

Chapter 7: Clouds and Aerosols

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1 Executive Summary

2
3 **A new framework is used, which separates forcing (instantaneous change in the radiative budget) and rapid adjustments (which modify the radiative budget indirectly through fast atmospheric and surface changes) from feedbacks (which operate through changes in climate variables that are mediated by a change in surface temperature).** This framework offers a clear distinction between the traditional concept of radiative forcing (RF) and the relatively new concept of adjusted forcing (AF). For aerosols one can further distinguish forcing processes arising from aerosol-radiation interactions (ari) and aerosol-cloud interactions (aci). [7.1, Figures 7.1 and 7.2]

10
11 **Clouds exert an average cooling influence on Earth, of about -17 W m^{-2} .** This is the net result of a greenhouse (infrared) warming due mainly to high clouds (about $+30 \text{ W m}^{-2}$) and a cooling effect from reflecting solar radiation contributed by all cloud types (about -47 W m^{-2}). This result is not new since AR4, but the important role of clouds in redistributing radiative fluxes vertically within the atmosphere is now better quantified. [7.2.1, 7.2.2, Figure 7.6]

16
17 **The net “clear-sky” feedback from water vapour and lapse rate changes together is *very likely* positive, that is, amplifies global climate changes.** We estimate a *very likely* (90%) range¹ for this feedback parameter, as traditionally defined, of 1.09 (0.91 to 1.27) $\text{W m}^{-2} \text{K}^{-1}$. The mean value and spread in climate models are essentially unchanged from AR4, but are now supported by stronger observational evidence and better process understanding of what determines relative humidity distributions. However the traditional view of a weak inherent climate sensitivity boosted by strong positive feedback from water vapour depends on the analysis framework; a valid alternative yields stronger intrinsic sensitivity, with a weak net clear-sky feedback. [7.2.4, Figure 7.8]

25
26 **The net radiative feedback due to all cloud types is *likely* (>66% chance) positive,** although a negative feedback (damping global climate changes) is still possible. We assign a *very likely* range of -0.2 to $1.4 \text{ W m}^{-2} \text{K}^{-1}$ for the cloud feedback parameter. This conclusion is reached by considering a plausible range for unknown contributions by processes yet to be accounted for, in addition to those occurring in current climate models. The cloud feedback remains the most uncertain radiative feedback in climate models. Observations alone do not currently provide a robust, direct constraint, but multiple lines of evidence now indicate positive feedback contributions from changes in both the height of high clouds and the horizontal distribution of clouds. Additional feedback from low cloud amount is also positive in most climate models, but that result is not well understood, nor effectively constrained by observations, so confidence in it is low. [7.2.4, Figures 7.9-7.10]

36
37 **Persistent contrails from aviation contribute a RF of $+0.02$ ($+0.01$ to $+0.03$) W m^{-2} for year 2010, and the combined contrail and contrail-cirrus AF from aviation is assessed to be $+0.05$ ($+0.02$ to $+0.15$) W m^{-2} .** This forcing can be much larger regionally but is very unlikely to produce observable regional effects on either the mean or diurnal range of surface temperature. [7.2.5]

41
42 **Climate-relevant aerosol properties result from aerosol sources and a number of atmospheric processes, which are considerably better understood than at time of the last IPCC assessment, but the representation of these processes varies greatly in global models.** For instance, the importance of new particle formation, the role of organics, and how mixing increases aerosol mass absorption efficiency are better appreciated. However, it remains unclear what level of sophistication in global aerosol and climate models is required to estimate aerosol-radiation and aerosol-cloud interactions to a sufficient accuracy. [7.3.1, 7.3.2, 7.3.3, Figures 7.11-7.14]

49
50 **Aerosol-radiation interactions result in rapid adjustments, which affect the stability of the atmosphere and cause changes in cloud dynamics that are distinct from any aerosol-cloud interaction.** Observations and detailed large eddy simulations show cloud cover decreases with absorbing aerosol embedded in the cloud layer, and increases when aerosols are above cloud. There is however limited evidence to gauge the relative importance of these two effects at the global scale. [7.3.4]

¹ Unless specified otherwise, all ranges for forcing and feedback parameters are 5 to 95% uncertainty intervals.

Aerosol-climate feedbacks occur mainly through changes in the source strength of natural aerosols or changes in the sink efficiency of natural and anthropogenic aerosols; a limited number of modelling studies have bracketed the feedback parameter within $\pm 0.2 \text{ W m}^{-2} \text{ K}^{-1}$ with a low level of confidence.

There is medium confidence for a weak DMS-CCN-cloud albedo feedback due to a weak sensitivity of CCN population to changes in DMS emissions. Although the limited evidence is for a rather weak aerosol-climate feedback at the global scale during the 21st century, regional effects on the aerosol may be important. [7.3.5]

Research continues to provide evidence for an aerosol influence on the microphysical properties of clouds. There is emerging evidence from observations and large eddy simulations that some of these influences may compensate and thus yield smaller net effects. However there is also evidence for strong aerosol influences that can play an important role in determining the cloud fraction and associated adjusted forcing. These two types of responses are evident in both liquid-only and mixed-phase stratiform clouds. We do not have a good understanding of the prevalence of these two responses. [7.4.2, 7.4.3, 7.4.4]

Research now emphasises the importance of considering the aerosol-cloud-precipitation system, as opposed to aerosol influences in individual clouds. This replaces earlier thinking that correctly identified aerosol-cloud microphysical processes but did not adequately address the multiple feedbacks that can occur within an evolving system. This new paradigm allows for a broader spectrum of responses in climate models and is preferable to earlier approaches that constrained climate model responses to be monotonic and relatively strong. [7.4.3, Figure 7.16]

The representation of aerosol-cloud processes in climate models continues to be a challenge. Aerosol and cloud variability at scales significantly smaller than those resolved in climate models, and the subtle responses of clouds to aerosol at these scales, mean that for the foreseeable future, climate models will continue to rely on parameterisations of aerosol-cloud interactions or other methods that represent subgrid variability. This implies large uncertainties for estimates of the forcings associated with aerosol-cloud interactions. [7.4, 7.5.2]

Cosmic rays enhance aerosol nucleation and cloud condensation nuclei production in the free troposphere, but the effect is too weak to have any climatic influence during a solar cycle or over the last century (*medium evidence, high agreement*). Changes in solar activity affect the flux of galactic cosmic rays impinging upon the Earth's atmosphere, which has been hypothesized to affect climate through changes in cloudiness. Based on available information, no robust association between changes in cosmic rays and cloudiness have been identified. In the event that such an association exists, it is very unlikely to be due to cosmic ray-induced nucleation of new aerosol particles. [7.3, 7.4.5]

The radiative forcing due to aerosol-radiation interactions (RFari) is assessed to be -0.4 (-0.7 to -0.1) W m^{-2} . This assessment is less negative than reported in AR4 because of a re-evaluation of aerosol absorption. The uncertainty estimate is also more robust and based on multiple lines of evidence from models, remotely sensed data, and ground based measurements. [7.5.1, Figure 7.17]

The RFari of different aerosol species (see Table 7.1) shows that aerosols have both negative and positive forcings. These estimates are not as robust as the total RFari as there is not as much supporting observational evidence. The dust RFari could be largely from natural and/or climate feedback effects and should not be considered solely as an anthropogenic forcing agent. [7.3.6, 7.5.1, Figure 7.18]

Table 7.1: The RFari of different aerosol species.

| Species | RFari (W m^{-2}) |
|---|--------------------------------|
| Sulphate | -0.4 (-0.6 to -0.2) |
| Black carbon from fossil fuel and biofuel | $+0.3$ ($+0.1$ to $+0.5$) |
| Nitrate | -0.13 (-0.23 to -0.03) |
| Biomass burning aerosol | $+0.0$ (-0.1 to $+0.1$) |
| Secondary organic aerosol | -0.08 (-0.28 to $+0.12$) |
| Primary organic matter | -0.05 (-0.09 to -0.01) |
| Dust (not necessarily anthropogenic) | -0.1 (-0.3 to $+0.1$) |

1
2 **The adjusted forcing due to aerosol-radiation interactions (AFari) is assessed to be -0.5 (-0.9 to -0.1) W m^{-2} .** The AF includes a rapid adjustment of cloud of -0.1 (-0.3 to $+0.1$) W m^{-2} . While there is robust
3 evidence for the existence of rapid adjustment of clouds in response to absorbing aerosols, these effects are
4 multiple and poorly represented in climate models, leading to a rather large uncertainty. There is the
5 potential of an additional positive contribution to AFari from cloud drop inclusions that is currently not
6 included in this assessment because of the low agreement among the few available studies. [7.3.4, 7.5.2]
7

8
9 **The RF from absorbing aerosol on snow and ice is assessed to be $+0.04$ ($+0.02$ to $+0.09$) W m^{-2} .** Unlike
10 in the last IPCC assessment, this estimate includes the effects on sea-ice, accounts for more physical
11 processes, and incorporates evidence from both models and observations. This RF has a 2-4 times larger
12 global mean surface temperature change per unit forcing and represents a significant forcing mechanism in
13 the Arctic and over Tibet. [7.3.4, 7.5.1]
14

15 **The total adjusted forcing due to aerosols (AFari+aci, excluding the effect of absorbing aerosol on
16 snow and ice) is assessed to be -0.9 (-1.5 to -0.3) W m^{-2} .** The AFari+aci estimate includes rapid
17 adjustments, such as changes to the cloud lifetime and aerosol microphysical effects on mixed-phase, ice and
18 convective clouds. This range was obtained by giving equal weight to satellite-based studies and estimates
19 from climate models and inverse studies grouped together, and includes a small correction term to be in line
20 with a 1750 reference period. It is consistent with multiple lines of evidence suggesting less negative
21 estimates for aerosol-cloud interactions than in AR4. [7.4, 7.5.2, 7.5.3, Figure 7.19]
22

23 **The radiative and adjusted forcings due to aerosol-cloud interactions (RFaci and AFaci) are assessed
24 to be -0.3 (-0.7 to 0) W m^{-2} and -0.4 (-0.9 to 0) W m^{-2} .** The AFaci is estimated as the residual between
25 AFari+aci and AFari. These ranges reflect our understanding of aerosol-cloud interactions in liquid and ice
26 clouds, in particular from observations and large-eddy simulations, which reveal some compensating effects.
27 However they remain somewhat inconsistent with available estimates from climate models, which do not
28 capture all of the relevant processes and tend to simulate more negative forcings. [7.5.2, 7.5.3, Figure 7.19]
29

30 **Multiple lines of evidence suggest that global mean precipitation increases in a warming climate, but at
31 a rate that is limited by the increase of radiant energy absorbed by the surface rather than by the more rapid
32 increase in atmospheric water vapour.** [7.6.2, 7.6.3]
33

34 **Regional precipitation changes will vary and are hard to predict, but a physical basis exists for
35 predicting some systematic aspects of the changes.** First, there is high confidence (high agreement,
36 medium evidence) that as climate warms wet regions will become wetter on average and dry regions drier,
37 although circulation changes will alter results in specific regions. Second, there is also high confidence (high
38 agreement, medium evidence) that short-duration (a few days or less) precipitation extremes as a whole will
39 increase more strongly with temperature than will the globally-averaged precipitation, but this will vary by
40 region according to local shifts in mean precipitation. Lack of understanding of soil moisture, clouds and
41 precipitation interactions, as well as controls on local circulations, limits confidence in projections of
42 precipitation changes on the scale of even the largest catchments. [7.6.2, 7.6.3, 7.6.5]
43

44 **There is a variety of evidence that rapid responses related to aerosol-cloud interactions impact the
45 spatial and temporal distribution of precipitation, but these effects are situation dependent and not
46 systematic.** There is very little evidence that aerosol-cloud interactions have a marked impact on the total
47 amount of precipitation, except in so far as they modify globally averaged surface temperatures. [7.6.4]
48

49 **Theory, model studies and observations suggest that some Solar Radiation Management (SRM)
50 methods, if realisable, could substantially offset a global temperature rise and some of its effects.** There
51 is medium confidence (*medium evidence, medium agreement*) that stratospheric aerosol SRM is scalable to
52 counter the RF and some of the climate effects expected from a twofold increase in CO_2 concentration.
53 Models cited in this assessment explored various source injection strategies and treated aerosol microphysics
54 differently; these variations produce significantly different estimates for the injection rate of chemicals
55 needed to generate the required RF. There is no consensus on whether a similarly large RF could be achieved
56 from cloud brightening SRM due to insufficient understanding of aerosol-cloud interactions. It does not

1 appear that land albedo change SRM can produce a large RF. Limited literature on other SRM methods
2 precludes their assessment. [7.4.3, 7.7.2, 7.7.3, 7.7.4]

3
4 **SRM would produce an inexact compensation for the RF by greenhouse gases.** There is very high
5 confidence that there would be residual regional differences in climate (e.g., temperature and rainfall) in
6 comparison to an unperturbed climate, however models consistently suggest that SRM would generally
7 reduce climate differences compared to a world with the same elevated greenhouse gas concentrations and
8 no SRM. [7.7.3, Figures 7.22-7.23]

9
10 **Numerous side effects and risks from SRM have been identified.** For example, there is high confidence
11 that SRM by stratospheric sulfate aerosols would increase polar stratospheric ozone depletion. Moreover, if
12 SRM were used to counter a large RF by greenhouse gases and then terminated, most of the warming that
13 had been offset would become evident within a few decades, and the rate of climate change would exceed
14 the rate that would have occurred in the absence of SRM. Other side effects have been identified, and there
15 would be other unanticipated or unexplored impacts. SRM will not compensate for ocean acidification from
16 increasing CO₂. [7.7.2, 7.7.3, 7.7.4, Figure 7.24]

7.1 Introduction

The atmosphere is mostly composed of gases, but also contains liquid and solid matter in the form of particles. It is usual to distinguish these particles according to their size, chemical composition, water content and sedimentation velocity into atmospheric aerosol particles, cloud particles and falling hydrometeors. Despite their small mass or volume fraction, particles in the atmosphere strongly influence the transfer of radiant energy and the spatial distribution of latent heating through the atmosphere, thereby influencing the weather and climate.

Clouds usually form in rising air, which expands adiabatically and cools until cloud formation occurs, usually through the nucleation or freezing of aerosol particles. Cloud particles are generally larger than aerosol particles and mostly composed of liquid water or ice. The evolution of a cloud is governed by the balance between a number of dynamical, radiative and microphysical processes. Cloud particles of sufficient size become falling hydrometeors, which are categorised as drizzle drops, raindrops, snow crystals, graupel and hailstones. Precipitation is an important, but difficult to comprehend, climate variable which is influenced by the distribution of moisture and cloudiness, and to a lesser extent by the concentrations and properties of aerosol particles.

Atmospheric aerosols are small solid and liquid particles suspended in the air that can be of natural or anthropogenic origin. They interact with solar radiation, through absorption and scattering, and to a lesser extent with terrestrial radiation, through absorption, scattering and emission. Aerosols can serve as cloud condensation nuclei (CCN) and ice nuclei (IN) upon which cloud droplets and ice crystals form. They also play a wider role in biogeochemical cycles in the Earth system, for instance by carrying nutrients to ocean ecosystems.

Cloud and aerosol amounts and properties are extremely variable in space and time. The short lifetime of cloud particles in subsaturated air creates relatively sharp cloud edges and fine-scale variations in cloud properties, which is much less typical of aerosol layers. While the distinction between aerosols and clouds is generally appropriate and useful, it is not always unambiguous, which can cause interpretational difficulties (e.g., Charlson et al., 2007; Koren et al., 2007).

Clouds respond to climate forcing mechanisms in multiple ways, and can alter both the net response of the system, for instance the globally-averaged surface temperature, but also modify the potency of the initial forcing. The representation of cloud processes in climate models has been recognised for decades as a continuing source of much of the uncertainty associated with our understanding of changes in the climate system (e.g., Arakawa, 1975; Arakawa, 2004; Bony et al., 2006; Cess et al., 1989; Charney, 1979; Randall et al., 2003), but cloud processes have never been systematically assessed by the IPCC before. Key issues include the representation of cumulus convection, cloud formation and dissipation processes. Inter-model differences in cloud feedbacks constitute by far the primary source of spread of both equilibrium and transient climate responses simulated by the CMIP3 climate models (Dufresne and Bony, 2008) despite the fact that most models simulate a near-neutral or positive cloud feedback (Randall et al., 2007 ; and later in this chapter).

Anthropogenic aerosols are responsible for a radiative forcing of climate change through their interaction with radiation and as a result of their interaction with clouds. Quantification of these effects is fraught with uncertainties (Haywood and Boucher, 2000; Lohmann and Feichter, 2005). While previous attempts to quantify the net anthropogenic radiative forcing from first principles have found that it is very likely or virtually certain to be negative, they could not rule out the possibility of positive values (Forster et al., 2007; Haywood and Schulz, 2007). Our inability to better quantify non-greenhouse-gas radiative forcings, and primarily those that result from aerosol-cloud interactions, underlie difficulties in constraining climate sensitivity from observations, even if we had a perfect knowledge of the temperature record (Stevens and Schwartz, 2012).

How clouds and aerosols contribute to climate change is conceptualized with the help of Figure 7.1, which also provides an overview of important terminological distinctions. *Forcings* associated with forcing agents such as greenhouse gases and aerosols act on global mean surface temperature through the global radiative (energy) budget. *Rapid forcing adjustments* (sometimes called rapid responses) arise when forcing agents, by

1 altering flows of energy internal to the system, affect cloud cover or other components of the climate system
2 and thereby alter the global budget indirectly. Because these adjustments do not operate through changes in
3 the global mean surface temperature, which are slowed by the massive heat capacity of the oceans, they are
4 generally rapid and most are thought to occur within a few weeks. *Feedbacks* are associated with changes in
5 climate variables that are mediated by a change in temperature; they contribute to amplify or damp the
6 changes to the global mean surface temperature via their impact on the radiative budget. Such a framework
7 offers a clear distinction between the traditional concept of radiative forcing (RF, defined as the
8 instantaneous radiative forcing with stratospheric adjustment only) and the relatively new concept of
9 adjusted forcing (AF, which includes other atmospheric and surface rapid adjustments) as introduced in
10 Chapter 1 and further discussed in Section 8.1. Figure 7.2 links the former terminology of aerosol direct,
11 semi-direct and indirect effects with the new terminology used in this Chapter and in Chapter 8.

12 [INSERT FIGURE 7.1 HERE]

13 **Figure 7.1:** Overview of feedback and forcing pathways involving clouds and aerosols. Forcing mechanisms are
14 represented by black arrows; forcing agents are boxes with grey shadows, rapid forcing adjustments (also called rapid
15 responses) are shown with brown arrows and feedbacks are other-coloured arrows. See text for further discussion.

16 [INSERT FIGURE 7.2 HERE]

17 **Figure 7.2:** Schematic of the new terminology used in this assessment report for aerosol-radiation and aerosol-cloud
18 interactions and how they relate to the terminology used in AR4. The radiative forcing from aerosol-radiation
19 interactions (abbreviated RFari) encompasses radiative effects from anthropogenic aerosols before any adjustment takes
20 place, and corresponds to what is usually referred to as the aerosol direct effect. Rapid adjustments induced by aerosol
21 radiative effects on the surface energy budget, the atmospheric profile and cloudiness contribute to the adjusted forcing
22 from aerosol-radiation interactions (abbreviated AFari). They include what has earlier been referred to as the semi-
23 direct effect. The radiative forcing from aerosol-cloud interactions (abbreviated RFaci) refers to the instantaneous effect
24 on cloud albedo due to changing concentrations of cloud condensation and ice nuclei. All subsequent changes to the
25 cloud lifetime and thermodynamics are rapid adjustments, which contribute to the adjusted forcing from aerosol-cloud
26 interactions (abbreviated AFaci).

27
28
29
30 For the first time in the IPCC WGI assessment reports, clouds and aerosols are discussed together in a single
31 chapter. Doing so allows us to assess, and place in context, recent developments in a large and growing area
32 of climate change research. In addition to assessing cloud feedbacks and aerosol forcings, which were
33 covered in previous assessment reports in a less unified manner, it becomes possible to assess understanding
34 of the multiple interactions among aerosols, clouds and precipitation and their relevance for climate and
35 climate change. Section 7.2 describes understanding of the role of clouds in climate change, and Section 7.3
36 discusses aerosol processes and properties. Section 7.4 covers aerosol-cloud interactions. Estimates of the
37 aerosol RF and AF due to aerosol-radiation and aerosol-cloud interactions are synthesised in Section 7.5.
38 Section 7.6 is devoted to precipitation processes. Finally Section 7.7 assesses solar radiation management
39 techniques aimed at cooling the planet as a number of these techniques rely on the modification of aerosols
40 and clouds.

41 7.2 Clouds

42 This section summarises our understanding of clouds in the current climate from observations and process
43 models; advances in the representation of cloud processes in climate models since AR4; assessment of cloud,
44 water vapour and lapse rate feedbacks; and the radiative forcing due to clouds induced by moisture released
45 by two anthropogenic processes (air traffic and irrigation). Aerosol-cloud interactions are assessed in Section
46 7.4. The fidelity of climate model simulations of clouds in the current climate is assessed in Chapter 9.

47 7.2.1 Clouds in the Present-Day Climate System

48 7.2.1.1 Cloud Formation, Cloud Types, and Cloud Climatology

49 To form a cloud, air must cool or moisten until it is sufficiently supersaturated to nucleate some of the
50 available condensation or freezing nuclei. Clouds may be composed of liquid water (possibly supercooled),
51 ice, or both (mixed phase). The nucleated cloud particles are initially very small, but grow by vapour
52 deposition. Other microphysical mechanisms dependent on the cloud phase (e.g., droplet collision and
53 coalescence for liquid clouds, riming and Wegener-Bergeron-Findeisen processes for mixed-phase clouds,
54
55
56
57
58

1 and snow formation for ice clouds) can produce a broader spectrum of particle sizes and types; turbulent
2 mixing produces further variations in cloud properties on scales from kilometres to less than a centimetre
3 (Brenguier, 1993). If and when some of the droplets or ice particles become large enough, these will fall out
4 of the cloud as precipitation.

5
6 Clouds form in diverse ways, including large-scale or orographically-driven ascent, convection in unstable
7 atmospheric layers, radiative or evaporative cooling, or turbulent mixing of a moist layer. Air circulations
8 often organise convection and associated clouds into coherent systems having scales from tens to thousands
9 of kilometres, such as cyclones or frontal systems. These represent a significant modelling and theoretical
10 challenge, since they are usually too large to represent in explicit cloud models (Section 7.2.2.1.1), but are
11 also not well resolved nor parameterised by most climate models; this gap however is beginning to close
12 (Section 7.2.2.1.2). Finally, clouds and cloud systems are organized by larger-scale circulations into different
13 regimes such as deep convection near the equator, subtropical marine stratocumulus, or midlatitude storm
14 tracks guided by the tropospheric westerly jets. Figure 7.3 shows a selection of widely occurring cloud
15 regimes schematically and as they might appear in a typical geostationary satellite image.

16 [INSERT FIGURE 7.3 HERE]

17 **Figure 7.3:** Diverse cloud regimes reflect diverse meteorology. (a) A visible-wavelength geostationary satellite image
18 shows (from top to bottom) expanses and long arcs of cloud associated with extratropical cyclones, subtropical coastal
19 stratocumulus near Baja California breaking up into shallow cumulus clouds in the central Pacific, and mesoscale
20 convective systems outlining the Pacific intertropical convergence zone or ITCZ. (b) A schematic section through a
21 typical warm front of an extratropical cyclone (see orange dots in panel (a)) showing multiple layers of upper-
22 tropospheric ice (cirrus) and mid-tropospheric water (altostratus) cloud upwind of the frontal zone, an extensive region
23 of nimbostratus associated with frontal uplift and turbulence-driven boundary layer cloud in the warm sector. (c) A
24 schematic section along the low-level trade wind flow from a subtropical west coast of a continent to the ITCZ (see red
25 dots in panel (a)), showing typical low-latitude cloud types, shallow stratocumulus above the cool waters of the oceanic
26 upwelling zone near the coast and trapped under a strong subsidence inversion, and shallow cumulus over warmer
27 waters further offshore transitioning to precipitating cumulonimbus cloud systems with extensive cirrus anvils
28 associated with rising air motions in the ITCZ.

29
30
31 Clouds cover about two thirds of the globe (Figure 7.4a,c). The midlatitude oceanic storm tracks and tropical
32 precipitation belts are particularly cloudy, while continental desert regions and the central subtropical oceans
33 are relatively cloud-free. Clouds are composed of liquid at temperatures above 0°C, ice below –38°C, and
34 either phase at intermediate temperatures (Figure 7.4b). While temperatures at any given altitude are warmer
35 in the tropics, clouds also extend higher there, such that ice cloud amounts are comparable to those at high
36 latitudes. Approximately 40% of the liquid clouds are at least lightly precipitating (Figure 7.4d), but this
37 fraction is much smaller for ice clouds. These fractions are sensitive to the criterion for defining
38 precipitation, especially for ice clouds.

39
40 In this chapter clouds above the 440 hPa pressure level are considered “high,” those below 680 hPa “low,”
41 and those in-between are considered mid-level, following Rossow and Schiffer (1991). Most high clouds
42 (mainly cirrus and deep cumulus outflows) occur near the equator and over tropical continents, but can also
43 be seen in the midlatitude storm track regions and over midlatitude continents in summer (Figure 7.5a,e);
44 they are produced by the storms generating most of the global rainfall in regions where tropospheric air
45 motion is upward, such that dynamical, rainfall and high-cloud fields closely resemble one another (Figure
46 7.5d, h). Mid-level clouds (Figure 7.5b,f), comprising a variety of types, are seen mainly in the storm tracks.
47 Low clouds (Figure 7.5c,g), including shallow cumulus and stratiform clouds, occur over essentially all
48 oceans but are most prevalent over cooler subtropical oceans where tropospheric air motion is downward.
49 They are less common over land, except at night and in winter.

50 [INSERT FIGURE 7.4 HERE]

51 **Figure 7.4:** (a) Annual mean cloud fractional occurrence (CloudSat/CALIPSO 2B-GEOPROF-LIDAR dataset for
52 2006–2011); (b) annual zonal mean liquid water path (blue shading, ocean only, O’Dell et al. (2008) microwave
53 radiometer dataset for 1988–2005; the 90% uncertainty range, assessed to be 70–150% of the plotted value, is
54 schematically indicated by the white error bar) and ice water path (grey shading, from CloudSat 2C-ICE dataset for
55 2006–2011; the 90% uncertainty range, assessed to be 50–200% of the plotted value, is schematically indicated by the
56 black error bar). (c-d) latitude-height sections of annual zonal mean cloud (including precipitation falling from cloud)
57 occurrence and precipitation (attenuation-corrected radar reflectivity >0 dBZ) occurrence; the latter has been doubled to
58

1 make use of a common colour scale (2B-GEOPROF-LIDAR dataset). The dashed curves show the annual-mean 0°C
2 and -38°C isotherms.

3 4 **[INSERT FIGURE 7.5 HERE]**

5 **Figure 7.5:** (a-d) DJF mean high, middle and low cloud cover from CloudSat/CALIPSO GEOPROF dataset (2006–
6 2011), and 500 hPa vertical pressure velocity (colours) and GPCP precipitation (magenta contours at 3 mm day⁻¹ in
7 dash and 7 mm day⁻¹ in solid); (e-h) same as (a-d), except for JJA. For low clouds, the CALIPSO-only GOCCP dataset
8 is used at locations where it indicates as larger fractional cloud cover, because the GEOPROF dataset removes some
9 clouds with tops at altitudes below 750 m. Low cloud amounts are likely underrepresented in regions of high cloud
10 (Chepfer et al., 2008), although not as severely as with earlier satellite technologies.

11 12 *7.2.1.2 Effects of Clouds on Earth’s Radiation Budget*

13
14 The effect of clouds on Earth’s present-day top-of-atmosphere (TOA) radiation budget, or cloud radiative
15 effect (CRE), can be inferred from satellite data by comparing upwelling radiation in cloudy and non-cloudy
16 conditions. Loeb et al. (2009) estimate that by enhancing the planetary albedo, clouds exert a global and
17 annual shortwave cloud radiative effect (SWCRE) of -47 W m⁻² and, by contributing to the greenhouse
18 effect, exert a mean longwave effect (LWCRE) of +30 W m⁻². Other published satellite estimates differ from
19 these by 10% or less (Loeb et al., 2009). The net global mean CRE of -17 W m⁻² implies a net cooling effect
20 of clouds on climate. Each of the individual longwave and shortwave CREs is large compared to the ~4 W
21 m⁻² radiative forcing of doubling CO₂. Thus clouds have the potential to cause significant climate feedback,
22 depending on how climate-sensitive are the properties governing these two effects (Section 7.2.4). Clouds
23 also exert a CRE at the surface, thus affecting the hydrological cycle (Section 7.6), though this aspect of
24 CRE has received less attention (see Section 7.2.2.2).

25 26 **[INSERT FIGURE 7.6 HERE]**

27 **Figure 7.6:** Distribution of annual-mean top of atmosphere (a) SWCRE, (b) LWCRE, (c) net CRE (2001–2010 average
28 from CERES-EBAF) and (d) precipitation (1981–2000 average from GPCP).

29
30 The regional patterns of annual-mean TOA CRE (Figure 7.6a-b) reflect those of the altitude-dependent cloud
31 distributions. High clouds, which are cold compared to the underlying surface, dominate patterns of
32 LWCRE, while the SWCRE is sensitive to optically thick clouds at all altitudes. Regions of deep, thick cloud
33 with large LWCRE and large negative SWCRE tend to accompany precipitation (Figure 7.6d), showing their
34 intimate connection with the hydrological cycle. The net CRE is negative over most of the globe and most
35 negative in regions of very extensive low-lying reflective stratus and stratocumulus cloud such as the
36 midlatitude and eastern subtropical oceans, where SWCRE is strong but LWCRE is weak. In these regions,
37 the spatial distribution of net CRE on seasonal timescales is strongly correlated with measures of low-level
38 stability or inversion strength (Klein and Hartmann, 1993; Williams et al., 2006; Wood and Bretherton,
39 2006; Zhang et al., 2010). However, models with comparably good simulations of this behaviour in the
40 current climate can produce a broad range of cloud responses to climate perturbations (Wyant et al., 2006).

41 42 **7.2.2 Cloud Processes**

43 44 *7.2.2.1 Modelling of Cloud Processes*

45
46 Cloud formation processes span scales from the submicron scale of cloud condensation nuclei to cloud
47 system scales of up to thousands of kilometres. This range of scales is impossible to resolve with numerical
48 simulations on computers, and is unlikely to become so for decades if ever. Nonetheless progress has been
49 made through a variety of modelling strategies.

50 51 *7.2.2.1.1 Explicit simulations in small domains*

52 High-resolution models in small domains have been widely used to simulate interactions of turbulence with
53 various types of cloud, e.g., cumulus, stratocumulus and cirrus. The grid spacing is chosen to be small
54 enough to resolve explicitly the dominant turbulent eddies that drive cloud heterogeneity, with the effects of
55 smaller-scale phenomena parameterized. This strategy is typically called large-eddy simulation (LES) when
56 boundary-layer eddies are resolved, and cloud-resolving modelling (CRM) when only deep cumulus motions
57 are well resolved. It is useful not only in simulating cloud and precipitation characteristics, but also in
58 understanding how turbulent circulations within clouds transport and process aerosols and chemical

1 constituents. Cloud microphysics, precipitation and aerosol interactions are treated with varying levels of
2 sophistication. Such models can be run in idealised settings, or with boundary conditions for specific
3 observed cases.

