1	Title:
2 3	Dominant role of mineral dust in cirrus cloud formation revealed by global- scale measurements
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24	Abstract:
25	Airborne mineral dust particles can act as natural seeds for cirrus clouds in the upper
26	troposphere. However, dust atmospheric abundance is unconstrained in cirrus-forming regions,
27	hampering our ability to predict these radiatively important clouds. Here, we present global-scale
28	measurements of dust aerosol abundance in the upper troposphere, and incorporate these into a
29	detailed cirrus formation model. We show that dust aerosol initiates cirrus clouds throughout the
30	extra-tropics in all seasons and dominates cirrus formation in the northern hemisphere (75-93%
31	of clouds seasonally). Using a global transport model with improved dust treatment, we also
32	explore which of Earth's deserts are the largest contributors of dust aerosol to cirrus-forming

regions. We find that the meteorological environment downstream of each emission region modulates dust atmospheric lifetime and transport efficiency to the upper troposphere, so that source contributions are disproportionate to emissions. Our findings establish the critical role of dust in Earth's climate system through the formation of cirrus cloud.

37 Main Text:

38 Interactions between aerosol and clouds are a poorly understood aspect of the climate system¹. Clouds strongly influence the balance of solar and terrestrial radiation that determines 39 40 air and surface temperatures. Mineral dust aerosol particles are principal players in initiating the formation of cirrus²⁻⁴, widespread high altitude clouds composed of ice that exert a net warming 41 effect on the planet⁵. Natural cloud seeding by mineral dust competes with other cirrus initiation 42 mechanisms, the dominant process then defining cirrus cloud coverage, vertical extent, 43 microphysical properties, and brightness. Mineral dust's potential to dictate cirrus properties 44 designate it a fundamentally important aerosol type for atmospheric radiative balance and global 45 climate⁶. 46

47 Approximately 1000-4000 Tg of dust aerosol is emitted annually from the world's arid 48 regions⁷⁻⁹, making it among the most abundant aerosol types in the atmosphere. Thick plumes 49 visible from space account for the overwhelming bulk of atmospheric dust aerosol. These low 50 altitude plumes (typically <6 km^{10,11}) reside at temperatures too warm for cirrus formation. A 51 tiny fraction of emitted dust particles are vertically transported to the cold upper troposphere 52 (UT). Even with very low relative abundance (~1 in 10^3 - 10^5 aerosol particles), dust and other 53 ice-nucleating particles (INP) may still effectively control cirrus cloud formation¹².

54 Cirrus are formed by nucleation of water into ice, a fleeting process that occurs at high 55 altitudes and is notoriously difficult to observe directly in the atmosphere⁴. Theoretical 56 treatments of cirrus formation are also challenging because global models must approximate the 57 small-scale (sub-grid) variability of aerosol and ambient conditions to which nucleation is 58 extremely sensitive^{11,13}.

59 Our ability to assess the impact of dust on the climate system and predict the effects of 60 future changes to dust emissions is hampered by a nearly universal lack of dust aerosol 61 abundance measurements in the cirrus-forming regions of the atmosphere. No large-scale dust 62 concentration measurements have been reported in the background UT. Widespread geographic

coverage of dust and other aerosol species can be achieved by some satellite sensors, though they 63 lack the sensitivity to detect dust once thick plumes dissipate $^{14-16}$, only a few are altitude-64 resolved, and absolute concentrations are generally not reported. 65

In our previous studies sampling cirrus clouds directly, we confirmed that mineral dust 66 particles were principally responsible for a limited set of northern hemispheric cirrus clouds². 67 Here we present wide-ranging measurements of dust aerosol in the background atmosphere and 68 investigate dust sources using global models. We assess dust's influence on cirrus by combining 69 70 highly sensitive measurements of dust aerosol, previous laboratory studies of dust INP activity, global simulations of air transport, and offline microphysical simulations of ice nucleation. 71

72

Mineral dust is lofted into the global upper troposphere

73 Mineral dust was sampled on a global scale during the NASA Atmospheric Tomography (ATom) airborne campaigns (Fig. 1). Continuous vertical profiling during north-south transects 74 of the Pacific and Atlantic Ocean basins generated two-dimensional aerosol curtains. The 75 Particle Analysis by Laser Mass Spectrometry (PALMS) instrument identified individual mineral 76 dust particles and other aerosol types from their chemical fingerprints. Dust mass and number 77 concentrations are derived by combining PALMS chemical information with size-resolved 78 aerosol number concentration measured concurrently¹⁷. Over a few minutes sample time, this 79 new, highly sensitive in situ measurement technique can detect dust aerosol at the minute 80 concentrations necessary to investigate its role in cirrus formation (Fig. S1). 81

In Fig. 2 and S2-S4 we present the airborne measurements as global-scale maps of 82 mineral dust in the background troposphere. Dust mass concentrations span several orders of 83 84 magnitude. Most atmospheric dust is emitted as intense plumes from a few arid regions of the 85 world (Fig. 1). As plumes dissipate into the background atmosphere, dust aerosol is transported globally and removed from the atmosphere by precipitation and surface deposition. Dust must 86 87 be lofted to reach the UT where cirrus form, with convective clouds providing a direct route. 88 However, during vertical transport dust and other aerosol are also efficiently scavenged by clouds and precipitation. From 2 to 12 km altitude, average dust mass concentrations decrease 89 by factors of 10-1000 due to dilution and removal by clouds (Fig. 2c-e). Vertical concentration 90 91 gradients vary across latitudes and are strongest over emission sources such as the tropical and

northern Atlantic basin. Polar profiles have shallower or even inverted altitude gradients due to
 weak convection, less precipitation, and low surface emissions¹⁸.

