 imes 10^{14} st 1.39$	1.30	$1.46 imes 10^{14} st 1.37$	0.55**
Total particle (5 nm $<$ D_p $<$ 120 nm) volume (mm ³)	16.58 * 1.14	18.03 * 1.85	1.09	8.62 * 1.72	0.52
Volatile particle volume (mm ³)	5.62 * 1.67	5.98 * 1.86	1.06	2.90 * 1.88	0.52
Non-volatile particle volume (mm ³)	10.65 * 1.11	12.03 * 1.85	1.13	5.42 * 1.73	0.51**
PSAP black-carbon-equivalent mass at 467 nm (mg)	17.12 * 1.12	14.48 * 1.24	0.85	7.24 * 1.33	0.42****
PSAP black-carbon-equivalent mass at 530 nm (mg)	16.85 * 1.12	12.89 * 1.36	0.76	6.79 * 1.33	0.40****
PSAP black-carbon-equivalent mass at 660 nm (mg)	16.15 * 1.13	16.07 * 1.16	0.99	6.01 * 1.39	0.37****
Carbon monoxide, CO (g)	5.99 ± 0.96	4.02 ± 0.54	0.67**	4.68 ± 1.27	0.78*
Nitrogen oxides, NO_x (g)	7.26 ± 0.50	7.60 ± 0.41	1.05	7.28 ± 0.33	1.00
Number of plume intercepts	4	5		10	

Summary of emissions indices measured in clear air at the medium-thrust cruise condition (fuel flow rate of $0.280 \pm 0.020 \text{ kg} \text{ s}^{-1}$) for the right inboard engine at altitudes of 9,140-10,970 m(30,000-36,000 ft). Particle emissions indices are reported as the geometric mean ± 1 g.s.d.; trace-gas emissions indices are reported as the arithmetic mean ± 1 arithmetic standard deviation (a.s.d.). D_p is the particle diameter; PSAP, particle soot absorption photometer. The ratios given are those of the emissions indices from the low-sulfur-content Jet A fuel or biofuel blend relative to the emissions indices from the medium-sulfur-content Jet A fuel. Ratios corresponding to significantly different means, as determined by Welch's unequal variances *t*-test, are set in bold: ****P < 0.001, ***P < 0.05, *P < 0.1.

of the sulfur mass in fuel is emitted as gaseous SO₂, which is photooxidized in the atmosphere to form small, secondary sulfate aerosols over longer timescales. Consequently, although our reported total and volatile emissions indices represent the near-field plume, they are a lower limit on the contribution of aircraft emissions to the atmospheric sulfur budget that is ultimately a small global source of pollution relative to terrestrial sources⁵.

We also tabulate emissions indices for the 50:50 blend of HEFA biojet fuel and low-sulfur-content Jet A (Table 1, Extended Data Tables 3–7). The emissions indices from burning the biofuel blend show that both volatile and non-volatile number emissions are reduced by roughly half compared to the Jet A fuels. Ratios of the emissions indices from the low-sulfur-content Jet A and biofuel blend fuels are given relative to the emissions indices from the medium-sulfur-content fuel, with those in bold denoting a statistically significant difference of means (P < 0.1). These large reductions are consistent across each of the investigated engine thrust conditions (Fig. 2), although the number emissions index reduction is less pronounced at the high-engine-thrust setting (about 25%) than at the low- and medium-thrust settings (about 50%). Ground-based tests on the DC-8 engines show a similar trend, albeit over a broader range of fuel flow rates and at sea level¹⁶. The differences in engine performance characteristics between surface tests and in-flight tests preclude a direct comparison of emissions indices, despite some promising approximation methods that use groundbased data to estimate emissions indices at cruise conditions^{24–26}. The greatest effect on emissions is associated with a reduction in black-carbon-equivalent mass, with the biofuel blend exhibiting emissions that are 30%–50% of those seen for the petroleum-based Jet A fuels.

Measured particle size distributions help to explain the differences between the reductions in number and mass (volume) emissions indices for the two fuels (Fig. 3). A pronounced decrease in both total and non-volatile particle number and volume associated with the biofuel blend is apparent, as is a slight shift towards smaller sizes in the mode peak diameter, by 3–5 nm for the number distributions and 9–12 nm for the volume distributions. This shift appears to be caused by a greater reduction in the number of larger, soot-mode aerosols, which serve as condensation nuclei for organic species and sulfuric acid. Because gas-to-particle condensation scales with particle size (more specifically, with diameter squared in the free molecular regime), the lower emission of the soot size mode from the biojet fuel blend diminishes the condensational sink, which in turn enhances the nucleation of new particles in a compensating manner.



Figure 2 | Summary of particle emissions indices at all thrust and cruise conditions. Geometric mean particle emissions indices (± 1 g.s.d.) for all thrust settings (shown as fuel flow rate ± 1 a.s.d.) and each fuel burned on the right inboard engine (no. 3) at altitudes of 9,140–10,970 m (30,000–36,000 ft). The ratio of the emissions indices for the 50:50

biofuel blend and the medium-sulfur-content Jet A fuel are labelled, with the number of asterisks denoting the statistical significance level of a difference in means test: ****P < 0.001, ***P < 0.01, **P < 0.05 (see Table 1 and Extended Data Tables 3–7).