4
5 Such models have come to play several key scientific roles. First, they can be directly compared to in-situ
6 and high-resolution remote sensing observations (e.g., Blossey et al., 2007; Fridlind et al., 2007; Stevens et
7 al., 2005a). Second, they have revealed important influences of small-scale interactions, turbulence,
8 entrainment and precipitation on cloud dynamics that must eventually be accounted for in parameterizations
9 (e.g., Ackerman et al., 2009; Derbyshire et al., 2004; Krueger et al., 1995; Kuang and Bretherton, 2006).
10 Third, they can be used to predict how cloud ensemble properties (such as cloud cover, depth, or radiative
11 effect) may respond to climate changes (e.g., Ackerman et al., 2003; Tompkins and Craig, 1998). Fourth,
12 they have become an important tool in testing and improving parameterizations of cloud-controlling
13 processes such as cumulus convection, turbulent mixing, small-scale horizontal cloud variability, and
14 aerosol-cloud interactions (Fletcher and Bretherton, 2010; Lock, 2009; Randall et al., 2003; Rio and
15 Hourdin, 2008; Stevens and Seifert, 2008), as well as the interplay between convection and large-scale
16 circulations (Kuang, 2008).

17
18 CRMs of deep convective cloud systems with horizontal resolutions of 2 km or finer (Bryan et al., 2003) can
19 reasonably characterise observed statistical properties of the cloud ensemble, including fractional area
20 coverage of cloud (Xu et al., 2002), vertical thermodynamic structure (Blossey et al., 2007), the distribution
21 of updraughts and downdraughts (Khairoutdinov et al., 2009), and organisation into mesoscale convective
22 systems (Grabowski et al., 1998). Finer grids (down to hundreds of meters) often give significantly better or
23 different results for individual storm characteristics such as vertical velocity or tracer transport, however.
24 Some cloud ensemble properties remain sensitive to CRM microphysical parameterization assumptions
25 regardless of resolution, particularly the vertical distribution and optical depth of clouds containing ice.

26
27 It is computationally demanding to run a CRM in a domain large enough to capture convective organisation
28 or perform regional forecasts. Some studies have created smaller regions of CRM-like resolution within
29 realistically forced regional-scale models (e.g., Boutle and Abel, 2012; Zhu et al., 2010; Zhu et al., 2012), a
30 special case of the common “nesting” approach for regional downscaling (see Section 9.6). One application
31 has been to orographic precipitation, both associated with extratropical cyclones (e.g., Garvert et al., 2005)
32 and with explicitly simulated cumulus convection (e.g., Hohenegger et al., 2008); better resolution of the
33 orography improves the simulation of precipitation initiation and wind drift of falling rain and snow between
34 watersheds.

35
36 LES of shallow cumulus cloud fields with horizontal grid spacing of about 100 m and vertical grid spacing
37 of about 40 m produces vertical profiles of cloud fraction, temperature, moisture and turbulent fluxes that
38 agree well with available observations (Siebesma et al., 2003), though the simulated precipitation efficiency
39 still shows some sensitivity to microphysical parameterizations (vanZanten et al., 2011). LES of
40 stratocumulus-topped boundary layers reproduces the turbulence statistics and vertical thermodynamic
41 structure well (e.g., Ackerman et al., 2009; Stevens et al., 2005a), and has been used to study the sensitivity
42 of stratocumulus properties to aerosols (e.g., Savic-Jovicic and Stevens, 2008; Xue et al., 2008) and
43 meteorological conditions. However, the simulated entrainment rate and cloud liquid water path are sensitive
44 to the underlying numerical algorithms, even with vertical grid spacings as small as 5 m, due to poor
45 resolution of the sharp capping inversion (Stevens et al., 2005b). On the other hand, using modern high-order
46 turbulence closure schemes, the statistics of boundary-layer cloud distributions can be reasonably simulated
47 even at CRM-like horizontal resolution of 1 km or larger (Cheng and Xu, 2008; Cheng and Xu, 2006).

48 49 *7.2.2.1.2 Cloud-resolving global models*

50 Since AR4, increasing computer power has led to three types of developments in global atmospheric models.
51 First, models have been run with resolution that is higher than in the past, but not sufficiently high that
52 cumulus clouds can be resolved. Second, models have been run with resolution high enough to resolve (or
53 “permit”) large individual cumulus clouds over the entire globe. In a third approach, the parameterizations of
54 global models have been replaced by embedded CRMs. The first approach is assessed in Chapter 9. The
55 other two approaches are discussed below.

1 Global cloud-resolving models (GCRMs) have been run with grid spacings as small as 3.5 km (Putman and
2 Suarez, 2011; Tomita et al., 2005). At present GCRMs can only be used for relatively short simulations of a
3 few simulated months to a year or two on the fastest supercomputers, but in the not-too distant future they
4 may provide climate projections. GCRMs provide a consistent way to couple convective circulations to
5 large-scale dynamics, but must still parameterize the effects of individual clouds, microphysics, and
6 boundary-layer circulations.

7
8 Because they avoid many of the biases associated with uncertain aspects of cumulus parameterization,
9 GCRMs better simulate many properties of convective circulations that are very challenging for
10 conventional GCMs, including the diurnal cycles of precipitation (Sato et al., 2009) and the Asian summer
11 monsoon (Oouchi et al., 2009). Inoue et al. (2010) showed that the cloudiness simulated by NICAM is in
12 good agreement with observations from CloudSat and CALIPSO, but the results are sensitive to the
13 parameterizations of turbulence and cloud microphysics (Iga et al., 2011; Inoue et al., 2010).

14
15 Heterogeneous multiscale methods, in which cloud-resolving models are embedded in each grid cell of a
16 larger scale model (Grabowski and Smolarkiewicz, 1999), have also been further developed as a way to
17 realise some of the advantages of GCRMs but at less cost. This approach has come to be known as
18 superparameterization, because the CRM effectively replaces some of the existing GCM parameterizations
19 (e.g., Khairoutdinov and Randall, 2001; Tao et al., 2009). Super-parameterized models occupy a middle
20 ground between “process models” and “climate models” (see Figure 7.7) in terms of both advantages and
21 cost.

22
23 Like GCRMs, super-parameterized models give more realistic simulations of the diurnal cycle of
24 precipitation (Khairoutdinov et al., 2005; Pritchard and Somerville, 2010) and the MJO (Benedict and
25 Randall, 2009) than most standard GCMs; they can also improve aspects of the Asian monsoon and the El
26 Niño-Southern Oscillation (DeMott et al., 2011; Stan et al., 2010). Moreover, because they also begin to
27 resolve cloud-scale circulations, both strategies provide a framework for studying aerosol-cloud interactions
28 that standard GCMs lack (Wang et al., 2011b).

30 [INSERT FIGURE 7.7 HERE]

31 **Figure 7.7:** Model and simulation strategy for representing the climate system and climate processes at different space
32 and time scales. Models are usually defined based on the range of spatial scales they represent, shown by square
33 brackets. The temporal scales that can be represented for a given model class can vary; for instance climate models can
34 be run for a few time steps, or can simulate millennia. The figure indicates the typical timescales for which a given
35 model is used. Computational power prevents one model from covering all time and space scales. Since the AR4 the
36 development of global cloud resolving models, and hybrid approaches such as the multi-scale modelling framework,
37 have helped fill the gap between climate system and climate process models.

39 7.2.2.2 *Recent Observational Advances*

40
41 New observations, and continuing support for existing observational networks have also led to advances in
42 understanding cloud processes since AR4. Observational constraints on long-term cloudiness changes or on
43 cloud feedbacks are discussed respectively in Sections 2.3.8 and 7.2.4.

44
45 In 2006, two new coordinated, downward-pointing active sensors, the cloud profiling radar (CPR) on the
46 CloudSat satellite and the CALIOP lidar on board the CALIPSO satellite were placed in orbit. These sensors
47 have significantly improved our ability to quantify cloud amounts, water content and precipitation globally
48 (see Figures 7.4 and 7.5), and complement the detection capabilities of passive multispectral sensors (e.g.,
49 Chan and Comiso, 2011). An international network of surface-based active cloud sensors now deployed at
50 several well-instrumented sites sampling a variety of climate regimes are being systematically compared
51 with climate models (Illingworth et al., 2007) and their parameterizations (Hogan et al., 2009; Neggers et al.,
52 2012). Finally, field programs continue to give insights into cloud processes such as tropical cumulus
53 convection over land (Lebel et al., 2010), cirrus formation (May et al., 2008) and aerosol-cloud-precipitation
54 interactions associated with low-lying, liquid water clouds (Rauber et al., 2007; Wood et al., 2011b).

55
56 The net downward flux of radiation at the surface, which is a key driver of the global hydrological cycle, is
57 sensitive to the vertical distribution of clouds. Through radiation budget measurements and cloud profiling it
58 has been estimated more accurately (Kato et al., 2011). Based on these observations, the global mean surface

1 longwave flux is about 10 W m^{-2} larger than the average in climate models, likely due to insufficient model-
2 simulated cloud cover or lower tropospheric moisture (Stephens et al., 2012).

3
4 Satellite cloud-observing capacities are reviewed by Stubenrauch et al. (2012). New sensors show more
5 clearly that low clouds are prevalent in nearly all types of convective system, to an extent not always
6 captured in models (Chepfer et al., 2008; Haynes et al., 2011; Naud et al., 2010). Cloud layers at different
7 levels overlap less often than often assumed in GCMs, especially over high-latitude continents (Mace et al.,
8 2009; Naud et al., 2008), and the common assumption that the radiative effects of precipitating ice can be
9 neglected is not necessarily warranted (Waliser et al., 2011). New observations have led to revised
10 treatments of overlap in some models, which significantly affects cloud radiative effects (Pincus et al.,
11 2006). Active sensors have also been useful in detecting low-lying Arctic clouds over sea ice (Kay et al.,
12 2008), improving our ability to test climate model simulations of the interaction between sea ice loss and
13 cloud cover (Kay et al., 2011).

14
15 Recent studies have confirmed that aircraft observations of unexpectedly numerous small ice crystals are
16 largely an artefact of crystal shattering (Korolev et al., 2011), helping to reconcile in-situ and satellite
17 observations. Satellite observations of ice water content have enabled model microphysical schemes to be
18 tested globally, revealing for example that the rate of conversion of cloud ice to precipitation varies
19 geographically in ways not consistent with current assumptions (Ma et al., 2012a) and that liquid water
20 clouds in CRMs generally begin to rain too early (Suzuki et al., 2011). Global datasets of low-lying liquid
21 cloud droplet number concentration based on passive observations at multiple wavelengths are being
22 improved (Bennartz, 2007; Quaas et al., 2006) and are gaining popularity as a metric for climate model
23 simulation of aerosol-cloud interactions (Wang et al., 2011b).

24 25 **7.2.3 Parameterization of Clouds in Climate Models**

26 27 *7.2.3.1 Challenges of Parameterization*

28
29 Cloud droplets or ice crystals form from vapour, evolve, collide, may grow to form precipitation, and
30 ultimately evaporate or fall out of the atmosphere. The representation of these processes in climate models is
31 particularly challenging, in part because some of the fundamental details of these microphysical processes
32 are poorly understood (particularly for ice- and mixed-phase clouds), and because the spatial heterogeneity
33 of the clouds and their turbulent properties occurs at scales significantly smaller than a GCM grid box.

34
35 Most CMIP5 climate model simulations use horizontal resolutions of 100–200 km in the atmosphere, with
36 vertical layers varying between 100 m near the surface to more than 1000 m aloft. Within regions of this size
37 in the real world, there is usually enormous subgrid-scale variability in cloud properties, associated with
38 variability in humidity, temperature and vertical motion (Figure 7.16). This variability must be accounted for
39 to accurately simulate cloud-radiation interaction, condensation, evaporation and precipitation, and other
40 cloud processes that crucially depend on how cloud condensate is distributed across each grid box (Cahalan
41 et al., 1994; Larson et al., 2001; Pincus and Klein, 2000).

42
43 The simulation of clouds in modern climate models involves several parameterizations that must closely
44 work together as a system. These include parameterization of turbulence and cumulus convection, cloud
45 fraction and microphysical processes (including their vertical overlap between different grid levels), and
46 aerosol and chemical transport. The system of parameterizations must balance simplicity, realism,
47 computational stability and efficiency. Each parameterization makes simplifying mathematical assumptions
48 about the subgrid variability within each grid cell; these assumptions vary significantly from model to model
49 and for pragmatic and historical reasons may not even be fully consistent across the parameterizations used
50 in one model. For example, clouds in a grid column may be assumed to be vertically stacked for the radiation
51 calculation, but not for calculating evaporation of precipitation. In summary, realistic simulation of clouds
52 and their response to climate change forms one of the greatest challenges of climate modelling.

53
54 Cloud process parameterization is important for specialized chemical-aerosol-climate models (see review by
55 Zhang, 2008) and for regional climate models as well as for CMIP5-class global models. A few of the former
56 models have added complexity in representing subgrid cloud variability and the cloud particle size
57 distribution (e.g., Jacobson, 2003).

7.2.3.2 *Recent Advances in Representing Cloud Microphysical Processes*

7.2.3.2.1 *Liquid clouds*

Recent development efforts have been focused on the introduction of more complex representations of microphysical processes, with the dual goals of coupling them better to atmospheric aerosols and linking them more consistently to the sub-grid variability assumed by the model for other calculations. For example, most AR4-era climate models used a single moment formulation for stratiform cloud water, predicting only the average cloud and rain water mass in each grid cell at a given time, diagnosing the droplet concentration using empirical relationships based on aerosol mass (e.g., Boucher and Lohmann, 1995; Menon et al., 2002), or altitude and proximity to land. Many were forced to employ an arbitrary lower bound on droplet concentration to reduce the aerosol radiative forcing (Hoose et al., 2009). Such formulations oversimplify microphysically-mediated cloud variations.

By contrast, more models participating in CMIP5 use two-moment schemes for liquid stratiform cloud. Some diagnose rain and snow number concentrations and mixing ratios (Morrison and Gettelman, 2008; Salzmann et al., 2010), allowing treatment of aerosol scavenging and the radiative effect of snow. Some models explicitly treat subgrid cloud water variability for calculating microphysical process rates (Morrison and Gettelman, 2008). Cloud droplet activation schemes now account more realistically for subadiabaticity, particle composition, and particle size (Abdul-Razzak and Ghan, 2000; Ghan et al., 2011; Liu et al., 2012). Despite such advances in internal consistency, a continuing weakness in traditional GCMs (and even in GCRMs and superparameterized models, albeit to a much lesser extent) is their inability to (fully) resolve cloud-scale vertical motions, to which microphysical processes are highly sensitive.

7.2.3.2.2 *Mixed-phase and ice clouds*

Ice treatments are following a similar path to those for liquid water, and face similar challenges but amplified by the greater complexity of ice processes. Many AR4 models predicted the condensed water amount in just two categories – cloud and precipitation – with a temperature-dependent partitioning between liquid and ice within either category. Although supersaturation with respect to ice is commonly observed at low temperatures, only one CMIP3 GCM (ECHAM) allowed ice supersaturation (Lohmann and Kärcher, 2002). Many climate models now include separate, physically-based equations for cloud liquid versus cloud ice, and for rain versus snow, allowing a more realistic treatment of mixed-phase processes and ice supersaturation (Gettelman et al., 2010; Liu et al., 2007; Salzmann et al., 2010; Tompkins et al., 2007; see also Section 7.4.4). These new schemes are tested in a single-column model against cases observed in field campaigns (e.g., Klein et al., 2009) or against satellite observations (e.g., Kay et al., 2012), and provide superior simulations of cloud structure than typical AR4-class parameterizations (Kay et al., 2012). New representations of the Wegener-Bergeron-Findeisen process in mixed-phase clouds (Lohmann and Hoose, 2009; Storelvmo et al., 2008b) compare the rate at which the pre-existing ice crystals deplete the water vapour (Korolev, 2007) with the condensation rate for liquid water driven by vertical updraught speed; these are not yet included in CMIP5 models. Climate models are increasingly representing detailed microphysics, including mixed-phase processes, inside convective clouds (Fowler and Randall, 2002; Lohmann, 2008; Song and Zhang, 2011). Such processes can influence storm characteristics like strength and electrification. We reiterate that microphysical detail may not translate to improved simulation if the dynamics is not commensurately well resolved.

7.2.3.3 *Recent Advances in Parameterizing Moist Turbulence and Convection*

Both the mean state and variability in climate models are sensitive to the parameterization of cumulus convection. Since AR4, the development of convective parameterization has been driven largely by rapidly growing use of process models, in particular LES and CRMs, to inform parameterization development (see Hourdin et al., 2012).

Accounting for greater or more state-dependent entrainment of air into deep cumulus updraughts has improved simulations of the Madden-Julian Oscillation (MJO), tropical convectively-coupled waves, and mean rainfall patterns in some models (Bechtold et al., 2008; Chikira and Sugiyama, 2010; Del Genio et al., 2012; Del Genio et al., 2007; Hohenegger and Bretherton, 2011; Kim et al., 2012; Mapes and Neale, 2011; Song and Zhang, 2009) but usually at the expense of other aspects of the simulation. Improved activation

1 criteria and parameterizations of cumulus momentum fluxes in another model improved ENSO and tropical
2 vertical temperature profiles (Neale et al., 2008; Richter and Rasch, 2008). Since AR4, more climate models
3 have adopted cumulus parameterizations that diagnose the expected vertical velocity in cumulus updrafts
4 (e.g., Chikira and Sugiyama, 2010; Donner et al., 2011; Park and Bretherton, 2009), in principle allowing
5 more complete representations of aerosol activation, cloud microphysical evolution, and gravity wave
6 generation by the convection.

7
8 Several global models now couple shallow cumulus convection more closely to moist boundary layer
9 turbulence (Couvreur et al., 2010; Neggers, 2009; Neggers et al., 2009; Siebesma et al., 2007) including cold
10 pools generated by nearby deep convection (Grandpeix and Lafore, 2010). Many of these efforts have led to
11 more accurate simulations of boundary-layer cloud radiative properties and vertical structure (e.g., Köhler et
12 al., 2011; Park and Bretherton, 2009), and have ameliorated the common problem of premature deep
13 convective initiation over land in one CMIP5 GCM (Rio et al., 2009).

14 7.2.3.4 *Recent Advances in Parameterizing Cloud Radiative Effects*

15
16 Some models have improved representation of subgrid cloud variability, which has important effects on grid-
17 mean radiative fluxes and precipitation fluxes, for example based on the use of probability density functions
18 of thermodynamic variables (Watanabe et al., 2009). Stochastic approaches for radiative transfer can account
19 for this variability in a computationally efficient way (Barker et al., 2008). New treatments of cloud overlap
20 have been motivated by new observations (Section 7.2.2.2).

21 7.2.4 *Cloud, Water-Vapour and Lapse Rate Feedbacks*

22
23 Climate feedbacks are a central concern for projecting the magnitude of climate change, because they
24 determine the sensitivity of global surface temperature to external forcing agents (see Chapter 9). Water
25 vapour, lapse rate and cloud feedbacks each involve moist atmospheric processes closely linked to clouds,
26 and in combination, produce most of the simulated climate feedback and most of its inter-model spread. The
27 strength of a radiative feedback can be expressed as its impact on the top-of-atmosphere net downward
28 radiative flux per degree of global surface temperature increase (or feedback parameter) and may be
29 compared with the basic “black-body” response of $-3.4 \text{ W m}^{-2} \text{ K}^{-1}$.

30 7.2.4.1 *Water-Vapour Response and Feedback*

31
32 As pointed out in previous reports (Randall et al., 2007, Section 8.6.3.1), physical arguments and models of
33 all types suggest global water vapour amounts increase in a warmer climate, leading to a positive feedback
34 via its enhanced greenhouse effect. The saturated water vapour mixing ratio (WVMR) increases nearly
35 exponentially and very rapidly with temperature, at 6–10% per degree near the surface, and even more
36 steeply aloft (up to 17% per degree) where air is colder. Mounting evidence continues to indicate that
37 changes in relative humidity in warmer climates will be nowhere near this rapid, at least in a global and
38 statistical sense, although new work highlights some robust changes that vary by region. Hence the WVMR
39 is expected to increase, on average, at a rate similar to the saturated WVMR.

40
41 Because global temperatures have been rising, the above arguments imply WVMR should be rising
42 accordingly, and multiple observing systems indeed show this (Section 2.3.6). One exception is that
43 meteorological station data suggest a plateauing of WVMR near the land surface over the last decade or so,
44 but humidity at this level exerts little greenhouse effect (Soden and Held, 2006) and is governed by
45 mechanisms different from those operating at upper levels (Joshi et al., 2008). A study challenging the water
46 vapour increase (Paltridge et al., 2009) used an older reanalysis product, whose trends are contradicted by
47 newer ones (Dessler and Davis, 2010) and by direct global observations which are considered more reliable
48 for trends (see Box 2.3). The study also reported decreasing relative humidity in data from Australian
49 radiosondes, but more complete studies show that region to be anomalous in this respect (Dai et al., 2011).
50 Thus reliable, large-scale, multi-decade trend data remain consistent with the expected global feedback.

51
52 Some studies have proposed that the response of upper-level humidity to natural fluctuations in the global
53 mean surface temperature is informative about the feedback. However, small changes to the global mean
54 (primarily from ENSO) involve geographically heterogeneous temperature change patterns, the responses to

1 which may be a poor analogue for global warming (Hurley and Galewsky, 2010b). Most climate models
2 reproduce these natural responses reasonably well (Dessler and Wong, 2009; Gettelman and Fu, 2008),
3 providing additional evidence that they at least represent the key processes.
4

5 The “last-saturation” concept approximates the WVMR of air by its saturation value when it was last in a
6 cloud (see Sherwood et al., 2010b for a review), which can be inferred from trajectory analysis. Studies since
7 the AR4 using a variety of models and observations (including concentrations of isotopically substituted
8 water vapour molecules) support this concept (Galewsky and Hurley, 2010; Sherwood and Meyer, 2006).
9 The concept has clarified what determines relative humidity in the subtropical upper troposphere and placed
10 the water-vapour feedback on firmer theoretical footing by directly linking actual and saturation WVMR
11 values (Hurley and Galewsky, 2010a).
12

13 In a warmer climate, an upward shift of the tropopause and poleward shift of the jets and associated climate
14 zones is expected (Sections 2.6.4 and 2.6.5) and simulated by most GCMs (Section 10.3.3.1). These changes
15 account, at least qualitatively, for robust regional changes in the relative humidity simulated in warmer
16 climate by GCMs, including decreases in the subtropical troposphere and tropical uppermost troposphere,
17 and increases near the extratropical tropopause and high latitudes (Sherwood et al., 2010a). This pattern may
18 be amplified however by non-uniform atmospheric temperature or wind changes (Hurley and Galewsky,
19 2010b). It is also the likely cause of most model-predicted changes in mid- and upper-level cloudiness
20 patterns (Sherwood et al., 2010a; Wetherald and Manabe, 1980; see also Section 7.2.4.3.3). Idealised CRM
21 simulations of warming climates also show upward shifts, with otherwise little change in mean relative
22 humidity (e.g., Kuang and Hartmann, 2007; Romps, 2011).
23

24 It remains unclear whether stratospheric water vapour contributes significantly to climate feedback.
25 Observations have shown decadal variations in stratospheric water vapour, which may have affected the
26 planetary radiation budget somewhat (Solomon et al., 2010) but do not have a clear relation to global
27 warming (Trenberth et al., 2007, Section 3.4.2.4). GCMs can simulate strong positive feedbacks from
28 stratospheric water vapour (Joshi et al., 2010), but with parameter settings producing unrealistic behaviour in
29 the present climate.
30

31 *7.2.4.2 Relationship Between Water Vapour and Lapse-Rate Feedbacks* 32

33 Lapse rates in the tropics should change roughly as predicted by a moist adiabat, due to the strong restoring
34 influence of convective heating. This restoring influence has now been directly inferred from satellite data
35 (Lebsock et al., 2010), and the near-constancy of tropical atmospheric stability and deep-convective
36 thresholds over recent decades is also now observable in sea surface temperature (SST) and deep convective
37 data (Johnson and Xie, 2010). The stronger warming of the atmosphere relative to the surface produces a
38 negative feedback on global temperature because the warmed system radiates more thermal emission to
39 space for a given increase in surface temperature than in the reference case where the lapse rate is fixed. This
40 feedback is not equally strong in all models, because lapse rates in middle and high latitudes change
41 differently (Dessler and Wong, 2009).
42

43 **[INSERT FIGURE 7.8 HERE]**

44 **Figure 7.8:** Clear-sky feedback parameters as predicted by [CMIP3] GCMs. Black points in the centre show the total
45 radiative response including the Planck response, with the Planck response and individual feedbacks from water vapour
46 and lapse rate shown to the right in red. On the left are the equivalent three parameters calculated in an alternative,
47 relative humidity-based framework. In this framework the Planck stabilization and each of the two feedbacks are all
48 weaker and more consistent among the models. [PLACEHOLDER FOR FINAL DRAFT: Current working draft of
49 figure taken from Held and Shell (2012); additional data from CMIP5 TBA].
50

51 As shown by Cess (1975) and discussed in the AR4 (Randall et al., 2007), models with a more negative
52 lapse-rate feedback tend to have a more positive water-vapour feedback. Cancellation between these is close
53 enough that their sum, or “clear-sky feedback,” has a 90% range in CMIP3 models of only +0.96 to +1.22 W
54 m⁻² K⁻¹ (based on a Gaussian fit to the data of Held and Shell (2012), see Figure 7.8). The physical reason
55 for this cancellation is that as long as water vapour infrared absorption bands are nearly saturated, outgoing
56 longwave radiation is determined by relative humidity (Ingram, 2010) which exhibits little systematic change
57 in any model (Section 7.2.4.1). In fact, Held and Shell (2012) and Ingram (2012a) argue that it makes more
58 sense physically to consider a reference climate perturbation in which relative humidity is held fixed, rather

1 than specific humidity as per the usual convention. In that framework, the climate is fundamentally less
2 stable but the water-vapour and lapse-rate feedbacks are each small; the large and partially compensating
3 conventional feedbacks are seen to have arisen from unphysical choices made in first setting out the
4 traditional framework.

5
6 There is some observational evidence suggesting significant departures from the adiabatic lapse rate in the
7 tropical upper troposphere (Section 2.2.4), which would be larger than simulated by models (Section 9.4.1).
8 Since the combined clear-sky feedback depends on relative humidity change, however, the imputed lapse-
9 rate variations would have little influence on the total feedback and might even slightly increase it (Ingram,
10 2012b). In summary, all sound evidence continues to support a strong, positive feedback (measured in the
11 traditional framework) from the combination of water-vapour and lapse-rate changes.

12 13 *7.2.4.3 Cloud Feedbacks*

14
15 The dominant source of spread among GCM climate sensitivities in AR4 was due to diverging cloud
16 feedbacks, particularly due to low clouds, and this continues to be true (Chapter 9). The net cloud feedback
17 also continues to be nearly zero or positive in all global models. Progress has been made since the AR4 in
18 understanding which simulated cloud changes underlie the positive feedbacks, and providing a stronger
19 theoretical and observational basis for key positive cloud feedbacks. There has also been progress in
20 quantifying cloud feedbacks, including separating the effects of different cloud types, using radiative-kernel
21 residual methods (Soden et al., 2008) and satellite simulators (see Section 9.2.2.3) (Zelinka et al., 2012a).

22
23 Until very recently cloud feedbacks have been diagnosed in models by differencing cloud radiative effects in
24 doubled CO₂ and control climates, normalized by the change in global mean surface temperature. Different
25 diagnosis methods do not always agree, and some simple methods can make positive cloud feedbacks look
26 negative (Soden and Held, 2006). Moreover, it is now recognised that some of the cloud changes are induced
27 directly by the atmospheric radiative effects of CO₂ independently of surface warming, and are therefore
28 rapid adjustments rather than feedbacks. Most of the published studies available for this assessment did not
29 separate these effects, but we do so where possible. It appears that these adjustments are sufficiently small in
30 most models (see Section 7.2.4.3.6) that general conclusions regarding feedbacks are not significantly
31 affected.

32
33 Four aspects of the global cloud response to greenhouse gas-induced climate change are distinguished here:
34 altitude increases of high-level clouds, effects of hydrological cycle and storm track changes on middle and
35 high clouds, changes in low-level cloud cover, and changes in high-latitude clouds. Finally, recent research
36 on the rapid cloud adjustments to CO₂ is assessed. Cloud changes cause both longwave (greenhouse
37 warming) and shortwave (reflective cooling) effects, which combine to give the overall cloud feedback or
38 forcing adjustment for global climate.

39 40 *7.2.4.3.1 Cloud altitude feedback mechanisms involving high-level clouds*

41 High clouds at low and middle latitudes exert little net top-of-atmosphere radiative effect in the current
42 climate due to near-compensation between their longwave and shortwave cloud radiative effects (Kiehl,
43 1994). Nonetheless, systematic changes in their height or optical thickness could produce a significant
44 radiative feedback by altering this balance.

45
46 The dominant driver of net longwave cloud feedback in models, and a dominant contributor to net positive
47 cloud feedback overall, appears to be a robust consequence of global warming: an increase in the height of
48 the tropopause and the main level at which the deepest convective clouds stop rising and cloudy air flows
49 outward, tentatively attributed in AR4 to the so-called “fixed anvil-temperature” mechanism (Hartmann and
50 Larson, 2002). According to this mechanism, the outflow level from tropical deep convective systems is
51 determined ultimately by the highest point at which water vapour emits significant infrared radiation; this
52 point is governed by the water vapour partial pressure, therefore occurring at a similar temperature (higher
53 altitude) in a warmer climate. A positive cloud altitude feedback results because the temperature difference
54 between the cloud and the surface increases, increasing the cloud’s greenhouse effect without necessarily
55 affecting its albedo. This occurs at all latitudes and has long been noted in model simulations (Cess et al.,
56 1990; Hansen et al., 1984). New studies confirm that this mechanism operates in GCMs, with a small
57 modification to account for lapse-rate changes, and accounts for most of the total longwave cloud feedback

1 in CMIP3 GCMs (Soden and Vecchi, 2011; Zelinka and Hartmann, 2010; Zelinka et al., 2012a). In some
2 GCMs the overall contributions from high cloud and from cloud height changes can be isolated (Figure 7.9)
3 with net (LW+SW) estimates for the feedback parameter of +0.14 (0.0 to +0.30) $\text{W m}^{-2} \text{K}^{-1}$ and +0.34 (+0.05
4 to +0.62) $\text{W m}^{-2} \text{K}^{-1}$, respectively. The increase in cloud height also occurs consistently in cloud-resolving
5 models (Kuang and Hartmann, 2007; Kubar et al., 2007).