94 Global models parameterize the detailed cloud processes and aerosol losses during convective transport. Uncertainties in these processes have led to highly variable UT aerosol 95 predictions¹¹, with surface-emitted aerosol often vastly over-estimated¹³. Surface-based and total 96 column techniques to evaluate modeled aerosol do not constrain UT dust concentrations because 97 the overwhelming majority of dust aerosol mass exists in the lower troposphere, below cirrus 98 altitudes. In Fig. 2 CESM/CARMA¹⁹ and GEOS/GOCART²⁰ dust simulations give context to 99 the dust encountered by the aircraft. Both models use revised aerosol convective transport 100 schemes (see Methods)^{21,22} that improve the accuracy of UT dust mass concentrations x10-100 101 (Fig. S5). Importantly, the global simulations of mineral dust abundance are now constrained by 102 wide-ranging in situ measurements that can resolve the minute quantities of dust present in the 103 104 background upper troposphere.

105

Certain deserts are more efficient at supplying dust to the UT

Dust emission rates and atmospheric concentrations from the world's most productive source regions have been investigated in many previous airborne and ground-based measurement campaigns, frequently targeting Northern Africa^{6,23-27}. Using the CESM/CARMA global model with an improved convective transport scheme, we explore how Earth's seven foremost dust emission zones (Fig. 1 black boxes) supply dust aerosol to the UT. We focus on cirrus-forming regions where supersaturated water vapor can nucleate ice onto dry particles, typically termed "deposition" nucleation (T<235 K, Fig. S6)⁴.

113 Transport of dust to cirrus-forming regions depends on the amount of dust emitted as well as the downwind meteorological environment. North African dust emissions, primarily from the 114 Saharan Desert, account for 60% of all emissions and dwarf other sources (Fig. 3, Table 1). 115 Infrequent deep convection over the Sahara limits direct dust transport into the UT (Fig. S7). 116 117 Once the main dust plume departs the African continent it slowly subsides and continues to experience minimal convection as it crosses the Atlantic through the Azores High. Most of the 118 plume eventually subsides into the marine boundary layer¹⁰, a high-loss environment where 119 aerosol is scavenged by rain and turbulent mixing to the ocean's surface. In contrast, central 120 121 Asian deserts emit only about 13% of global dust mass, yet through most of the year their

contributions to UT dust are larger than North Africa (Fig. 3). After dust is lofted to several km
 altitude by frontal systems, further vertical transport into the UT is driven by a variety of
 mechanisms^{28,29} such as dry convection, synoptic scale ascent, orographic uplift, and interactions
 with the Tibetan Plateau and the Asian Summer Monsoon^{30,31}. Asian dust in the UT is then
 transported across the Pacific via the sub-tropical jet (Fig. S7)^{28,30}.

Figure 3 summarizes seasonal patterns of UT dust contributions from the seven emission 127 zones. Surprisingly, North Africa does not overwhelm other dust sources. Asian emissions 128 129 compete annually with North Africa, contributing 59-73% of dust to the northern hemisphere 130 (NH) UT during boreal summer, despite Saharan emissions at their annual maximum. Middle Eastern desert inputs are smaller but more consistent seasonally. North America is a minor 131 contributor, both globally and to the NH. Australia, South Africa, and South America deserts are 132 the southern hemisphere's (SH) major sources and contribute 92.3% of the UT dust but account 133 134 for only 4.3% of global UT dust by mass. All contribute similarly to the UT in the SH and have 135 similar seasonal cycles. Tropical UT dust sources are the most varied, with SH emissions 136 dominating during austral summer and NH taking over during other seasons. Source 137 contributions to UT dust have a strong seasonal variability that is not commensurate with emission cycles (lines, Fig. 3), highlighting how downwind environments help modulate vertical 138 transport. 139

The Fig. 3 pie charts summarize how UT dust contributions from each zone do not 140 141 always scale with their annual emissions. Asia's contributions are disproportionally high and 142 North Africa's are low so that each accounts for about 40% of global UT dust. We compare the relative efficiency of dust transport into cirrus-forming regions by normalizing UT 143 concentrations to annual emissions for each zone in Table 1. Relative to North Africa, all other 144 NH zones have higher vertical transport efficiencies, with Asian dust emissions being the most 145 efficient. On a per-emissions basis, about 10 times as much Asian dust reaches cirrus-forming 146 regions compared to North African dust. This amplified Asian transport efficiency relative to 147 North Africa is larger than the estimated model biases for those regions (see Supplemental 148 material) and is consistent with a recent trajectory analysis 28 . In contrast the three SH dust 149 emission zones show less variability, with South America seeding the UT most efficiently. In 150 both hemispheres the largest source is the least efficient at transporting dust to cirrus-forming 151 regions. The well-known north-south emissions gradient in dust $(NH/SH = 7.5)^{7}$ is amplified in 152

UT concentrations (NH/SH = 22), giving the NH a considerably higher potential to induce cirrus
 formation by mineral dust aerosol.

Projections of future dust emission changes are highly uncertain, and even the sign is unknown^{1,32}. The underlying meteorological drivers for dust emission are different across emissions zones, as are their regional responses to climate change forcings³³. In particular, Asian emissions increasing relative to Africa³⁴ would further amplify the disproportionally high influence of Asian dust on cirrus.

