

Satellites Show Aerosol's Impact on Summer Arctic Cloud Freezing

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Abstract

Arctic aerosols affect cloud properties and climate. However, the magnitudes and mechanisms are uncertain, as are how aerosol-cloud relationships might change in a rapidly warming environment. We assessed some of the complex relationships between aerosols, surface, and meteorology in the relatively pristine summertime Arctic and quantified resulting impacts on clouds using CloudSat/CALIPSO data, AIRS relative humidity and temperature, plus MERRA-2 aerosol and meteorological reanalysis products. In line with previous studies, dust aerosol layers over the summertime Arctic sea ice are associated with icier clouds. However, summertime dust-associated cloud glaciation is uncommon at temperatures $>-10^{\circ}\text{C}$ and not likely at lower altitudes in the summer. We use DMS concentrations as a proxy for marine new particle formation. When DMS is elevated, open ocean clouds near the surface (0.6-1.5 km) are up to 12% more prevalent. These findings allow us to make some educated guesses about where key processes are occurring, such as ice nucleation from dust, and to more effectively prioritize aircraft targeting in future field campaigns, such as ARCSIX.

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Lauren Zamora^{1,2}, Ralph Kahn²

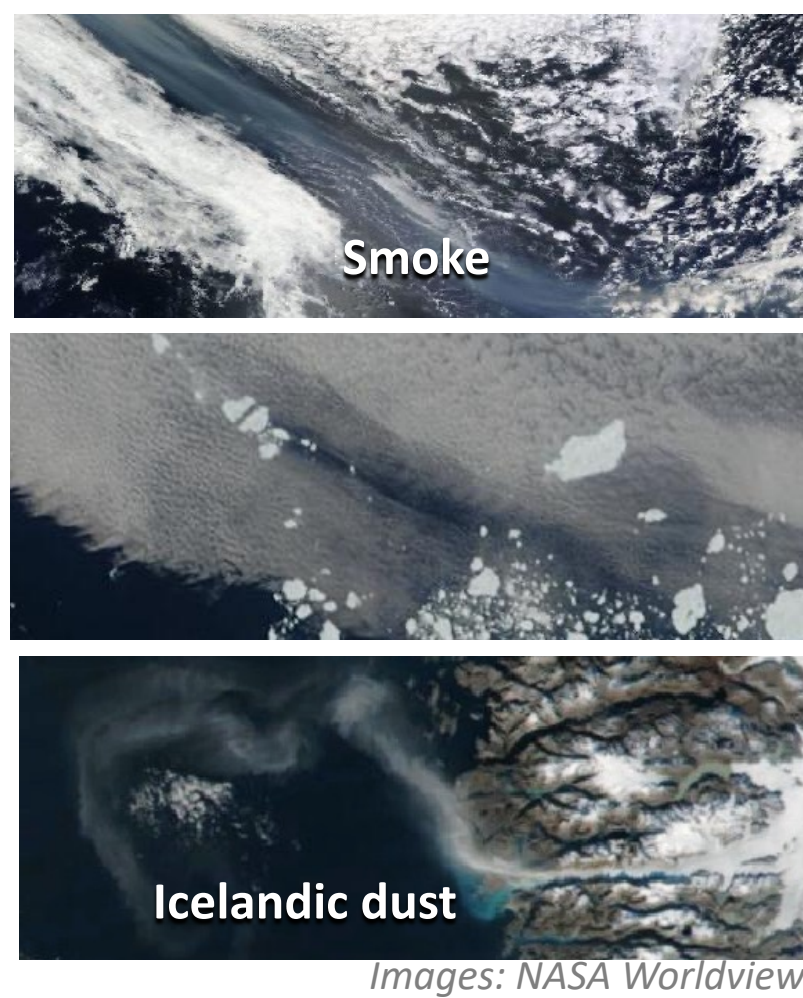
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Background

Arctic aerosols affect cloud properties and climate, and others have shown Arctic aerosols are associated with cloud glaciation^{1,2,3}. However, the magnitudes and mechanisms are uncertain, as are how aerosol-cloud relationships might change in a rapidly warming environment. We assessed some of the complex relationships between aerosols, surface, and meteorology in the relatively pristine summertime Arctic and quantified resulting impacts on clouds.