6 7 **[INSERT FIGURE 7.9 HERE]**

8 **Figure 7.9:** Cloud feedback parameters as predicted by GCMs. Total feedback shown at left by black symbols, broken
9 out into infrared and visible components in red and blue, respectively (Zelinka et al., 2012a; Zelinka et al., 2012b).
10 Centre panel shows components attributable to clouds in different height ranges (see text); values reported by Soden
11 and Vecchi (2011) do not conform exactly to this definition but are shown for comparison, with their “mixed” category
12 assigned “medium”. Right panel shows components attributable to different cloud property changes (not available from
13 all studies).

14
15 The observational record allows us to verify various elements of the above mechanism. First, the global
16 tropopause is rising as expected (Chapter 2). Second, cloud heights change as predicted with regional,
17 seasonal and interannual changes in near-tropopause temperature structure (Eitzen et al., 2009; Xu et al.,
18 2007; Zelinka and Hartmann, 2010), although the response may be affected by changes in stratospheric
19 circulation (Chae and Sherwood, 2010; Eitzen et al., 2009). Third, Norris et al. (2012) report that two
20 satellite datasets show a rising trend in high clouds since 1979 roughly consistent with the mechanism.
21 Davies and Molloy (2012) report a downward height trend among all clouds in a shorter record from a
22 satellite that measures heights more directly, but this is likely an artefact of instrument problems early in that
23 record (Evan and Norris, 2012).

24 25 *7.2.4.3.2 Feedback mechanisms involving the amount of middle and high cloud*

26 Nearly all GCMs simulate a reduced middle and high cloud amount in warmer climates in low- and mid-
27 latitudes, especially in the subtropics (Trenberth and Fasullo, 2009; Zelinka and Hartmann, 2010). In global
28 average, these cloudiness reductions cause a positive shortwave and negative longwave feedback; the latter
29 almost completely cancels the altitude response (Figure 7.9), which may explain why the important role of
30 cloud altitude feedbacks was not fully appreciated until recently. The net effect of changes in amount of all
31 cloud types is a positive feedback, but this comes mainly from the changes low clouds (see the following
32 section), implying a near-cancellation of LW and SW effects for the mid- and high-level amount changes.

33
34 Changes in predicted cloud cover are geographically correlated with simulated subtropical drying (Meehl et al.,
35 2007), suggesting that they are partly tied to large-scale circulation changes including the poleward shifts
36 found in most models (Sherwood et al., 2010a; Wetherald and Manabe, 1980). Bender et al. (2012) and
37 Norris et al. (2012) also report latitudinal variations in observed cloudiness trends since 1979 consistent with
38 the poleward shift of the circulation indicated in several other observables (see Chapter 2) and simulated
39 (albeit with weaker amplitude) by most GCMs (Yin, 2005). This shifts clouds to latitudes of weaker sunlight,
40 decreasing the planetary albedo. The impact of the observed shift appears to be significant and would imply
41 a strong positive feedback if it were due to global warming (Bender et al., 2012). Recent studies call into
42 question how much of the observed shifts are temperature- or ozone-driven however (Chapter 10.3), so the
43 true amount of positive feedback coming from poleward shifts remains highly uncertain but could be
44 underestimated by GCMs.

45
46 Tselioudis and Rossow (2006) predict reduced cloud cover in the storm-track itself in warmer climates based
47 on observed present-day relationships with meteorological variables combined with model-simulated
48 changes to those driving variables. In agreement with the above analysis, they found that this reduction
49 contributed little to net cloud feedback. The upward mass flux in deep clouds should decrease in a warmer
50 climate (Section 7.6) which might contribute to cloudiness decreases in convective regions (e.g., the ITCZ).
51 Most CMIP3 GCMs produce too little storm-track cloud in the southern hemisphere compared to nearly-
52 overcast conditions in reality. This bias could also imply that positive feedback from poleward shifts is
53 greater than in GCMs, or that the negative feedback many GCMs simulate in these regions from increasing
54 cloud amount may be unrealistic (Trenberth and Fasullo, 2010).

55
56 Changing coverage of thin cirrus clouds could in principle exert a significant feedback due to the net
57 warming effect of these clouds (e.g., Rondanelli and Lindzen, 2010), but no mechanistic argument for a

1 particular change one way or the other has been advanced, and responses in GCMs so far are small (Zelinka
2 et al., 2012b).

3 4 *7.2.4.3.3 Feedback mechanisms involving low clouds*

5 Differences in the response of low clouds to a warming are responsible for most of the spread in model-
6 based estimates of equilibrium climate sensitivity (Bony and Dufresne, 2005). Since the AR4 this finding has
7 withstood further scrutiny (e.g., Soden and Vecchi, 2011; Webb et al., 2012), holds in CMIP5 models (Vial
8 et al., 2012), and has been shown to apply also to the transient climate response (e.g., Dufresne and Bony,
9 2008). This ‘low-cloud problem’ is most potent over tropical and sub-tropical oceans (Williams and
10 Tselioudis, 2007; Williams and Webb, 2009; Xu et al., 2010), although not exclusively so (Trenberth and
11 Fasullo, 2010), and is usually associated with the representation of shallow cumulus or stratocumulus clouds
12 (Williams and Tselioudis, 2007; Williams and Webb, 2009; Xu et al., 2010). Because the low-cloud problem
13 emerges in a variety of idealized model formulations (Medeiros et al., 2008; Zhang and Bretherton, 2008), or
14 conditioned on a particular dynamical state (Bony et al., 2004), it appears to be attributable to how cloud,
15 convective and boundary layer processes are parameterized in GCMs.

16
17 The modelled response of low clouds does not appear to be dominated by a single feedback mechanism, but
18 rather the net effect of several potentially competing mechanisms as elucidated in LES and GCM sensitivity
19 studies (e.g., Bretherton et al., 2012; Zhang et al., 2012a; Zhang and Bretherton, 2008). Starting with some
20 proposed negative feedback mechanisms, it has been argued that in a warmer climate, low clouds will be: (i)
21 horizontally more extensive, because changes in the lapse rate of temperature also modify the lower-
22 tropospheric stability (Miller, 1997); (ii) optically thicker, because adiabatic ascent is accompanied by a
23 larger condensation rate (Somerville and Remer, 1984); and (iii) vertically more extensive, in response to a
24 weakening of the tropical overturning circulation (Caldwell and Bretherton, 2009). While these mechanisms
25 may play some role in subtropical low cloud feedbacks, none of them appears dominant. Regarding the first
26 mechanism, model fidelity to relationships observed to hold for the present climate fails to constrain the
27 response of models to climate change (Section 7.2.1.2; see also Webb and Lock, 2012). The second
28 mechanism, discussed briefly in the next section, appears to be a small effect. The third mechanism cannot
29 yet be ruled out, but does not appear to be the dominant factor in determining subtropical cloud changes in
30 GCMs (Bony and Dufresne, 2005; Zhang and Bretherton, 2008).

31
32 Since the AR4, several new positive feedback mechanisms have been proposed, most associated with the
33 marine boundary layer clouds thought to be at the core of the ‘cloud problem.’ These include the ideas that:
34 warming-induced changes in the absolute humidity lapse rate change the energetics of mixing in ways that
35 demand a reduction in cloud amount or thickness (Bretherton et al., 2012; Brient and Bony, 2012; Lock,
36 2009); energetic constraints prevent the surface evaporation from increasing with warming at a rate sufficient
37 to balance expected changes in dry air entrainment thereby reducing the supply of moisture to form clouds
38 (Rieck et al., 2012; Webb and Lock, 2012); and that increased concentrations of greenhouse gases reduce the
39 radiative cooling that drives stratiform cloud layers and thereby the cloud amount (Bretherton et al., 2012;
40 Caldwell and Bretherton, 2009; Stevens and Brenguier, 2009). These mechanisms could explain why models
41 consistently produce positive low-cloud feedbacks. Among CFMIP GCMs, the feedback parameter ranges
42 from +0.09 to +0.63 W m⁻² K⁻¹, and is largely associated with a reduction in low-cloud amount, albeit with
43 considerable variability in the spatial structure of these changes (Webb et al., 2012). One
44 “superparameterized” GCM (Section 7.2.2.1) simulates a negative low-cloud feedback (Wyant et al., 2006;
45 Wyant et al., 2009), but that model’s representation of low clouds was worse than some conventional GCMs.

46
47 The tendency of both GCMs and process models to produce these positive feedback effects suggests that the
48 feedback contribution from changes in low clouds is positive. However, the well-known deficiencies in
49 model representation of low clouds globally, the diversity of model results, the lack of reliable observational
50 constraints, and the tentative nature of the suggested mechanisms leaves us with low overall confidence in
51 the sign of the low-cloud feedback contribution.

52 53 *7.2.4.3.4 Feedbacks involving changes in cloud optical depth*

54 It has long been suggested that cloud water content could increase in a warmer climate simply due to the
55 availability of more vapour for condensation in a warmer atmosphere, yielding a negative feedback
56 (Paltridge, 1980; Somerville and Remer, 1984), but this argument ignores the physics of crucial cloud-

1 regulating processes like precipitation formation and turbulence. Observational evidence discounting a large
2 effect of this kind was reported in AR4 (Randall et al., 2007).

3
4 The global-mean net feedback from cloud optical depth changes in CFMIP models (Figure 7.9) scatters
5 around zero but is slightly positive on average. Optical depths tend to reduce slightly at low and middle
6 latitudes, but increase consistently at latitudes poleward of 50°, yielding a positive longwave feedback that
7 slightly outweighs the negative shortwave feedback. These latitude-dependent optical depth changes may be
8 attributed to phase changes at high latitudes and greater poleward moisture transport (Vavrus et al., 2009),
9 and possible to poleward shifts of the circulation.

10
11 The phase change contribution arises because at mixed-phase temperatures of -40°C – 0°C , cloud ice particles
12 are typically several-fold larger than cloud water drops (e.g., Mitchell et al., 2010), so a given mass of cloud
13 ice reflects less sunlight than the same mass of cloud water droplets. As climate warms, the shift from ice to
14 liquid clouds would raise albedos for a given water mass, and the slower fall speeds of liquid could increase
15 the water mass. The resulting negative cloud feedback appears in GCMs (Senior and Mitchell, 1993), with a
16 magnitude that correlates with the simulated amount of cloud ice in mixed-phase clouds (Tsushima et al.,
17 2006). The key physics is however not adequately represented in climate models, so this particular feedback
18 mechanism is highly uncertain.

19 20 *7.2.4.3.5 Feedback from Arctic cloud interactions with sea ice*

21 Arctic clouds, despite their low altitude, have a net warming effect at the surface in the present climate
22 because their downward emission of infrared radiation over the year outweighs their reflection of sunlight
23 during the short summer season. They also cool the atmosphere, however, so their effect on the energy
24 balance of the whole system is ambiguous and depends on the details of the vertical cloud distribution and
25 the impact of cloud radiative interactions on ice cover (Palm et al., 2010).

26
27 Low cloud amount over the Arctic oceans varies inversely with sea ice amount (open water producing more
28 cloud) as now confirmed since AR4 by visual cloud reports (Eastman and Warren, 2010) and lidar and radar
29 observations (Kay and Gettelman, 2009; Palm et al., 2010). The observed effect is weak in boreal summer,
30 when the melting sea-ice is at a similar temperature to open water and stable boundary layers with extensive
31 low cloud are common over both surfaces, and strongest in boreal autumn when cold air flowing over
32 regions of open water stimulates cloud formation by boundary-layer convection (Kay and Gettelman, 2009;
33 Vavrus et al., 2011). Kay et al. (2011) show that a GCM can represent this seasonal sensitivity of low cloud
34 to open water, but doing so depends on the details of how boundary-layer clouds are parameterized. Vavrus
35 et al. (2009) show that in a global warming scenario, GCMs simulate more Arctic low-level cloud in all
36 seasons, but especially during autumn and winter when open water and very thin sea ice increase
37 considerably, increasing upward moisture transport to the clouds.

38
39 A negative Arctic cloud feedback was suggested by Liu et al. (2008) on the basis that recent observed
40 surface warming has been greater in less cloudy areas, but this argument was not tested in a climate model
41 and does not control for the large correlated effects of weather on both clouds and surface temperature.
42 Gagen et al. (2011) present tree-ring evidence that summertime Arctic cloud cover was negatively correlated
43 with Arctic temperatures over the last millennium, which is consistent with the conclusions of the above
44 studies assuming there was less ice during warmer periods. Gagen et al. (2011) presented this as evidence of
45 a negative shortwave cloud feedback, but the year-round longwave feedback would likely offset this as noted
46 above. Note that there are pitfalls to using natural climate variations to infer cloud feedbacks in any case (see
47 Section 7.2.4.3.7).

48 49 *7.2.4.3.6 Rapid adjustment of clouds to a CO₂ change*

50 Gregory and Webb (2008) partitioned the response of top-of-atmosphere radiation in GCMs to an
51 instantaneous doubling of CO₂ into a ‘rapid’ (sub-seasonal) adjustment in which the land surface,
52 atmospheric circulations and clouds respond to the radiative effect of the CO₂ increase, and an ‘SST-
53 mediated’ response that develops more slowly as the oceans warm. This distinction is important not only to
54 help understand what is going on in the models, but because the presence rapid adjustments would cause
55 clouds to respond slightly differently to a transient climate change (in which SST changes have not caught
56 up to CO₂ changes) or to a climate change caused by other forcings, than they would to the same warming at

1 equilibrium driven by CO₂. There is also a rapid adjustment of the precipitation field, discussed in Section
2 7.6.3.

3
4 Gregory and Webb (2008) found that in some climate models, rapid adjustment of clouds can have
5 comparable top-of-atmosphere radiative effects to the ensuing SST-mediated cloud changes. Andrews and
6 Forster (2008) found that this behaviour was exceptional, and that on average, rapid cloud adjustments in a
7 suite of climate models causes less than 20% of their equilibrium radiative feedback; most of these come
8 from low cloud (Colman and McAvaney, 2011). Webb et al. (2012) and Andrews et al. (2012), however,
9 each showed using different model ensembles that adjustments contribute half the uncertainty that feedbacks
10 do, for example 2.2 K uncertainty from feedbacks as compared to 1.08 K from adjustments in the latter
11 study. Thus a better understanding of rapid adjustments could narrow the uncertainty in climate projections.

12 7.2.4.3.7 *Observational constraints on global cloud feedback*

13 A number of studies since AR4 have attempted to constrain cloud feedback (or total climate sensitivity) from
14 observations; here we discuss those using modern cloud, radiation or other measurements. Section 12.5
15 discusses those based on past temperature data and forcing proxies.

16
17 One approach is to seek observable aspects of present-day cloud behaviour that reveal cloud feedback.
18 Varying parameters in a GCM sometimes produces changes in cloud feedback that correlate with the
19 properties of cloud simulated for the present day, but this depends on the GCM (Yokohata et al., 2010), and
20 the resulting relationships do not hold across multiple models such as those from CMIP3 (Collins et al.,
21 2011; Gettelman et al., 2012a). Among the AR4 models, net cloud feedback is strongly correlated with mid-
22 latitude relative humidity (Volodin, 2008) and with characteristics of the southern-hemisphere storm track
23 (Trenberth and Fasullo, 2010); if valid either regression relation would imply a relatively strong positive
24 cloud feedback in reality, but no mechanism has been proposed to explain or validate these empirical
25 relationships and such apparent skill can easily arise fortuitously (Klocke et al., 2011). Likewise, Clement et
26 al. (2009) found realistic decadal variations of low cloud over the North Pacific in only one model
27 (HadGEM1) and argued that the relatively strong cloud feedback in this model should therefore be regarded
28 as more likely, but provided no evidence for such a link. Chang and Coakley (2007) examined midlatitude
29 maritime clouds and found cloud thinning with increasing temperature, consistent with a positive feedback,
30 while Gordon and Norris (2010) found the opposite result following a methodology that tried to isolate
31 thermal and advective effects. In summary, there is no evidence of a robust link between any of the noted
32 observables and the global feedback, though some apparent connections are tantalising and are being further
33 studied.
34

35
36 Several studies have attempted to derive global climate sensitivity from interannual relationships between
37 global-mean observations of top-of-atmosphere radiation and surface temperature. One problem with this is
38 the different spatial character of interannual and long-term warming; another is that the methodology can be
39 confounded by cloud variations not caused by those of surface temperature (Spencer and Braswell, 2008). A
40 range of climate sensitivities has been inferred based on such analyses (Forster and Gregory, 2006; Lindzen
41 and Choi, 2011). Crucially, however, among different GCMs there is no correlation between the interannual
42 and long-term cloud-temperature relationships (Dessler, 2010), contradicting the basic assumption of these
43 methods. Many but not all AOGCMs predict relationships that are consistent with observations (Dessler,
44 2010). More recently there is interest in relating the time-lagged correlations of cloud and temperature to
45 feedback processes (Spencer and Braswell, 2010) but again these relationships appear to reveal only a
46 model's ability to simulate ENSO or other modes of interannual variability properly, which are not obviously
47 informative about the cloud feedback on long-term global warming (Dessler, 2011).
48

49 While a number of studies have proposed methods to infer the long-term cloud feedback from observed
50 variability, for a method to be accepted it should have a sound physical basis and be shown to work
51 consistently when applied to different climate models. No method yet proposed passes both tests. Moreover,
52 some model studies show that the response of global cloud radiative effect to a global warming is sensitive to
53 relatively subtle details in the geographic warming pattern, such as the slight hemispheric asymmetry due to
54 the lag of southern ocean warming relative to northern latitudes (Senior and Mitchell, 2000; Yokohata et al.,
55 2008). Cloud responses to specified uniform ocean warming without CO₂ increases are not the same as those
56 to CO₂-induced global warming simulated with more realistic oceans (Ringer et al., 2006), partly because of
57 rapid adjustments (Section 7.2.4.3.6) and because low clouds also feed back tightly to the underlying surface

(Caldwell and Bretherton, 2009). Simulated cloud feedbacks also differ significantly between colder and warmer climates in some models (Crucifix, 2006; Yoshimori et al., 2009). These sensitivities highlight the challenges facing any attempt to infer long-term cloud feedbacks from simple data analyses.

7.2.4.4 Feedback Synthesis

Together, the water-vapour, lapse-rate and cloud feedbacks are the principal determinants of climate sensitivity. The water-vapour and lapse-rate feedbacks should be thought of as a single phenomenon rather than in isolation. To estimate a 90% probability range for that feedback, we double the variance of GCM results about the mean to account for possible common errors among models, to arrive at +1.09 (+0.91 to +1.27) $\text{W m}^{-2} \text{K}^{-1}$, similar to AR4. Values in this range are supported by a steadily growing body of observational evidence, model tests, and physical reasoning. Key aspects of the responses of water vapour and clouds to climate warming now appear to be constrained by large-scale dynamical mechanisms that are not sensitive to poorly-represented small-scale processes, and as such, are more credible.

Several cloud feedback mechanisms now appear consistently in GCMs, summarised in Figure 7.10, most supported by other lines of evidence. They nearly all act in a positive direction. First, high clouds are expected to rise in altitude and thereby exert a stronger greenhouse effect in warmer climates. This altitude feedback mechanism is well understood, has theoretical and observational support, occurs consistently in GCMs and CRMs, and explains about half of the mean positive cloud feedback in GCMs. Second, middle and high-level cloud cover tends to decrease in warmer climates even within the ITCZ, decreasing the albedo more than it increases the greenhouse effect thus adding positive feedback. This cannot be tested observationally but is consistent with anticipated changes to atmospheric water transport. Third, observations and most models suggest storm tracks shift poleward in a warmer climate, drying the subtropics and moistening the high latitudes, which causes further positive feedback via a net shift of cloud cover to latitudes that receive less sunshine. Finally, most GCMs also predict that low cloud amount decreases, especially in the subtropics, another source of positive feedback though one that differs significantly among models and lacks a well-accepted theoretical basis. Over middle and high latitudes, GCMs suggest warming-induced transitions from ice to water clouds may cause clouds to become more opaque, but this appears to have a small net radiative effect.

[INSERT FIGURE 7.10 HERE]

Figure 7.10: Robust cloud responses to greenhouse warming (those simulated by most models and possessing some kind of independent support or understanding). Key climatological features (tropopause, freezing level, circulations) are shown in grey. Changes anticipated in a warmer climate are shown in red (if contributing positive feedback) or brown (if contributing little or ambiguously to feedback); no robust mechanisms contribute negative feedback. Changes include rising high cloud tops and melting level, and increased polar cloud cover and/or optical thickness (high confidence); broadening of the Hadley Cell and/or poleward migration of storm tracks, and narrowing of the ITCZ (medium confidence); and reduced low-cloud amount and/or optical thickness (low confidence). Confidence assessments are based on degree of GCM consensus, strength of independent lines of evidence from observations or process models, and degree of basic understanding.

Currently, neither cloud process models (CRMs and LES) nor observations provide clear evidence to contradict or confirm any of the above mechanisms, except as noted. In some cases these models show stronger low-cloud feedbacks than GCMs, but each model type has limitations, and some GCM biases suggest positive feedbacks are underestimated. For low clouds, cloud process models suggest a variety of potentially opposing response mechanisms that may account for the current spread of GCM feedbacks. In summary we find no evidence to contradict either the cloud or water-vapour-lapse-rate feedback ranges shown by current GCMs, although the many uncertainties mean that true feedback could still lie outside these ranges.

Based on the above synthesis of cloud behaviour, the net radiative feedback due to all cloud types is judged *likely* (>66% chance) to be positive. This is reasoned probabilistically as follows. First, since evidence from observations and process models is mixed as to whether GCM cloud feedback is too strong or too weak overall, and since the positive feedback found in GCMs comes mostly from mechanisms now supported by other lines of evidence, the central (most likely) estimate of the total cloud feedback is taken as the mean from GCMs (+0.8 $\text{W m}^{-2} \text{K}^{-1}$). Second, since there is no accepted basis to discredit individual GCMs, the probability distribution of the true feedback must be at least as broad as the distribution of GCM results.

1 Third, since feedback mechanisms are probably missing from GCMs (particularly involving thin high clouds
2 or low clouds) and some CRMs suggest feedbacks outside the range in GCMs, the probable range of the
3 feedback must be broader than its spread in GCMs. We estimate the likely range of this feedback by
4 doubling the spread (quadrupling the variance) about the mean of the data in Figure 7.9, that is assuming an
5 uncertainty 170% as large as that encapsulated in the GCM range added to it in quadrature, and assuming
6 Gaussian errors. This yields a 90% (very likely) range of -0.2 to $1.4 \text{ W m}^{-2} \text{ K}^{-1}$, with a 16% probability of a
7 negative feedback.

8
9 Note that the assessment of feedbacks in this chapter is independent of constraints on climate sensitivity
10 from observed trends or palaeoclimate information discussed in Section 12.4.

11 7.2.5 *Anthropogenic Sources of Moisture and Cloudiness*

12
13 Human activity can be a source of additional cloudiness through specific processes involving a source of
14 water vapour in the atmosphere. We discuss here the impact of aviation and irrigation on water vapour and
15 cloudiness. The impact of water vapour sources from combustion at the Earth's surface is thought to be
16 negligible. Changes to the hydrological cycle because of land use change are not discussed.

17 7.2.5.1 *Contrails and Contrail-Induced Cirrus*

18
19
20
21 Aviation jet engines emit hot moist air, which can form line shaped persistent condensation trails (contrails)
22 in environments that are supersaturated with respect to ice and colder than -40°C . The contrails are
23 composed of ice crystals that are typically smaller than those of background cirrus (Frömming et al., 2011;
24 Heymsfield et al., 2010). Their effect on longwave radiation dominates over their shortwave effect
25 (Burkhardt and Kärcher, 2011; Rap et al., 2010a; Stuber and Forster, 2007) but models disagree on the
26 relative importance of the two effects. Contrails have been observed to spread into large cirrus sheets which
27 may persist for several hours, and observational studies confirm their overall positive net RF impact
28 (Haywood et al., 2009). Aerosol emitted within the aircraft exhaust may also affect high-level cloudiness.
29 This last effect is classified as an aerosol-cloud interaction and is discussed in Section 7.4.4. Rap et al.
30 (2010b) confirmed the assessment that aviation contrails are very unlikely, at current levels of coverage, to
31 have an observable effect on surface temperature or diurnal temperature range.

32
33 Forster et al. (2007) estimated the 2005 RF from contrails as $+0.01$ (-0.007 to $+0.02$) W m^{-2} , and quoted
34 Sausen et al. (2005) to update the 2000 forcing for aviation-induced cirrus (including linear contrails) to
35 $+0.03$ ($+0.01$ to $+0.08$) W m^{-2} . Lee et al. (2009) scaled these estimates upward 18% to account for revised
36 fuel use estimates, propulsive efficiency and flight routes for year 2005. Kärcher et al. (2010) obtain a range
37 of $+0.008$ to $+0.020 \text{ W m}^{-2}$ for contrails in the year 2000. Traffic distance has further increased by 22%
38 between 2005 and 2010, so that overall we adopt a RF estimate of $+0.02$ ($+0.01$ to $+0.03$) W m^{-2} for
39 persistent (linear) contrails for 2010.

40
41 Satellite estimates of contrail-induced cirrus forcing estimates (e.g., Boucher, 1999) may unintentionally
42 include cirrus changes not directly caused by aviation. Schumann and Graf (2012) constrained their model
43 with observations of the diurnal cycle of contrails and cirrus in a region with high air traffic relative to a
44 region with little air traffic, and estimated an AF of $+0.045$ to $+0.075 \text{ W m}^{-2}$ for contrails and contrail-
45 induced cirrus in 2006, but their model has a large SW contribution, suggesting larger estimates are possible.
46 An alternative approach was taken by Burkhardt and Kärcher (2011), who estimated for year 2002 a global
47 RF of $+0.03 \text{ W m}^{-2}$ from contrails and contrail cirrus within a climate model (Burkhardt and Kärcher, 2009),
48 after compensating for reduced background cirrus cloudiness in the main traffic areas. Combining these
49 estimates with uncertainties on spreading rate, optical depth, ice particle shape and radiative transfer
50 (Markowicz and Witek, 2011) and accounting for the ongoing increase in air traffic, we assess a combined
51 contrail and contrail-induced cirrus AF for year 2010 to be $+0.05$ ($+0.02$ to $+0.15$) W m^{-2} .

52 7.2.5.2 *Irrigation-Induced Cloudiness*

53
54
55 Boucher et al. (2004) estimated a global AF due to water vapour from irrigation in the range of $+0.03$ to
56 $+0.10 \text{ W m}^{-2}$ but the net climate effect was dominated by the evaporative cooling at the surface and by
57 atmospheric thermal responses to low-level humidification. Regional surface cooling was confirmed by a

1 number of more recent regional and global studies (Kueppers et al., 2007; Lobell et al., 2009). The resulting
2 increase in water vapour may induce a small enhancement in precipitation downwind of the major irrigation
3 areas (Puma and Cook, 2010), as well as some regional circulation patterns (Kueppers et al., 2007). Sacks et
4 al. (2009) reported a 0.001 increase in cloud fraction over land (0.002 over irrigated land). This suggests an
5 AF no more negative than -0.1 W m^{-2} with very low confidence.

7.3 Aerosols

9 The section assesses the role of aerosol particles (aerosols) in the climate system, focusing on aerosol
10 processes and properties, as well as other factors, that influence aerosol-radiation and aerosol-cloud
11 interactions. Processes directly relevant to aerosol-cloud interactions are discussed in Section 7.4, and
12 estimates of aerosol radiative and adjusted forcings are assessed in Section 7.5. The time evolution of
13 aerosols and their forcing are discussed in Chapters 2 and 8, with Chapter 8 also covering changes in natural
14 volcanic aerosols.

7.3.1 Aerosols in the Present-Day Climate System

7.3.1.1 Aerosol Formation and Aerosol Types

20 Atmospheric aerosols, whether natural or anthropogenic, originate from two different pathways: emissions of
21 primary particulate matter and formation of secondary particulate matter from gaseous precursors (Figure
22 7.11). The main constituents of the atmospheric aerosol are inorganic species (such as sulphate, nitrate, sea
23 salt), organic species (also termed organic carbon or organic aerosol), black carbon (BC, a distinct type of
24 carbonaceous material formed from the incomplete combustion of fossil and biomass based fuels), and
25 mineral species (mostly desert dust). Dust, sea spray and BC are introduced into the atmosphere as primary
26 particles, whereas non-sea-salt sulphate and nitrate are predominantly from secondary aerosol formation
27 processes. Organic aerosol (OA) has both significant primary and secondary sources. In the present-day
28 atmosphere, the majority of BC, sulphate and nitrate come from anthropogenic sources, whereas sea spray
29 and most mineral aerosol is of natural origin. Atmospheric OA is influenced by both natural and
30 anthropogenic sources of volatile organic compounds. Emission rates of aerosols and aerosol precursors are
31 summarized in Table 7.2. The characteristics and role of the main aerosol species are listed in Table 7.2.

[INSERT FIGURE 7.11 HERE]

34 **Figure 7.11:** Overview of atmospheric aerosol processes and environmental variables influencing aerosol-radiation and
35 aerosol-cloud interactions. Gas-phase processes and variables are highlighted in red while particulate-phase processes
36 and variables appear in green. Although this figure shows a linear chain of processes from aerosols to forcings, it is
37 increasingly recognized that aerosols and clouds form a coupled system with two-way interactions.

[INSERT TABLE 7.2 HERE]

40 **Table 7.2:** Global and regional anthropogenic and natural emissions important for atmospheric aerosols. For the
41 anthropogenic components the maximum and minimum values from available inventories are presented according to
42 Granier et al. (2011). Units are Tg yr^{-1} except for BVOCs in TgC yr^{-1} and DMS in TgS yr^{-1} . Dust and sea-spray
43 estimates span the range in the historical CMIP5 simulations. SOA range is taken from Spracklen et al. (2011). BVOC
44 range from Arneeth et al. (2008).

[INSERT TABLE 7.3 HERE]

47 **Table 7.3:** Key aerosol properties of the main aerosol species in the troposphere. Brown carbon is a particular type of
48 OA but is treated here as an additional component because it is light absorbing. The estimate of aerosol burdens and
49 lifetimes in the troposphere are based on the AeroCom models, except for primary biological aerosol particles (PBAP),
50 which are treated by analogy to other coarse mode aerosol types.

7.3.1.2 Aerosol Climatology

54 Since the AR4, new and improved observational aerosol datasets have emerged. A number of field
55 experiments have taken place such as the Intercontinental Chemical Transport Experiment (INTEX,
56 Bergstrom et al., 2010; Logan et al., 2010), African Monsoon Multidisciplinary Analysis (AMMA, Hansell
57 et al., 2010; Jeong et al., 2008), Integrated Campaign for Aerosols, gases and Radiation Budget (ICARB,
58 2008 and references therein), Megacity Impact on Regional and Global Environments field experiment

(MILAGRO, Paredes-Miranda et al., 2009), Geostationary Earth Radiation Budget Inter-comparisons of Longwave and Shortwave (GERBILS, Christopher et al., 2009), Research of the Composition of the Troposphere from Aircraft and Satellites (ARCTAS, Lyapustin et al., 2010), the Amazonian Aerosol Characterization Experiment 2008 (AMAZA-08, Martin et al., 2010a) and Atmospheric Brown Clouds (ABC, Engling and Gelencsér, 2010; Nakajima et al., 2007), which have improved our understanding of regional aerosol properties.