160 Assessing dust's role in global cirrus cloud formation

We now evaluate the ability of dust ice-nucleating particles (INP) to generate cirrus. Climate models are unable to explicitly simulate cirrus ice nucleation because model grid cells are typically much larger (10's of km) than individual clouds, and long time steps (~1 hr) do not directly capture the transient water vapor, temperature, and INP fluctuations that lead to cloud formation. Instead, most models rely on sub-grid parameterizations of these highly sensitive variables^{35,36}. In the real atmosphere, temperature and water vapor can have gradients across meter-length scales, and cloud formation occurs over seconds to minutes.

We employ a detailed microphysical model of cirrus ice nucleation at high time 168 resolution, initialized by ATom dust and water vapor measurements (Fig. 4). Starting at the 169 aircraft position, future air temperatures are predicted from 10-day forward trajectories 170 171 calculated from the NOAA Global Forecast Systems model. High-frequency perturbations are superimposed on trajectory temperatures to simulate UT gravity waves not resolved by the 172 model (Fig. S8), thereby providing a realistic ensemble of future temperatures that aircraft-173 174 sampled air parcels will experience. The detailed freezing model is run every hour along each 175 future trajectory. These simulations probe the role of dust aerosol in forming in situ cirrus, i.e., clouds that are not directly associated with convective storms yet account for a large fraction of 176 UT cirrus³⁷. Cirrus simulations do not include freezing during convective transport, where dust 177 178 aerosol is also likely to nucleate ice.

The cirrus simulations incorporate the new dust measurements (Fig. S1), which provide a critical constraint and thereby enable a realistic assessment of cirrus formation in the background atmosphere. Dust particles induce ice formation by deposition heterogeneous nucleation⁴, and background aerosol freeze homogeneously as aqueous solution droplets¹². The competition

between these processes depends on dust abundance, size, and the cooling rate that drives supersaturation of water vapor with respect to ice (S_{ice}). In cases of rapid cooling or in dust-poor environments, dust aerosol has little influence on cirrus cloud properties. Not all dust particles are efficient INP⁴, and the active fraction is extremely sensitive to particle size and S_{ice} (Fig. S9, S10). We employ a recent parameterization of desert dust ice nucleation efficiency based on dozens of controlled cloud formation experiments³⁸.

Within the 10-day simulation period, air parcels from 2759 trajectories experienced cirrus 189 190 formation events at some point. Heterogeneous nucleation on mineral dust aerosol was solely 191 responsible for 71% of cirrus clouds, and the rest were formed by homogeneous freezing (Fig. 5). The relative contribution of dust to ice nucleation is fairly constant above 195 K (Fig. 5a). 192 The two competing nucleation mechanisms act to distribute the available water vapor into 193 drastically different ice crystals populations (Fig. 5b). Homogeneous freezing of the relatively 194 195 abundant aqueous aerosols distributes the condensable water vapor into numerous small ice crystals and generates cirrus with high solar reflectivity. Conversely, most dust-induced cirrus 196 197 initially contain fewer (and given the same water vapor, larger) ice crystals and are optically thinner³⁷, thereby exerting a smaller radiative effect. If we consider only the most radiatively 198 relevant cirrus having abundant ice crystals ($N_i > 10 L^{-1}$), dust still initiates about half (47%) of 199 cirrus globally and 72% in the NH extra-tropics. 200

201 Simulations that artificially suppressed nucleation on dust reveal that dust aerosol changes cloud properties in two important ways (Fig. 5b). First, dust increases cirrus occurrence 202 203 by 42% (from 1945 to 2759 cases). Second, dust inhibits the formation of homogeneously nucleated, more reflective clouds in 59% of the homogeneous cases (1153 out of 1945). Global 204 modeling studies suggest that cloud radiative effects and climate sensitivity depend strongly on 205 ice sedimentation rates that scale with crystal size^{39,40}. If dust nucleation were not considered 206 when simulating cirrus formation, reduced sedimentation rates will alter estimates of cloud 207 radiative effects, UT temperatures, and surface temperature responses to greenhouse gas 208 emissions. 209

Air parcel trajectories span much of the global upper troposphere (Fig. S11). Dust
dominates cirrus formation in the northern extra-tropics (seasonally 75-93% of cirrus cases, Fig.
5c). Dust-induced cirrus occurs throughout the year in the NH (Fig. S12), with the highest

observed dust concentrations in springtime leading to an increasingly dominant effect on cirrus.
Despite generally lower dust abundance in the SH extra-tropics, dust impact on cirrus formation
is still significant (58-71% of cirrus cases). Tropical and subtropical air parcels experienced
fewer freezing events, and despite strong dust emissions within those latitudes, very low UT
concentrations reduced dust's influence on cirrus (34-63%). The tropical tropopause layer (TTL)
above 13 km altitude is minimally sampled by air trajectories (Fig. S11), and therefore dust's
influence on TTL cirrus cannot be fully resolved by this analysis.

220 The analysis confirms that dust aerosol concentration is the principal limiting factor to 221 the cirrus formation mechanism across all seasons and measurement regions (Fig S13). Seasonal differences in temperature, RH, and atmospheric vertical motion add variability to the dust-222 induced cirrus fraction. Heterogeneous nucleation becomes the dominant cirrus-forming 223 mechanism in regions where median dust concentrations are above $\sim 10 L^{-1}$, and at 100 L⁻¹ nearly 224 all *in situ* cirrus are formed by dust aerosol only. The cirrus studied here appear more sensitive 225 to dust aerosol than higher altitude TTL cirrus⁴¹, which required greater dust concentrations to be 226 227 similarly affected.