METHODS

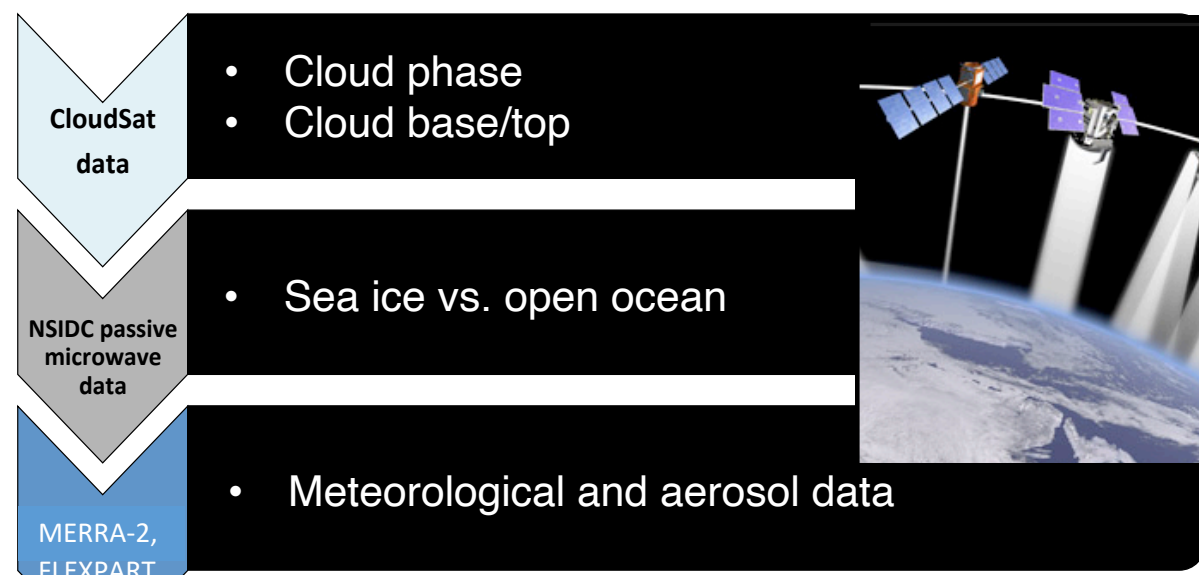
We used CloudSat/CALIPSO data, AIRS RH and T, plus MERRA-2 aerosol and meteorological reanalysis products. These resources provide immense amounts of long-term vertically-resolved information that we can stratify into fine meteorological bins to:



Control for co-varying meteorology

Isolate roles of surface and meteorology

Identify role of aerosol subtype



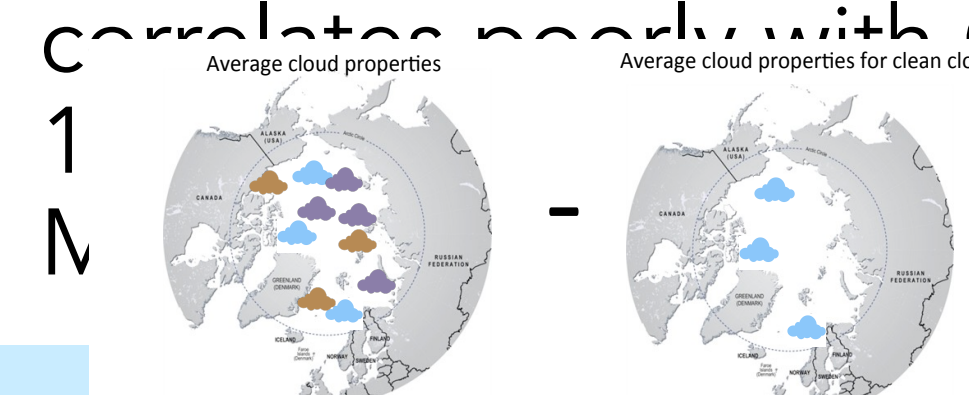
Cloud observation criteria:
• Cloud base > 0.6 km
• Clean clouds: lower 25% of aerosol data in top 5 km

Cloud "iciness" scale:
• 100% for icy clouds
• 50% for mixed phase
• 0% for liquid clouds