Figure 3 | Size distributions of particle emissions at high-thrust and cruise conditions. Geometric mean (*1 g.s.d.) size distributions of particle number emissions index (EI_N; left) and volume emissions index (EI_V; right) for the total (top) and non-volatile particle fraction (bottom) measured at the high-thrust condition, behind the no. 3 engine. Black

This competition manifests as the size-dependent number emissions reductions observed in Fig. 3.

We find that blending petroleum-based fuels with a HEFA biojet fuel reduces the volatile and non-volatile particle emissions by 50%-70% at atmospheric cruise conditions. However, the emissions indices for soot particle numbers remain in the soot-rich regime (around 10^{14} kg⁻¹) for the biofuel blend. Theoretical calculations³ suggest that the initial number of contrail ice particles scales linearly with soot number emissions index in the soot-rich regime and that ambient and ultrafine particles are unlikely to contribute to contrail formation.

Understanding the implications of these findings for future aviation-related effects on upper tropospheric clouds is complicated by the idea that, despite these potential reductions in the number of ice crystals, the frequency and ice mass of contrails may actually increase, owing to the 8% increase in the hydrogen content of the biofuel blend compared to petroleum-based fuel (Extended Data Table 1). However, it remains to be seen whether this increased water vapour in the early plume is relevant to contrail evolution after the vortex mixing phase, where the supersaturation of ambient water vapour with respect to ice is the primary driver of the persistence and ice mass of contrails²⁷. Future modelling studies should explore the extent to which the ice mass concentration of contrails (and hence longwave forcing) might increase, as well as the changes in the optical depth, lifetime and coverage of clouds and the overall radiative forcing associated with reduced ice number concentrations^{28–30}. A critical first step is the determination of the number and size of engine exhaust particles at cruise conditions, for which data have so far been non-existent for engines burning biojet fuel blends³ and sparse for conventional, petroleum-based fuels². This work provides key microphysical parameters relating to aerosols, which are necessary for transportation and climate modelling efforts to constrain future aviation-related impacts on the environment as the aviation

squares and green diamonds show data for the medium-sulfur-content Jet A fuel and the 50:50 biofuel blend, respectively. Solid lines are log-normal fits and the shaded area represents the difference between the two curves. The geometric mean diameter $D_{\rm g}$ for each fit is noted in the legend; all fit parameters are given in Extended Data Tables 8 and 9.

industry considers transitioning towards more widespread adoption of biofuels.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Information Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to R.H.M. (richard.h.moore@nasa.gov).

METHODS

Source aircraft and engines. The NASA DC-8 (Tail Number N817NA) and its four CFM56-2-C1 engines have previously been described in detail^{16,31–34}. The engines were installed on the airframe in 1986 and are maintained in accordance with the CFM56-2C and Douglas DC-8 maintenance manuals and inspection task cards. With a maximum take-off thrust of 97,860 N and an in-flight maximum cruise thrust of 22,150 N, the CFM56-2-C1 is the first high-bypass (6.0 bypass ratio) engine of the CFM56 family^{16,32,35}. At cruise, the maximum pressure ratio is 31.3 and the cruise bucket-specific fuel consumption is 68.4 g N⁻¹hr⁻¹ (refs 16, 32, 35). Given the use of the aircraft as a NASA flying laboratory, the annual flight hours for the aircraft and engines are typically low (<1,000 h) as compared to commercial aircraft.

Biofuel blending and ACCESS fuel properties. Blending of conventional jet fuels with biojet fuels is relevant for the aviation fuel supply over the next few decades. For example, the European Advanced Biofuels Flightpath aims to enable around 1.8 million tonnes (2 million tons) of sustainable aviation biofuels to be in use by 2020³⁶. Meanwhile, the US Farm-to-Fly initiative will accelerate biofuel development, and the US Federal Aviation Administration (FAA) targets annual production of 4.5 billion litres (1 billion gallons) of drop-in renewable jet fuel by 2018³⁷. For comparison, the FAA target is roughly 6% of the US domestic production of jet fuel and 1% of the global jet fuel consumption expected for 2018^{38,39}. Here we examine the emissions from three different fuels: a low-sulfur-content Jet A fuel, a medium-sulfur-content Jet A fuel, and a 50:50 (by volume) blend of the lowsulfur-content Jet A fuel and an HEFA biojet fuel. The medium-sulfur-content Jet A fuel was created by doping the low-sulfur-content Jet A fuel that was delivered from the refinery with a small aliquot of tetrahydrothiophene to increase the sulfur content of the fuel to 416 ± 37 p.p.m.m. without changing its other properties. Typical sulfur concentrations in petroleum-based jet fuels are of the order of several hundred parts per million by mass.

The main difference between the HEFA fuel and traditional petroleum-based fuels is that the former contains no sulfur or aromatic species, whereas traditional jet fuels typically have aromatic contents of 18%–25%. In addition to strict standards related to fuel density, viscosity and freezing behaviour that affect safety of flight, fuel aromatics are limited to below 25% to limit solvent deterioration of nitrile elastomers; meanwhile, a minimum aromatic content of 8% was established to swell the elastomer seals in some current fuel systems^{40–42}. For this reason, only 50:50 (by volume) blends of HEFA and petroleum-based fuels are currently certified for flight. All investigated fuels were tested before use and conform to flight worthiness specifications outlined by ASTM^{43,44}; the ACCESS fuel properties are summarized in Extended Data Table 1.