228 Observational studies support the cirrus simulation results. Simulated occurrence 229 frequencies of non-convective cirrus qualitatively match the latitude dependence of cirrus detected by satellite⁴², with sub-TTL cirrus maxima between 40 and 70°. Satellite and direct 230 sampling studies infer the competition between homogeneous and heterogeneous nucleation 231 mechanisms from cirrus properties. In broad sampling studies at similar altitudes to ATom, high 232 ice crystal concentrations ($N_i > 500 L^{-1}$) indicative of young clouds formed by homogeneous 233 freezing were rarely observed^{41,43}, <1% and <10%. A large fraction (40-90%) of cirrus below 234 the TTL were very thin ($N_i < 15 L^{-1}$), consistent with Fig. 5. In another extensive study, a high 235 fraction of non-convective cirrus were consistent with heterogeneous nucleation or a 236 combination of both mechanisms³⁷. Some observations indicate higher levels or broader regions 237 of clear-sky S_{ice} in the $SH^{44,45}$, consistent with homogeneous freezing being more common than 238 in the NH. Similarly, satellite sensors observe higher N_i in mid and high latitudes during winter 239 when regional dust emissions are lower^{46,47}. However, other analyses of water vapor or cirrus 240 particle concentrations suggest that systematic differences in NH-SH S_{ice} are still unclear^{48,49}. 241

This study demonstrates that mineral dust aerosols convectively lofted to the UT are 242 abundant enough to initiate cirrus cloud formation during all seasons throughout the extra-243 tropics, including the SH. Our approach applied to the ATom cases provides the most 244 quantitative analysis to date of dust's influence on cirrus formation in remote regions. Non-dust 245 INP candidates are less well constrained in their UT abundance and nucleation efficiency. 246 Although we consider dust to be the most important (abundant and active) INP type in the UT, 247 other particle types may contribute seasonally or regionally. Furthermore, we do not consider 248 249 heterogeneous nucleation mechanisms that occur at warmer temperatures, such as immersion freezing, where dust is also effective. In this way the analysis here defines a lower limit to the 250 influence of dust heterogeneous nucleation on the occurrence and properties of global cirrus. 251

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267 **Author contributions:**

KDF wrote the paper with contributions from all authors. KDF, GPS, CAB, AK, CJW, DMM,

GSD, and TB collected airborne data. PY and KHR provided CESM-CARMA model results.

HB, ASD, and PRC provided GEOS model results. ER provided forward trajectory results. EJJ

271 provided cirrus model results.

272 **Competing interests:**

273 The authors declare no competing interests.

Tables: **Table 1.** Annual dust surface emissions and average UT concentrations sourced from each emission $zone^{a}$.

	Surface emissions			UT concentrations		
Dust Emission Zone	Total emissions (Tg yr ⁻¹)	% of total	Relative to largest hemispheric source	Average UT conc. (ng m ⁻³)	% of total	Relative to largest hemispheric source and relative to emissions [‡]
Northern hemisphere sources						
1. North Africa	1338	59.8%	≡ 1	2.06	39.8%	≡ 1
2. Asia	298	13.3%	0.22 (0.17 - 0.29)	2.03	39.1%	9.7 (1.8 - 18.3)
3. Middle East	280	12.5%	0.21 (0.15 – 0.34)	0.76	14.7%	2.8 (0.9 - 6.6)
4. North America	71	2.5%	0.04 (0.03 - 0.05)	0.11	2.1%	2.6 (0.3 – 7.6)
Southern hemisphere sources						
5. Australia	107	4.8%	= 1	0.09	1.8%	≡ 1
6. South Africa	85	3.8%	0.80 (0.67 - 0.98)	0.06	1.1%	1.5 (0.5 - 4.4)
7. South America	71	3.2%	0.66(0.45 - 0.87)	0.07	1.4%	3.4 (0.9 – 16.2)

^a Data are from 2014-2018 simulations using the revised CESM/CARMA model.

²⁷⁷ [†]Annual average of monthly ratios (monthly ranges in parentheses)

[‡]The relative effectiveness of emissions from each zone contributing to UT dust, calculated for
each month as (UT conc. from zone / UT conc. from largest source) / (emissions from zone /
emissions from largest source), then averaged (monthly range in parentheses). Values are
normalized to the largest hemispheric emission source.

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283 Figure Legends:

Fig. 1. Global-scale airborne sampling of mineral dust aerosol during four NASA ATom
campaigns. Flight tracks are colored by seasonal deployment and shaded by altitude from 0.2 to
13 km (light to dark). Continents are colored by satellite-derived land type

(https://modis.gsfc.nasa.gov/data/dataprod/mod12.php), where brown denotes arid regions that
 are the predominant sources of mineral dust to the atmosphere. Black boxes encompass seven
 principal dust emission zones.

Fig. 2. Dust aerosol measurements and simulations during ATom1 in Aug, 2016. Airborne *in situ* measurements over the Pacific basin (a,c), Atlantic basin (b,d), and polar regions for all
 longitudes (e) are compared to two models. Flight tracks (top panels) are colored by measured
 dust mass concentration on a log scale. Background shading shows simulated CESM/CARMA
 dust concentrations, plotted as vertical curtains at the aircraft location. Lines delineate five

latitude bands (polar/mid-latitudes/tropics, see Methods). In the lower panels, measured dust
concentrations (solid) are compared to the CESM/CARMA (long dash) and GEOS/GOCART
(short dash) simulations for each latitude band. See Fig. S2-S4 for other seasons and variability.
Several of the planet's largest dust features were captured, including the intense North African
plume observed over the mid-Atlantic a few days after emission (b) and again >10 days
downwind over the Pacific Ocean at similar altitudes (a). See Supplementary Information for
discussion of spatial and seasonal patterns.