THE CALIPSO LIDAR V5 product

prevalence on ECI structures, lidar and slope,

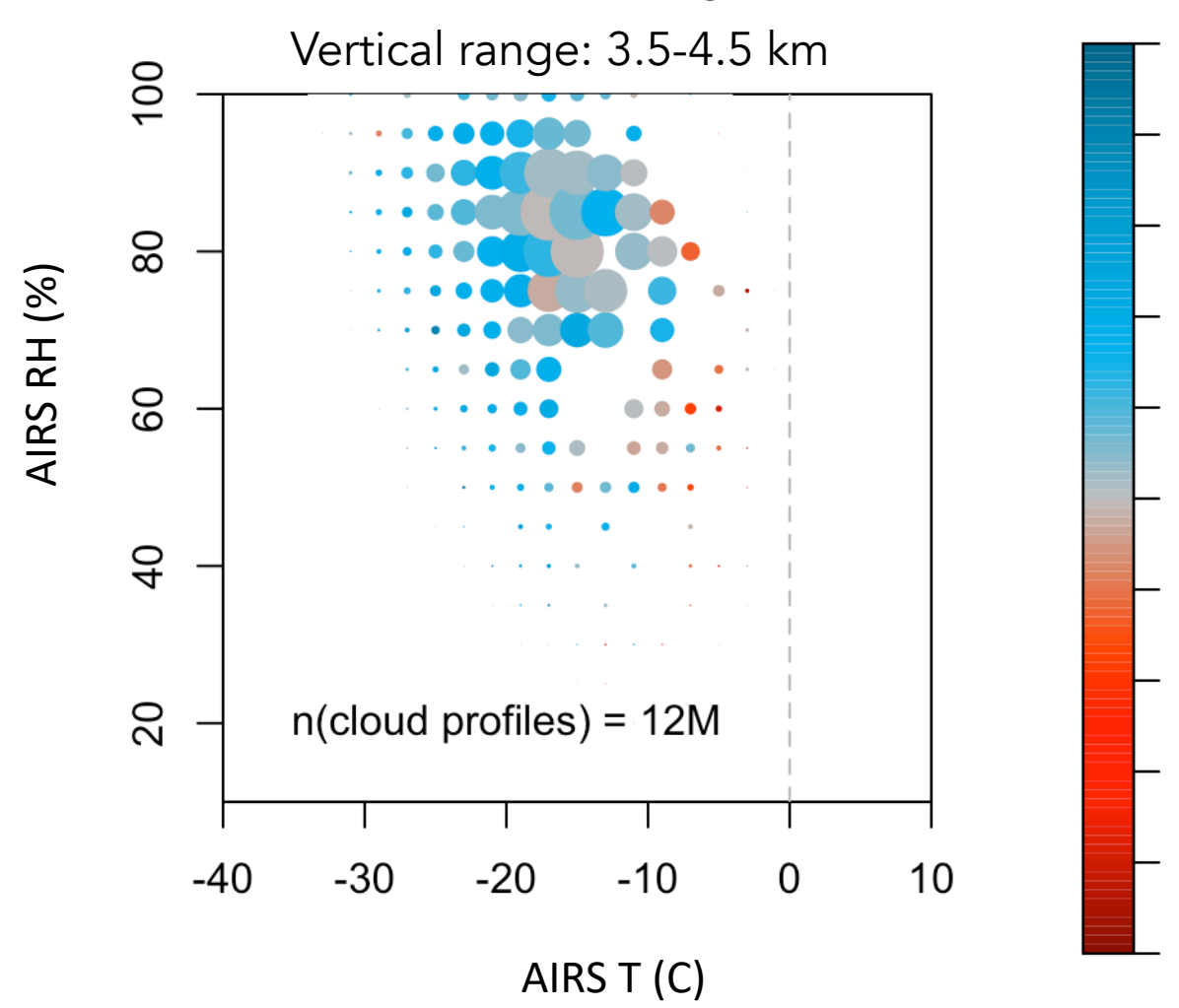
MERRA-2 black carbon (OC), and mineral dust validated in Zamora et al., 2022. MERRA-2 correlates poorly with CALIPSO data below 4 km