Particle and trace gas measurements. Measurements were made from the NASA HU-25 Falcon aircraft using the NASA Langley Aerosol Research Group (LARGE) suite of in situ instruments, which have been used for numerous airborne research campaigns, as described elsewhere¹⁸. Similarly, the instruments onboard the DLR Falcon 20 and NRC T-33 aircraft are well characterized with long flight heritage^{21,45}. Results from the following instruments were used here. Total particle number concentration $(D_p > 5 \text{ nm})$ was measured by a custom condensation particle counter (CPC) onboard the DLR Falcon 20. Fine particle number concentration ($D_{\rm p}$ > 10 nm) was measured on the NRC T-33 by a TSI 7610 CPC, and on the HU-25 by a pair of TSI 3010 CPCs; one of the HU-25 CPCs measured the non-volatile particle fraction after sample treatment with a thermal denuder at 350 °C, and the other measured the total (volatile and non-volatile) aerosol number. Similarly, total and non-volatile particle size distributions were measured by a TSI nano Scanning Mobility Particle Sizer (nanoSMPS; 3936, 3085, 3025A) for thermally undenuded and denuded sample streams, respectively. Because the nanoSMPS requires 45 s to complete a size upscan, a dual lag chamber system was used, where two cylinders were charged at high flow rate (191 min⁻¹) while the aircraft was sampling the exhaust plume. These lag chambers were then sampled by the nanoSMPS at 1.51 min⁻¹. The upper portion of the size distribution $(D_{\rm p}>85\,{\rm nm})$ was measured by an ultrahigh-sensitivity aerosol size spectrometer (UHSAS; Droplet Measurement Technologies), which was calibrated using soot particles from a Mini-CAST soot generator that were size-classified with a differential mobility analyser⁴⁶. The nanoSMPS size distributions were integrated to yield the total and non-volatile particle volume. Finally, particle black-carbonequivalent masses at three different optical wavelengths were measured by a Radiance Research Particle Soot Absorption Photometer (PSAP), assuming a mass absorption coefficient (MAC) of 7.5 m²g⁻¹ at 550 nm (ref. 47). This MAC was corrected for each of the PSAP wavelengths assuming an inverse wavelength dependence to yield MACs of $8.83 \text{ m}^2 \text{ g}^{-1}$, $7.78 \text{ m}^2 \text{ g}^{-1}$ and $6.25 \text{ m}^2 \text{ g}^{-1}$ at wavelengths of 467 nm, 530 nm and 660 nm, respectively. The PSAP mass concentrations were corrected for filter scattering artefacts following ref. 48, assuming a single scattering albedo (SSA) of 0.1, which is consistent with Mie theory calculations using the measured size distribution and a black-carbon refractive index of 1.95 - 0.75i (ref. 47). The PSAP correction is weakly sensitive to this assumed SSA and varies by only 1% over the SSA range of 0.03–0.3.

Carbon dioxide (CO₂), carbon monoxide (CO) and nitrogen oxide (NO_x) concentrations were measured using Los Gatos Research instruments. CO and CO₂ were measured via cavity-enhanced absorption. To measure NO_x, the sample stream was mixed with excess ozone to convert NO to NO₂, which was measured via cavity ring-down spectroscopy.

Calculation of emissions indices. As the Falcon sampling probe traverses the DC-8 engine exhaust it encounters varying concentrations of particles and trace gases owing to the spatial inhomogeneity of the plume. Making sense of these time-varying quantities requires that we normalize them to the rate of fuel burned by the engines to arrive at an averaged emissions index across the plume. The emissions index (EI) of particle or trace-gas species X is determined as

$$EI_{X} = \frac{\Delta X \times S(X)}{\Delta CO_{2}} (EI_{CO_{2}})$$

where ΔX and ΔCO_2 are the dilution-corrected, background-subtracted peak areas of the measured concentrations of species X and CO_2 at standard temperature and pressure (STP), respectively. S(X) is a unit-conversion scaling factor for particle concentrations (number or mass per air volume at STP) or trace-gas concentrations (parts per million by volume), defined as

$$S(X) = \begin{cases} V_{\rm m}/M_{\rm CO_2} & \text{for particles} \\ M_{\rm X}/M_{\rm CO_2} & \text{for trace gases} \end{cases}$$

where $V_{\rm m}$ is the molar volume of ideal gas at STP (22.41 mol⁻¹), and $M_{\rm X}$ and $M_{\rm CO_2}$ are the molar masses of species X and CO₂, respectively. EI_{CO2} is the emissions index of CO₂, assuming the carbon content in the fuel is constant and is completely converted to CO₂, and is defined as

$$EI_{CO_2} = \frac{RT}{PV_m} \frac{M_{CO_2}}{(M_C + \alpha M_H)} \approx 3,160 \text{ g kg}^{-1}$$

where *R* is the ideal gas constant, *T* is the temperature at STP (273.15 K), *P* is the pressure at STP (1 atm), α is the hydrogen-to-carbon molar ratio of the fuel, and $M_{\rm C}$ and $M_{\rm H}$ are the molar masses of carbon and hydrogen, respectively. Values of α calculated from Extended Data Table 1 for the Jet A fuels and the blended fuel are 1.92 and 2.07, respectively; however, for this analysis, we assume a constant value of 1.92 for simplicity. This assumption introduces insignificant error (about 1%) into the reported emissions indices compared to the measurement uncertainties.