Long-term aerosol mass concentrations are also measured more systematically at the surface by global and regional networks (see Chapter 2), and there are institutional efforts to improve the coordination and quality assurance of the measurements. A survey of the main aerosol types can be constructed from such measurements (e.g., Jimenez et al., 2009; Figure 7.12). Such analyses show a wide spatial variability in aerosol concentrations, dominant aerosol type, and aerosol composition worldwide. Mineral dust dominates the aerosol mass over some continental regions with relatively higher concentrations in urban S. Asia, accounting for about 35% of the total aerosol mass with diameter smaller than 10 μm . In the rural U.S. and South America, OC contributes the largest fraction to the atmospheric aerosol (i.e., 20% or more), while in other areas of the world the OC fraction ranks second or third with a mean of about 16%. Sulfate normally accounts for about 10–30% by mass, except for the areas in rural Africa, urban Oceania and South America with less than about 10%. The mass fractions of nitrate and ammonium are only around 6% and 4% on average, respectively. In most areas, BC fractions are less than 5% of the aerosol mass, although this percentage may be larger (about 12%) in South America, urban Africa, urban Europe, South-East and East Asia and urban Oceania due to the larger impact of combustion sources. Sea-salt aerosols can be dominant at oceanic remote sites with 50-70% of aerosol mass.

[INSERT FIGURE 7.12 HERE]

Figure 7.12: Bar chart plots summarizing the annual, seasonal or monthly mean mass surface concentration ($\mu\text{g m}^{-3}$) of seven major aerosol components for particles with diameter smaller than 10 μm , from various rural and urban sites in six continental areas of the world with at least an entire year of data, and two marine sites. For each location, the panels represent the median, the 25–75 percentiles (box), and the 10–90 percentiles (whiskers) for each aerosol component. These include: 1) urban North America. (Chow et al., 1993; Ito et al., 2004; Kim et al., 2000; Liu et al., 2005; Malm and Schichtel, 2004; Sawant et al., 2004); rural North America (Chow et al., 1993; Liu et al., 2005; Malm and Schichtel, 2004; Malm et al., 1994); 2) marine northern hemisphere Atlantic Ocean (Ovadnevaite et al., 2011; Rinaldi et al., 2009); 3) urban Europe (Hueglin et al., 2005; Lenschow et al., 2001; Lodhi et al., 2009; Lonati et al., 2005; Perez et al., 2008; Putaud et al., 2004; Querol et al., 2006; Querol et al., 2008; Querol et al., 2001; Querol et al., 2004; Rodriguez et al., 2002; Rodriguez et al., 2004; Roosli et al., 2001; Viana et al., 2006; Viana et al., 2007; Yin and Harrison, 2008); rural Europe (Gullu et al., 2000; Hueglin et al., 2005; Kocak et al., 2007; Putaud et al., 2004; Puxbaum et al., 2004; Querol et al., 2009; Querol et al., 2001; Querol et al., 2004; Rodriguez et al., 2002; Rodriguez et al., 2004; Salvador et al., 2007; Theodosi et al., 2010; Viana et al., 2008; Yin and Harrison, 2008; Yttri, 2007); 4) high Asia, with altitude larger than 1680 m. (Carrico et al., 2003; Decesari et al., 2010; Ming et al., 2007a; Qu et al., 2008; Ram et al., 2010; Rastogi and Sarin, 2005; Rengarajan et al., 2007; Shresth et al., 2000; Zhang et al., 2001; Zhang et al., 2012c; Zhang et al., 2008); urban South Asia (Chakraborty and Gupta, 2010; Khare and Baruah, 2010; Kumar et al., 2007; Lodhi et al., 2009; Raman et al., 2010; Rastogi and Sarin, 2005; Safai et al., 2010; Stone et al., 2010); urban China (Cheng et al., 2000; Hagler et al., 2006; Oanh et al., 2006; Wang et al., 2003; Wang et al., 2005b; Wang et al., 2006; Xiao and Liu, 2004; Yao et al., 2002; Ye et al., 2003; Zhang et al., 2002; Zhang et al., 2012c; Zhang et al., 2011); rural China (Hagler et al., 2006; Hu et al., 2002; Zhang et al., 2012c) [PANEL MISSING]; urban China (Cheng et al., 2000; Hagler et al., 2006; Oanh et al., 2006; Wang et al., 2003; Wang et al., 2005b; Wang et al., 2006; Xiao and Liu, 2004; Yao et al., 2002; Ye et al., 2003; Zhang et al., 2002; Zhang et al., 2012c; Zhang et al., 2011); South-East and East Asia (Han et al., 2008; Khan et al., 2010; Kim et al., 2007; Lee and Kang, 2001; Oanh et al., 2006); 5) South America (Artaxo et al., 1998; Artaxo et al., 2002; Bourotte et al., 2007; Celis et al., 2004; Fuzzi et al., 2007; Gioda et al., 2011; Mariani and Mello, 2007; Martin et al., 2010b; Morales et al., 1998; Souza et al., 2010); 6) urban Africa (Favez et al., 2008; Mkoma, 2008; Mkoma et al., 2009b); rural Africa (Maenhaut et al., 1996; Mkoma, 2008; Mkoma et al., 2009a; Mkoma et al., 2009b; Nyanganyura et al., 2007; Weinstein et al., 2010); 7) marine southern hemisphere Indian Ocean (Rinaldi et al., 2011; Sciare et al., 2009); 8) urban Oceania (Chan et al., 1997; Maenhaut et al., 2000; Radhi et al., 2010; Wang et al., 2005a; Wang and Shooter, 2001).

Aerosol optical depth (AOD), which is related to the column-integrated aerosol amount, is measured by the Aerosol Robotic Network (AERONET, Holben et al., 1998; Holben et al., 2001), other ground-based networks, and a number of satellite-based sensors. Retrievals from aerosol-dedicated instruments such as MODIS (Kleidman et al., 2012; Levy et al., 2010; Remer et al., 2005), MISR (Kahn et al., 2005; Kahn et al., 2007) and POLDER/PARASOL (Tanré et al., 2011) are used preferentially to less specialised instrumentation such as AVHRR (Geogdzhayev et al., 2002; Jeong and Li, 2005), TOMS (Torres et al.,

1998; Torres et al., 2002) and ATSR/AATSR although the latter are useful because of their longer measurement records (see Chapter 2). While each AOD retrieval shows some skill against more accurate AERONET measurements, there are still large differences in regional and seasonal patterns because of differences in sampling, cloud screening and treatment of the surface reflectivity (Kokhanovsky et al., 2010). The global but incomplete satellite measurements can be combined with information from global aerosol models through data assimilation techniques (e.g., Benedetti et al., 2009; Figure 7.13a). Due to the heterogeneity in their sources, their short lifetime and the dependence of sinks on the meteorology, aerosol distributions show large variations on daily, seasonal and interannual scales.

[INSERT FIGURE 7.13 HERE]

Figure 7.13: a) spatial distribution of the AOD (unitless) from the MACC model with assimilation of MODIS AOD (Benedetti et al., 2009; Morcrette et al., 2009) for the year 2010; b) to e) latitudinal vertical cross-sections of the aerosol extinction coefficient (km^{-1}) for four longitudinal bands (180°W to 120°W , 120°W to 60°W , 20°W to 40°E , and 60°E to 120°E) from the CALIOP instrument (Winker et al., 2009). The black circles show the extinction scale height for 90% (top) and 63% (bottom) of the average AOD.

Spaceborne lidars, such as on the CALIPSO mission (Winker et al., 2009), now provide a climatology of the aerosol extinction coefficient (Figure 7.13b-d), highlighting that over most regions the majority of the optically-active aerosol resides in the lowest 1–2 km. There is less information available on the vertical profile of aerosol number and mass concentrations, although a number of field experiments involving research aircraft have measured aerosol concentrations. In particular BC mixing ratios have been measured during the ARCTAS (Jacob et al., 2010), ARCPAC (Warneke et al., 2010), A-FORCE and HIPPO1 (Schwarz et al., 2006; Schwarz et al., 2010) campaigns (Figure 7.14). Commercial aircraft also report total aerosol number concentrations (Brenninkmeijer et al., 2007).

[INSERT FIGURE 7.14 HERE]

Figure 7.14: Comparison of profiles of BC mass mixing ratios (ng kg^{-1}) as measured by airborne SP2 instruments during the ARCTAS (Jacob et al., 2010), HIPPO1 (Schwarz et al., 2006; Schwarz et al., 2010) and A-FORCE (Oshima et al., 2012) campaigns and as simulated by a range of AeroCom models. The model values are averages for the month corresponding to each field campaign.

7.3.2 Aerosol Sources and Processes

7.3.2.1 Aerosol Sources

Sea spray particles comprise sea salt and primary marine matter, and are produced at the sea surface by bubble bursting induced mostly by breaking waves. The effective emission flux of sea spray particles to the atmosphere depends on the surface wind speed and atmospheric stability, and to a lesser extent on the temperature and composition of the sea water. Our understanding of sea spray emissions has increased substantially since AR4; however, process-based estimates of the total mass, number size distribution and chemical composition of emitted sea spray particles continue to have large uncertainties (de Leeuw et al., 2011; Table 7.2). It has been shown that sea spray can contain primary organic matter, preferentially in the submicronic size range, the amount of which depends on biological activity in ocean waters (Facchini et al., 2008). Uncertainty in the source translates into a large uncertainty in the natural level of aerosol number concentration in the marine atmosphere which, unlike the aerosol optical depth and sea spray mass concentrations, is more difficult to constrain from space observations (Jaeglé et al., 2011).

Dust particles are produced mainly by disintegration of aggregates following creeping and saltation of larger soil particles over desert and other arid surfaces (Kok, 2011; Zhao et al., 2006). The magnitude of dust emissions to the atmosphere depends on the surface wind speed and many soil-related factors such as its texture, moisture and vegetation cover. The range of estimates for the global dust emissions spans a factor of about 5 (Huneeus et al., 2011; Table 7.2). The fraction of dust coming from anthropogenic sources remain ill quantified although some recent satellite observations suggest it could be 20 to 25% (Ginoux et al., 2012a; Ginoux et al., 2012b).

Primary biological aerosol particles (PBAP) include bacteria, pollen, fungal spores, lichen, viruses and fragments of animals and plants (Després et al., 2012). Most of these particles are emitted in the coarse mode (Pöschl et al., 2010) and the contribution to the accumulation mode is likely to be very small. There are only

1 a few estimates of the global flux of PBAP and these are poorly constrained (Burrows et al., 2009; Heald and
2 Spracklen, 2009; see Table 7.2).

3
4 The main natural aerosol precursors are dimethylsulfide (DMS) emitted by the oceans and biogenic volatile
5 organic compounds (BVOC) emitted mainly by the terrestrial biosphere. BVOC emissions depend on the
6 amount and type of vegetation, temperature, radiation and several environmental factors such as the ambient
7 CO₂ concentration (Grote and Niinemets, 2008; Pacifico et al., 2009). While speciated BVOC emission
8 inventories have been derived for some continental regions, global emission inventories or schemes are
9 available only for isoprene and some monoterpenes (Guenther et al., 2006; Muller et al., 2008). The total
10 global BVOC emissions have large uncertainties, despite the apparent convergence in different model-based
11 estimates (Arneth et al., 2008).

12
13 The contribution of secondary organic aerosol (SOA) to the total organic aerosol is larger than previously
14 thought, but the split between primary organic aerosol (POA) and SOA has remained somewhat ambiguous
15 due to atmospheric transformation processes affecting both these components (Jimenez et al., 2009;
16 Robinson et al., 2007). Globally, most of atmospheric SOA is expected to originate from biogenic sources,
17 even though anthropogenic sources could be equally important at northern midlatitudes (De Gouw and
18 Jimenez, 2009; Lin et al., 2012). Recent studies suggest that the SOA formation from BVOCs may be
19 enhanced substantially by anthropogenic pollution due to i) high nitrogen oxide concentrations enhancing
20 BVOC oxidation, and ii) high anthropogenic POA concentrations that facilitate transformation of VOCs to
21 the particle phase (Carlton et al., 2010; Heald et al., 2011; Spracklen et al., 2011). The uncertainty range of
22 atmospheric SOA formation has decreased since AR4 but is still larger than a factor of 5 (Farina et al., 2010;
23 Hallquist et al., 2009; Heald et al., 2010; Spracklen et al., 2011; Table 7.2).

24
25 Anthropogenic sources of aerosols (BC, POA) and aerosol precursors (sulphur dioxide, ammonia, nitric acid
26 and VOCs) are known from emission inventories (Table 7.2). They are generally better constrained than
27 natural sources, exceptions being anthropogenic sources of BC, which are likely to be underestimated (Bond
28 et al., 2012), and anthropogenic emissions of some VOCs, fly-ash and dust which are still poorly known.
29 Since AR4, remote sensing by satellites has been increasingly used to constrain natural and anthropogenic
30 aerosol and aerosol precursor emissions (e.g., Dubovik et al., 2008; Huneus et al., 2012; Jaeglé et al., 2011).

31 32 7.3.2.2 *Aerosol Processes*

33
34 New particle formation is the process by which low-volatility vapors nucleate into stable embryos, which
35 under certain condensable vapor regimes, can grow rapidly to produce nanometre-sized aerosol particles.
36 Since AR4, substantial progress in our understanding of atmospheric nucleation and new particle formation
37 has been made (Zhang et al., 2012b). Multiple lines of evidence indicate that while sulfuric acid is the main
38 driver of nucleation (Kerminen et al., 2010; Sipila et al., 2010), the nucleation rate is affected by ammonia
39 and amines (Kirkby et al., 2011; Kurten et al., 2008; Smith et al., 2010; Yu et al., 2012) as well as low-
40 volatile organic vapors (Metzger et al., 2010; Paasonen et al., 2010; Wang et al., 2010a). Neutral nucleation
41 pathways are very likely to dominate over ion-induced nucleation in continental boundary layers, but the
42 situation might be different higher up in the atmosphere (Hirsikko et al., 2011; Kazil et al., 2010).

43
44 Condensation is the main process transferring low-volatile vapors between the gas phase and aerosol
45 particles, and also the dominant process for growth to larger sizes. The growth of the smallest particles
46 depends crucially on the condensation of organic vapors (Riipinen et al., 2011), and is therefore tied strongly
47 with atmospheric SOA formation discussed in Section 7.3.3.1. The treatment of condensation of semi-
48 volatile compounds, such as ammonium nitrate and most organic vapors, remains a challenge in climate
49 modeling.

50
51 Small aerosol particles collide with one other and stick together. This process, termed Brownian coagulation,
52 affects the aerosol mixing state and is the main sink for the smallest aerosol particles. In locations where
53 aerosol lifetimes are long and amounts of condensable vapors are low, such as in the stratosphere and large
54 parts of the free troposphere, coagulation can contribute significantly to aerosol growth.

55
56 Since AR4, observations of atmospheric nucleation and subsequent growth of nucleated particles to larger
57 sizes have been increasingly reported in different atmospheric environments (Kulmala and Kerminen, 2008;

1 Manninen et al., 2010; O'Dowd et al., 2010). Global model studies suggest that after their growth in the
2 atmosphere, nucleated particles have a large impact on the cloud condensation nuclei (CCN) concentrations
3 (Merikanto et al., 2009; Pierce and Adams, 2009b; Spracklen et al., 2008; Yu and Luo, 2009) and potentially
4 also on cloud droplet number concentrations and aerosol-cloud interactions (Kazil et al., 2010; Makkonen et
5 al., 2012; Wang and Penner, 2009). Quantifying the climatic effects caused by atmospheric nucleation
6 remain, however, very difficult because these effects depend in a non-linear way on the nucleation rate, the
7 survival probability of nucleated particles against coagulation, the growth of these particles to larger sizes,
8 and the number concentration and size distribution of primary particle emissions (Chang et al., 2009;
9 Reddington et al., 2011; Spracklen et al., 2010).

10
11 Cloud processing affects the number concentration, composition, size and mixing state of atmospheric
12 aerosol particles via aqueous-phase chemistry taking place inside clouds, via altering aerosol precursor
13 chemistry around and below clouds, and via different aerosol-cloud hydrometeor interactions. A more
14 detailed discussion on aerosol-cloud interactions will be presented in Section 7.4.

15
16 The understanding and modelling of aerosol sinks has seen less progress since AR4. Improved dry
17 deposition models, which depend on the particle size as well as the roughness properties of the surface, have
18 been developed and are increasingly being used in global aerosol models (Feng, 2008; Kerkweg et al., 2006;
19 Petroff and Zhang, 2010). For the largest particles in the coarse mode, it is important to consider
20 sedimentation throughout the atmosphere and its role in dry deposition at the surface. The uncertainty in the
21 estimate of wet deposition (in-cloud and below-cloud scavenging) is controlled by the uncertainties in the
22 prediction of the amount of precipitation, the size and to some extent the chemical composition of particles.
23 For insoluble primary particles like BC and dust, wet deposition also depends strongly on their degree of
24 mixing with soluble compounds. Parameterization of wet deposition of aerosols remains a key source of
25 uncertainty in aerosol models, which affects the vertical distribution and long-range transport of aerosols
26 (Lee et al., 2011; Vignati et al., 2010), with further impact on model estimates of aerosol forcings. It should
27 be noted however that, when the aerosol source term is known, aerosol sinks can be adjusted in global
28 aerosol models by constraining aerosol concentrations against observations, which helps to limit the impact
29 of the very large uncertainties on aerosol sources and sink processes.

30 31 **7.3.3 Progresses and Gaps in Understanding Climate Relevant Aerosol Properties**

32
33 The climate effects of atmospheric aerosol particles depend on their atmospheric distribution, along with
34 their hygroscopicity, optical properties and ability to act as CCN and ice nuclei (IN). Key quantities for
35 aerosol optical and cloud forming properties are the particle number size distribution, chemical composition,
36 mixing state and shape. These properties are determined by a complex interplay between their sources,
37 atmospheric transformation processes and their removal from the atmosphere (Section 7.3.2 and Figure
38 7.11). Since AR4, measurement of some of the key aerosols properties has been greatly improved in
39 laboratory and field experiments using advanced instrumentation, which allows for instance the analysis of
40 individual particles. These experimental studies have in turn stimulated improvement in the model
41 representations of the aerosol physical, chemical and optical properties. We focus our assessment on some of
42 the key issues where there has been progress since AR4.

43 44 **7.3.3.1 Chemical Composition and Mixing State**

45
46 Research on the climate impacts of aerosols has moved further away from the simple paradigm of externally-
47 mixed sulphate, BC and biomass burning aerosols. Although the role of inorganic aerosols as an important
48 anthropogenic contributor to aerosol-radiation and aerosol-cloud interactions has not been questioned, BC
49 and organics have received increasing attention.

50
51 The physical properties of BC (strongly light-absorbing, refractory with a vaporization temperature near 400
52 K, aggregate in morphology, insoluble in most organic solvents) allow a strict definition at least in principle
53 (Bond et al., 2012). Direct measurement of individual BC-containing particles is possible with a laser-
54 induced incandescence (also called SP2, Gao et al., 2007; Moteki and Kondo, 2010; Schwarz et al., 2008b),
55 which has enabled accurate measurements of the size of BC cores, as well as total BC mass concentrations.
56 Condensation of gas-phase compounds on BC and coagulation with other particles alter the mixing state of
57 BC (e.g., Adachi et al., 2010; Li et al., 2003; Pósfai et al., 2003), which can make internally mixed BC in

1 polluted urban air on a timescale of about 12 h (McMeeking et al., 2010; Moteki et al., 2007). The resulting
2 BC-containing particles could become hygroscopic, which can lead to reduced lifetime and atmospheric
3 loading (Stier et al., 2006). Koch et al. (2009b) and Schwarz et al. (2010) used airborne measurements to
4 evaluate AeroCom model simulations of the vertical distribution of BC aerosol in many regions, and found
5 that most models simulate too much BC in the upper troposphere (Schwarz et al., 2010, see Figure 7.14).

6
7 Formation processes of OA remain highly uncertain, which is a major weakness in the present understanding
8 and model representation of atmospheric aerosols (Hallquist et al., 2009; Kanakidou et al., 2005).

9 Measurements by aerosol mass spectrometers have provided some insights into sources and atmospheric
10 processing of OA (Lanz et al., 2007; Ulbrich et al., 2009; Zhang et al., 2005a). Measurements at continental
11 midlatitudes including urban and rural/remote air suggest that the majority of SOA is likely to be oxygenated
12 OA (Zhang et al., 2007a; Zhang et al., 2005b). Measurements within and downstream of urban air indicate
13 that under most circumstances SOA substantially contributes to the total OA mass (de Gouw et al., 2005;
14 Volkamer et al., 2006; Zhang et al., 2007a). Some of the OA is light absorbing and can be referred to as
15 brown carbon (BrC, Andreae and Gelencser, 2006). The absorption properties of BrC can be attributed to
16 water soluble organic compounds and humic-like substances (Graber and Rudich, 2006; Kirchstetter et al.,
17 2004), although they are poorly quantified (Alexander et al., 2008; Flowers et al., 2010).

18
19 There is a large range in the complexity with which OA is represented in global aerosol models. Some
20 complex, yet still parameterized, chemical schemes have been developed recently which account for
21 multigenerational oxidation (Donahue et al., 2011; Jimenez et al., 2009; Robinson et al., 2007). Some
22 regional and global models use semi-empirical schemes, where semi- or non-volatile organic compounds
23 (SVOC) are produced from parent VOCs by oxidation processes and partitioned between the aerosol and gas
24 phases (Fan et al., 2005; Heald et al., 2005; Russell and Allen, 2005; Tsigaridis and Kanakidou, 2003). The
25 representation of SOA in many of the models used in CMIP5 is still very crude in that the source terms are
26 prescribed and/or the models ignore the complex chemical and aging processes (Hennigan et al., 2009), but
27 the impact of these simplifications on aerosol forcing estimates is unclear.

28
29 There are multiple observations that show co-existence of external and internal mixtures relatively soon after
30 emission (Hara et al., 2003; Schwarz et al., 2008a; Twohy and Anderson, 2008). Generally, in biomass
31 burning aerosol, organic compounds and black carbon were frequently found to be internally mixed with
32 ammonium, nitrate, and sulphate (Deboudt et al., 2010; Pratt and Prather, 2010). Studies over urban
33 locations revealed that as much as 90% of the particles are internally mixed with secondary inorganic species
34 (Bi et al., 2011). Likewise there is evidence of internal mixing between dust and biomass burning aerosols
35 when these aerosol types age together (Hand et al., 2010). In sea-spray the primary organic matter is mixed
36 with the sea salt with increasing percentage mass contribution in submicron sizes with decreasing particle
37 size. Studies have shown that the state of mixing can alter particle hygroscopicity and hence their ability to
38 act as CCN (Wex et al., 2010).

39 40 7.3.3.2 *Size Distribution and Optical Properties*

41
42 Aerosol size distribution is one of a key parameter determining both the aerosol spectral optical and CCN
43 properties. Since AR4 much effort has been put on measuring and simulating the aerosol number rather than
44 volume size distribution. For instance, number size distributions in the submicron range (30-500 nm) were
45 measured at 24 sites in Europe for two years (Asmi et al., 2011), although systematic measurements of this
46 parameter are still limited in other regions. Validation studies show agreement between column-averaged
47 volume size distribution from sun photometer measurements (Dubovik et al., 2006) and direct in situ
48 (surface as well as aircraft-based) measurements (Gerasopoulos et al., 2007; Haywood et al., 2011; Radhi et
49 al., 2010; Smirnov et al., 2011), but these inversion products have not been comprehensively validated.

50
51 The aerosol extinction coefficient depends on aerosol size distribution, aerosol refractive index and mixing
52 state. Humidification of internally-mixed aerosols as the RH increases, further influences their light
53 scattering and absorption properties through changes in particle shape, size, and refractive index (Frenay et
54 al., 2010). Lidars provide further measurements of the vertical profile of aerosol extinction coefficient. Yu et
55 al. (2010) and Koffi et al. (2012) found that global aerosol models tend to have a positive bias on the aerosol
56 extinction scale height in some (but not all) regions, resulting in an overestimate of aerosol concentrations
57 above 6 km. The vertical integral of the extinction coefficient is the aerosol optical depth or AOD. It is a key

ingredient to estimate aerosol radiative effects (see Section 7.3.4) and can be measured to a good accuracy at particular locations from sunphotometry. Satellite retrievals of AOD are more uncertain. While error estimates exist for individual measurements (Kleidman et al., 2012; Remer et al., 2005), the uncertainty on the global-mean AOD can only be obtained from the spread between different satellite retrievals (see Section 7.3.1.2) or model simulations. This remains a source of uncertainty when estimating aerosol-radiation interactions, more so for the anthropogenic AOD which is more difficult to constrain from observations (Su et al., 2012).

Aerosol absorption is another key climate-relevant aerosol property. Earlier in-situ methods to measure absorption suffered from important uncertainties (Moosmüller et al., 2009). Recent measurements using photo-acoustic methods and laser-induced incandescence methods are more accurate but remain sparse. Coating of soluble material over a primary aerosol such as a BC or dust core enhance the BC mass absorption efficiency by up to a factor of 2 (Bond and Bergstrom, 2006; Cross et al., 2010), with typical values of about 8–20 m² g⁻¹ at a wavelength of 532 nm. Absorption AOD can be retrieved from sunphotometer measurements in situations where AOD is larger than 0.2. Koch et al. (2009b) used AERONET retrievals of aerosol AOD to show that most AeroCom models underestimate absorption in many regions, but there remain representativeness issues when comparing point observations to a model climatology.

7.3.3.3 CCN Properties of Atmospheric Aerosols

A subset of aerosol particles acts as CCN. The ability of an aerosol particle to take up water and subsequently activate is determined by its size and composition. Whether these particles eventually become cloud droplets or not, understanding the water vapour uptake of aerosol is an important step in assessing RFari. Common CCN in the atmosphere are composed of sea salt, sulphates and sulphuric acid, nitrate and some organics (Table 7.3). CCN activity of inorganic aerosols is relatively well understood. Lately most attention has been paid to the CCN activity of mixed organic/inorganic aerosols (e.g., King et al., 2010; Prisle et al., 2010). Uncertainties in our current understanding of CCN properties are associated with SOA (Good et al., 2010), mainly because OA is still poorly characterized (Jimenez et al., 2009). For SOA it is not clear how important surface tension effects and bulk-to-surface partitioning are, and if the water activity coefficient changes significantly as a function of the solute concentration (Good et al., 2010; Prisle et al., 2008). The size of the CCN has been found to be more important than their chemical composition at two continental locations as larger particles are more readily activated than smaller particles because they require a lower critical supersaturation (Dusek et al., 2006; Ervens et al., 2007). However, the chemical composition may be important in other locations such as the marine environment, where primary organic particles (hydrogels) have been shown to be exceptionally good CCN (Ovadnevaite et al., 2011).

The bulk hygroscopicity parameter κ (Petters and Kreidenweis, 2007), has been introduced to provide a concise way to describe how effectively an aerosol particle functions as a CCN. It can be measured experimentally and is increasingly being used as a way to characterize aerosol properties. Pringle et al. (2010) used surface and aircraft measurements to evaluate the values of the hygroscopicity parameter simulated by a global aerosol model, and found generally good agreement. When the aerosol is dominated by organics, discrepancies between values of κ obtained directly from both CCN activity measurements and sub-saturated particle water uptake measurements have been observed in some instances (e.g., Irwin et al., 2010; Prenni et al., 2007; Roberts et al., 2010), whereas in other studies closure has been obtained (e.g., Duplissy et al., 2008; Rose et al., 2011). Adsorption theory (Kumar et al., 2011) replaces κ -theory for CCN activation for insoluble particles (e.g., dust) while alternative theories are still required for explanation of primary marine organics that seem to have peculiar gel-like properties (Ovadnevaite et al., 2011).

7.3.3.4 IN Properties of Aerosols

Aerosols that act as IN are solid substances at atmospheric temperatures and supersaturations. Mineral dust and primary bioaerosols such as bacteria, fungal spores and pollen, are typically known as good IN. Conflicting evidence has been presented for the ability of BC, organic and biomass burning particles to act as IN. Four heterogeneous ice nucleation modes are distinguished in the literature. Immersion freezing refers to freezing that is initiated from within a cloud droplet, condensation freezing refers to freezing during droplet formation, and contact freezing is initiated when an IN collides with a supercooled cloud droplet. Deposition

nucleation refers to the direct deposition of vapour onto IN. Lidar observations revealed that liquid cloud droplets are present before ice crystals form via heterogeneous freezing mechanisms (Ansmann et al., 2008; de Boer et al., 2011) indicating that deposition nucleation does not seem to be important for mixed-phase clouds.

Laboratory measurements of the ice activity of various aerosols is summarized in terms of their ice onset temperatures versus relative humidities, for both deposition/condensation nucleation and immersion freezing, in Figure 7.15. The reported nucleation onset points refer to different fractions of ice nucleating particles due to different detection thresholds of measurement methods. Bioaerosols initiate immersion freezing at the highest temperatures and are considered as very efficient IN, but their concentration in the upper troposphere is relatively low (Hoose et al., 2010a). Remote sensing observations confirm that ice clouds in air containing dust can be found at significantly warmer temperatures than in dust-free conditions (Choi et al., 2010; Sassen et al., 2003; Seifert et al., 2010). Laboratory results indicate that, in comparison with natural IN such as mineral dust and biological particles, soot initiates ice only at the coldest temperatures (Hoose and Möhler, 2012). As in-situ observations indicate an enrichment of BC in atmospheric ice particle residuals in tropospheric mixed phase clouds (Cozic et al., 2008; Targino et al., 2009; Twohy et al., 2010), there must be some mechanism for BC to enter ice clouds, but that does not necessarily mean that BC acted as IN. Since BC has anthropogenic sources its increase since pre-industrial times may have caused changes to the lifetime of mixed-phase clouds, as discussed in Section 7.4, and thus to AFaci as summarized in Section 7.5.

IN can either be bare or mixed with other substances. As bare particles age in the atmosphere, they acquire liquid surface coatings by condensing soluble species and water vapour or by scavenging soluble particles, which may transform IN from deposition or contact nuclei into possible immersion nuclei. A change from contact to immersion freezing implies an activation at colder temperatures with consequences for the lifetime of mixed-phase clouds and AFaci (Section 7.5).

[INSERT FIGURE 7.15 HERE]

Figure 7.15: The onset temperatures and relative humidities for deposition/condensation freezing and immersion freezing for **bioaerosols** (Ahern et al., 2007; Diehl et al., 2001; Iannone et al., 2011; Kanji et al., 2011; Mohler et al., 2008; Mortazavi et al., 2008; von Blohn et al., 2005; Yankofsky et al., 1981), **mineral dusts** (Archuleta et al., 2005; Bundke et al., 2008; Connolly et al., 2009; Cziczo et al., 2009a; Field et al., 2006; Kanji and Abbatt, 2006; Kanji et al., 2011; Knopf and Koop, 2006; Koehler et al., 2010; Kulkarni and Dobbie, 2010; Lüönd et al., 2010; Mohler et al., 2006; Murray et al., 2011; Niedermeier et al., 2010; Niemand et al., 2012; Roberts and Hallett, 1968; Salam et al., 2006; Schaller and Fukuta, 1979; Welti et al., 2009; Zimmermann et al., 2008), **organics** (Baustian et al., 2010; Kanji et al., 2008; Petters et al., 2009; Prenni et al., 2007; Shilling et al., 2006; Wagner et al., 2010; 2011; Wang and Knopf, 2011; Zobrist et al., 2007), **solid ammonium sulphate** (Abbatt et al., 2006; Baustian et al., 2010; Mangold et al., 2005; Shilling et al., 2006; Wise et al., 2009; 2010) and **BC (soot)** (Crawford et al., 2011; DeMott, 1990; DeMott et al., 1999; Diehl and Mitra, 1998; Dymarska et al., 2006; Fornea et al., 2009; Gorbunov et al., 2001; Kanji et al., 2011; Mohler et al., 2005), from a compilation of experimental data of sub- and super-micron aerosol particles in the literature (for references see supplementary material). The large range of observed ice nucleation onset conditions is due to different experimental setups, particle sizes, activated fractions and chemical composition. Only those IN species for which at least three papers exist are shown. The solid line refers to saturation with respect to liquid water and the dashed line refers to the homogeneous freezing of solution droplets after Koop et al. (2000). Adapted from Hoose and Möhler (2012).