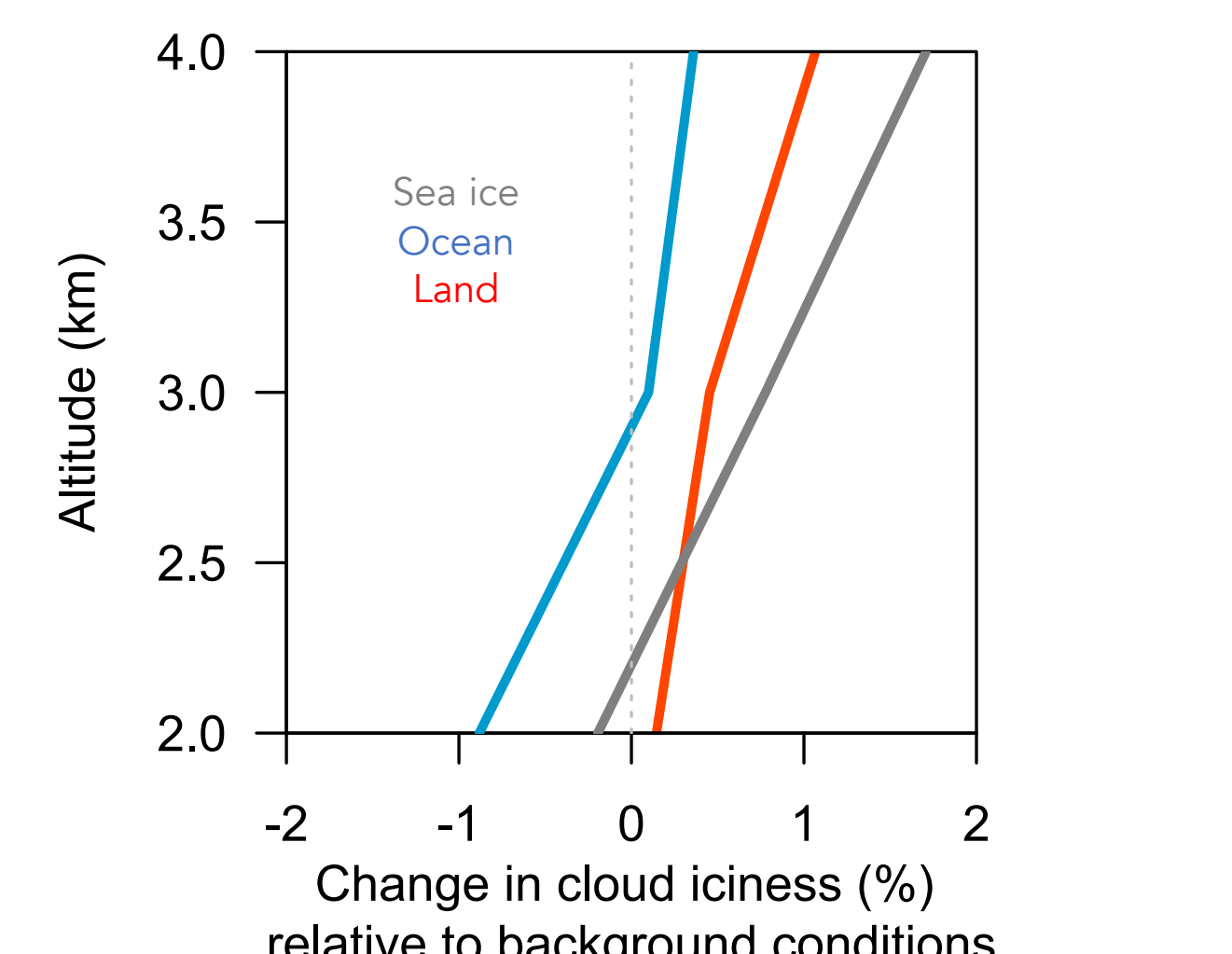
Aerosol net indirect effect on system-wide cloud properties

Preliminary Results

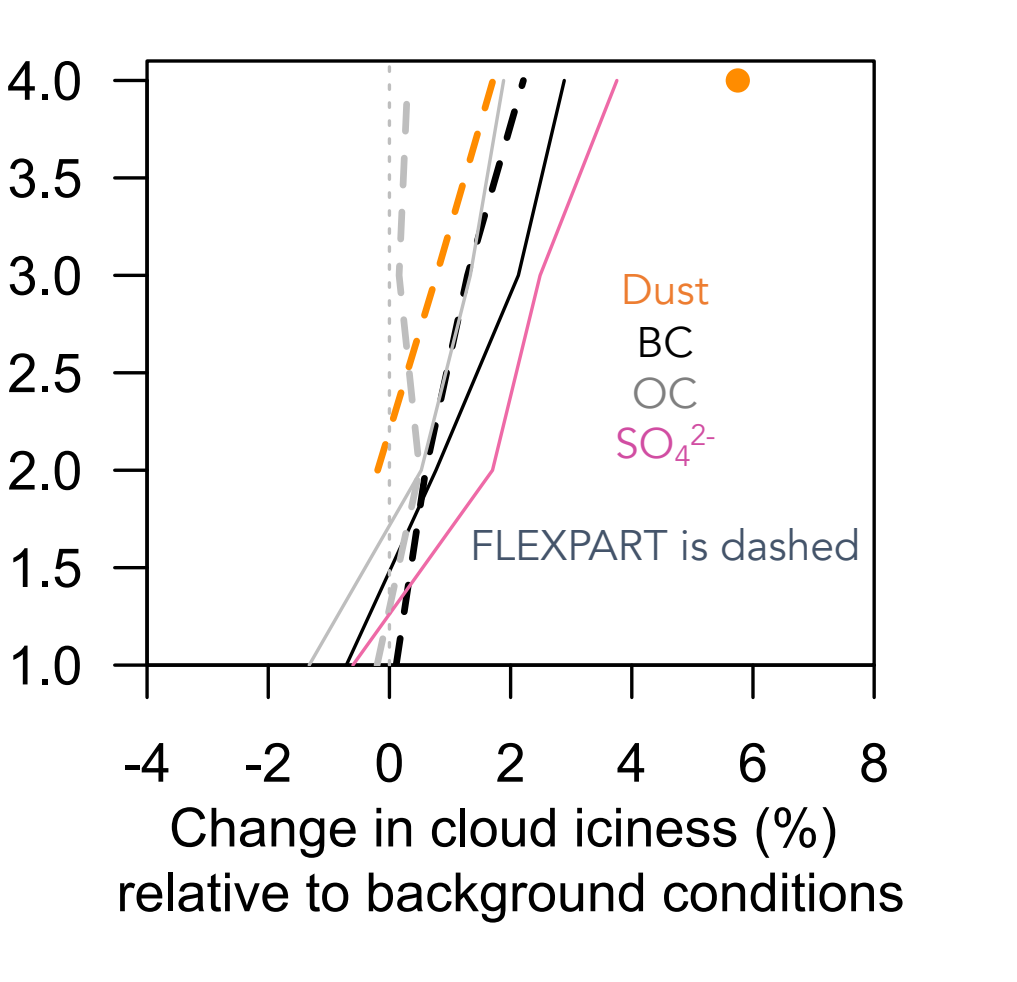
Change in mean cloud iciness (%) at all FLEXPART dust levels, relative to background dust levels