The emissions indices reported here are tabulated as geometric means # 1 g.s.d. for particle emissions indices and as arithmetic means ± 1 a.s.d. for the trace-gas emissions indices. The statistical significance of the difference in means is assessed using Welch's unequal variances *t*-test.

Particle sampling and transmission loss corrections. The size-dependent inlet aspiration efficiency of the NASA HU-25 HIMIL inlet and particle losses to the sampling lines are estimated using the Particle Loss Calculator⁴⁹, which accounts for diffusional, inertial and sedimentation losses. Particles were sampled at a flow rate of $37 \, l \, min^{-1}$ through a 4.35-mm tube that was assumed to be oriented parallel to the air flow around the aircraft (mean air velocity of about 200 m s^{-1}). In reality, the inlet tube is shrouded, which ensures parallel sampling streamlines, but which reduces the local air velocity by an unknown amount. However, the sensitivity of the inlet aspiration efficiency to airspeed is negligible for the ultrafine particles sampled during ACCESS. The sample then passes through approximately 0.34 m of tubing with an inner diameter of 4.35 mm and approximately 4 m of tubing with an inner diameter of 7.9 mm before being sampled by the instrumentation in the cabin of the chase plane. We find that the uncertainty associated with particles larger than 20 nm in diameter is within 10%, but that large corrections (about 40%) are necessary at the lowest particle sizes (diameters of <10 nm). Given uncertainties associated with the CPC detection efficiency curves, we choose not to apply these corrections to the tabulated integrated number, volume and mass emission indices (Table 1 and Extended Data Tables 3-7), following ref. 33. Instead we focus on the differences in the sampled emissions indices across fuel types and engine powers, for which the sampling characteristics and efficiency should be nearly the same. On the basis of the measured mean size distributions, we estimate the uncertainty associated with neglecting these corrections on the tabulated number emissions indices to be 7%-9% and on the volume and mass emissions indices to be around 3%. The variability in the measured data (as expressed by the g.s.d. about the geometric mean) is typically much larger than 10%. Log-normal size distributions discussed in the next section have been corrected for these sampling and transmission losses.

Particle size distribution log-normal fits. Particle number and volume size distributions (5–300 nm diameter) measured by the SMPS and UHSAS were fitted using a single-mode log-normal function of the form

$$\frac{\mathrm{d}(\mathrm{EI}_i)}{\mathrm{d}[\log(D_\mathrm{p})]} = \frac{i}{\sqrt{2\pi}\,\log(\sigma_{\mathrm{g},i})} \exp\left\{-\frac{[\log(D_\mathrm{p}) - \log(D_{\mathrm{g},i})]^2}{2[\log(\sigma_{\mathrm{g},i})]^2}\right\}$$

where *i* is the total particle number (*N*) or volume (*V*), EI_{*i*} is the corresponding emissions index, D_p is the dry particle diameter, $D_{g,i}$ is the geometric mean diameter and $\sigma_{g,i}$ is the g.s.d. Fit coefficients for the number and volume size distributions are given in Extended Data Tables 8 and 9. All size distributions have been corrected for particle sampling and transmission losses.

Sample size. No statistical methods were used to predetermine sample size. Data availability. The datasets generated during and/or analysed during this study are available in the NASA Aeronautics Flight Projects repository at http://aero-fp. larc.nasa.gov/ and from the corresponding author on reasonable request.

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Extended Data Table 1 | Mean fuel properties (± 1 a.s.d.) for each of the three fuels investigated

Fuel Property	Medium-sulfur- content Jet A fuel	Low-sulfur- content Jet A fuel	50:50 HEFA:low-sulfur- content Jet A fuel blend	Jet A Specification Range [†]
Sulfur (p.p.m.m.)	416 ± 37	22 ± 13	11 ± 3	< 3000
Aromatics (per cent by volume)	21.1 ± 0.7	21.4 ± 1.4	12.9 ± 1.2	8–25
Hydrogen Content (per cent by mass)	13.6 ± 0.3	13.8 ± 0.2	14.7 ± 0.2	> 13.4
Naphthalenes (per cent by volume)	0.68 ± 0.04	0.68 ± 0.05	0.4 ± 0.0	< 3.0
Density (kg m ⁻³)	809.2 ± 1.8	810 ± 0.5	787.4 ± 2.5	775-840
Net Heat of Combustion (MJ kg ⁻¹)	43.14 ± 0.05	43.15 ± 0.06	43.52 ± 0.04	>42.8
Number of Samples Tested	5	4	5	

[†]ASTM D1655 (ref. 43) and D7566 (ref. 44).