7.3.4 Aerosol-Radiation Interactions (ARI)

7.3.4.1 Aerosol-Radiation Interactions

Direct radiative effect (DRE) is the change in radiative flux caused by the combined direct effect of anthropogenic and natural aerosols. The DRE results from well-understood physics and is close to being an observable quantity, yet our knowledge of aerosol and environmental characteristics needed to quantify the DRE at the global scale remains incomplete (Anderson et al., 2005; Jaeglé et al., 2011; Satheesh and Moorthy, 2005). The DRE requires knowledge of the spectrally-varying aerosol extinction coefficient, single scattering albedo and phase function, which can in principle be estimated from the aerosol size distribution, shape, chemical composition, hygroscopicity, and mixing state (Figure 7.11). Radiative properties of the surface, atmospheric molecules and clouds also influence the DRE. In the solar spectrum and cloud-free conditions, the DRE is typically negative at the top-of-atmosphere but gets weaker, and can become positive with increasing aerosol absorption, decreasing upscatter fraction, or increasing albedo of the underlying

1 surface. DRE is weaker in cloudy conditions, except where the cloud layer is thin or where aerosols are
2 located above or between clouds (e.g., Chand et al., 2009). The DRE at the surface is negative and can be
3 much stronger than the DRE at the top-of-atmosphere over regions where aerosols are absorbing (Li et al.,
4 2010). In the longwave spectrum, top-of-atmosphere DRE is generally positive and mainly exerted by
5 coarse-mode aerosols such as desert dust (Reddy et al., 2005), and stratospheric aerosols (Jacobson, 2001).

6
7 There have been many measurement-based estimates of the DRE (Bauer et al., 2011; Bergamo et al., 2008;
8 Di Biagio et al., 2010; Yu et al., 2006) although some studies involve some degree of modelling. Observed
9 and calculated shortwave radiative fluxes agree within measurement uncertainty when aerosol properties are
10 known (e.g., Osborne et al., 2011). Global observational estimates of the DRE rely on satellite remote
11 sensing of aerosol properties and/or measurements of the Earth's radiative budget (Chen et al., 2011; Kahn,
12 2012). Estimates of shortwave cloud-free top-of-atmosphere DRE over the ocean range from -4 to -6 W m^{-2}
13 on a global, annual average, mainly contributed by sea spray (Bellouin et al., 2005; Loeb and Manalo-Smith,
14 2005; Myhre et al., 2007; Yu et al., 2006). However, DRE can reach tens of W m^{-2} locally. Estimates over
15 land are more difficult as the surface is less well characterised (Chen et al., 2009; Jethva et al., 2009) despite
16 recent progress in aerosol inversion algorithms (e.g., Dubovik et al., 2011). Attempts to estimate the DRE in
17 cloudy-sky remain elusive (e.g., Peters et al., 2011a) although passive and active remote sensing of aerosols
18 over clouds is now possible (de Graaf et al., 2012; Omar et al., 2009; Torres et al., 2007; Waquet et al.,
19 2009). Notable areas of positive top-of-atmosphere DRE exerted by absorbing aerosols include the Arctic
20 over ice surfaces (Stone et al., 2008) and seasonally off the shore of Namibia over stratocumulus clouds.
21 While AOD and aerosol size are relatively well constrained, uncertainties in the aerosol single-scattering
22 albedo (Loeb and Su, 2010) and vertical profile (e.g., Zarzycki and Bond, 2010) contribute significantly to
23 the overall uncertainties in DRE.

24 25 7.3.4.2 *Rapid Adjustments to ARI*

26
27 Aerosol-radiation interactions give rise to rapid adjustments, which are particularly pronounced for
28 absorbing aerosols, where the associated cloud changes are often referred to as the semi-direct effect (see
29 Figure 7.2). The adjusted forcing (AF) from aerosol-radiation interactions is quantified in Section 7.5; only
30 the corresponding processes governing rapid adjustments are discussed here. Impacts on precipitation are
31 discussed in Section 7.6.

32
33 Since AR4, additional studies have found correlations between cloud cover and absorbing aerosols (e.g.,
34 Brioude et al., 2009; Wilcox, 2010), and eddy-resolving, regional and global scale modelling studies have
35 helped confirm a causal link. Relationships between cloud and aerosol reveal a more complicated picture
36 than initially anticipated (e.g., Ghan et al., 2012; Hill and Dobbie, 2008; Koch and Del Genio, 2010; Sakaeda
37 et al., 2011; Zhuang et al., 2010).

38
39 Absorbing aerosols modify atmospheric stability. The effect of this on cloud cover depends on the height of
40 the aerosol relative to the cloud and the type of cloud (Allen and Sherwood, 2010; Koch and Del Genio,
41 2010; Yoshimori and Broccoli, 2008). Aerosol also reduces the downwelling solar radiation at the surface.
42 Together the changes in atmospheric stability and reduction in surface fluxes provide a means for aerosols to
43 significantly modify the fraction of surface-forced clouds (Feingold et al., 2005; Sakaeda et al., 2011). These
44 changes may also affect precipitation as discussed in Section 7.6.2.

45
46 Cloud cover is expected to decrease if absorbing aerosol is embedded in the cloud layer. This has been
47 observed (Koren et al., 2004) and simulated (e.g., Feingold et al., 2005) for clouds over the Amazon forest in
48 the presence of smoke aerosols. In the stratocumulus regime, absorbing aerosol above cloud-top strengthens
49 the temperature inversion, reduces entrainment and tends to enhance cloudiness. Satellite observations
50 (Wilcox, 2010) and modelling (Johnson et al., 2004) of marine stratocumulus show a thickening of the cloud
51 layer beneath layers of absorbing smoke aerosol, which induces a local negative forcing. The responses of
52 other cloud types, such as those associated with deep convection, are not well determined.

53
54 Absorbing aerosols embedded in cloud drops enhances their absorption, which can affect the dissipation of
55 cloud. The contribution to R_{Fari} is small (Ghan et al., 2012; Stier et al., 2007), and there is contradictory
56 evidence regarding the magnitude of the cloud dissipation effect influencing A_{Fari} (Bond et al., 2012;
57 Feingold et al., 2005; Ghan et al., 2012; Jacobson, 2012).

1
2 Global forcing estimates are necessarily based on global models (see Section 7.5.2), although the accuracy of
3 GCMs in this regard is limited by their ability to represent low cloud controlling processes with fidelity. This
4 is an area of concern as discussed in Section 7.2, and limits confidence in these estimates.

6 **7.3.5 *Aerosol Responses to Climate Change and Feedback***

7
8 The climate drivers of changes in aerosols can be split into physical changes (temperature, humidity,
9 precipitation, soil moisture, solar radiation, wind speed, sea ice extent, etc), chemical changes (availability of
10 oxidants) and biological changes (vegetation cover and properties, plankton abundance and speciation, etc).
11 The response of aerosols to climate change may constitute a feedback loop whereby climate processes
12 amplify or dampen the initial perturbation (Carslaw et al., 2010; Raes et al., 2010). We assess here the
13 relevance and strength of aerosol-climate feedbacks in the context of future climate change scenarios.

15 **7.3.5.1 *Changes in Sea Spray and Mineral Dust***

16
17 Concentrations in sea spray will respond to changes in surface wind speed, precipitation and to the expected
18 decrease in sea ice cover (Struthers et al., 2011). There is no agreement among climate models about the
19 balance of effects, with estimates ranging from an overall 19% reduction in global sea salt burden from the
20 present-day to year 2100 (Liao et al., 2006), to little sensitivity (Mahowald et al., 2006a), or a sizeable
21 increase (Bellouin et al., 2011; Jones et al., 2007). In particular there is little understanding of how surface
22 wind speed may change over the ocean in a warmer climate, and observed recent changes (e.g., Young et al.,
23 2011; Chapter 2) may not be indicative of future changes. Given that sea spray particles comprise a
24 significant fraction of CCN concentrations over the oceans, such large changes will feed back on climate
25 through changes in cloud droplet number (Korhonen et al., 2010a).

26
27 Studies of the effects of climate change on dust loadings give a wide range of results. Woodward et al.
28 (2005) found a factor of 3 increase in dust burden in 2100 relative to present-day because of a large increase
29 in bare soil fraction. A few studies reported moderate (–10 to –20%) increases or decreases (e.g., Jacobson
30 and Streets, 2009; Liao et al., 2009; Tegen et al., 2004). Mahowald et al. (2006b) found a 60% decrease
31 under double CO₂ concentration. The large range reflects different responses of the atmosphere and
32 vegetation cover to climate change forcings and results in low confidence in these predictions.

34 **7.3.5.2 *Changes in Sulfate, Ammonium and Nitrate Aerosols***

35
36 The DMS-sulfate-cloud-climate feedback loop could operate in numerous ways through changes in
37 temperature, absorbed solar radiation, mixed layer depth and nutrient recycling, sea-ice extent, wind speed,
38 shift in marine ecosystems due to ocean acidification and climate change, atmospheric processing of DMS
39 into CCN, and no study has included all the relevant effects. However two decades of research have yielded
40 important insights into this complex, coupled system (Ayers and Caine, 2007; Carslaw et al., 2010). There
41 is now medium agreement between observations and Earth System model simulations for a weak feedback
42 due to a weak sensitivity of the CCN population to changes in DMS emissions (Carslaw et al., 2010; Quinn
43 and Bates, 2011; Woodhouse et al., 2010). Parameterisations of oceanic DMS production lack robust
44 mechanistic justification (Halloran et al., 2010) and as a result the sensitivity to ocean acidification and
45 climate change remains uncertain (Bopp et al., 2004; Cameron-Smith et al., 2011; Kim et al., 2010).

46
47 In the atmosphere chemical production of sulfate increases with temperature (Aw and Kleeman, 2003;
48 Dawson et al., 2007; Kleeman, 2008), but most studies to date predict a small (0 to 9%) reduction in global
49 sulfate burden, mainly because of future increases in precipitation (Liao et al., 2006; Pye et al., 2009;
50 Racherla and Adams, 2006; Unger et al., 2006). However Rae et al. (2007) found a small increase in global
51 sulfate burden from 2000–2100 because the simulated future precipitation was reduced in regions of high
52 sulphate abundance.

53
54 Changes in temperature have a large impact on nitrate aerosol formation through the Clausius-Clapeyron
55 relation for HNO₃. There is some agreement among global aerosol models that climate change alone will
56 contribute to a decrease in the nitrate concentrations (Bellouin et al., 2011; Liao et al., 2006; Pye et al., 2009;
57 Racherla and Adams, 2006) with the exception of Bauer et al. (2007) who found little change in nitrate for

1 year 2030. It should be noted however that changes in precursor emissions are likely to increase nitrate
2 concentrations in the future (Bellouin et al., 2011). Besides the changes in meteorological parameters,
3 climate change can also influence ammonium formation by changing concentrations of sulfate and nitrate,
4 but the effect of climate change alone was found to be small (Pye et al., 2009).

5 6 7.3.5.3 *Changes in Carbonaceous Aerosols*

7
8 There is evidence that future climate change could lead to increases in the occurrence of wildfires because of
9 changes in fuel availability, readiness of the fuel to burn and ignition sources (Kloster et al., 2010; Marlon et
10 al., 2008; Mouillot et al., 2006; Pechony and Shindell, 2010). However vegetation dynamics may also play a
11 role that is not well understood. Increased fire occurrence would increase aerosol emissions, but decrease
12 BVOC emissions. This could lead to a small positive or negative net radiative effect and feedback (Carslaw
13 et al., 2010).

14
15 A large fraction of secondary organic carbon aerosol forms from the oxidation of isoprene, sesquiterpenes
16 and monoterpenes from biogenic sources. Emissions from vegetation can increase in a warmer atmosphere,
17 everything else being constant (Guenther et al., 2006). Global aerosol models simulate an increase in
18 isoprene emissions of 22 to 55% by 2100 in response to temperature change (Heald et al., 2008; Liao et al.,
19 2006; Sanderson et al., 2003) and a change in global SOA burden of -6% to +11% through the climate-
20 induced changes in aerosol processes and removal rates (Heald et al., 2008; Liao et al., 2006; Tsigaridis and
21 Kanakidou, 2007). Increasing CO₂ concentrations are believed to inhibit BVOC emissions (Arneft et al.,
22 2007) which could offset the temperature effect and adds significant uncertainty to future emissions. Future
23 changes in vegetation cover, whether they are natural or anthropogenic, also introduce large uncertainty in
24 emissions (Lathière et al., 2010). There is little understanding on how the marine source of organic aerosol
25 may change with climate, notwithstanding the large range of emission estimates for the present day (Carslaw
26 et al., 2010).

27 28 7.3.5.4 *Synthesis*

29
30 The emissions, properties and concentrations of aerosols or aerosol precursors could respond significantly to
31 climate change, but there is little consistency across studies in the magnitude or sign of this response. The
32 lack of consistency arises mostly from our limited understanding of processes governing the source of
33 natural aerosols and the complex interplay of aerosols with the hydrological cycle. The feedback parameter
34 as a result of the future changes in emissions of natural aerosols is mostly bracketed within $\pm 0.1 \text{ W m}^{-2} \text{ K}^{-1}$
35 (Carslaw et al., 2010). With respect to anthropogenic aerosols, Liao et al. (2009) showed a significant
36 positive feedback (feedback parameter of +0.04 to +0.15 $\text{W m}^{-2} \text{ K}^{-1}$ on a global mean basis) while Bellouin
37 et al. (2011) simulated a smaller feedback of -0.02 to -0.08 $\text{W m}^{-2} \text{ K}^{-1}$. Overall we assess that models
38 simulate relatively small feedback parameters (i.e., within $\pm 0.2 \text{ W m}^{-2} \text{ K}^{-1}$) with very low confidence,
39 however regional effects on the aerosol may be important.

40 41 7.4 **Aerosol-Cloud Interactions**

42 43 7.4.1 *Introduction*

44
45 This section assesses our understanding of aerosol-cloud-precipitation interactions, emphasizing the ways in
46 which anthropogenic aerosols may be affecting the distribution and radiative properties of clouds and
47 precipitation. The idea that anthropogenic aerosols are changing cloudiness, thus contributing a substantial
48 forcing to the climate system, has been addressed to varying degrees in all of the previous IPCC assessment
49 reports.

50
51 Since AR4, research has continued to articulate new pathways through which the aerosol may affect the
52 radiative properties of clouds, as well as the intensity and spatial patterns of precipitation (e.g., Rosenfeld et
53 al., 2008). Progress can be identified on four fronts: (i) global-scale modelling has advanced in its ability to
54 represent a greater diversity of aerosol-cloud interactions, and with greater internal consistency; (ii)
55 observational studies continue to document strong local correlations between CCN proxies and clouds or
56 precipitation, but have become more quantitative and are increasingly identifying the methodological
57 challenges associated with such correlations; (iii) regional-scale modelling is increasingly being used to

1 assess regional influences of aerosol on cloud field properties and precipitation; (iv) fine-scale modelling
2 studies have begun to be used more widely, and among other things have shown how turbulent mixing, cloud
3 and regional-scale circulations may buffer the effects of aerosol perturbations.

4
5 This section focuses on the microphysics of aerosol-cloud interactions in liquid and mixed-phase clouds.
6 Their radiative implications are quantified in Section 7.5. This section also includes discussion of aerosol
7 influences on light precipitation in shallow clouds but defers discussion of aerosol effects on precipitation
8 from mixed-phase clouds to Section 7.6.

10 *7.4.1.1 Overview and Classification of Hypothesized Aerosol-Cloud Interactions*

11
12 Denman et al. (2007) catalogued several possible pathways via which the aerosol might affect clouds. Given
13 the number of possible aerosol-cloud interactions, and the difficulty of isolating them individually, we see
14 little value in attempting to assess each effect in isolation, especially since modelling studies suggest that the
15 effects may interact and compensate (Morrison and Grabowski, 2011; Stevens and Feingold, 2009). Instead,
16 we group all radiative consequences of aerosol-cloud interactions into two broad categories: the immediate
17 impact in the absence of macrophysical changes to clouds, denoted “radiative forcing due to aerosol-cloud
18 interactions” or RFaci, and the final result including follow-on impacts of macrophysical responses to the
19 initial change, denoted “adjusted forcing due to aerosol-cloud interactions”, or AFaci (Figure 7.2). RFaci
20 represents the classical “Twomey (1977)” or cloud albedo effect whereby greater CCN numbers increase the
21 droplet surface area, but is extended to include changes in the breadth of the size distribution (Section 7.4.2).
22 AFaci additionally accounts for any secondary effects that result as clouds adjust to the rapid changes in their
23 environment accompanying an aerosol perturbation (Figure 7.1), such as lifetime effects, wherein cloud
24 macrostructure adjusts to changes in cloud microstructure (Albrecht, 1989; Liou and Ou, 1989; Pincus and
25 Baker, 1994). AFaci does not, however, include adjustments associated with aerosol-radiation interactions.
26 Although AFaci subsumes RFaci, we retain an estimate of RFaci in Section 7.5 for continuity with prior
27 assessments, and because it is better understood than the model-dependent effects determining AFaci.
28 Possible contributions to the AFaci from liquid clouds are discussed in Section 7.4.3, separately from those
29 associated with adjustments by ice or mixed phase clouds (Section 7.4.4). Figure 7.16 shows a schematic of
30 many of the processes to be discussed in Sections 7.4, 7.5, and 7.6.

32 **[INSERT FIGURE 7.16 HERE]**

33 **Figure 7.16:** Schematic depicting the myriad aerosol-cloud-precipitation related processes occurring within a typical
34 GCM grid box. The schematic conveys the importance of considering aerosol-cloud-precipitation processes as part of
35 an interactive system encompassing a large range of spatiotemporal scales. Cloud types include low-level stratocumulus
36 and cumulus where research focuses on droplet activation, mixing, cloud scavenging, and new particle formation; ice-
37 phase cirrus clouds where a key issue is homogeneous freezing; and deep convective clouds where some of the key
38 questions relate to aerosol influences on liquid, ice, and liquid-ice pathways for precipitation formation, cold pool
39 formation, and scavenging. These processes influence the short- and longwave forcing of the system and hence climate.

41 *7.4.1.2 Advances and Challenges in Observing Aerosol-Cloud Interactions*

42
43 Since AR4, progress has been made in understanding how measurement artefacts affect retrievals of both
44 aerosol (Jeong and Li, 2010; Kahn et al., 2005; Tanré et al., 1996; Tanré et al., 1997) and cloud properties
45 (Platnick et al., 2003) in broken cloud fields. Two key issues are that measurements classified as “cloud-
46 free” may not be, and that aerosol measured in the vicinity of clouds is significantly different than it would
47 be were the cloud field, and its proximate cause (high humidity), not present (e.g., Loeb and Schuster, 2008).
48 The latter results from humidification effects on aerosol optical properties (Charlson et al., 2007; Su et al.,
49 2008; Tackett and Di Girolamo, 2009; Twohy et al., 2009), contamination by undetectable cloud fragments
50 (Koren et al., 2007) and the remote effects of radiation scattered by cloud edges on aerosol retrieval (Várnai
51 and Marshak, 2009; Wen et al., 2007).

52
53 While passive satellite retrievals are unable to distinguish aerosol layers above or below clouds from those
54 intermingling with the cloud field, newer active space-based remote sensing has begun to address this
55 problem (Anderson et al., 2005; Huffman et al., 2007; Stephens et al., 2002). Spectral polarization and multi-
56 angular measurements can discriminate between cloud droplets and aerosol particles and thus improve
57 estimates of aerosol loading and absorption (Deuzé et al., 2001; Mishchenko et al., 2007). Field studies
58 (Raubert et al., 2007; Vogelmann et al., 2012; Wood et al., 2011b) and laboratory investigations (e.g.,

1 Stratmann et al., 2009) of aerosol-cloud interactions also continue to make important contributions to our
2 understanding of how aerosols impact cloud processes, and how clouds in turn modify aerosols. The latter
3 occurs along a number of pathways including aqueous chemistry, which adds non-volatile mass to drops
4 (e.g., Schwartz and Freiberg, 1981), coalescence scavenging, whereby drop collision-coalescence diminishes
5 the drop (and aerosol) number concentration (Hudson, 1993), new particle formation in the vicinity of clouds
6 (Clarke et al., 1999) and aerosol removal by rain (see also Section 7.3). As a result, our understanding of the
7 distribution and properties of the aerosol in the vicinity of clouds continues to improve apace with an
8 appreciation of the limits of this understanding (Anderson et al., 2009).

9
10 The observational challenge of inferring causality from correlation remains a large and limiting one. Because
11 the aerosol is a strong function of air-mass history and origin, and is strongly influenced by cloud and
12 precipitation processes (Anderson et al., 2009; Clarke et al., 1999; Petters et al., 2006), and both are affected
13 by meteorology (Engström and Ekman, 2010), correlations between the aerosol and cloud, or precipitation,
14 cannot be taken as generally indicating a cloud response to the aerosol (e.g., Painemal and Zuidema, 2010).
15 Furthermore, attempts to control for other important factors (air-mass history or cloud dynamical processes)
16 are limited by a lack of understanding of cloud controlling factors in the first place (Anderson et al., 2009;
17 Siebesma et al., 2009; Stevens and Brenguier, 2009). These problems greatly undermine confidence in
18 observationally-based inferences of aerosol effects on clouds and precipitation.

19 20 7.4.1.3 *Advances and Challenges in Modelling Aerosol-Cloud Interactions*

21
22 Fine-scale models, capable of resolving cloud-scale circulations have greatly advanced as a tool for testing
23 the physical mechanisms proposed to govern aerosol-cloud-precipitation interactions (Ackerman et al., 2009;
24 vanZanten et al., 2011). A general finding from explicit numerical simulations of clouds is that various
25 aerosol impact mechanisms tend to be mediated (and often buffered) by interactions across scales not
26 included in the idealized models that gave rise to the original idea (Stevens and Feingold, 2009). Specific
27 examples involve the interplay between the drop-size distribution and mixing processes that determine cloud
28 macrostructure (Ackerman et al., 2004; Bretherton et al., 2007; Small et al., 2009; Stevens et al., 1998;
29 Wood, 2007), or the dependence of precipitation development in stratiform clouds on details of the vertical
30 structure of the cloud (Wood, 2007). As a result it is likely that the physical system is less sensitive to
31 aerosol perturbations than are large-scale models, which do not represent all of these compensating
32 processes.

33
34 At larger scales, regional models allow for a much broader range of scale interactions and timescales long
35 enough for robust features to emerge (Bangert et al., 2011; Seifert et al., 2012). Regional models allow
36 exploration of aerosol-cloud interactions in the context of non-idealized meteorology, variability in land
37 surface, and diurnal/monthly cycles (Tao et al., 2012). These advantages must be weighed against their
38 inability to resolve the fine-scale cloud processes discussed above. Regional models have brought to light the
39 possibility of aerosol gradients manifesting changes in circulation patterns via numerous mechanisms
40 including gradients in heating rates (Lau et al., 2006; Section 7.3.4.2), changes in the radiative properties of
41 cloud anvils (van den Heever et al., 2011; see Section 7.6), or changes in the spatial distribution of
42 precipitation (Lee, 2012; Section 7.6).

43
44 The representation of aerosol effects in global models has also advanced. Most global models now represent
45 an increasing number of hypothesized aerosol-cloud interactions, and are undergoing evaluation through
46 comparisons to data and to other models (Quaas et al., 2009). Historically, aerosol-cloud interactions in
47 climate models have largely been introduced based on simple constructs (e.g., Albrecht, 1989; Pincus and
48 Baker, 1994; Twomey, 1977). There has been significant progress on droplet activation (e.g., Ghan et al.,
49 2011) and ice nucleation (Barahona and Nenes, 2008; DeMott et al., 2010) parameterizations, however these
50 still depend heavily on unresolved (climate model) physics such as updraught velocity. Similarly,
51 parameterizations of aerosol influences on cloud amount cannot account for known non-monotonic
52 responses (Section 7.4.3.2). Global models are now beginning to represent effects in convective, ice and
53 mixed-phase clouds (e.g., Lohmann, 2008; Section 7.6; Song and Zhang, 2011). In addition
54 “superparameterisation” approaches (Section 7.2.2.1.2) hold promise for treating aerosol-cloud interactions
55 more comprehensively, with recent results supporting the notion that aerosol forcing is smaller than
56 simulated by standard climate models (Wang et al., 2011b; see Section 7.5.3). Nevertheless, we caution that
57 for both liquid-only and mixed-phase clouds, the inability of GCMs to resolve cloud-scale updraught

1 velocities and associated cooling rates is a pervasive concern for aerosol-cloud interactions, even for models
2 with grid sizes on the order of kilometres.

3
4 Although advances have been considerable, the challenges remain formidable. The representation of clouds
5 in large-scale models remains primitive (Section 7.2.3) and even if large-scale models were able to represent
6 clouds with greater fidelity, fine-scale modelling suggests that the outcome of an aerosol perturbation
7 depends on the details of the interaction of clouds, turbulence, radiation and precipitation processes on a
8 range of scales not represented by large-scale models (vanZanten et al., 2011). For this reason it is not
9 surprising that large-scale models exhibit a range of manifestations of aerosol-cloud interactions, which
10 limits quantitative inference (Quaas et al., 2009). This emphasises the need to incorporate into climate
11 models the lessons learned from cloud-scale models, in a physically consistent way.

12 *7.4.1.4 Combined Modelling and Observational Approaches*

13
14 Combined approaches, which attempt to maximize the respective advantage of models and data, are
15 beginning to add to understanding of aerosol-cloud interactions. These include inversions of the observed
16 historical record using large-scale modelling studies, but also the use of reanalysis and chemical transport
17 models to help interpret satellite records (Chameides et al., 2002; Koren et al., 2010; Mauger and Norris,
18 2010), field study data to help constrain fine-scale modelling studies (Ackerman et al., 2009; vanZanten et
19 al., 2011), or satellite/surface-based climatologies to constrain large-scale modelling (Quaas et al., 2009).

20 **7.4.2 Radiative Forcing due to Aerosol-Cloud Interactions (RFaci)**

21 *7.4.2.1 The Physical Basis*

22
23
24 The cloud albedo effect (Twomey, 1977), or RFaci, is the mechanism by which an increase in aerosol
25 number concentration leads to an increase in liquid cloud albedo (reflectance of incoming solar radiation) by
26 increasing the cloud droplet number concentration and hence increasing total droplet surface area, with the
27 liquid water content and cloud geometrical thickness held fixed. Although only the change in the droplet
28 concentration is considered in the original concept of the cloud albedo effect, a change in the shape of the
29 droplet size distribution that is directly induced by the aerosols, may also play a role (e.g., Feingold et al.,
30 1997; Liu and Daum, 2002). In the Arctic, anthropogenic aerosols may influence the longwave emissivity of
31 thin liquid clouds and generate a positive forcing at the surface (Garrett and Zhao, 2006; Lubin and
32 Vogelmann, 2006). This line of research is at a stage of relative infancy and is not covered further in this
33 assessment.

34
35
36 The physical basis of RFaci is fairly well understood, with research since AR4 generally reinforcing earlier
37 work. Detailed in-situ aircraft observations show that droplet concentrations observed just above the cloud
38 base generally agree with those predicted given the aerosol and updraught observed below the cloud (e.g.,
39 Fountoukis et al., 2007). Vertical profiles of cloud effective radius also agree with those predicted by models
40 that take into account the effect of entrainment (Lu et al., 2008), although uncertainties still remain in
41 estimating the shape of the droplet size distribution (Brenquier et al., 2011), and the degree of non-
42 adiabaticity within clouds. Multi-dimensional radiative transfer calculations have also been applied to
43 estimate cloud albedo instead of using the traditional two-stream approximation to find that the latter could
44 overestimate the albedo effect under certain conditions (Duda et al., 1996; Zuidema et al., 2008).

45 *7.4.2.2 Observational Evidence for Aerosol-Cloud Interactions*

46
47
48 At low AOD (i.e., less than about 0.3) there is ample observational evidence for increases in aerosol
49 resulting in an increase in drop concentration and decrease in drop size (for constant liquid water) but
50 uncertainties remain regarding the magnitude of this effect, and its sensitivity to spatial averaging. Based on
51 simple metrics, there is a large range of physically plausible responses, with aircraft measurements (e.g.,
52 Hegg et al., 2012; Lu et al., 2007; Lu et al., 2008; Twohy et al., 2005) tending to show stronger responses
53 than satellite-derived responses (McComiskey and Feingold, 2008; Nakajima and Schulz, 2009). At high
54 AOD, droplet concentration tends to saturate and, if the aerosol is absorbing, there may be reductions in
55 droplet concentration and cloudiness (Koren et al., 2008). This effect is part of AFaci (Section 7.4.3.2).

1 Since clouds experience rapid adjustments to liquid water and depth when influenced by aerosol
2 perturbations, RF_{aci} is a hypothetical construct. Hence while an aerosol influence on cloud microphysics is
3 observable, the radiative forcing associated with the “constant liquid water RF_{aci}” is generally not (Section
4 7.4.3.2).

6 7.4.2.3 *Advances in Process Level Understanding*

7
8 At the heart of the albedo effect lie two fundamental issues. The first is droplet activation and its sensitivity
9 to aerosol and meteorological parameters. The primary controls on droplet concentration are the aerosol
10 number concentration (particularly at diameters greater than about 60 nm) and cooling rate (proportional to
11 updraught velocity). Aerosol size distribution can play an important role under high aerosol loadings,
12 whereas aerosol composition tends to be much less important, except perhaps under very polluted conditions
13 and low updraught velocities (e.g., Ervens et al., 2005). The relative unimportance of composition is partially
14 because aging makes particles more hygroscopic, but also because of the self-regulation of the
15 supersaturation field that determines drop activation (e.g., higher hygroscopicity, which initially favours
16 activation, leads to higher water vapour uptake, which then lowers supersaturation and suppresses
17 activation). The second issue is the presence of condensed water that strongly determines how much energy
18 can be reflected; there is no RF_{aci} unless clouds are present. Simple arguments show that in a relative sense
19 the amount of reflected energy is approximately two-and-a-half times more sensitive to changes in the liquid
20 water path than to changes in droplet concentration (Boers and Mitchell, 1994). Since both these parameters
21 experience similar ranges of variability, the magnitude of aerosol-cloud related forcing rests mostly on
22 dynamical factors such as turbulent strength and entrainment that control cloud amount, and a few key
23 aerosol parameters such as aerosol number concentration and size distribution, and to a much lesser extent,
24 composition.