Fig. 3. Annual cycle of UT dust sourced from each desert emission zone from the revised
 CESM/CARMA model. (a-g) Shading indicates each source's fractional contribution to UT dust
 mass as a function of latitude (left axis) and season. Annual averages are listed as percent. Lines
 (right axis) compare fractional UT concentrations (black) and surface emissions (orange) within
 the northern (solid) and southern (dash) hemispheres. Pie charts (h) are global annual averages
 of surface emissions and UT dust contributions from each emission zone.

Fig. 4. Predicting cirrus formation by combining in situ measurements with cloud-aerosol 308 309 simulations. (1) Dust aerosol, water vapor, and temperature are measured from aircraft during continuous vertical profiling, creating a global-scale map of initial conditions. (2) The future 310 movement of each sampled air parcel is calculated from meteorological wind fields. Air parcels 311 that ascend are cooled, whereby cirrus formation becomes possible. (3) Detailed microphysical 312 simulations of cirrus formation are performed along each air parcel's future trajectory to 313 investigate the competition between dust heterogeneous nucleation and homogeneous freezing of 314 background aqueous aerosol (see Methods). Illustration courtesy of K. Bogan, CIRES, 315 University of Colorado Boulder. 316

Fig. 5. Mineral dust's role in global cirrus cloud formation evaluated from all ATom 317 deployments. a) Heterogeneous nucleation on mineral dust (brown curves) dominates the 318 freezing mode. Homogeneous freezing on aqueous aerosol (green) can occur where dust 319 concentrations are very low or where rapid cooling produces highly super-saturated conditions 320 quickly. Solid lines denote all cirrus freezing events, and dotted lines are for cirrus with ice 321 concentrations N_i >10 L⁻¹ (see text). b) Dust-induced freezing generates more frequent clouds 322 with lower ice crystal concentrations. If dust heterogeneous nucleation is suppressed (blue) 323 cirrus clouds with higher ice concentrations form instead. c) The latitudes for cirrus freezing 324 325 events are compiled for all ATom cases (left axes). Seasonal distributions are shown in Fig. S12. Gray shading shows the sample space (number of 1-hr data points) for all air parcel trajectories. 326 Measured dust concentrations (right axis) are averaged for the five latitude ranges in Fig. 2. 327

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454 Methods:

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456 **The NASA Atmospheric Tomography Mission (ATom) airborne sampling campaigns**

During the NASA ATom mission between 2016 and 2018, the NASA DC8 flying 457 research laboratory executed four global-scale sampling campaigns over the Pacific and Atlantic 458 Ocean basins⁵⁰. ATom was designed to evaluate satellite measurements and global models of 459 gas-phase and aerosol species. The vast majority of sampling time occurred in the remote 460 atmosphere, 100's to 1000's of km from continental emission sources. Flights spanned latitudes 461 from about 86 °S to 82 °N. The aircraft executed constant vertical profiles from 0.2 to 13 km 462 altitude, with 5-15 minute legs of constant altitude at the top and bottom of each profile. 463 Despite its broad spatial coverage and multiple deployments, the ATom data set does not 464 constitute a climatology. An analysis of the representativeness of ATom sampling is pending⁵¹. 465 We analyze measurements from all four aircraft deployments, ATom1 (Jul 29 – Aug 23, 2016), 466 ATom2 (Jan 26 - Feb 21, 2017), ATom3 (Sept 28 - Oct 27, 2017), and ATom4 (Apr 24 - May 467 21, 2018). We exclude low altitude flight segments near airports and the Aug 23, 2016 transit 468

- flight over the continental US. In Fig. 2 and elsewhere we average dust concentrations across broad latitude bands, defined as polar (> 60°), mid-latitudes (27° - 60°), or tropical/subtropical
- 471 (<27°) for all deployments. Longitude 71°W separates Pacific and Atlantic Ocean basins.
- 472 ATom aircraft data for each flight are publically available at
- 473 https://daac.ornl.gov/ATOM/guides/ATom_merge.html.
- 474

475 Aerosol concentration measurements

Size-resolved aerosol concentration from $0.003-4.8 \ \mu m^{52}$ was measured by combining 476 two nucleation-mode aerosol size spectrometers (NMASS)⁵³, a Droplet Measurement 477 Technologies Ultra High Sensitivity Aerosol Spectrometer (UHSAS)⁵⁴, and a TSI, Incorporated 478 Laser Aerosol Spectrometer (LAS). The UHSAS and LAS optical spectrometers encompass the 479 mineral dust size range reported here, $D>0.1-4.8 \,\mu\text{m}$. Particle diameters were derived from 480 optical spectrometer calibrations using ammonium sulfate, which has a refractive index that is 481 also appropriate for many mineral dusts and most other tropospheric particle types^{17,52,54}. 482 Sample flows were 0.06 and 0.1 lpm, respectively. 483