Extended Data Table 2 | Summary of cruise emissions index tables

	Engine thrust condition (fuel flow in kg s ⁻¹)						
	High	Medium	Low				
	(0.373±0.025)	(0.280±0.020)	(0.231±0.017)				
Engine #2	Extended Data Table 3	Extended Data Table 5	Extended Data Table 6				
Engine #3	Extended Data Table 4	Table 1 in main text	Extended Data Table 7				

Extended Data Table 3 | Emissions indices for no. 2 engine under high-thrust and cruise conditions

	Medium-sulfur-content Jet A fuel	Low-sulfur-content 50:5 Jet A fuel		50:50 HEFA:low-sul Jet A fuel bl	:50 HEFA:low-sulfur-content Jet A fuel blend	
Emissions index (per kg of fuel)	Emissions index	Emissions index	Ratio	Emissions index	Ratio	
Total particle (D _p > 5 nm) number	-	3.30×10 ¹⁵ * 1.04	-	1.97×10 ¹⁵ * 1.48	-	
Ultrafine particle (5 < D _n < 10 nm) number	-	1.81×10 ¹⁵ * 1.08	-	1.08×10 ¹⁵ * 1.99	-	
Fine particle ($D_n > 10 \text{ nm}$) number	1.43×10 ¹⁵ * 1.13	1.57×10 ¹⁵	1.10	1.02×10 ¹⁵ * 1.16	0.71***	
Volatile fine particle number	7.58×10 ¹⁴ * 1.27	6.53×10 ¹⁴	0.86	5.06×10 ¹⁴ * 1.09	0.67*	
Non-volatile fine particle number	6.59×10 ¹⁴ * 1.08	9.17×10 ¹⁴	1.39	5.03×10 ¹⁴ * 1.30	0.76**	
Total particle (5 < D_p < 120 nm) volume (mm ³)	68.23 * 1.31	78.80	1.16	37.77 * 1.15	0.55	
Volatile particle volume (mm ³)	20.07 * 1.12	29.71	1.48	15.79 * 1.20	0.79	
Non-volatile particle volume (mm ³)	46.02 * 1.55	49.09	1.07	21.65 * 1.21	0.47	
PSAP black-carbon-equivalent mass at 467 nm (mg)	85.13 * 1.12	70.49	0.83	38.94 * 1.37	0.46****	
PSAP black-carbon-equivalent mass at 530 nm (mg)	82.40 * 1.12	67.94	0.82	37.55 * 1.39	0.46****	
PSAP black-carbon-equivalent mass at 660 nm (mg)	75.38 * 1.12	66.43	0.88	36.84 * 1.39	0.49***	
Carbon monoxide, CO (g)	5.62 ± 0.95	3.57	0.64	4.43 ± 1.17	0.79	
Nitrogen oxides, NO _x (g)	9.67 ± 0.71	-	-	9.61 ± 0.80	0.99	
Number of plume intercepts	4	1		7		

Emissions indices were measured in clear air at a single-engine fuel flow rate of 0.373 ± 0.025 kg s⁻¹. Data are for the left inboard engine (no. 2) at altitudes of 9,140-10,970 m (30,000-36,000 ft). Significance level: ****P < 0.001, ***P < 0.001, **P < 0.05, *P < 0.1. Particle emissions indices are reported as the geometric mean ± 1 g.s.d.; trace-gas emissions indices are reported as the arithmetic mean ± 1 a.s.d.

Extended Data Table 4 | Emissions indices for no. 3 engine under high-thrust and cruise conditions

	Medium-sulfur-content Jet A fuel	Low-sulfur-content 50:50 HEFA:low-sulf Jet A fuel Jet A fuel ble		lfur-content end	
Emissions index (per kg of fuel)	Emissions index	Emissions index	Ratio	Emissions index	Ratio
Total particle (D _n > 5 nm) number	-	2.11×10 ¹⁵ * 1.23	-	1.62×10 ¹⁵ * 1.44	-
Ultrafine particle (5 < D _n < 10 nm) number	-	1.16×10 ¹⁵ * 1.45	-	9.15×10 ¹⁴ * 1.12	-
Fine particle ($D_n > 10 \text{ nm}$) number	1.07×10 ¹⁵ * 1.13	1.34×10 ¹⁵	1.25	7.87×10 ¹⁴ * 1.19	0.74**
Volatile fine particle number	5.81×10 ¹⁴ * 1.07	4.97×10 ¹⁴	0.86	3.98×10 ¹⁴ * 1.06	0.69****
Non-volatile fine particle number	4.86×10 ¹⁴ * 1.24	8.43×10 ¹⁴	1.74	3.81×10 ¹⁴ * 1.39	0.78
Total particle (5 < D_p < 120 nm) volume (mm ³)	44.30 * 1.14	50.01	1.13	23.32 * 1.14	0.53*
Volatile particle volume (mm ³)	9.51 * 1.54	13.53	1.42	9.39 * 1.54	0.99
Non-volatile particle volume (mm ³)	32.81 * 1.34	36.48	1.11	13.27 * 1.06	0.40
PSAP black-carbon-equivalent mass at 467 nm (mg)	54.00 * 1.20	53.55	0.99	20.84 * 1.07	0.39***
PSAP black-carbon-equivalent mass at 530 nm (mg)	52.38 * 1.20	51.77	0.99	20.00 * 1.07	0.38***
PSAP black-carbon-equivalent mass at 660 nm (mg)	50.95 * 1.20	50.10	0.98	19.54 * 1.07	0.38***
Carbon Monoxide, CO (g)	5.18 ± 0.72	3.82	0.74	4.70 ± 1.20	0.91
Nitrogen Oxides, NO _x (g)	9.45 ± 0.49	9.77	1.03	9.03 ± 0.58	0.96
Number of Plume Intercepts	4	1		5	