Change in FLEXPART dust-associated cloud iciness over different surface types



Change in cloud iciness relative to background aerosol conditions over sea ice



Data are binned into similar meteorological groups

FLEXPART dust aerosols are associated with icier clouds at some altitudes; effects vary with surface type

Other aerosol types are also associated with icier clouds. Large disagreement between modeled aerosols leads to uncertainty in aerosol effects

Average CloudSat/CALIPSO cloud fraction (CF)

Sea ice

Open ocean

Altitude (km)

CF change (% of clean CF)

CF change (absolute, %)

Altitude (km)

Altitude (km)

Altitude (km)

Altitude (km)

Altitude (km)

Altitude (km)

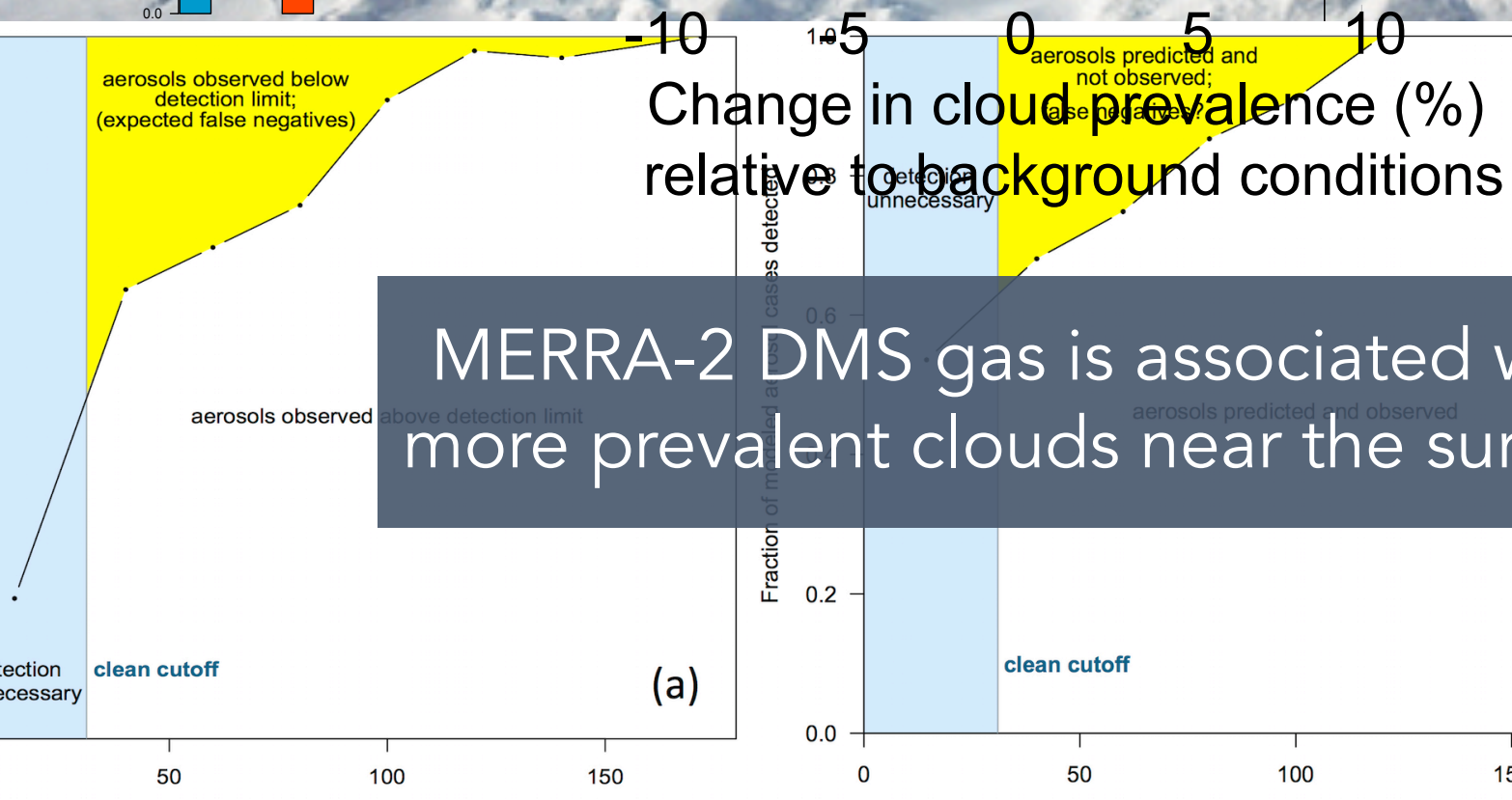
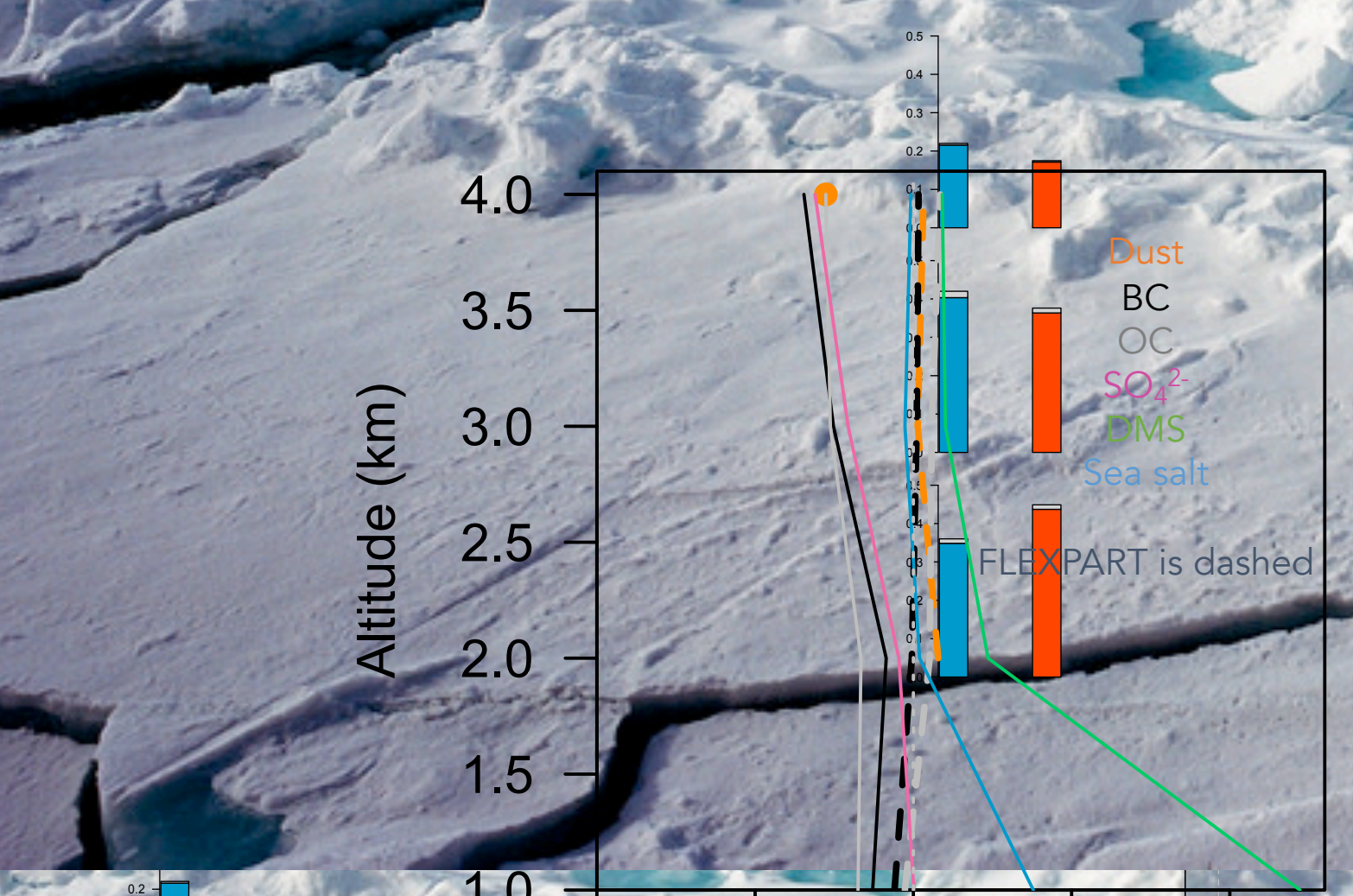
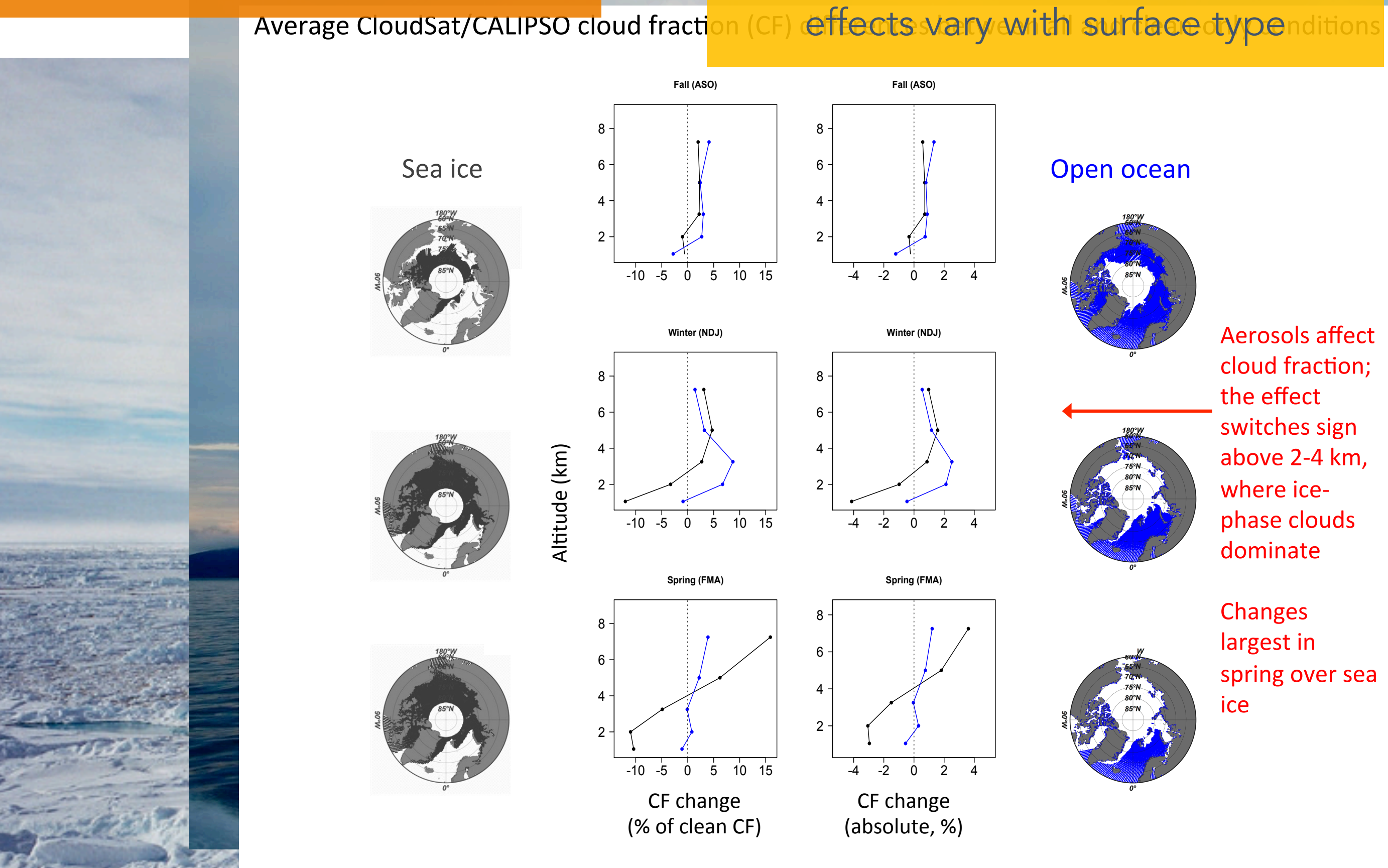
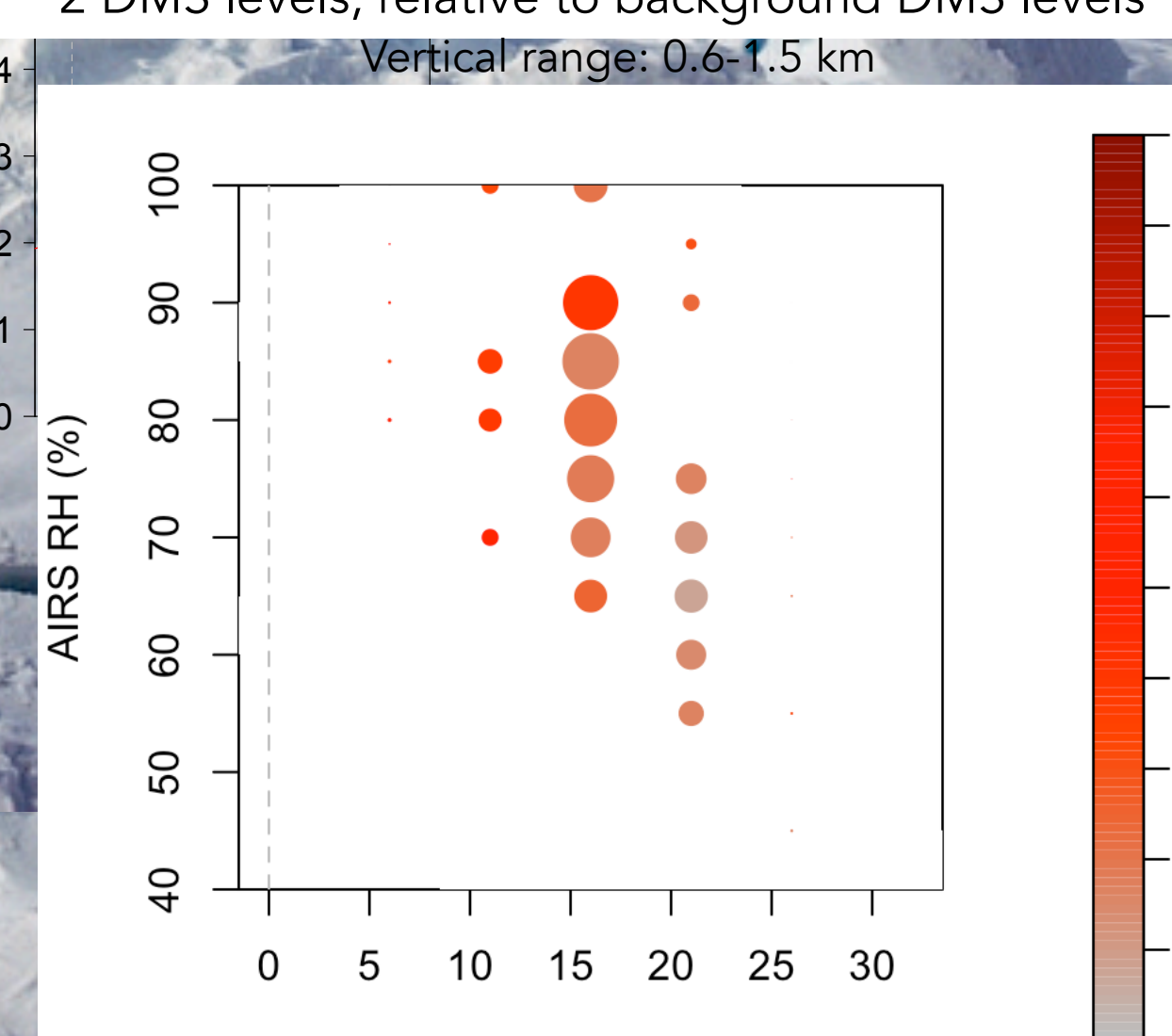