Air was isokinetically sampled using a forward-facing, shrouded, diffuser inlet designed 484 by the University of Hawaii. This inlet transmits aerosol with approximately 100% efficiency 485 for sizes up to $\sim 4 \,\mu m$ aerodynamic diameter at low altitude, whereby efficiency drops to 50% at 486 sizes greater or equal to 5.0 μ m (~3 and \geq 3.2 μ m at 12 km)⁵⁵. In characterizing the aircraft inlet, 487 the authors reported the 50% cutpoint sizes as lower diameter limits due to limited large particle 488 statistics in their reference measurement, and they acknowledged that actual efficiencies could be 489 higher⁵⁵. A reliable transmission efficiency curve cannot be generated. Therefore, while it is 490 likely that some undersampling occurred at the large end of the reported size range (D>3-4 µm), 491 due to the ambiguity in the aircraft inlet inefficiency no attempt was made to adjust measured 492 dust concentrations⁵². Supermicron particle number concentrations were scaled to account for 493 sub-isokinetic sampling and losses in instrument tubing, with typical scaling factors of x0.97-494 1.04 for 1 μ m particles and x3.0-3.4 for 4 μ m particles⁵². Cloudy flight segments are excluded at 495 496 1 sec resolution based on data from a wing-mounted cloud particle probe.

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498 Mineral dust aerosol measurements using PALMS

The NOAA Particle Analysis by Laser Mass Spectrometry (PALMS) airborne singleparticle mass spectrometer instrument has been described in detail previously^{56,57}. PALMS characterizes the size and chemical composition of individual aerosol particles from about 0.1–5 μ m in diameter. Single-particle mass spectra are post-processed to classify each particle into a compositional type and that are then counted to determine their relative abundance^{17,58}.

Mineral dust particles are identified as spectra having multiple crustal metal signatures 504 such as silicon, aluminum, iron, and calcium, and often with trace amounts of alkalis, barium, 505 506 tin, antimony, or lanthanides. The dust particle class is very diverse, with several different 507 composition sub-types representing a wide variety of mineralogies. Typical mineral dust spectra have distinctive signatures and are easily differentiated from other particle types that also contain 508 metals, such as biomass burning, alkali salts, meteoric, and oil combustion. Dust classification is 509 further refined using a cluster analysis routine that groups similar spectra⁵⁸. Manual sorting of 510 clusters and reclassification of a minor fraction of the dust spectra further reduces 511 misclassification error, estimated to be $<5\%^{17}$. 512

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514 Mineral dust quantification using PALMS and aerosol size spectrometers

PALMS and other single-particle mass spectrometer instruments do not measure aerosol 515 concentrations directly because unlike commercial size spectrometers, particle detection 516 efficiency is a strong and variable function of size. Recently, we developed a method to combine 517 PALMS size-resolved composition with size-resolved absolute concentration measured over the 518 same size range by optical particle spectrometers instruments¹⁷. Briefly, the fractional 519 abundance of mineral dust within a size range measured by PALMS is multiplied by the absolute 520 concentration within that size range measured by aerosol size spectrometers⁵² to derive number, 521 surface area, volume, and mass concentration of mineral dust aerosol from 0.1 to 4.8 µm 522 geometric diameter. Note that the large size limit is approximate since aircraft inlet sampling 523 524 efficiency becomes increasingly less certain above about ~3-4 µm aerodynamic diameter (see previous sections)^{52,55}. Particle spectrometer number distributions are converted to volume 525 assuming spherical shape, and mineral dust density is prescribed as 2.5 g cm⁻³ to convert volume 526 to mass. Concentrations are reported at ambient temperature and pressure conditions, except 527 standard concentrations are reported (1013 mbar, 273.15 K) are reported for Fig. 2 and S2-S4. 528

529	PALMS and the aerosol size spectrometers sampled continuously during flight. Mass
530	concentration products are averaged every 3 minutes of sampling time. Principal uncertainties in
531	the derived dust mass and number concentrations include the statistical sampling of aerosol and
532	the volume concentration measurement from size spectrometers. Estimated statistical
533	uncertainties from PALMS statistical sampling and identification of mineral dust are 50% at 0.01
534	μ g m ⁻³ and 25% at 1 μ g m ^{-3 17} . Uncertainties in aerosol size spectrometer concentrations over 3
535	minutes are estimated as 7-17% for number and 30-131% in volume and principally due to
536	atmospheric variability of sparse D>0.5 μ m aerosol ⁵² . We estimate the LOD as the minimum
537	concentration for 1 dust particle detected per sample time. For 3 min samples, typical mass
538	LODs are ~0.0001-0.01 μ g m ⁻³ (Fig. S1B) above 3 km altitude. Sample times for dust number
539	concentrations were typically 6 min but up to 24 min for flight segments at constant altitude
540	(range <1 km). Typical LODs in the UT are ~0.5-10 L^{-1} (Fig. S1A). Number concentrations are
541	reported for the size range D=0.18 to 4.8 μ m.

542 Cirrus-forming regions may contain mineral dust particles that are too large to sample by 543 the aircraft inlet and *in situ* instruments. However, dust number concentrations for large sizes 544 will be relatively low. By extrapolating the number distributions from ~4 to 25 μ m diameter 545 using a log-linear function (Fig. S10a), we estimate that large, unsampled dust particles 546 contribute <7% of the total dust number concentration for all ATom cases, yielding a negligible 547 change (<1%) in cirrus initiation statistics.

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Airborne measurements of gases and temperature

550 Water vapor was measured by the Diode Laser Hygrometer (DLH), an external-path 551 diode laser absorption instrument which has flown on the DC-8 and numerous other airborne 552 platforms^{59,60}. DLH water vapor measurements have been intercompared and validated under 553 conditions relevant to this study⁶¹. The temporal resolution of these measurements is greater than 554 20 Hz; the spatial resolution is approximately 10 m in the horizontal and 1 m in the vertical.