Emissions indices were measured in clear air at a single-engine fuel flow rate of 0.373 ± 0.025 kg s⁻¹. Data are for the right inboard engine (no. 3) at altitudes of 9,140-10,970 m (30,000-36,000 ft). Significance level: ****P < 0.001, ***P < 0.01, **P < 0.05, *P < 0.1. Particle emissions indices are reported as the geometric mean ± 1 g.s.d.; trace-gas emissions indices are reported as the arithmetic mean ± 1 a.s.d.

Extended Data Table 5 | Emissions indices for no. 2 engine under medium-thrust and cruise conditions

	Medium-sulfur-content Low-sulfur-content Jet A fuel Jet A fuel		ntent	50:50 HEFA:low-sulfur-content Jet A fuel blend	
Emissions index (per kg of fuel)	Emissions index	Emissions index	Ratio	Emissions index	Ratio
Total particle ($D_p > 5$ nm) number	-	3.25×10 ¹⁵ * 1.09	-	2.77×10 ¹⁵ * 1.24	-
Ultrafine particle (5 < D _p < 10 nm) number	-	2.26×10 ¹⁵ * 1.10	-	2.23×10 ¹⁵ * 1.24	-
Fine particle (D _p > 10 nm) number	9.97×10 ¹⁴ * 1.14	1.12×10 ¹⁵ * 1.28	1.12	6.01×10 ¹⁴ * 1.35	0.60*
Volatile fine particle number	5.65×10 ¹⁴ * 1.08	4.96×10 ¹⁴ * 1.39	0.88	3.14×10 ¹⁴ * 1.51	0.56**
Non-volatile fine particle number	4.28×10 ¹⁴ * 1.26	6.13×10 ¹⁴ * 1.23	1.43	2.79×10 ¹⁴ * 1.27	0.65
Total particle (5 < D _p < 120 nm) volume (mm ³)	30.65 * 1.06	37.96 * 1.39	1.24	16.37 * 1.54	0.53*
Volatile particle volume (mm ³)	10.30 * 1.30	15.02 * 1.35	1.46	6.08 * 2.10	0.59
Non-volatile particle volume (mm ³)	20.03 * 1.05	22.75 * 1.44	1.14	9.83 * 1.32	0.49**
PSAP black-carbon-equivalent mass at 467 nm (mg)	41.29 * 1.18	42.40 * 1.13	1.03	17.09 * 1.07	0.41***
PSAP black-carbon-equivalent mass at 530 nm (mg)	39.56 * 1.18	41.07 * 1.13	1.04	16.79 * 1.10	0.42***
PSAP black-carbon-equivalent mass at 660 nm (mg)	37.88 * 1.20	41.45 * 1.18	1.09	17.51 * 1.16	0.46***
Carbon Monoxide, CO (g)	6.06 ± 0.66	4.06 ± 0.41	0.67***	5.12 ± 1.42	0.85
Nitrogen Oxides, NO _x (g)	7.53 ± 0.21	7.67 ± 0.58	1.02	7.35 ± 1.44	0.98
Number of Plume Intercepts	4	8		7	

Emissions indices were measured in clear air at a single-engine fuel flow rate of 0.280 ± 0.020 kg s⁻¹. Data are for the left inboard engine (no. 2) at altitudes of 9,140-10,970 m (30,000-36,000 ft). Significance level: ****P < 0.001, ***P < 0.01, **P < 0.05, *P < 0.1. Particle emissions indices are reported as the geometric mean * 1 g.s.d.; trace-gas emissions indices are reported as the arithmetic mean \pm 1 a.s.d.

Extended Data Table 6 | Emissions indices for no. 2 engine under low-thrust and cruise conditions