25 7.4.3 *Forcing Associated with Adjustments in Liquid Clouds (AF_{aci})*

26 7.4.3.1 *The Physical Basis for Adjustments in Liquid Clouds*

27
28 The adjustments giving rise to AF_{aci} are multi-faceted and are often associated with changes in cloud
29 lifetime or cloud water, previously referred to as ‘lifetime’ effects (Figure 7.2). However this old
30 nomenclature is misleading because it assumes a relationship between cloud lifetime and cloud amount or
31 water content. Moreover, the effect of the aerosol on cloud amount may have nothing to do with cloud
32 lifetime per se (e.g., Pincus and Baker, 1994).

33
34 The traditional view has been that adjustment effects associated with aerosol-cloud-precipitation interactions
35 will add to the initial albedo increase by increasing cloud amount. The chain of reasoning involves three
36 steps: first that droplet concentrations depend on the number of available CCN; second that precipitation
37 development is regulated by the droplet concentration; and third that the development of precipitation
38 reduces cloud amount (Stevens and Feingold, 2009).

39
40 Of the three steps the first has ample support in both observations and theory (Section 7.4.2). More
41 problematic are the second two links in the chain of reasoning. The physical basis for a pervasive, positive
42 dependence of cloud amount on the available CCN is weak. Although increased droplet concentrations
43 inhibit the initial development of precipitation in single clouds and stratocumulus (see Section 7.4.3.2.1), it is
44 not clear that such an effect operates in an evolving, broken cloud field. In the trade-cumulus regime, some
45 modelling studies suggest the opposite, with increased aerosol concentrations actually promoting the
46 development of deeper clouds and invigorating precipitation (Stevens and Seifert, 2008; see discussion of
47 similar responses in deep convective clouds in Section 7.6). Others have shown alternating cycles of larger
48 and smaller cloud water in both aerosol-perturbed stratocumulus (Sandu et al., 2008) and trade cumulus (Lee
49 et al., 2012), pointing to the important role of environmental feedback. Although the original studies that
50 hypothesized cloud amount effects (Albrecht, 1989; Liou and Ou, 1989) are often taken as demonstrative of
51 this point, there is limited unambiguous observational evidence (exceptions to be given below). Many
52 climate models assume such an effect *a priori*, which likely influences their forcing estimates.

53 7.4.3.2 *Observational Evidence of Adjustments in Liquid Clouds*

1 Since AFaci subsumes RFaci, and because RFaci is hard to observe, we now discuss observations of aerosol-
2 perturbed cloud fields more generally.

3 4 7.4.3.2.1 *Stratocumulus*

5 The cloud albedo effect is best manifested in so-called ship tracks, which are bright lines of clouds behind
6 ships. As shown during the Monterey Area Ship Track (MAST) experiment, many ship tracks are
7 characterized by an increase in the droplet concentration resulting from the increase in aerosol number
8 concentration and an absence of drizzle size drops, which leads to a decrease in the droplet radius and an
9 increase in the cloud albedo (Durkee et al., 2000), all else equal. Coakley and Walsh (2002) showed that
10 cloud water responses can be of different sign. This is supported by more recent shiptrack analyses based on
11 the new A-Train satellites (Christensen and Stephens, 2011); liquid water decreases weakly (−6%) in
12 overcast clouds in response to ship-track intrusions but increases significantly (39%) in precipitating, broken
13 stratocumulus clouds where aerosol intrusions result in larger cloud fraction. Adjustments are therefore key
14 to understanding radiative response. A-Train satellites studies of long-term degassing of low-lying volcanic
15 aerosol on stratocumulus point to smaller drop sizes but ambiguous changes in cloud fraction and cloud
16 water (Gasso, 2008). The global radiative forcing of visible ship tracks has been estimated from satellite and
17 found to be insignificant at about -0.5 mW m^{-2} (Schreier et al., 2007), although there is some concern that
18 this analysis may not have identified all shiptracks. Nevertheless, this result is supported by two years of data
19 downwind of shipping lanes that were unable to distinguish aerosol influences from meteorological
20 influences on cloud microphysical or macrophysical properties (Peters et al., 2011b). Counter evidence of
21 shiptracks significantly increasing the cloud fraction and albedo of broken cloud scenes are also emerging
22 (Goren and Rosenfeld, 2012).

23
24 The development of precipitation in stratocumulus, whether due to aerosol or meteorological influence can,
25 in some instances, change a highly reflective closed-cellular cloud field to a weakly reflective broken open-
26 cellular field (Comstock et al., 2005; Savic-Jovicic and Stevens, 2008; Sharon et al., 2006; Stevens et al.,
27 2005b; vanZanten et al., 2005; Wang and Feingold, 2009a). In some cases, compact regions (pockets) of
28 open-cellular convection become surrounded by regions of closed-cellular convection. It is, however,
29 noteworthy that observed precipitation rates can be similar in both open and closed-cell environments (Wood
30 et al., 2011a). The lack of any apparent difference in the large-scale environment of the open cells, versus the
31 surrounding closed cellular convection, suggests the potential for multiple equilibria (Baker and Charlson,
32 1990; Feingold et al., 2010). Therefore in the stratocumulus regime, the onset of precipitation may lead to a
33 chain of events that leads to a large-scale reduction of cloudiness in agreement with Liou and Ou (1989) and
34 Albrecht (1989). The transition may be bidirectional: ship tracks passing through open-cell regions also
35 appear to revert the cloud field to a closed-cell regime inducing a potentially strong AFaci (Christensen and
36 Stephens, 2011; Goren and Rosenfeld, 2012; Wang et al., 2011a).

37 38 7.4.3.2.2 *Trade-cumulus*

39 Precipitation from shallow convective clouds proves difficult to observe in trade cumuli, as the clouds are
40 small, and not easily observed by space-based remote sensing techniques (Stephens et al., 2008). A recent A-
41 Train satellite study of trade cumuli influenced by aerosol associated with slow volcanic degassing points to
42 smaller droplet size, decreased precipitation efficiency, increased cloud amount, and higher cloud tops (Yuan
43 et al., 2011). Other studies show that in the trade cumulus regime cloudiness tends to increase with
44 precipitation amount, most likely because processes that favour precipitation development also favour clouds
45 (Nuijens et al., 2009) and because precipitating trade cumuli tend to regenerate through colliding outflows
46 (Zuidema et al., 2012).

47
48 While observationally-based assessments of aerosol-cloud interactions have a long history, a more recent
49 development is assessment of the ability of detailed models to reproduce the radiative fingerprints of aerosol-
50 cloud interactions in cumulus cloud fields (Schmidt et al., 2009). This involves comparison between
51 measurements of the irradiance field with the same field calculated in a fine-scale model that represents
52 aerosol-cloud interactions. Such an approach identifies key forcing parameters (e.g., cloud-field properties,
53 aerosol hygroscopicity and absorption) and provides the link between aerosol-cloud interactions and AFaci.

54 55 7.4.3.3 *Advances in Process Level Understanding*

1 Since AR4, there has been progress towards understanding some basic processes relevant to AFaci. One
2 question is how susceptible precipitation is to droplet concentration, and by inference to the available
3 aerosol. Some studies point to the drop effective radius as a threshold indicator of the onset of drizzle
4 (Rosenfeld and Gutman, 1994; Wang et al., 2011a). Others focus on the sensitivity of the conversion of
5 cloud-water to rain-water (autoconversion) and find that it scales with the *square* of the inverse of the droplet
6 concentration (Khairoutdinov and Kogan, 2000). However theoretical work that incorporates a fuller
7 description of rain formation processes suggests that this strongly over-estimates the sensitivity of rain
8 formation in shallow clouds (Stevens and Seifert, 2008), and that rain formation scales with approximately
9 the inverse *square-root* of the droplet concentration (Kostinski, 2008; Seifert and Stevens, 2010).
10 Observational studies also suggest a weaker scaling law (approximately the inverse of drop concentration;
11 Comstock et al., 2005; Pawlowska and Brenguier, 2003; vanZanten et al., 2005). Note that thicker, liquid
12 clouds generate rain via accretion of cloud drops by raindrops, a process that is relatively insensitive to
13 droplet concentration, and therefore to aerosol perturbations (e.g., Khairoutdinov and Kogan, 2000).

14
15 The balance of evidence suggests that autoconversion likely scales with approximately the inverse square
16 root of droplet concentration and that liquid water path has significantly more leverage over precipitation
17 than does droplet concentration. Some of the effects that reduce and even eliminate the sensitivity of rain
18 formation to the autoconversion process have begun to be incorporated in parameterizations used by large-
19 scale models (Posselt and Lohmann, 2009).

20
21 Small-scale studies (Ackerman et al., 2004; Small et al., 2009; Xue et al., 2008) and A-Train satellite
22 observations (Christensen and Stephens, 2011; Lebsock et al., 2008) tend to confirm two responses of the
23 cloud liquid water to increasing aerosol. Under clean conditions when clouds are prone to precipitation, an
24 increase in the aerosol tends to increase cloud amount. Under non-precipitating conditions, clouds tend to
25 thin in response to increasing aerosol. Treatment of the subtlety of these responses and associated detail in
26 small-scale cloud processes is not currently feasible in GCMs, nor will it be in the foreseeable future.

27
28 Since AR4, cloud resolving model simulation has begun to stress the importance of scale interactions when
29 addressing aerosol-cloud interactions. Large model domains (order 100 km) allow mesoscale circulations to
30 develop in response to changes in the aerosol. These dynamical responses may have a significant impact on
31 cloud morphology and radiative forcing. Examples include the dramatic changes in cloud morphology
32 associated with changes in cellular structure discussed above and the cloud-free shadows that appear
33 alongside ship tracks (Wang and Feingold, 2009b). Similar examples of large-scale changes in circulation
34 associated with aerosol, and associated influence on precipitation are discussed in Section 7.6.4. These
35 underscore the imprudence of applying simplistic rules for aerosol-cloud-precipitation interactions in GCMs.

36 37 *7.4.3.4 Advances in and Insights Gained from Large-Scale Modelling Studies*

38
39 Regional models are increasingly including representation of aerosol-cloud interactions using sophisticated
40 microphysical models (Bangert et al., 2011; Seifert et al., 2012; Yang et al., 2011) and offer an important
41 middle ground between fine-scale, idealized simulations and GCMs. Some of these models are operational
42 weather forecast models that undergo regular evaluation. Yang et al. (2011) show improved simulations of
43 stratocumulus fields when aerosol-cloud interactions are introduced. Regional models are increasingly being
44 used in conjunction with satellite observations to provide the meteorological context for aerosol-cloud
45 interactions (see Section 7.4.1.4) with some (e.g., Painemal and Zuidema, 2010) suggesting that droplet
46 concentration differences, are primarily driven by synoptic scale influences rather than aerosol.

47
48 It appears increasingly likely that cloud adjustments, which almost certainly exist locally, vary from one
49 cloud regime to the next. As such it requires models to correctly represent the distribution of cloud regimes
50 evincing such effects, and knowledge of how such effects manifest themselves across these regimes. Because
51 diverse effects offer the possibility of compensating one another, at least globally, it seems possible that
52 AFaci may be less important than previously thought (Stevens and Feingold, 2009).

53 54 *7.4.4 Adjustments in Cold Clouds*

55 56 *7.4.4.1 The Physical Basis for Adjustments in Cold Clouds*

1 Mixed-phase clouds, containing both liquid water and ice particles, exist at temperatures between 0°C and –
2 38°C. At warmer temperatures ice melts rapidly, whereas at colder temperatures liquid water freezes
3 homogeneously. The formation of ice in mixed-phase clouds depends on heterogeneous freezing, initiated by
4 ice nuclei (IN; Section 7.3.3.4 and Figure 7.15), usually insoluble aerosol particles. In spite of their very low
5 concentrations (on the order of 1 l^{-1}), IN have an important influence on mixed-phase clouds. Mineral dust
6 particles have been identified as good IN but far less is known about the IN ability of other aerosol types,
7 and their preferred modes of nucleation. For example, the ice nucleating ability of black carbon particles
8 remains controversial (Hoose and Möhler, 2012). In the case of cirrus (ice-only) clouds forming via
9 homogeneous freezing at temperatures colder than –38°C, soluble matter can hinder glaciation by depressing
10 the freezing temperature of super-cooled drops (e.g., Baker and Peter, 2008; Girard et al., 2004). Hence
11 anthropogenic perturbations to the aerosol have the potential to affect glaciation, water and ice optical
12 properties, and their radiative effect.

13
14 Because the equilibrium vapour pressure with respect to ice is lower than that with respect to liquid, the
15 initiation of ice in a supercooled liquid cloud will cause vapour to diffuse rapidly toward ice particles at the
16 expense of the liquid water (Wegener-Bergeron-Findeisen process; e.g., Hudson et al., 2010; Verheggen et
17 al., 2007). This favours the depositional growth of large ice crystals, which may sediment away from the
18 water-saturated region of the atmosphere, influencing the subsequent evolution of the cloud. Hence
19 anthropogenic perturbations to the IN can influence the rate at which ice forms, which in turn may regulate
20 cloud amount (Lohmann, 2002a; Storelvmo et al., 2011; see also Section 7.2.3.2.2) and upper tropospheric
21 humidity.

22
23 Finally, the ice-phase provides enthalpy to the environment, which influences cloud dynamics, as well as
24 alternate, complex pathways for precipitation to develop (e.g., Zubler et al., 2011 and Section 7.6).

25 26 7.4.4.2 *Observations of Aerosol Effects on Arctic Ice and Mixed-Phase Stratiform Clouds*

27
28 Arctic mixed-phase clouds have received a great deal of attention since AR4, with major field programs
29 conducted in 2004 (Verlinde et al., 2007) and 2009 (Brock et al., 2011; Jacob et al., 2010; McFarquhar et al.,
30 2011), in addition to long-term monitoring at Barrow, Alaska (Shupe et al., 2008) and analysis of earlier
31 field experiments (Uttal et al., 2002). Mixed-phase Arctic clouds persist for extended periods of time (days
32 and even weeks; Zuidema et al., 2005), in spite of the inherent instability of the ice-water mix. We focus
33 here on the role of the aerosol and refer to Section 7.2.3.2.2 for a discussion of meteorological aspects. In
34 spite of their low concentrations, IN have an important influence on cloud persistence, with clouds tending to
35 glaciate and disappear rapidly when IN concentrations are relatively high and/or updraught velocities too
36 small to sustain a liquid water layer (e.g., Ovchinnikov et al., 2011). The details of the heterogeneous ice-
37 nucleation mechanism remain controversial but there is increasing evidence that ice forms in Arctic stratus
38 via the liquid phase (immersion freezing) so that the CCN population also plays an important role (de Boer et
39 al., 2011; Lance et al., 2011). If ice indeed forms via the liquid phase this represents a self-regulating
40 feedback that helps sustain the clouds: as ice forms, water is depleted, which restricts further ice formation
41 and competition for water vapour via the Wegener-Bergeron-Findeisen process.

42 43 7.4.4.3 *Advances in Process Level Understanding*

44
45 Since AR4 research on ice-microphysical processes has been very active as evidenced by activity in the
46 abovementioned field experiments. The persistence of some mixed-phase stratiform clouds has prompted
47 efforts to explain this phenomenon in a theoretical framework (Korolev and Field, 2008). Cloud persistence
48 may require a level of understanding of very detailed processes. For example, ice particle growth by vapour
49 diffusion depends strongly on particle habit (Harrington et al., 2009), the details of which may have similar
50 influence on glaciation times to the choice of different ice nucleation mechanism (Ervens et al., 2011). A
51 recent review (Morrison et al., 2012) discusses the myriad processes that create a resilient mixed-phase cloud
52 system, invoking the ideas of “buffering” seen in liquid clouds (Stevens and Feingold, 2009). Bistability has
53 also been observed in the mixed-phase Arctic cloud system; the resilient cloud state is sometimes interrupted
54 by a cloud-free state (Stramler et al., 2011), however there is much uncertainty regarding the meteorological
55 and microphysical conditions determining which of these states is preferred.

1 Significant effort has been expended on heterogeneous freezing parameterizations employed in cloud or
2 larger-scale models. Korolev (2007) developed a theoretically based parameterization of the Wegener-
3 Bergeron-Findeisen process that has lately been employed in different GCMs (Lohmann and Hoose, 2009;
4 Storelvmo et al., 2008b). Other parameterizations remain empirical (e.g., DeMott et al., 2010; Gettelman et
5 al., 2010; Hoose et al., 2008; Lohmann and Diehl, 2006; Phillips et al., 2008; Salzmann et al., 2010;
6 Storelvmo et al., 2008a), although some recent work attempts to represent the processes explicitly (Jacobson,
7 2003) or ground the development of parameterisations in concepts derived from classical nucleation theory
8 (Hoose et al., 2010b). The details of how these processes are treated have important implications for tropical
9 anvils (Fan et al., 2010).

10
11 Ice nucleation in cirrus clouds (at temperatures less than -35°C) depends crucially on the cloud updraught
12 velocity and hence the supersaturation with respect to ice. For homogeneous nucleation, the onset relative
13 humidities have been parameterized using results of parcel model simulations (e.g., Barahona and Nenes,
14 2009; Sassen and Dodd, 1988), airborne measurements in cirrus or wave clouds (Heymsfield et al., 1998;
15 Heymsfield and Miloshevich, 1995), extensions of classical homogeneous ice nucleation theory
16 (Khvorostyanov and Sassen, 1998; Khvorostyanov and Curry, 2009), and data from laboratory
17 measurements (e.g., Bertram et al., 2000; Friedman et al., 2011; Koop et al., 2000; Magee et al., 2006;
18 Mohler et al., 2003). If IN are present, then heterogeneous nucleation is the preferred freezing pathway
19 because it occurs at lower onset relative humidities (or higher onset temperatures) than homogeneous
20 nucleation. The onset relative humidities (or temperatures) for heterogeneous nucleation depend on the type
21 and size of the involved ice nuclei (Figure 7.15 and Section 7.3.3.4).

22 23 *7.4.4.4 Advances in and Insights Gained from Large-Scale Modelling Studies*

24
25 Since the AR4 mixed-phase and ice clouds have received significant attention, with effort on representation
26 of both heterogeneous (mixed-phase clouds) and homogeneous (cirrus) freezing processes. The physics of
27 cirrus clouds usually only involve ice-phase microphysical processes and are somewhat simpler than mixed-
28 phase clouds. Nevertheless, representation of aerosol-cloud interactions in mixed-phase and ice clouds is
29 considerably less advanced than that involving liquid-only clouds.

30
31 For realistic concentrations of bacteria, biological particles have been found to play a negligible role for
32 heterogeneous freezing in GCMs (Diehl and Wurzler, 2010; Hoose et al., 2010a; 2010b; Phillips et al., 2009;
33 Sesartic et al., 2011). However some investigators (Ariya et al., 2009; Sun et al., 2010) argue that biological
34 particles even in low concentrations may still be important because they can trigger secondary ice production
35 via ice multiplication. Anthropogenic changes to the biosphere could conceivably also reduce the prevalence
36 of biological IN. Our poor understanding of the climatology and lifecycle of aerosol particles that can serve
37 as IN complicates attempts to assess what constitutes an anthropogenic perturbation to the IN population, let
38 alone the effect of such a perturbation. BC can impact background cirrus by affecting ice nucleation
39 properties but its effect remains uncertain (Kärcher et al., 2007). The numerous GCM studies that have
40 evaluated AFaci for ice clouds are summarised in Section 7.5.

41 42 *7.4.5 Impact of Cosmic Rays on Aerosols and Clouds*

43
44 High solar activity leads to variations in the strength and three-dimensional structure of the heliosphere,
45 which reduces the flux of galactic cosmic rays (GCR) impinging upon the Earth's atmosphere by increasing
46 the deflection of low energy GCR. As GCR is the primary source of atmospheric ionization, it has been
47 suggested that GCR may act to amplify relative small variations in solar activity into climatologically
48 significant effects (Ney, 1959), via a hypothesised relationship between ionization and cloudiness (e.g.,
49 Dickinson, 1975; Kirkby, 2007). There have been many studies aiming to test this hypothesis since AR4,
50 which fall in two categories: i) studies that seek to establish a causal relationship between cosmic rays and
51 aerosols/clouds by looking at correlations between the two quantities on timescales of days to decades, and
52 ii) studies that test through observations or modelling one of the physical mechanisms that have been put
53 forward. We assess these two categories of studies in the next two sections.

54 55 *7.4.5.1 Correlations Between Cosmic Rays and Properties of Aerosols and Clouds*

1 Many empirical relationships have been reported between GCR or cosmogenic isotope archives and some
2 aspects of the climate system (e.g., Bond et al., 2001; Dengel et al., 2009; Ram and Stolz, 1999). The forcing
3 from changes in total solar irradiance alone does not seem to account for these observations, implying the
4 existence of an amplifying mechanism such as the hypothesized GCR-cloud link. We focus here on observed
5 relationships between GCR and aerosol and cloud properties. Such relationships have focused on decadal
6 variations in GCR induced by the 11-year solar cycle, shorter variations associated with the quasi-periodic
7 oscillation in solar activity centred on 1.68 years or sudden and large variations known as Forbush decrease
8 events. It should be noted that GCR co-vary with other solar parameters such as solar and UV irradiance,
9 which makes any attribution of cloud changes to GCR problematic (Laken et al., 2011).

10
11 Some studies have shown co-variation between GCR and low-level cloud cover using global satellite data
12 over periods of typically 5–10 years (Marsh and Svensmark, 2000; Pallé Bagó and Butler, 2000). Such
13 correlations have not proved to be robust when extending the time period under consideration (Agee et al.,
14 2012), restricting the analysis to particular cloud types (Kernthaler et al., 1999) or locations (Udelhofen and
15 Cess, 2001; Usoskin and Kovaltsov, 2008). The purported correlations have also been attributed to ENSO
16 variability (Farrar, 2000; Laken et al., 2012) and artefacts of the satellite data cannot be ruled out (Pallé,
17 2005). Statistically significant, but weak, correlations between diffuse fraction and cosmic rays have been
18 found at some locations in the UK over the 1951 to 2000 period (Harrison and Stephenson, 2006). Harrison
19 (2008) also found a unique 1.68-year periodicity in surface radiation for two different UK sites between
20 1978 and 1990, potentially indicative of a cosmic ray effect. Svensmark et al. (2009) found large global
21 reductions in the aerosol Ångström exponent from AERONET, liquid water path from SSM/I, and cloud
22 cover from MODIS and ISCCP after large Forbush decreases, but these results were not corroborated by
23 other studies who found no statistically significant links between GCR and clouds at the global scale
24 (Čalogović et al., 2010; Kristjánsson et al., 2008; Laken and Čalogović, 2011). Although some studies found
25 small but significant positive correlations between GCR and high- and mid-altitude clouds (Laken et al.,
26 2010; Rohs et al., 2010), these variations were very weak, and the results were highly sensitive to how the
27 Forbush events were selected and composited (Laken et al., 2009).

28 29 *7.4.5.2 Physical Mechanisms Linking Cosmic Rays to Cloudiness*

30
31 The most widely studied mechanism proposed to explain the possible link between GCR and cloudiness is
32 the “ion-aerosol clear air” mechanism, in which atmospheric ions produced by GCR facilitate aerosol
33 nucleation and growth ultimately impacting CCN concentrations and cloud properties (Carslaw et al., 2002;
34 Usoskin and Kovaltsov, 2008). The variability of atmospheric ionization rates due to GCR changes can be
35 considered relatively well quantified (Bazilevskaya et al., 2008), whereas resulting changes in aerosol
36 nucleation rates are very poorly known (Enghoff and Svensmark, 2008; Kazil et al., 2008). The Cosmics
37 Leaving Outdoor Droplets (CLOUD) experiment at CERN indicates that GCR-induced ionization enhances
38 water–sulphuric acid nucleation in the middle and upper troposphere, but is very unlikely to give a
39 significant contribution to nucleation taking place in the continental boundary layer (Kirkby et al., 2011).
40 Field measurements qualitatively support this view but cannot provide any firm conclusion on the role of
41 ions due to the scarcity and other limitations of free-troposphere measurements (Arnold, 2006; Mirme et al.,
42 2010), and due to difficulties in separating GCR-induced nucleation from other nucleation pathways in
43 continental boundary layers (Hirsikko et al., 2011). If strong enough, the signal from GCR-induced
44 nucleation should be detectable at the Earth’s surface because a big fraction of CCN in the global boundary
45 layer is expected to originate from nucleation taking place in the free troposphere (Merikanto et al., 2009).
46 Based on surface aerosol measurements at one site, Kulmala et al. (2010) found no connection between GCR
47 and new particle formation or any other aerosol property over a solar cycle (1996–2008). Our understanding
48 of the “ion-aerosol clear air” mechanism as a whole relies on a few model investigations that simulate GCR
49 changes over a solar cycle (Kazil et al., 2012; Pierce and Adams, 2009a; Snow-Kropla et al., 2011) or during
50 strong Forbush decreases (Bondo et al., 2010; Snow-Kropla et al., 2011). Although all model studies found a
51 detectable connection between GCR variations and either CCN changes or column aerosol properties, the
52 response appears to be too weak to cause a significant radiative effect because GCR are unable to effectively
53 raise CCN and droplet concentrations (Kazil et al., 2012).

54
55 A second pathway linking GCR to cloudiness has been proposed through the global electric circuit (GEC). A
56 small direct current is able to flow vertically between the ionosphere (maintained at approximately 250 kV
57 by thunderstorms and electrified clouds) and the Earth’s surface over fair-weather regions because of GCR-

1 induced atmospheric ionization. Charges can accumulate at the upper and lower cloud boundaries as a result
2 of the effective scavenging of ions by cloud droplets (Tinsley, 2000). This creates conductivity gradients at
3 the cloud edges (Nicoll and Harrison, 2010), and may influence droplet-droplet collision (Khain et al., 2004),
4 cloud droplet-particle collisions (Tinsley, 2000), and cloud droplet formation processes (Harrison and
5 Ambaum, 2008). These microphysical effects may potentially influence cloud properties both directly and
6 indirectly. Although Harrison and Ambaum (2010) observed a small reduction in downward LW radiation
7 which they associated with variations in surface current density, supporting observations are extremely
8 limited. Our current understanding of the relationship between cloud properties and the GEC remains very
9 low, and there is no evidence yet that associated cloud processes could be of climatic significance.

11 7.4.5.3 *Synthesis*

12 Although there is some evidence that ionization from cosmic rays may enhance aerosol nucleation in the free
13 troposphere, there is medium evidence and high agreement that the cosmic ray-ionization mechanism is too
14 weak to influence global concentrations of CCN or their change over the last century or during a solar cycle
15 in any climatically significant way. The lack of trend in the cosmic ray intensity over the last 50 years (Agee
16 et al., 2012; McCracken and Beer, 2007) provides another strong argument against the hypothesis of a major
17 contribution of cosmic rays to ongoing climate change.

20 7.5 Radiative Forcing and Adjusted Forcing by Anthropogenic Aerosols

21 In this section, aerosol forcing estimates are synthesized and updated from AR4. As depicted in Figure 7.2,
22 RF refers to the instantaneous radiative forcing either due to aerosol-radiation interactions (ari), formerly
23 known as the direct aerosol forcing, or aerosol-cloud interactions (aci), formerly known as the indirect
24 aerosol forcing. AF refers to the adjusted forcing (also referred to as radiative flux perturbation) and can be
25 estimated from Hansen-style experiments with fixed sea-surface temperatures (see Sections 7.1 and 8.1). It
26 includes rapid adjustments, such as changes to the cloud lifetime, changes in lapse-rate due to absorbing
27 aerosols and aerosol microphysical effects on mixed-phase, ice and convective clouds. AF can again be
28 caused by ari, aci or the sum of ari and aci.

29 Chapter 2 of AR4 (Forster et al., 2007) assessed RFari to be $-0.5 \pm 0.4 \text{ W m}^{-2}$ and broke this down into
30 components associated with several species. Land albedo changes associated with black carbon (BC) on
31 snow were assessed to be $+0.1 \pm 0.1 \text{ W m}^{-2}$. The RFaci was assessed to be -0.70 W m^{-2} with a -1.8 to -0.3
32 W m^{-2} uncertainty range. These uncertainty estimates were based on a combination of model results and
33 remote observations. The semi-direct effect and other aerosol indirect effects were assessed in Chapter 7
34 (Denman et al., 2007) to contribute additional uncertainty. The combined total aerosol forcing was given as
35 two distinct ranges: -0.2 to -2.3 W m^{-2} from models and a -0.1 to -1.7 W m^{-2} range from inverse estimates.

36 For consistency with Chapter 8, all quoted ranges represent a 5% to 95% uncertainty range and we evaluate
37 the forcings between 1750 and ~2010. Note that for several aerosol species (such as biomass burning) this
38 does not quite equate to the anthropogenic effect as emissions started to be influenced by humans before the
39 Industrial Revolution. Many studies estimate aerosol forcings between 1850 and present day and conversion
40 to a forcing between 1750 and present-day contributes to increase the uncertainty (Bellouin et al., 2008).

45 7.5.1 *Estimates of RF and AF from Aerosol-Radiation Interactions (RFari and AFari)*

47 7.5.1.1 *Radiative Forcing and Adjusted Forcing from All Aerosols*

48 Building on our understanding of aerosol processes and their radiative effects (Section 7.3), this section
49 assesses RFari and AFari. The subsection additionally assesses the forcings from absorbing aerosol (BC and
50 dust) on both snow and ice.

51 Observations can give useful constraints to aspects of the global RFari estimates (Section 7.3.4, Anderson et
52 al., 2005; Kahn, 2012). Estimates of RFari are either taken from global aerosol models directly (Schulz et al.,
53 2006) or based mostly on observations, but using supplemental information from models (Loeb and Su,
54 2010; Myhre, 2009). Three studies (Bellouin et al., 2008; Myhre, 2009; Zhao et al., 2008) improved satellite-
55 based RFari estimate over those quoted in AR4. The most complete analysis of the globally-averaged all-sky
56 RFari estimate over those quoted in AR4. The most complete analysis of the globally-averaged all-sky
57

1 preindustrial-to-present RF (Myhre, 2009) estimated an RFari of $-0.3 \pm 0.2 \text{ W m}^{-2}$. There is medium
2 confidence in this best estimate, but the uncertainty range is likely underestimated (Loeb and Su, 2010).

3
4 Lohmann et al. (2010) found RFari in five GCMs ranging from -0.1 to -0.4 W m^{-2} . A second phase of
5 AeroCom model results gives an RFari median of -0.39 W m^{-2} , with a model range of about -0.1 to -0.6 W
6 m^{-2} , after their forcings for 1850-2000 had been scaled by emissions to represent 1750–2010 changes
7 (Myhre et al., 2012). Figure 7.17 shows the zonal mean total RFari for AeroCom phase II models and the
8 1750–2010 RFari from all models are shown in Figure 7.19. Robust features are the maximum negative RF
9 around 10°N – 50°N , in the region with highest aerosol concentrations and a positive RF at high latitudes due
10 to the higher surface albedo there.