- 555 The Meteorological Measurement System⁶² measures ambient dynamic temperature from 556 open wire platinum sensor. Accurate and science quality static temperature is indirectly derived 557 from 3D wind aero-dynamic calibration using aircraft induced maneuvers.
- 558
- 559 Global simulations of mineral dust aerosol using CESM/CARMA
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We use the NSF/NCAR Community Earth System Model (CESM) coupled with the 560 Community Aerosol and Radiation Model for Atmospheres (CARMA) sectional aerosol model 561 19,21,63,64 . CARMA tracks 20 discrete size bins from 0.1 μ m to 17.4 μ m in diameter for internally 562 mixed aerosols including mineral dust, sea-salt, organic material, black carbon and sulfate. 563 Integrated dust mass concentration is calculated between 0.1 and 4.5 µm for comparison with 564 PALMS dust mass concentrations (Fig 2, S2 through S5). The model runs at 1.9x2.5 degree 565 resolution and 56 vertical layers from the surface to ~45 km. At every time step, 566 CESM/CARMA is nudged to the Goddard Earth Observing System model v.5 (GEOS5) 567 dynamic meteorological wind and temperature fields. To compare the modeled dust mass 568 concentrations with ATom measurements (Fig. 2, S2-S5) we extract model output at the aircraft 569 location and time. 570

The dust emission source function depends on particle size and surface wind speed $^{65-67}$. 571 Some dust sources such as co-emission with biomass burning smoke⁶⁸ and resuspension from 572 glaciers^{18,69} are not specifically included. CARMA simulates several dust removal processes at 573 the sub-grid level. Dust can be scavenged below cloud by raindrops. Also, dust can be activated 574 575 as cloud condensation nuclei, then subsequently removed both at and above the cloud base. The parameterization of deep convection in the CESM model assumes a stationary state of an 576 577 ensemble of convective plumes, for which the transport of water, temperature and momentum has been widely validated⁷⁰. Yu et al.²¹ revised CESM/CARMA convective removal processes 578 based on the methodology of previous studies^{71,72}. Aerosol removal was constrained by 579 comparing modeled vertical distributions of sea-salt and black carbon against the ATom datasets. 580 In the present study, we found that simulations with a tuning parameter ($ract = 0.02 \text{ hPa}^{-1}$) in the 581 activation term of Yu et al. best matched ATom mineral dust mass concentration measurements. 582 583 For the source apportionment runs (Fig 3, Table 1, S6, S7) we conducted 8 simulations, in each of which only a single source region emitted dust (7 boxed regions in Fig. 1, plus the global 584 remainder that accounted for <0.1% of total emissions and UT concentrations). The simulations 585 cover 2014 to 2018, preceded by a 3-year spin-up. Monthly and annual mean model output are 586 used to determine the dust UT contributions. Dust emissions and atmospheric concentrations are 587 calculated for the entire CARMA size range. 588

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590 Global simulations of mineral dust aerosol using GEOS/GOCART

591 Dust is simulated in the Goddard Chemistry, Aerosol, Radiation, and Transport (GOCART) aerosol module^{20,73,74} in the Goddard Earth Observing System model v.5 (GEOS) 592 framework.⁷⁵. GOCART simulates BC, organic carbon, sulfate, nitrate, ammonium, dust, and sea 593 salt. For the ATom comparisons, GEOS/GOCART was run at a global ~50-km horizontal 594 resolution with 72 vertical pressure layers from the surface up to 0.01 mbar (~85 km). The 595 dynamical time step was 450 s. The model was run in 'replay mode', which resets the model 596 dynamical state every 6 h to a balanced state provided by the Modern-era Reanalysis for 597 Research and Applications v.2 atmospheric reanalysis. A 2.5-yr simulation was conducted from 598 the beginning of 2016 to cover all of the ATom missions. The first half of 2016 was used as a 599 spin-up period. For model-measurement comparisons, dust mass concentrations are extracted 600 from the model domain at the time and location of the aircraft using a linear interpolation method 601 in both space and time. 602

The emission of dust particles is calculated online based on the GEOS dynamic 603 604 meteorological fields including surface wind and soil moisture using a topographic-based algorithm⁶⁵. Simulated dust size bins are 0.2–2, 2–3.6, 3.6–6.0, 6.0–12, and 12–20 µm dry 605 optical diameter. For comparison to PALMS dust mass concentrations (Fig 2, S2-S5), integrated 606 607 mass is calculated by summing the first two bins, 0.2-3.6 µm, using densities of 2500 and 2650 kg m^{-3} for the first and second bin, respectively. Dust particles are removed by dry and wet 608 deposition via impaction collection of rain/cloud droplets on dust particles. Table S1 describes 609 the treatment of dust wet scavenging in GEOS for the previous baseline case (R7), a sensitivity 610 experiment (R9), and the revised treatment (R23) that was recently formulated using ATom 611 biomass burning aerosol data²². 612

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614 Model evaluation using ATom measurements

The reported dust mass concentrations characterize the atmosphere at the time and location of measurement. Large-scale models should extract simulated dust concentrations that are co-located with aircraft sampling. To assess the skill of CESM/CARMA and GEOS/GOCART to reproduce the PALMS mineral dust observations, we calculated the mean log bias and centered root mean squared log error (CRMSLE) of the models following the method of Schill et al.²² (Fig. S5). A mean log bias of 1 means the model is biased by an order of magnitude $(10^1 = 10)$. This analysis can emphasize disagreement at low concentrations since

relative errors can be large near detection limits.CRMSLE represents the width of the errors
around the mean log bias and is analogous to the standard deviations of the errors if they are
normally distributed.