	Medium-sulfur-content Jet A fuel	Low-sulfur-con Jet A fuel	ent 50:50 HEFA:low-sulfur- Jet A fuel blend		fur-content end
Emissions index (per kg of fuel)	Emissions index	Emissions index Ratio		Emissions index	Ratio
Total particle ($D_n > 5 \text{ nm}$) number	-	-	-	-	-
Ultrafine particle (5 < D_p < 10 nm) number	-	-	-	-	-
Fine particle ($D_n > 10 \text{ nm}$) number	8.61×10 ¹⁴ * 1.27	-	-	4.39×10 ¹⁴ * 1.04	0.51***
Volatile fine particle number	5.19×10 ¹⁴ * 1.23	-	-	2.48×10 ¹⁴ * 1.00	0.48***
Non-volatile fine particle number	3.39×10 ¹⁴ * 1.36	-	-	1.90×10 ¹⁴ * 1.09	0.56***
Total particle (5 < $D_n < 120 \text{ nm}$) volume (mm ³)	20.79 * 1.19	-	-	13.13	0.63
Volatile particle volume (mm ³)	4.16 * 2.18	-	-	2.13	0.51
Non-volatile particle volume (mm ³)	15.61 * 1.00	-	-	11.00	0.70
PSAP black-carbon-equivalent mass at 467 nm (mg)	32.85 * 1.07	-	-	9.53	0.29
PSAP black-carbon-equivalent mass at 530 nm (mg)	31.90 * 1.04	-	-	7.86	0.25
PSAP black-carbon-equivalent mass at 660 nm (mg)	32.03 * 1.09	-	-	9.88	0.31
Carbon Monoxide, CO (g)	8.17 ± 0.68	-	-	6.33 ± 0.25	0.77***
Nitrogen Oxides, NO _x (g)	6.82 ± 0.20	-	-	6.72 ± 0.12	0.98
Number of Plume Intercepts	5	0		2	

Emissions indices were measured in clear air at a single-engine fuel flow rate of 0.231 ± 0.017 kg s⁻¹. Data are for the left inboard engine (no. 2) at altitudes of 9,140-10,970 m (30,000-36,000 ft). Significance level: ****P < 0.001, ***P < 0.01, **P < 0.05, *P < 0.1. Particle emissions indices are reported as the geometric mean * 1 g.s.d.; trace-gas emissions indices are reported as the arithmetic mean ± 1 a.s.d.

Extended Data Table 7 | Emissions indices for no. 3 engine under low-thrust and cruise conditions

	Medium-sulfur-content Jet A fuel	Low-sulfur-content Jet A fuel		50:50 HEFA:low-sulfur-content Jet A fuel blend	
Emissions index (per kg of fuel)	Emissions index	Emissions index	Ratio	Emissions index	Ratio
Total particle (D _n > 5 nm) number	-	-	-	-	-
Ultrafine particle (5 < D _n < 10 nm) number	-	-	-	-	-
Fine particle (D _p > 10 nm) number	4.33×10 ¹⁴ * 1.33	-	-	2.26×10 ¹⁴ * 1.03	0.52**
Volatile fine particle number	2.59×10 ¹⁴ * 1.27	-	-	1.48×10 ¹⁴ * 1.03	0.57**
Non-volatile fine particle number	1.73×10 ¹⁴ * 1.43	-	-	7.76×10 ¹³ * 1.11	0.45**
Total particle (5 < D_p < 120 nm) volume (mm ³)	9.46 * 1.14	-	-	4.54 * 1.46	0.48*
Volatile particle volume (mm ³)	6.48 * 1.00	-	-	1.19 * 1.05	0.18
Non-volatile particle volume (mm ³)	5.97 * 1.39	-	-	2.33 * 1.25	0.39
PSAP black-carbon-equivalent mass at 467 nm (mg)	13.44 * 1.02	-	-	4.21 * 1.08	0.31****
PSAP black-carbon-equivalent mass at 530 nm (mg)	12.73 * 1.03	-	-	4.06 * 1.14	0.32****
PSAP black-carbon-equivalent mass at 660 nm (mg)	13.07 * 1.09	-	-	4.08 * 1.29	0.31***
Carbon Monoxide, CO (g)	8.14 ± 0.65	-	-	8.50 ± 1.72	1.04
Nitrogen Oxides, NO _x (g)	6.43 ± 0.36	-	-	6.19 ± 0.32	0.96
Number of Plume Intercepts	4	0		4	

Emissions indices were measured in clear air at a single-engine fuel flow rate of 0.231 ± 0.017 kg s⁻¹. Data are for the left inboard engine (no. 3) at altitudes of 9,140-10,970 m (30,000-36,000 ft). Significance level: ****P < 0.001, **P < 0.01, **P < 0.05, *P < 0.1. Particle emissions indices are reported as the geometric mean * 1 g.s.d.; trace-gas emissions indices are reported as the arithmetic mean ± 1 a.s.d.