11
12 Remote sensing data, observations of fine-mode aerosol properties and a better knowledge of bulk aerosol
13 optical properties make the estimate of total RFari more robust than the RF for individual species (see
14 Forster et al., 2007). The best estimate of total RFari is taken to be -0.4 W m^{-2} , based on combining the
15 AeroCom median with the refined satellite-based estimates. The methodology of Loeb and Su (2010) is
16 combined with an uncertainty analysis of aerosol optical properties to give an overall assessment for the
17 RFari of -0.4 W m^{-2} , with a 5-95% range of -0.7 to -0.1 W m^{-2} . This is also consistent with the uncertainty
18 analysis of Bellouin et al. (2012) and Ma et al. (2012b). Uncertainties in the radiative transfer also contribute
19 to the overall uncertainty (Randles et al., 2012; Stier et al., 2012), while uncertainties in the vertical
20 distribution are important for absorbing aerosols (Samset et al., 2012; Section 7.3.3).

21
22 The assessed RFari of $-0.4 \pm 0.3 \text{ W m}^{-2}$ has a smaller magnitude and smaller range than in AR4. There is
23 also increased confidence in this assessment due to more robust satellite-based estimates and their better
24 agreement with models.

25 26 **[INSERT FIGURE 7.17 HERE]**

27 **Figure 7.17:** Annual zonal mean RFari (in W m^{-2}) due to all anthropogenic aerosols from the different AeroCom II
28 models. No adjustment for missing species in certain models has been applied. The multi-model mean is shown with a
29 black solid line. Adapted from Myhre et al. (2012).

30
31 AFari adds the radiative effects from rapid adjustments onto RFari. Studies have evaluated the rapid
32 adjustments separately as a semi-direct effect (see Section 7.3.5) and/or the AFari has been directly
33 evaluated. Rapid adjustments are principally caused by cloud changes. There is high confidence that the
34 local heating caused by absorbing aerosols can cause cloudiness to increase or decrease depending on their
35 conditions. However, there is low confidence in determining the sign and magnitude of the rapid adjustments
36 at the global scale as current models differ in their responses and are known to inadequately represent some
37 of the important relevant cloud processes (see Section 7.3.5). Existing estimates of AFari nevertheless rely
38 on such global models. Five GCMs were analysed for RFari and AFari in Lohmann et al. (2010). Their rapid
39 adjustments ranged from -0.3 to $+0.1 \text{ W m}^{-2}$. In a further study, Takemura and Uchida (2011) found a rapid
40 adjustment of $+0.06 \text{ W m}^{-2}$. The sensitivity analysis of Ghan et al. (2012) found a -0.1 W m^{-2} to $+0.1 \text{ W m}^{-2}$
41 range over model variants, where an improved aging of the mixing state led to small negative rapid
42 adjustment of around -0.1 W m^{-2} . Bond et al. (2012) assessed scaled RF and efficacy estimates from seven
43 earlier studies focusing on BC and found a range of rapid adjustments between -0.2 W m^{-2} and -0.01 W m^{-2} .
44 There is a potential additional rapid adjustment term from the effect of cloud drop inclusions (see Section
45 7.3.4.2). Based on the Ghan et al. (2011) and Jacobson (2012), Bond et al. (2012) estimate an additional
46 AFari of $+0.2 \text{ W m}^{-2}$, with an uncertainty range of -0.1 to $+0.7 \text{ W m}^{-2}$, however there is low evidence for
47 this effect and we do not include it in our assessment. Overall a best estimate for the rapid adjustment is
48 taken to be -0.1 W m^{-2} , with a 5% to 95% uncertainty range of -0.3 to $+0.1 \text{ W m}^{-2}$. The uncertainties are
49 added in quadrature to the estimate of RFari to give an assessment for AFari of $-0.5 \pm 0.4 \text{ W m}^{-2}$.

50 51 *7.5.1.2 Radiative Forcing by Species*

52
53 AeroComII studies have calculated aerosol distributions using 1850 and 2000 simulations with the same
54 meteorology to isolate RFari for individual aerosol types (SO_4 , BC fossil-fuel, OC fossil-fuel, biomass
55 burning or BB, SOA, NO_3). Unless otherwise noted in the text below the best estimate and 5–95% ranges for
56 individual types quoted in Figure 7.18 are solely based on the AeroComII range (Myhre et al., 2012) and the
57 estimates have been scaled by emissions to derive 1750–2010 RFari values. Note that although global

1 numbers are presented here these RF estimates all exhibit large regional variations and individual aerosol
2 species can contribute significantly to regional climate change despite rather small RF estimates (e.g., Wang
3 et al., 2010b).

4
5 For sulphate AeroComII models give a RF median and 5–95% uncertainty range of $-0.35 \pm 0.2 \text{ W m}^{-2}$, for
6 the 1850–2000 period and $-0.37 \pm 0.2 \text{ W m}^{-2}$ for the 1750–2010 period. This estimate and uncertainty range
7 are consistent with the AR4 estimate of $-0.4 \pm 0.2 \text{ W m}^{-2}$, which is retained as the best estimate for AR5.

8
9 RF from BC is evaluated in different ways in the literature. The BC RF in this report is from fossil fuel and
10 biofuel sources, while open burning sources are attached to the BB aerosol, which also includes other
11 organic species (see Section 7.3.2). The radiative effect from all anthropogenic plus natural sources of BC
12 derived using an analysis of AERONET level 2.0 (e.g., Ramanathan and Carmichael, 2008) is likely biased
13 high due to the inability of AERONET to measure small optical depths (Section 7.3.4, Bond et al., 2012).
14 Bond et al. (2012) analysed BC forcing across 18 model variants, including 8 AeroComII models, using and
15 improved AERONET scaling and accounting for model bias in the BC vertical profile. Scaling for
16 absorption AOD in the models to account for a low bias in emissions (see Section 7.3.2) typically led to 50%
17 or larger underestimates in RF, whilst BC at too high altitudes in models led to a RF overestimate of around
18 15% (Zarzycki and Bond, 2010; Section 7.3.3). Bond et al. (2012) also performed an uncertainty analysis,
19 principally of the scaling approach and forcing efficiency with each contributing roughly a $\pm 40\%$ (2 standard
20 deviation) uncertainty. RFs from Bond et al. (2012) were assessed to be $+0.17$, $+0.12$ and $+0.15 \text{ W m}^{-2}$ from
21 fossil fuel, biofuel and open burning sources, respectively. AeroComII models give a BC RFari from fossil
22 fuel and biofuel of $+0.19 \pm 0.013 \text{ W m}^{-2}$ for the 1850–2000 period and $+0.24 \pm 0.19 \text{ W m}^{-2}$ for the 1750–
23 2010 period (Myhre et al., 2012). The Bond et al. (2012) scaling is used to derive the best estimate of BC
24 RFari of $+0.3 \text{ W m}^{-2}$ and a $\pm 0.2 \text{ W m}^{-2}$ uncertainty range is adopted, based on both the AeroComII range and
25 the Bond et al. (2012) bottom-up uncertainty analysis.

26
27 Biomass burning aerosol RFari, which includes both BC and OA species, is estimated from AeroComII
28 models as $-0.01 \pm 0.08 \text{ W m}^{-2}$, and an estimate of $+0.0$ (-0.1 to $+0.1$) W m^{-2} is adopted. Note here that the
29 BB RFari is close to zero, but consists of rather strong RFari from BC and OA as shown in Figure 7.18. The
30 SOA RFari estimate is $-0.08 \pm 0.2 \text{ W m}^{-2}$ and the POM from fossil fuel estimate is $-0.05 \pm 0.04 \text{ W m}^{-2}$. For
31 OA from natural burning and for SOA the natural radiative effects can be an order of magnitude larger than
32 the RF (see Sections 7.3.2, Section 7.3.3, and O'Donnell et al., 2011) and they could thus contribute to
33 climate feedback (see Section 7.3.6).

34
35 The AeroComII RF estimate is used for nitrate aerosol, giving an RFari of $-0.13 \pm 0.1 \text{ W m}^{-2}$, but comprises
36 a relatively large 1850 to 1750 correction term. This is in good agreement with earlier estimates (e.g., Adams
37 et al., 2001; Bauer et al., 2007; Myhre et al., 2009). In the AeroComII models ammonium aerosol is included
38 within the sulphate and nitrate estimates.

39
40 Anthropogenic sources of mineral aerosols can result from changes in land use and water use or climate
41 change. Estimates of the RF from anthropogenic mineral aerosols are highly uncertain, because natural and
42 anthropogenic sources of mineral aerosols are often located close to each other (Ginoux et al., 2012a;
43 Mahowald et al., 2009). Using a compilation of observations of dust records over the 20th century with
44 model simulations, Mahowald et al. (2010) deduced a 1750–2000 change in mineral aerosol RFari including
45 both natural and anthropogenic changes of $-0.14 \pm 0.11 \text{ W m}^{-2}$. This is consistent within the AR4 estimate of
46 $-0.1 \pm 0.2 \text{ W m}^{-2}$ (Forster et al., 2007) which is retained here. Note that some of this forcing could be due to
47 feedback processes (see Section 7.3.6).

48 [INSERT FIGURE 7.18 HERE]

49 **Figure 7.18:** Mean (solid line), median (dashed line), one standard deviation (box) and full (min-max) range (whiskers)
50 for RFari (in W m^{-2}) from different aerosol types from AeroCom II models. The forcings are for the 1850 to 2000
51 period. Adapted from Myhre et al. (2012).

52 7.5.1.3 Absorbing Aerosol on Snow and Sea-Ice

53
54 Forster et al. (2007) estimated the RF for surface albedo changes from BC deposited on snow to be $+0.10 \pm$
55 0.10 W m^{-2} , with a low level of understanding, based largely on studies from Hansen and Nazarenko (2004)

1 and Jacobson (2004). Since AR4, observations of BC in snow have been conducted using several different
2 measurement techniques (e.g., Doherty et al., 2010; Forsström et al., 2009; Huang et al., 2011; Kaspari et al.,
3 2011; McConnell et al., 2007; Ming et al., 2009; Xu et al., 2009), providing data with which to constrain
4 models. Laboratory measurements have confirmed the albedo reduction due to BC in snow (Hadley and
5 Kirchstetter, 2012). The albedo effects of non-BC constituents have also been investigated but not rigorously
6 quantified. Global modelling studies since AR4 have quantified present-day radiative effects from BC on
7 snow of +0.02 to +0.08 W m⁻² (Flanner et al., 2009; Flanner et al., 2007; Hansen et al., 2007; Koch et al.,
8 2009a; Rypdal et al., 2009; Skeie et al., 2011; Wang et al., 2011c). These studies apply different BC
9 emission inventories and atmospheric aerosol representations, include forcing from different combinations of
10 terrestrial snow, sea-ice, and snow on sea-ice, and include different indirect effects such as snow grain size
11 evolution and melt-induced accumulation of impurities at the snow surface, observed on Tibetan glaciers
12 (Xu et al., 2012). The forcing operates mostly on terrestrial snow and is largest during March–May, when
13 boreal snow and ice are exposed to strong insolation (Flanner et al., 2007). All climate change studies
14 exploring this forcing find that it warms the Arctic. Estimates of the change in global mean surface
15 temperature per unit forcing are 1.7–4.5 times greater for snow and sea-ice forcing than for CO₂ forcing
16 (Bellouin and Boucher, 2010; Flanner et al., 2009; Flanner et al., 2007; Hansen and Nazarenko, 2004;
17 Hansen et al., 2005). The Koch et al. (2009a) estimate is not included in this range due to the low signal to
18 noise ratio in their study. The greater response of global-mean temperature occurs primarily because all of
19 the forcing energy is deposited directly into the cryosphere, whose evolution drives a positive albedo
20 feedback on climate. Key sources of forcing uncertainty include BC concentrations in snow and ice, BC
21 mixing state and optical properties, snow and ice coverage and patchiness, co-presence of other light-
22 absorbing particles in the snow pack, snow effective grain size and its influence on albedo perturbation, the
23 masking of snow surfaces by clouds and vegetation, and the accumulation of BC at the top of snowpack
24 caused by melting and sublimation. Bond et al. (2012) derive a 1750–2010 snow and sea-ice RF estimate of
25 +0.046 (+0.015 to +0.094) W m⁻² for BC by 1) considering forcing ranges from all relevant global studies, 2)
26 accounting for biases caused by a) modelled Arctic BC-in-snow concentrations using measurements from
27 Doherty et al. (2010), and b) exclusion of soil dust, which reduces BC forcing by approximately 20%, 3)
28 combining in quadrature individual uncertainty terms from Flanner et al. (2007) plus that originating from
29 the co-presence of dust, and 4) scaling the present-day radiative contributions from BB, biofuel, and fossil
30 fuel BC emissions according to their 1750–2010 changes. Here, we adopt an estimate of +0.04 (+0.02 to
31 +0.09) W m⁻² and note that the surface temperature change is roughly three (two to four) times more
32 responsive to this RF relative to CO₂.

33 7.5.2 Estimates of RF and AF from Aerosol-Cloud Interactions (RF_{aci} and AF_{aci})

34 RF_{aci} and AF_{aci} can be estimated in different ways. Both refer to changes in TOA radiation since pre-
35 industrial times but whether it is estimated from changes in SW radiation, net radiation or cloud radiative
36 forcing varies among publications. In this section and in Figure 7.19 we refer to estimates of the change in
37 net TOA radiation whenever possible. If that estimate is not available we use, in order of preference, the
38 change in net SW radiation, the change in net cloud forcing and last, the change in SW cloud forcing. Given
39 that the LW RF_{aci} is negligible and that RF_{aci} is limited to cloudy skies, the different ways to estimate
40 RF_{aci} are comparable. However, AF_{aci} has contributions from clear-sky conditions and LW radiation,
41 especially if aerosol effects on mixed-phase and ice clouds are considered or if the atmospheric circulation
42 responds to changes in the SW forcing.

43
44
45 Despite the consolidation in our understanding of the physical basis of the indirect forcing there still remain
46 large uncertainties in quantification, because of the difficulties in representing relevant aerosol properties,
47 clouds, and aerosol-cloud interactions in climate models as discussed in Section 7.4. Ensemble-mean
48 globally-averaged model estimates of RF_{aci} have thus remained rather constant over time (Figure 7.19) and
49 amount to roughly -1 W m⁻². This estimate is obtained from the average over all published estimates,
50 treating each of them as equal (taking one value for the best estimate per model and paper). The -1 W m⁻²
51 estimate is slightly more negative than the estimate of RF_{aci} in AR4 where the reported value of -0.7 W m⁻²
52 was obtained from putting more emphasis of the newest results at that time (Forster et al., 2007). Whereas
53 early models used offline three-dimensional sulphate fields, state-of-the art GCMs have their own aerosol
54 schemes and consider sea salt, mineral dust and carbonaceous aerosols in addition to sulphate. There does
55 not seem to be a systematic tendency for models that use a parameterization based on cloud parcel models
56 instead of empirical relationships between the aerosol mass/number concentration and the cloud droplet

1 number concentration to simulate a larger or smaller RFaci. Sensitivity studies show that RFaci is larger if
2 the background aerosol concentration is low (Chen and Penner, 2005) as this increases the cloud
3 susceptibility. As shown by Storelvmo et al. (2009) different empirical relationships that are used to bypass
4 cloud activation can cause a difference of 1.3 W m^{-2} in RFaci. RFaci depends strongly on the assumed
5 minimum droplet concentration because that determines the susceptibility of the cloud (Hoose et al., 2009).
6 Models that use autoconversion rates of cloud droplets to form rain drops which depend inversely on the
7 droplet concentration automatically simulate an increase in cloud lifetime and cloud liquid water. In small-
8 scale studies no increase in cloud lifetime is found because smaller droplets also evaporate more readily
9 (Jiang et al., 2006), a process that is not yet considered in GCMs. AFaci tends to be less negative if changes
10 to the cloud droplet size distribution (dispersion) are considered (e.g., Rotstayn and Liu, 2005) or if a
11 prognostic equation for precipitation is introduced (Posselt and Lohmann, 2009) because that shifts the
12 emphasis from autoconversion to accretion between cloud droplet and rain drops in better agreement with
13 what can be inferred from observations and large-eddy simulations (Wood, 2005).

14
15 Recent estimates of RFaci using satellite-based observations are systematically weaker (-0.4 W m^{-2} in
16 Figure 7.19) than the estimates, which were derived from GCM calculations based on parameterization of
17 physical processes or in-situ observations (Lohmann and Lesins, 2002; Quaas et al., 2008; Quaas et al.,
18 2009). The satellite-based data show lower susceptibility of cloud effective radius or droplet number
19 concentration to aerosol optical depth or number concentration than do in-situ observations or detailed cloud
20 parcel model calculations (McComiskey and Feingold, 2008), leading to the differences in the estimate of
21 RFaci. This result is at least partly due to scale-related averaging biases in satellite retrievals (McComiskey
22 and Feingold, 2012). It is also generally difficult to separate the RFaci from the rapid cloud response and
23 meteorological effects in both observations and fully coupled numerical model calculations (e.g., George and
24 Wood, 2010; Lohmann et al., 2010; Mauger and Norris, 2010; Painemal and Zuidema, 2010). Penner et al.
25 (2011) have recently questioned the applicability of present day (satellite-based) observations of cloud-
26 aerosol susceptibility to present-day minus preindustrial calculations in climate models. Their work suggests
27 that RFaci may be significantly higher than suggested by Quaas et al. (2008).

28
29 There is conflicting evidence for the importance of AFaci associated with cirrus, ranging from a statistically
30 significant impact on cirrus coverage (Hendricks et al., 2005) to a very small effect (Liu et al., 2009). Penner
31 et al. (2009) obtained a rather large (unadjusted) forcing of anthropogenic ice-forming aerosol on upper
32 tropospheric clouds; however, they ignored potential compensating effects on lower lying clouds. The
33 climate impact of anthropogenic lead-containing mineral dust particles, among the most efficient ice-forming
34 substances, has been investigated. In the extreme scenario in which 100% of ice-forming mineral dust
35 particles in cirrus clouds contained lead, up to $+0.8 \text{ W m}^{-2}$ more longwave radiation was emitted to space as
36 compared to pure mineral dust particles (Cziczo et al., 2009b). A new study based on two GCMs and
37 different ways to deduce AFaci on cirrus clouds estimates that effect to be $+0.27 \pm 0.1 \text{ W m}^{-2}$ (Gettelman et
38 al., 2012b), which partially offsets the total AFaci.

39
40 A complementary approach to GCM-based estimates of AF/RF is to infer AF/RF as a residual using the
41 observed temperature record over land, and estimates of the ocean heat uptake and the evolution of
42 greenhouse gas and solar RF (Anderson et al., 2003; Hegerl et al., 2007). These approaches, called inverse
43 estimates, normally involve models of intermediate complexity. The only inverse study that obtained RFaci
44 bracketed it to be between 0 and -1.2 W m^{-2} (Knutti et al., 2002). Both RFaci estimates constrained by
45 satellite observations and the ensemble-mean model estimate of -1 W m^{-2} fit into this range.

46
47 In conclusion, there are different rapid adjustments that contribute to the AF through changes in cloud
48 condensate, cloud cover, cloud phase or precipitation efficiency (Section 7.4). These adjustments can either
49 increase or decrease RFaci. While adjustments in both directions occur in CRMs (Section 7.4), climate
50 models do not always include the processes leading to compensation effects and positive adjustments. For
51 this reason we defer our assessment of RFaci and AFaci to Section 7.5.3.

52 **7.5.3 Estimates of AF from Combined Aerosol-Radiation and Aerosol-Cloud Interactions (AFari+aci)**

53
54
55 The first GCM estimates of AFari+aci only included aci in liquid stratiform clouds. Its mean value of -1.5 W
56 m^{-2} is only slightly less negative than the mean AFaci suggesting that ari play a secondary role once aci are
57 considered. AFari+aci is much more variable if secondary processes (+SEC) such as aci on mixed-phase or

convective clouds are additionally included in GCMs (Figure 7.19). The model average to -1.4 W m^{-2} is dominated by a few large negative estimates. Thus, the median value of -1.1 W m^{-2} may be more meaningful. The spread between the GCMs studies that also include aci in mixed-phase clouds (+MPC) can be explained by differences in the frequency of glaciation of supercooled clouds. If more IN are available in present-day conditions, supercooled clouds glaciate more readily and precipitate (see Section 7.4.4.1). In these cases an additional cooling stems from more LW radiation being emitted to space. If on the contrary more IN become coated with soluble material and become less efficient, supercooled clouds remain longer in the atmosphere, which enhances AFaci of liquid clouds but leads to a small positive LW effect that slightly compensates for the SW cooling.

Regional and global models systematically misrepresent the distribution of clouds, and cloud processes, especially those for shallow maritime clouds. One persistent shortcoming of global models is the tendency to only treat aerosol-cloud interactions in terms of stratiform but not convective clouds. In fact most GCMs neglect the radiative effect of convective clouds entirely. Recent efforts to consistently address both types of cloud representations represent a significant advance in large scale-modelling (Jacobson, 2003; Lohmann, 2008; Suzuki et al., 2008). Nonetheless our understanding of aerosol-cloud interactions is incomplete, and what is well-understood, is incompletely represented in large-scale models. For these reasons, and because lifetime effects depend critically on the interplay of uncertainly parameterized physical processes, global model-based estimates of lifetime effects remain uncertain. This is reflected in the large spread of climate model estimates of AFari+aci that also consider aci in convective clouds (+CNV). One of the most reliable estimate in the +CNV category may be the estimate of -1.1 W m^{-2} to the farthest right because it is obtained from a multi-scale modelling framework approach (Wang et al., 2011b), which resolves convection. Using satellite retrievals to estimate AFari+aci leads to the least negative average AFari+aci of -0.7 W m^{-2} (Figure 7.19).

An inverse estimate that is obtained purely from energy balance arguments bounds AFari+aci (including unknown residuals that are assumed to be small) since 1950 to be between -0.7 to -1.5 W m^{-2} (Murphy et al., 2009), which is the same range as obtained from all inverse estimates of AFari+aci (Figure 7.19).

We produce a best estimate and 5–95% uncertainty range for AFari+aci in the following way. The global CMIP5 models and inverse estimates are grouped together and a bootstrapping method is used to estimate a mean and standard deviation of -0.98 and 0.32 W m^{-2} . Processing the satellite-based estimates in the same way leads to a mean and standard deviation of -0.73 and 0.30 W m^{-2} , respectively, with the caveat that the method applied to a small sample may underestimate the variance. We combine these two estimates into a best estimate and a 5 to 95% range for RFari+aci of -0.9 (-1.5 to -0.3) W m^{-2} , accounting for a small 1850 to 1750 correction term. This estimate is significantly lower than the AR4 estimate but is consistent with several new lines of evidence. Firstly the best estimate is close to the estimate of -1.1 W m^{-2} from the multi-scale modelling framework approach (Wang et al., 2011b). Secondly climate model studies that include missing processes such as aci on mixed-phase clouds or by putting more emphasis on accretion instead of autoconversion by treating rain as a prognostic variable arrive at smaller AFari+aci estimates than those which ignore them. Thirdly those AFari+aci studies taking satellite data into account, and therefore are observationally based, yield the least negative forcing. Fourthly the average AFari+aci from the CMIP5/ACCMIP models also amounts to about -1 W m^{-2} (see Table 7.5).

The AFaci is estimated as the residual between AFari+aci and AFari. We further assume that AFari and AFaci are additive and uncertainties are independent, which yields to our assessment of AFaci of -0.4 (-0.9 to 0) W m^{-2} . Models indicate that RFaci is less than AFaci, which implies an estimate of -0.3 (-0.7 to 0) W m^{-2} , again less negative than the AR4 estimate for the same reasons as given above.

[INSERT FIGURE 7.19 HERE]

Figure 7.19: Upper panel: GCM, satellite and inverse estimates of RFari, RFaci, AFaci and AFari+aci. Each symbol represents the best estimate per model and paper as detailed in Table 7.4. The RFaci studies are divided into those from GCMs published prior to TAR, AR4 and AR5, those including satellite data (SAT) and the inverse estimate (INV). AFaci and AFari+aci studies from GCMs on liquid stratiform clouds are also divided into those published prior to AR4 and AR5 and from the CMIP5/ACCMIP models. GCM estimates that include adjustments beyond aci in liquid stratiform clouds are marked +MPC when including aci in mixed-phase clouds and are marked +CNV when including aci in convective clouds. For RFaci from inverse estimates the range instead of the best estimate is given because it is only one study.

Lower panel: Box whisker plots of GCM, satellite and inverse estimates of RFari, RFaci, AFaci and AFari+aci. They are grouped into RFari from CMIP5/ACCMIP GCMs, RFaci from GCMs (TAR, AR4, AR5) and satellites (SAT), AFaci from GCMs, AFari+aci from GCMs taking aci only on liquid stratiform clouds (AR4+AR5) and including secondary processes (aci on mixed-phase or convective clouds) into account (+SEC), AFari+aci studies from the CMIP5 models, satellites (SAT) and inverse estimates of AFari+aci. Displayed are the averages (cross sign), median values (middle line), 33% and 67% percentiles (box boundaries) and 5% and 95% percentiles (ends of vertical lines) except for the inverse estimates, which is an expert assessment of the combined estimate of multiple inverse estimates. References for the individual estimates are provided in Table 7.4.

[INSERT TABLE 7.4 HERE]

Table 7.4: List of references for each category of estimates displayed in Figure 7.19.

[INSERT TABLE 7.5 HERE]

Table 7.5: Estimates of aerosol AF (in $W m^{-2}$) in some of the CMIP5 models. The AF are estimated from fixed-SST experiments using atmosphere-only version of the models listed. Different models include different aerosol effects.

7.6 Links to Precipitation

7.6.1 Introduction

Precipitation is limited by the availability of moisture and energy. Energy, which is required to evaporate water from the surface, is provided by the differential heating, through radiative processes, of the surface relative to the atmosphere. Evaporation cools the surface and, by making water available for condensation (which returns to the surface as precipitation), warms the atmosphere by an equal amount (Figure 7.20; Trenberth et al., 2009). The oceans provide a huge reservoir of moisture, but without the energetic imbalances set by radiative processes, there would be no driving force for the hydrological cycle. These basic constraints are most powerful when applied globally, but also help define how the structure of atmospheric circulations moderates their influence on regional scales (Neelin and Held, 1987; Raymond et al., 2009).

[INSERT FIGURE 7.20 HERE]

Figure 7.20: Illustration of major drivers affecting precipitation. Radiative drivers cool the atmosphere, warm the surface, and thereby provide the energy for evaporation and condensation/precipitation. Circulations organize and distribute precipitation. The ability of changes in the position or strength of circulation features to change the distribution of precipitation is referred to as a dynamic effect, the ability of warmer circulations to transport more water vapour and thereby change the amount of precipitation is referred to as a thermodynamic effect. The immediate effect of warming, greenhouse gases, clouds and aerosols on precipitation is indicated by blue or red if their change over the 20th century is thought to have changed a precipitation driver in a way that will increase, respectively decrease, precipitation. Grey indicates that changes are unknown or have multiple effects of different sign.

Precipitation fields simulated by the current generation of climate models still exhibit substantial biases (Liepert and Previdi, 2012; Stephens et al., 2010; Chapter 9). Nonetheless, it is possible to identify robust features in simulated response of precipitation to changes in precipitation drivers. In almost every case the Clausius-Clapeyron relation, which governs water phase changes, underlies these robust features.

7.6.2 The Effects of Global Warming on Large Scale Precipitation Trends

Atmospheric specific humidity is strongly expected to increase by about $7.5\% K^{-1}$ in the column average (see Section 7.2.4.1). Precipitation however increases much less rapidly with temperature because the net evaporation is energetically limited. This limitation arises because the spectroscopic properties of water vapour demand that the fractional increase in its ability to absorb (and emit) infrared radiation must increase less rapidly than the fractional increase in concentration (Stephens and Hu, 2010).

Globally-averaged precipitation increases with global mean surface temperature at about $2\% K^{-1}$ (Held and Soden (2006) and Richter and Xie (2008); for the CMIP3 models analysed by Andrews et al. (2009) the response is $1.4\text{--}3.4\% K^{-1}$), but not uniformly, as atmospheric circulations tend to increase the precipitation in the wettest latitudes, and dry latitudes may even see a precipitation decrease (Allen and Ingram, 2002; Held and Soden, 2006). This tendency of wet regions to get wetter and dry regions to get drier is moderated by a

1 weakening in the tropical overturning circulation (Figure 7.20). On smaller scales the signal is less clear due
2 to model-specific regional circulation shifts (Neelin et al., 2006), but there is some evidence that the sub-
3 tropical dry zones are expanding, both as a result of the tropical convergence zones narrowing (Chou et al.,
4 2009; Neelin et al., 2006), and the storm tracks moving poleward (Allen et al., 2012) and strengthening
5 (O’Gorman and Schneider, 2008).

6
7 The “wet-get-wetter” and “dry-get-drier” response that is evident at large scales over oceans, can be
8 understood as a simple consequence of the sharp increase in atmospheric water vapour combined with the
9 result that changes in atmospheric circulations are significantly smaller in a relative sense (Held and Soden,
10 2006). Wet regions are wet because they import moisture from dry regions, increasingly so with warmer
11 temperatures. This thermodynamic effect is evident in trends in of 20th century precipitation (Allan and
12 Soden, 2007; Zhang et al., 2007b; see Section 2.3.1), and has withstood additional analysis and scrutiny
13 since AR4 (Chou et al., 2009; Muller and O’Gorman, 2011; Seager et al., 2010), increasing our confidence in
14 the effect.

15
16 The wet-get-wetter response is mitigated by the anticipated reduction in the overturning rate of the
17 atmosphere (which can be understood partly as a consequence of the change in the dry static stability of the
18 atmosphere with warming, Figure 7.20, see also Section 7.2). This argument is less robust in dry regions
19 because of the opposing effects of increased horizontal vapour divergence by the circulation and gains from
20 local surface evaporation, primarily over the ocean (Bony et al., 2012; Chou et al., 2009), but can still be
21 used to understand the GCM land responses to some extent (Muller and O’Gorman, 2011). The non-uniform
22 nature of surface warming (e.g., land areas warm more and differently than neighbouring oceans, Joshi et al.,
23 2008) induces regional circulation shifts that affect precipitation trends. Low understanding of soil-moisture-
24 precipitation feedbacks, whose sign remains uncertain, also complicates local responses to warming
25 (Hohenegger et al., 2009). In summary, the effect of warming on precipitation on the scale of individual
26 catchments remains highly uncertain, even though some broad-scale responses are robust and relatively well
27 understood.

28 29 **7.6.3 Radiative Forcing of the Hydrological Cycle**

30
31 As much as half of the total change in precipitation can be related directly to the radiative effects of the CO₂,
32 irrespective of whether it causes any surface warming (Andrews et al., 2009; Bony et al., 2012). These
33 hydrological adjustments to radiative forcing are rapid, and can be linked to rapid adjustments of clouds
34 (Wyant et al., 2012; also Section 7.2).

35 36 **[INSERT FIGURE 7.21 HERE]**

37 **Figure 7.21:** Illustration of the response of the large-scale overturning to increasing CO₂ concentrations (adapted from
38 Bony et al., 2012). Approximately half of the response is evident before any warming is felt, but additional warming
39 continues to slow down the circulation and adds linearly to the rapid adjustment. The rapid adjustment is different over
40 land and ocean, with the increase in CO₂ initially causing an intensification of the circulation over land. The robustness
41 of the result is illustrated by the common behaviour of 15 CMIP5 models, irrespective of the details of their
42 configuration.