In some environments, PALMS observed zero dust particles for a significant fraction of data samples. Excluding these samples from the log bias analysis would artificially skew the biases to lower values. Therefore, for samples with zero dust particles, the measured mass concentration is approximated as half the LOD (see Fig. S1).

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630 Forward trajectories of air transport

Forward trajectories were calculated using the Traj3D model^{76,77} run with National
Centers for Environmental Predictions (NCEP) Global Forecast System (GFS) 0.25°x0.25°
resolution meteorology (https://www.ncdc.noaa.gov/data-access/model-data/modeldatasets/global-forcast-system-gfs). Trajectories were initialized along the flight track each
minute and run forward for 10 days with a time resolution of one hour.

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637 **Cirrus ice nucleation simulations**

Our objective is to investigate the impact of mineral dust particles on cirrus formation and initial crystal concentrations. We simulate the competition between homogeneous freezing of aqueous aerosols and heterogeneous ice nucleation on dust particles using a box model. After nucleation, processes such as differential sedimentation, entrainment, and aggregation conspire to reduce ice concentrations as cirrus evolve^{78–81}. Simulation of these subsequent processes is beyond the scope of the current study. See Supplementary Information for a discussion of nucleation mechanisms and contributions by non-dust INP types.

As an air parcel cools and the supersaturation over ice (Sice) increases above about 1.1, 645 646 heterogeneous nucleation on dust particles produces the first ice crystals. If the concentration of ice crystals nucleated on dust particles is sufficient and the cooling is slow enough, then 647 depositional growth of these ice crystals will deplete vapor and halt the rising S_{ice} before the 648 threshold for homogeneous freezing (Sice=1.45-1.70, depending on temperature) is reached. In 649 650 this case, the ice concentration will be determined by the abundance of dust heterogeneous nuclei. If the dust concentration is sufficiently low and cooling is sufficiently rapid, then Sice will 651 continue to rise after heterogeneous ice nucleation on dust has occurred, and homogeneous 652

freezing will ultimately dominate the ice concentration. The ice concentration just after
nucleation will generally be the maximum value over the lifecycle of the cirrus cloud. To
represent a cloud we track the nucleation and growth of hundreds of individual ice crystals⁸² and
associated changes in water vapor. This Lagrangian approach avoids the numerical diffusion
associated with growth/sublimation of ice crystals in an Eulerian (bin) model.

Aqueous aerosols are represented by a log-normal size distribution with a mode radius of 658 0.015 μ m, a standard deviation of 2 and a concentration of 100 cm⁻³. Results are not strongly 659 sensitive to the aqueous aerosol size distribution nor their concentration. Homogeneous freezing 660 rates are calculated using a water activity parameterization⁸³. For representation of 661 heterogeneous nucleation on dust particles, we use the ice nucleation active site (INAS) density 662 approach^{41,84} with a measurement-based parameterization of dust active site density that depends 663 on ice saturation ratio and temperature ³⁸. A fundamental assumption is that the ice nucleation 664 activity of UT dust is accurately represented by the laboratory experiments on desert dust 665 samples. Dust particles can accumulate coatings of soluble material during atmospheric aging, 666 and some laboratory studies indicate that thick coatings reduce nucleation efficiency^{85,86}. 667 668 However, preliminary analysis with PALMS indicates that for these and similar laboratory experiments, dust samples were subjected to coatings that are likely much thicker (and possibly 669 670 more deactivating) than experienced by typical atmospheric dust particles. Mineral dust size distributions are taken from the ATom measurements (see Fig. S10). Dust particles are depleted 671 672 from the size distribution as heterogeneous nucleation takes place.

673Cooling rate is a critical factor controlling the competition between heterogeneous and674homogeneous nucleation and the concentration of ice crystals 87,88 . We use a parameterized wave675spectrum 82 to superimpose high-frequency temperature perturbations to simulate gravity waves676that typically define UT cooling rates. The wave amplitudes were adjusted based on recent677analyses of superpressure balloon measurements 89,90 .

678The parcel-model ice nucleation simulations are initiated for all aircraft sample points679with T \leq 235 K and where $S_{ice}>0.9$ at any point along the 10-day forward trajectory, yielding 2759680separate cases from 23590 total calculated trajectories. Mixing of air parcels is not considered in681this analysis. If more than 90% of the dust particles are depleted by ice nucleation, and the S_{ice} 682has decreased to less than 1.03, we assume no further nucleation will take place. As S_{ice} rises683with decreasing temperature, dust heterogeneous nucleation always precedes homogeneous

- freezing, resulting in many clouds being formed by a combination of both mechanisms.
- 685 However, in nearly all mixed cases homogeneous freezing produced many more ice crystals and
- therefore dominated the cirrus properties. The dominant nucleation mechanism is defined here
- as that which generated a higher ice crystal concentration.

688 **Data availability:**

- In situ data and model output for this study are publically available at
- 690 <u>https://doi.org/10.3334/ORNLDAAC/2006</u>. ATom aircraft data are publically available at
- 691 https://doi.org/10.3334/ORNLDAAC/1925.

692 Code availability:

- 693 Code for the CESM model is publically available at http://www.cesm.ucar.edu/models/cesm1.0/.
- 694 Code for the GEOS model is publically available at https://gmao.gsfc.nasa.gov/GEOS_systems/.
- 695

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