Figure 22: Based on CALIPSO Arctic profiles under non-cloudy conditions, we compare a) the expected fraction and b) possible maximum fraction of false negatives (aerosol present but not detected) for the combustion tracer, black carbon (BC; ng C m⁻³). The expected fraction of false negatives in panel a) was determined by comparing bin-out-of-cloud 2000 AMT₃₅₅ and -B BC concentrations with the fraction of the total number of samples between 1-5 km that had converted backscatter values (Mm⁻¹ sr⁻¹) above the CALIPSO clear-sky nighttime backscatter detection limit from

Change in mean cloud prevalence (%) at all MERRA-2 DMS levels, relative to background DMS levels



As might be expected if small particles were acting as CCN, DMS-related cloud prevalence effects appear to be higher in less stable and more humid environments

Preliminary Conclusions

In line with previous studies, dust aerosol layers over the summertime Arctic sea ice are associated with icier clouds, at least between 2.5-4.5 km; at cold conditions present at 3.5-4.5 km, up to ~6% clouds transition to an icier phase when dust levels are elevated relative to background levels. However, summertime dust-associated cloud glaciation is uncommon at temperatures > -10 °C and not likely at lower altitudes in the summer.

DMS concentrations as a proxy for new particle formation. When elevated, open ocean clouds near surface (0.6-1.5 km) are up to 12% prevalent.

The system is very noisy; results only on statistically large samples, and necessarily to individual clouds. Work ongoing and will benefit from further model validation and refinement.

Findings allow us to make some educated guesses about where key processes are occurring, such as ice nucleation from dust, and to more effectively prioritize aircraft targeting in field campaigns, such as ARCSIX.



We will apply results to upcoming ARCSIX aircraft campaign, summer 2024

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