Extended Data Table 8	Fit coefficients for the number	er size distribution
	,	

Medium-sulfur-content Jet A fuel		t A fuel	50:50 HEFA:low-sulfur-content Jet A fuel blend			
Size Distribution Parameter	N (kg ⁻¹)	D _{g,N} (nm)	$\sigma_{{\mathbf{g}},{\mathbf{N}}}$	N (kg ⁻¹)	D _{g,N} (nm)	$\sigma_{\mathbf{g},\mathbf{N}}$
Total Particle Number, d(El _N)	/d[log(D _n)]:					
Engine #2:						
High Engine Thrust	(1.50±0.03)×10 ¹⁵	29.4±0.4	1.79 ± 0.02	(1.09±0.02)×10 ¹⁵	26.8±0.3	1.74 ± 0.02
Medium Engine Thrust	$(1.07 \pm 0.02) \times 10^{15}$	24.7±0.4	1.78 ± 0.03	$(6.32 \pm 0.13) \times 10^{14}$	24.0±0.3	1.69 ± 0.02
Low Engine Thrust	$(8.28\pm0.21)\times10^{14}$	24.8±0.4	1.72 ± 0.03	$(6.07 \pm 0.67) \times 10^{14}$	19.3±1.8	2.09±0.21
Engine #3:						
High Engine Thrust	(1.14±0.02)×10 ¹⁵	27.8±0.3	1.76 ± 0.02	(8.77±0.12)×10 ¹⁴	25.1±0.2	1.71 ± 0.01
Medium Engine Thrust	$(7.01 \pm 0.17) \times 10^{14}$	23.3±0.3	1.69 ± 0.02	$(3.50\pm0.08)\times10^{14}$	21.5±0.3	1.73 ± 0.03
Low Engine Thrust	$(4.68 \pm 0.07) \times 10^{14}$	23.0±0.2	1.66 ± 0.02	$(2.11 \pm 0.25) \times 10^{14}$	18.6±0.9	1.43 ± 0.08
Non-Volatile Particle Number	, d(El _N)/d[log(D _p)]:					
Engine #2:						
High Engine Thrust	(7.64±0.15)×10 ¹⁴	35.3±0.4	1.72 ± 0.02	(5.41±0.15)×10 ¹⁴	28.7±0.5	1.75 ± 0.03
Medium Engine Thrust	(5.00±0.14)×10 ¹⁴	29.7±0.5	1.64 ± 0.03	(2.62±0.07)×10 ¹⁴	27.8±0.4	1.71±0.02
Low Engine Thrust	$(4.50\pm0.23)\times10^{14}$	25.5±0.9	1.86±0.07	$(4.15\pm0.63)\times10^{14}$	20.9±2.4	2.03±0.27
Engine #3:						
High Engine Thrust	(6.30±0.13)×10 ¹⁴	32.5±0.4	1.71±0.02	$(3.94 \pm 0.12) \times 10^{14}$	28.0±0.5	1.68 ± 0.03
Medium Engine Thrust	$(3.18\pm0.14)\times10^{14}$	27.0±0.7	1.63 ± 0.04	$(1.78 \pm 0.06) \times 10^{14}$	26.3±0.5	1.68 ± 0.03
Low Engine Thrust	$(2.82\pm0.08)\times10^{14}$	23.5±0.5	1.73±0.03	$(1.09\pm0.08)\times10^{14}$	23.4±0.8	1.58 ± 0.06

Log-normal fit coefficients (± 1 a.s.d.) to the geometric mean particle number size distributions measured behind the left and right inboard engines (no. 2 and no. 3, respectively) at altitudes of 9,140–10,970 m (30,000–36,000 ft).



Extended Data Table 9 | Fit coefficients for the volume size distribution

	Medium-sulfur-content Jet A fuel		50:50 HEFA:low-sulfur-content Jet A fuel b			
Size Distribution Parameter	V (mm ³ kg ⁻¹)	D _{g,V} (nm)	$\sigma_{\rm g,V}$	V (mm ³ kg ⁻¹)	D _{g,V} (nm)	$\sigma_{g,V}$
Total particle volume, d(El _V)/	d[log(D _p)]:					
Engine #2:	F					
High Engine Thrust	66.8±2.4	65.7±1.2	1.57 ± 0.03	34.4±1.1	57.2±1.0	1.61±0.03
Medium Engine Thrust	28.5±0.8	57.5±0.8	1.58±0.02	14.7±0.5	52.6±0.5	1.60 ± 0.03
Low Engine Thrust	19.9±0.6	51.9±0.8	1.59±0.02	13.0±2.3	61.6±6.0	1.67±0.19
Engine #3:						
High Engine Thrust	34.4±1.0	60.8±1.0	1.61±0.03	21.9±0.6	53.3±0.7	1.54 ± 0.02
Medium Engine Thrust	14.4±0.6	50.4±1.0	1.51±0.03	6.7±0.5	51.9±2.2	1.73±0.08
Low Engine Thrust	8.9±0.3	48.7±0.7	1.55 ± 0.02	3.3±0.5	49.1±5.0	1.92±0.18
Non-volatile particle volume,	d(El _V)/d[log(D _p)]:				
Engine #2:						
High Engine Thrust	44.1±1.1	68.3±0.7	1.46 ± 0.02	20.4±0.6	61.2±0.8	1.54 ± 0.02
Medium Engine Thrust	19.6±1.1	60.9±1.8	1.57 ± 0.05	9.7±0.5	61.5±1.6	1.66 ± 0.05
Low Engine Thrust	15.8±0.8	64.3±1.8	1.64 ± 0.05	8.9±2.9	58±12	1.77±0.31
Engine #3:						
High Engine Thrust	37.8±1.4	73.0±1.5	1.63 ± 0.04	14.5±0.5	60.5±1.3	1.65 ± 0.04
Medium Engine Thrust	12.6±1.5	66.7±5.2	1.80 ± 0.16	6.0 ± 0.5	61.8±3.4	1.85±0.11
Low Engine Thrust	6.1±0.3	51.8±1.1	1.56 ± 0.03	1.9±0.2	44.9±1.8	1.57±0.07

Log-normal fit coefficients (±1 a.s.d.) to the geometric mean particle volume size distributions measured behind the left and right inboard engines (no. 2 and no. 3, respectively) at altitudes of 9,140–10,970 m (30,000–36,000 ft).