43
44 In the absence of a compensating temperature change, an increase in any well-mixed greenhouse gas reduces
45 the net radiative cooling rate of the atmosphere. This reduces the rainfall rate and the strength of the
46 overturning circulation. The dynamic effects are similar to those that result from the effect of atmospheric
47 warming on the lapse rate, which also reduces the strength of the overturning circulation (e.g., Section 7.6.2)
48 and are robustly evident over a wide range of models and model configurations (Bony et al., 2012; see also
49 Figure 7.21). These circulation changes are more pronounced over the ocean, because asymmetries in the
50 land-sea response to changing concentrations of greenhouse gases (Joshi et al., 2008) amplify the maritime
51 and dampen or even reverse the terrestrial signal (Bony et al., 2012; Wyant et al., 2012).

52
53 The response of the hydrological cycle to the partitioning of radiation between the surface and atmosphere
54 partly explains why forcings having the same AF do not produce the same precipitation responses. Increased
55 solar forcing does not strongly affect the net atmospheric cooling, and thus the hydrological cycle feels only
56 the associated warming with little rapid adjustment (Andrews et al., 2009), despite a small atmospheric
57 absorption of additional solar radiation (Takahashi, 2009). As a result, solar radiation management schemes

1 that maintain a constant surface temperature will lead to a reduction in globally averaged precipitation and
2 potentially substantially different regional distributions (Schmidt et al., 2012b; Section 7.7).

3
4 Changes in cloud radiative effects, and the radiative forcing from aerosol particles are also effective in
5 changing the profile of radiative heating, and thus influence the hydrological cycle. Cloud effects can be
6 large, but have been less studied than those of aerosols. To the extent that aerosol particles only scatter
7 radiation, their global effect is expected to be analogous to that associated with changes in the solar
8 irradiance. Absorbing aerosols act more like CO₂ however, as they heat and stabilize the troposphere thereby
9 reducing precipitation in a way that partially offsets their overall warming effect (Andrews et al., 2009). The
10 impact of non-well-mixed precipitation drivers such as clouds, aerosols and tropospheric ozone can depend
11 sensitively on their height and geographic distribution (Allen et al., 2012; Ming et al., 2010; Shindell et al.,
12 2012b), because of the impact of spatially non-uniform heating on atmospheric circulations of all scales, and
13 associated adjustments of cloud cover which can further alter the circulations (Section 7.2).

14 15 **7.6.4 Aerosol-Cloud Interactions**

16
17 In contrast to aerosol-radiation interactions, aerosol-cloud interactions do not directly influence the radiant
18 streams of energy that drive the climate system, and thus are not expected to have any influence on the
19 global mean precipitation rate. But because such interactions affect the microphysical structure of clouds
20 they may influence how precipitation is distributed.

21
22 There is a growing body of literature suggesting that aerosol-cloud interactions affect the dynamics of
23 individual convective clouds and storms, but in a variety of different ways that may be situation dependent,
24 and remain poorly understood. An increase in cloud condensation nuclei may affect the cloud microphysical
25 development thereby changing where and how much ice forms in convective clouds (Khain et al., 2005;
26 Rosenfeld and Woodley, 2001), or how much cloud water is available for detrainment and evaporation after
27 shallow convection (Stevens and Seifert, 2008). These changes tend to affect the vertical distribution of
28 latent heating in ways that would support the invigoration of convection (Tao et al., 2012). Some support for
29 these ideas is provided by observations (Devasthale et al., 2005; Koren et al., 2010; Koren et al., 2008), but
30 the interpretation of the observational evidence is difficult for reasons discussed extensively in Section
31 7.4.1.2. Modelling studies suggest that aerosol-cloud interactions accompanying increased aerosol loading
32 may either invigorate or suppress storms, depending on environmental factors such as vertical wind shear
33 (Fan et al., 2009). For the more studied cases in which invigoration is present, the degree of invigoration
34 depends sensitively on a number of factors, including the representation of cloud and aerosol microphysics
35 (Ekman et al., 2011), as well as the types of clouds and storms being simulated (Khain, 2009; Seifert and
36 Beheng, 2006). Studies that have looked at the effect of aerosol-cloud interactions in more of a climate
37 context, wherein convection is allowed to equilibrate with its forcing, find that invigoration effects are minor
38 (van den Heever et al., 2011) or negligible (Morrison and Grabowski, 2011), suggesting that convective
39 intensity is also limited by energetic constraints (e.g., Chou et al., 2009; Raymond et al., 2009).

40
41 Because precipitation development in clouds is a time dependent process, which proceeds at rates that are
42 partly determined by the cloud microphysical structure (Seifert and Zängl, 2010), aerosol-cloud interactions
43 may lead to shifts in topographic precipitation to the leeward side of mountains when precipitation is
44 suppressed, or to the windward side in cases when it is more readily initiated. Orographic clouds show a
45 reduction in the annual precipitation over topographical barriers downwind of major urban areas in some
46 studies (Givati and Rosenfeld, 2004; Jirak and Cotton, 2006) but not in others (Halfon et al., 2009). Even in
47 cases where effects are reported, the interpretation of the data remains controversial (Alpert et al., 2008;
48 Levin and Cotton, 2009).

49
50 The idea that precipitation can respond in a variety of different, often compensating, ways to a microphysical
51 perturbation from the ambient aerosol (as also discussed in Section 7.4) receives support from studies using
52 high-resolution numerical weather prediction methods. Although large aerosol perturbations may affect the
53 day-to-day predictions of summer season rainfall over Germany, no systematic effect of aerosol-cloud
54 interactions on precipitation is evident (Seifert et al., 2012). Likewise, although weekly cycles in aerosol
55 properties are readily apparent, a robust effect on precipitation or storm intensity has proven difficult to
56 detect (Barnet et al., 2009; Bäumer et al., 2008; Stjern, 2011; Tuttle and Carbone, 2011; Yuter et al., 2012).

7.6.5 *The Physical Basis for Changes in Precipitation Extremes*

The physical basis for an effect of aerosol-cloud interactions on precipitation extremes was discussed in the previous section. Here we briefly discuss process understanding of the effect of warming on precipitation extremes; observed trends supporting these conclusions are presented in Section 2.7.

Precipitation within individual storms is expected to increase with the available moisture content in the atmosphere rather than with the global precipitation (Allen and Ingram, 2002; Held and Soden, 2006). Global models provide robust support for these ideas, particularly in the extra-tropics (O’Gorman and Schneider, 2009), although the increase in the surface humidity, where the air is warmer, proves to be a better indicator of changes in extremes than changes in column humidity. When these effects are combined with a weakening of the overturning mass circulation, individual storms are expected to become more intense. On this issue global model results are less robust (Kharin et al., 2007; O’Gorman and Schneider, 2009). However, experiments designed to mimic the conditions of the tropical atmosphere under climate change using large-eddy simulation (Romps, 2011) and cloud-resolving models (Muller et al., 2011) provide support for the thermodynamic scaling based on surface saturation humidity, for which precipitation extremes increase by 7–8% K⁻¹.

Inferences from observed variations in particular regions suggest a dependence on temperature that might be twice as large as the thermodynamic arguments predict (Lenderink et al., 2011; Lenderink and Van Meijgaard, 2008) and may be different at short time scales (e.g., an hour or less) than at daily or storm-duration time scales (Haerter et al., 2010; Hardwick Jones et al., 2010). It is possible that in a warmer climate individual storms are deeper and drive more convergence (Chou et al., 2009; Romps, 2011; Sugiyama et al., 2010); however, a problem with regional observational approaches to this question is that the atmosphere is typically more convectively unstable during regionally hotter weather, exaggerating the effect (e.g., Haerter and Berg, 2009) relative to that in a globally warmer climate where convective instability should not differ much from today (Section 7.2.4.2). From this combination of evidence we conclude that while it is very likely that precipitation extremes will increase with warming, by how much remains uncertain and may vary with time scale.

Following the AR4 studies have continued to show that extremes in precipitation are associated with the coincidence of particular weather patterns (e.g., Lavers et al., 2011). We currently lack an adequate basic understanding of what controls the return time and persistence of such rare events.

7.7 **Solar Radiation Management and Related Techniques**

7.7.1 *Introduction*

Geoengineering is the deliberate large-scale intervention in the Earth system to counter undesirable impacts of climate change on the planet (e.g., Keith, 2000; Royal Society, 2009). One class of proposed geoengineering methods is Solar Radiation Management (SRM), which aims to counter the warming associated with increasing greenhouse gas concentrations by reducing the amount of solar energy absorbed by the climate system. A related technique seeks to deliberately decrease the greenhouse effect in the climate system by altering high-level cloudiness. Another class of geoengineering methods, known as Carbon Dioxide Reduction (CDR), is discussed in Chapter 6. The potential impacts of SRM are discussed in Section 19.5.4 of the WGII AR5. Some of the issues relating to cost, implementation, governance, ethics, economics, laws, and politics are assessed in Chapter 6 of the WGIII contribution to AR5. Here we assess proposed geoengineering methods that have the potential to influence components of the energy budget by at least a few tenths of a W m⁻² in the global mean without presuming their technological feasibility. It should be noted that no technology for SRM has been fully developed and can be considered ready for large-scale deployment. The physical underpinnings of specific methods are discussed in Section 7.7.2 followed by an assessment of their impact on climate in Section 7.7.3.

7.7.2 *Assessment of Proposed SRM Methods*

7.7.2.1 *Stratospheric Aerosols*

1 Observations of the aftermaths of major volcanic eruptions such as Pinatubo (June, 1991) demonstrate that
2 an increase in the amount of stratospheric aerosols cools the planet. Some proposed SRM techniques
3 increase stratospheric aerosol in a similar way to explosive volcanic eruptions (Budyko, 1974; Crutzen,
4 2006). Most stratospheric aerosol SRM research to date has explored the possibility of forming sulphuric
5 acid aerosols by injecting sulphur containing gases into the stratosphere, Other particles composed of black
6 carbon (Crutzen, 2006; Keith, 2010; Kravitz et al., 2012b) or metal oxides (Keith, 2010) have been
7 suggested with the potential for larger RFs, but they are not assessed further here because the literature on
8 them is too limited.

9
10 The RF from stratospheric aerosols will depend on the injection strategy: chemical species (gaseous SO₂,
11 H₂SO₄ or sprayed aerosols), location(s), rate, and frequency of injection. The injection strategy effects
12 particle size - an important factor influencing the mass scattering efficiency, and sedimentation rate. These
13 properties are important to SRM, changing the reflectivity of the aerosol and its loss rate. Particle size may
14 also have other physiochemical effects (some mentioned below). The eventual aerosol size distribution is
15 controlled by the balance between new particle formation, deposition of vapour on pre-existing particles,
16 evaporation of particles, coagulation, and sedimentation that changes as particles grow larger. Models that
17 include more comprehensive representations of aerosol microphysics (English et al., 2012; Heckendorn et
18 al., 2009; Pierce et al., 2010) found smaller aerosol burdens, larger particles and weaker RF than the earlier
19 studies that assumed a fixed particle size (e.g., Rasch et al., 2008a). This is because coagulation and
20 condensation of sulphuric acid on pre-existing particles both contribute to increase particle size when the
21 injection rate is increased. Pierce et al. (2010) suggested that the injection of H₂SO₄, a condensable vapour,
22 from an aircraft could produce larger forcings with lower sulfur loadings than SO₂ injection. Recent
23 modelling studies agree that emissions of sulfate aerosol precursors of 10 MtS yr⁻¹ or greater would be
24 needed to achieve forcing stronger than -4 W m⁻² (estimated as the mean of experiments reported by
25 Heckendorn et al. (2009), Niemeier et al. (2011) and Pierce et al. (2010)).

26
27 A variety of potential side effects have been identified. Tilmes et al. (2009; 2008) found that SRM by
28 stratospheric aerosols would lead to an increase of chemical ozone loss for the Arctic, a delay of the recovery
29 of the Antarctic ozone hole by 30–70 years, changes in ozone abundance (column ozone depletion of 3–10%
30 in polar latitudes and increases of 3–5% in the tropics) and discernable shifts in tropopause altitude (lifting
31 by 1 km) from radiative heating associated with the ozone changes. The change in ozone could increase UV
32 radiation reaching the surface, although attenuation by the aerosols may partially compensate (Vogelmann et
33 al., 1992). A decrease in direct radiation and increase in diffuse radiation reaching the Earth's surface could
34 increase photosynthesis in terrestrial ecosystems (Mercado et al., 2009; see Chapter 6) and impact some solar
35 energy technologies (see WGII AR5, Section 19.5.4). Kravitz et al. (2009) explored rain acidification from
36 stratospheric geoengineering and concluded that acid deposition rates would be orders of magnitude lower
37 than those that impact most ecosystems. These impacts remain poorly quantified and there may be additional
38 ones, for instance on tropospheric photochemistry and cirrus cloud microphysics.

39 40 7.7.2.2 *Cloud Brightening*

41
42 Boundary layer clouds act to cool the planet, and relatively small changes in cloud albedo or extent can have
43 profound effects on the Earth's radiation budget (e.g., Slingo, 1990). Examples of cloud brightening include
44 shiptracks produced in marine stratocumulus clouds by emissions of particles from ships and changes in
45 trade cumulus cloud properties produced by a relatively weak but continuous volcanic eruption of SO₂ (Yuan
46 et al., 2011), as discussed in Section 7.4.3.2. Latham (1990) suggested that it might be possible to
47 deliberately increase cloud albedo by introducing additional sea salt particles into the marine boundary layer,
48 to act as cloud condensation nuclei and “brighten” clouds through the aerosol-cloud interaction first
49 described by Twomey (1974) and discussed in Section 7.4.

50
51 Changes in cloud morphology (e.g., from open to closed cells) or switching from low to high liquid water
52 content clouds have the potential to create large AF locally. Marine stratocumulus clouds (with relatively
53 weak precipitation) are thought to be an optimal cloud type for brightening because of their relatively low
54 values of droplet concentration and the longer lifetime of sea-salt particles in non-precipitating
55 environments. However these clouds occupy a relatively small fraction of the planet and large local RF (30
56 to 100 W m⁻²) would be required to produce a globally-averaged RF of the order of -1 to -5 W m⁻².

1 Latham et al. (2008), Jones et al. (2009) and Rasch et al. (2009) explored the impact of changing droplet
2 concentration without an explicit examination of the complex interactions that govern the way aerosols affect
3 clouds. Their studies show that about -1 W m^{-2} of AF could be produced by increasing droplet concentration
4 over about 5% of the ocean surface area in marine stratocumulus regions. Forcing as strong as -4 W m^{-2} was
5 produced by increasing droplet concentration over 75% of the ocean surface, although it is not clear whether
6 such increases are achievable. There is less agreement among models when a more complete treatment of
7 aerosol-cloud interactions is considered. Korhonen et al. (2010b) concluded that increases in droplet
8 concentration would be much smaller than previous studies had assumed. In contrast, Partanen et al. (2012)
9 used the same seeding mechanism in a different model to find much larger increases in droplet
10 concentration, with an AF as strong as -5 W m^{-2} . Wang et al. (2011a) used very high resolution simulations
11 to demonstrate that cloud brightening is sensitive to cloud-dynamical feedbacks that are missing in current
12 global models. They also explored aerosol injection strategies and concluded that rates initially proposed for
13 cloud seeding would be insufficient to produce droplet concentration values needed for large-scale cloud
14 brightening. These studies highlight the limitations of our understanding of aerosol-cloud interactions, and
15 the resulting uncertainty in estimates of the AF that can be produced with cloud brightening SRM.

16
17 Changes in clouds due to aerosols are often swamped by meteorological variability (Wood et al., 2011b)
18 making detection of aerosol effects on clouds extremely difficult. Total AF from anthropogenic aerosols is
19 assessed to be -0.9 (-1.5 to -0.3) W m^{-2} in Section 7.5, that is at least 5 times smaller than the AF that
20 climate models assume is achievable with SRM.

21
22 While aerosol seeding over oceans was originally proposed to brighten clouds, Hill and Ming (2012) and
23 Partanen et al. (2012) found that aerosol-radiation interactions (the aerosol direct effect) could also be
24 important, adding 30 to 50% to the AF from aerosol-cloud interactions, sometimes dominating the forcing in
25 certain regions.

26 27 7.7.2.3 Surface Albedo Changes

28
29 Planetary albedo might also be increased by engineering local changes to the albedo of urban areas,
30 croplands, grasslands, deserts and the ocean surfaces.

31
32 Hamwey (2007) estimated the potential RF from whitening roofs and pavements at -0.17 W m^{-2} and
33 calculations by Akbari et al. (2012) suggest a global-mean equilibrium cooling between 0.01 to 0.07°C .
34 However, more detailed radiative transfer calculations (Lenton and Vaughan, 2009; Oleson et al., 2010),
35 overestimation of the surface area of the built environment that would be available for whitening, and
36 possible atmospheric feedbacks (Jacobson and Ten Hoeve, 2012) suggest that these are somewhat
37 overestimated.

38
39 The surface albedo of the world's grassland and/or cropland could be modified by replacing native species
40 with other natural or bioengineered species (Hamwey, 2007). A 25% increase in grassland albedo could give
41 an RF of -0.5 W m^{-2} (Lenton and Vaughan, 2009), with the maximum effect in summer mid-latitudes.
42 (Doughty et al., 2011; Ridgwell et al., 2009). Ridgwell et al. (2009) estimated a global-mean surface cooling
43 of 0.11°C for a $+0.04$ increase in cropland albedo. Potential effects in low-latitude regions are a reduction in
44 soil moisture, cloud cover and precipitation. The possibility for increasing crop and grassland albedo across a
45 wide variety of species remains largely non-quantified. Irvine et al. (2011) tested the impact of increasing
46 desert albedo up to 0.80 in a single model. This cooled surface temperature by -1.1°C (versus -0.22°C and $-$
47 0.11°C for their largest crop and urban albedo change).

48
49 The low albedo and large extent of ocean surfaces mean that only a small increase in albedo could be
50 sufficient to offset several W m^{-2} of RF by greenhouse gases. Methods have been proposed (Evans et al.,
51 2010; Seitz, 2011) to increase the fraction of the oceans covered with foam which is known to be very
52 reflective (Whitlock et al., 1982). Neither the extent of foam generation and persistence required for a
53 significant climate impact, nor the impact of artificial foam on the world's ocean (including ocean biology,
54 air-sea fluxes of latent heat, sensible heat, trace gases, aerosols, ocean circulation, and surface emissivity)
55 have been assessed quantitatively in the peer-reviewed literature.

7.7.2.4 Cirrus Thinning

Thin high cirrus clouds are relatively transparent to incoming sunlight, but absorb some of the infrared energy leaving the Earth's surface. Their overall radiative effect is to warm the climate (see Section 7.2.1.2). Reducing the coverage or longwave opacity of these clouds would therefore produce a negative AF. Cirrus cloud coverage and optical thickness are affected by ice crystal fall speeds, which depend on ice crystal size. Mitchell and Finnegan (2009) suggested seeding the upper troposphere with efficient IN that would trigger ice nucleation at higher temperatures than in the unperturbed atmosphere. The engineered ice crystals would grow preferentially and sediment out, resulting in thinner cirrus and reducing the water vapour available for normal particle formation. This strategy depends upon details of the cirrus nucleation processes that remain poorly understood (Section 7.4.4), but climate model simulations suggest that small changes in ice nucleation and the subsequent competition of ice particles for water vapour responses could produce responses in AF of order -5 W m^{-2} (e.g., Lohmann et al., 2008).

7.7.3 Climate Response to SRM Methods

The many ways that SRM can be implemented make quantitative model evaluation and intercomparison difficult, motivating idealised experiments to explore climate model responses. One such experiment consists of an artificial reduction of the total solar irradiance, which approximates the radiative impact of space mirrors (Early, 1989), but represents the radiative impact of other SRM methods less accurately. This results in a global cooling. Balancing the RF from a CO_2 increase with a reduction in the total solar irradiance results in a residual surface temperature change that is generally positive at mid- and high-latitudes, especially over continents, and generally negative in the Tropics (Bala et al., 2008; Kravitz et al., 2012a; Lunt et al., 2008; Schmidt et al., 2012b; Figures 7.22a-b and 7.23a-d). This is a robust prediction of climate models, and can be understood in terms of the difference between the relatively uniform longwave forcing (associated with greenhouse gases) and the latitudinally and seasonally varying shortwave forcing (from SRM). The residual changes in surface temperature are nevertheless much smaller than the warming due to the CO_2 RF alone. These features can be seen in Figure 7.23a-d showing results from the recent Geoengineering Model Intercomparison Project (Kravitz et al., 2011). The substantial warming from $4\times\text{CO}_2$ AF at high latitudes (4°C – 18°C) is reduced substantially through idealized SRM leaving residual changes, particularly a warming of 0°C – 3°C near the winter pole. While increasing CO_2 concentrations lead to a positive RF that warms the entire troposphere, SRM produces a negative RF that tends to cool the surface. The combination of RF used to balance the globally-averaged surface temperature increases atmospheric stability, reducing the evaporative flux, leading to a reduction in global precipitation rate as seen in Figures 7.22c-d and Figures 7.23e-g (Andrews et al., 2010; Bala et al., 2008; Schmidt et al., 2012b) and discussed in Section 7.6 Other factors can also modulate the climate response in evaporation, soil moisture and precipitation, such as the enhanced closure of stomata with increased CO_2 concentrations. Although models agree on the globally and annually averaged impact on precipitation, there is less agreement in the spatial pattern of the changes but some residual patterns in the idealized experiments are quite robust, for example the $\sim 5\%$ reduction in precipitation over SE Asia and the Pacific Warm Pool in JJA. Precipitation changes are substantially more varied between model simulations using more complex strategies for SRM as described below.

[INSERT FIGURE 7.22 HERE]

Figure 7.22: Zonally- and annually-averaged change in surface air temperature ($^\circ\text{C}$) for (a) an abrupt $4 \times \text{CO}_2$ experiment and (b) the GeoMIP G1 experiment for 11 coupled atmosphere-ocean general circulation models. (c) and (d) Same as (a) and (b) but for the change in precipitation (mm day^{-1}). In the GeoMIP G1 experiment (Kravitz et al., 2011) an abrupt fourfold increase in CO_2 concentration is balanced by a reduction in total solar irradiance to produce a top of atmosphere flux imbalance of less than $\pm 0.1 \text{ W m}^{-2}$ during the first 10 years of the simulation. All changes are relative to the pre-industrial control experiment and averaged over years 21–50.

Idealized experiments have also been conducted in which solar radiation is reduced only over the ocean (e.g., to approximate the effects of marine cloud seeding or increased sea foam). Other studies have explored albedo changes over land (e.g., to represent a surface albedo increase through plant albedo or brightening desert regions) and over particular latitudinal bands (Caldeira and Wood, 2008) or regions (Irvine et al., 2010; Irvine et al., 2011). Bala et al. (2010) found that although global-mean precipitation might decrease, precipitation and runoff over land could increase because of the RF gradient between ocean and land. In

1 contrast, increasing the albedo of desert areas may result in reduction in rainfall over the Indian and Sahel
2 regions (Irvine et al., 2010).

3
4 **[INSERT FIGURE 7.23 HERE]**

5 **Figure 7.23:** Multi-model mean of the change in surface air temperature (°C) averaged over December, January and
6 February months for (a) the abrupt $4 \times \text{CO}_2$ simulation and (b) the GeoMIP G1 experiment. (c-d) same as (a-b) but for
7 the June, July and August months. (e-h) same as (a-d) but for the change in precipitation (mm day^{-1}). All changes are
8 relative to the pre-industrial control experiment and averaged over years 21-50. Stippling denotes agreement on the sign
9 of the temperature anomaly in at least 75% of the models in panels.

10
11 High CO_2 concentrations from anthropogenic emissions will persist in the atmosphere for a long time (for
12 more than a thousand years in the absence of CDR). If SRM were used to counter positive forcing, it would
13 be needed as long as the CO_2 concentrations were high. If greenhouse gas concentrations increase, then the
14 scale of SRM to offset the resulting warming would need to increase, with amplifying residual effects from
15 increasingly imperfect compensation. Figure 7.24 shows the globally averaged warming and precipitation
16 changes associated with a $1\% \text{ yr}^{-1}$ CO_2 increase (GeoMIP experiment G2), with and without SRM, followed
17 by termination of SRM at year 50. While the rate of temperature change varies by a factor of 2–3 between
18 models, all GeoMIP models return to temperature levels consistent with CO_2 forcing within one to two
19 decades. The associated very rapid warming would have significant impact on ecosystem and human
20 adaptation. Other strategies could be considered that use SRM with mitigation to try to avoid transitions
21 across climate thresholds or to keep temperatures below a threshold in combination with greenhouse gas
22 emissions reductions. Such scenarios would not necessarily be subject to the “termination effect” as
23 described above and SRM might be used for shorter times, perhaps less than a century (Smith and Rasch,
24 2012; Wigley, 2006).

25
26 **[INSERT FIGURE 7.24 HERE]**

27 **Figure 7.24:** Timeseries of surface temperature (°C, left) and precipitation change (mm day^{-1} , right) for GeoMIP
28 experiment G2, relative to each model’s $1 \times \text{CO}_2$ reference simulation. Solid lines are simulations using SRM to
29 balance a $1\% \text{ yr}^{-1}$ increase in CO_2 concentration until year 50 after which SRM is stopped. Dashed lines are for $1\% \text{ CO}_2$
30 increase simulations with no SRM.

31
32 Several modelling studies (Jones et al., 2010; Rasch et al., 2008b; Robock et al., 2008) examined the model
33 response to SRM in less idealized scenarios with stratospheric aerosol SRM when the forcing is sufficient to
34 counter the effects of a doubling CO_2 on globally-averaged surface temperature. The results show that
35 changes in temperature, winds and precipitation are similar to those obtained in idealized studies described
36 above, with a residual warming at high latitudes, and a reduction in global-mean precipitation. Similar
37 responses have been observed following volcanic eruptions like Pinatubo (Trenberth and Dai, 2007). Each of
38 these studies used a somewhat different experimental design, and there are significant disagreements in
39 regional responses that may be due to the experimental design, model differences, or larger differences in
40 forcing produced by different representation of the stratospheric aerosol production, transport, and loss
41 processes.

42
43 Jones et al. (2009), Rasch et al. (2009), and Hill and Ming (2012) used coupled ocean/atmosphere/sea ice
44 models to assess the climate impacts of cloud brightening due to droplet concentration changes. A global
45 cooling was achieved, in spite of the heterogeneous RF associated with cooling by subtropical and
46 midlatitude cloud systems in all three models, but temperature and precipitation signatures that differed
47 substantially. One common feature was the cooling over the seeded regions (the marine stratocumulus
48 regions) and a warmer North Pacific adjacent to a cooler northwestern Canada, a feature reminiscent of La
49 Niña.

50
51 **7.7.4 Synthesis**

52
53 Multiple, consistent and independent lines of evidence (from theory, model studies and observations) suggest
54 that some SRM strategies may be able to counter a portion of the global warming due to anthropogenic
55 greenhouse gases. In particular SRM by stratospheric aerosols appears to be scalable to counter at least a
56 twofold increase in CO_2 concentration, but models disagree on the amount of material that would need to be
57 injected to achieve this. SRM by marine cloud brightening has received somewhat less attention, and issues
58 are less well understood because of the high level of uncertainty about aerosol-cloud interactions, and there

1 is as yet no consensus on whether it will be possible to achieve a significant RF. It does not appear that SRM
2 strategies that change land albedo (whiter rooftops and paving, vegetation changes) can produce sufficient
3 RF to counter the warming expected from a doubling of CO₂. Much less attention has been devoted to other
4 SRM strategies, so it is not yet possible to assess their efficacy, scalability, viability and consequences.

5
6 There is high agreement among models that the compensation between greenhouse gas warming and SRM
7 cooling is imprecise and that some regional impacts would remain, but the models disagree on the details.
8 Models consistently indicate that, if feasible, a geoengineered climate with SRM and high atmospheric CO₂
9 levels would be much closer to 20th century climate than a world with elevated CO₂ concentrations and no
10 SRM (Lunt et al., 2008; Moreno-Cruz et al., 2011). A wide range of side effects has been identified for SRM
11 methods. Some are common to all SRM methods (e.g., a global decrease in precipitation for no temperature
12 change in the global-mean). Others are particular to a specific SRM method (e.g., the high latitude
13 stratospheric ozone depletion from stratospheric aerosol SRM). SRM will not address the issue of ocean
14 acidification from increasing CO₂ and may have other impacts on the climate system. If SRM were
15 terminated at a high CO₂ concentration, the global average temperature would rapidly (within a decade or
16 two) approach temperatures consistent with that CO₂ forcing, severely stressing ecosystem and human
17 adaptation.

18
19 **[START FAQ 7.1 HERE]**

20 21 **FAQ 7.1: How do Clouds Affect Climate and Climate Change?**

22
23 *Clouds strongly affect the current climate, but observations alone cannot yet tell us how they will affect a*
24 *future, warmer climate. Comprehensive prediction of clouds requires a global climate model. Such models*
25 *produce cloud fields that roughly resemble those observed, but are far from perfect. Climate models vary in*
26 *how they predict clouds will change in a warmer climate, but based on all available evidence, it seems likely*
27 *that the net cloud-climate feedback is positive, and will not significantly limit global warming.*

28
29 Since the 1970s, scientists have recognised the critical importance of cloudiness for climate change. Clouds
30 affect the climate system in a variety of ways. They produce precipitation (rain and snow) that is necessary
31 for most life on land. They warm the atmosphere as water vapour condenses and releases latent heat. They
32 strongly affect the flows of both solar and infrared radiation through the atmosphere. Finally, they are
33 intimately associated with powerful vertical motions that can carry air from near the surface to the upper
34 troposphere in less than an hour. These strong, vertical currents carry energy, moisture, momentum, and
35 various chemical constituents, including aerosol particles. For decades, climate scientists have been using
36 observations to study how clouds change with the daily weather, with the seasonal cycle, and with year-to-
37 year changes, such as those associated with El Niño.

38
39 Each of the various cloud processes has the potential to change as the climate state evolves. Cloud feedbacks
40 are of intense interest in the context of anthropogenic climate change. Any change in a cloud process that is
41 caused by climate change—and in turn influences climate—represents a cloud-climate feedback. Because
42 clouds interact so strongly with both infrared and visible radiation, relative changes in cloudiness of only a
43 few per cent can have a potent effect on the climate system.

44
45 Many types of possible cloud-climate feedbacks have been identified. Broadly speaking, they would occur
46 through changes in cloud amount, cloud-top height, and/or cloud optical properties. We still are not sure
47 what types of cloud feedbacks will actually occur, nor how significant they will be for climate change.
48 Nevertheless, all the models used for the past two IPCC assessments produce cloud feedbacks that either
49 enhance anthropogenic greenhouse warming or have little overall effect. Feedbacks are not “put into” these
50 models, but are an aspect of the solutions to the model equations describing the atmosphere and other parts
51 of the climate system. The differences in the strength of the cloud feedbacks produced by these models
52 explain why they react more or less sensitively to changes in greenhouse gas concentrations.

53
54 Low clouds reflect a lot of solar radiation back to space, but have only a weak effect on the infrared radiation
55 emitted by the Earth. As a result, they tend to cool the Earth in the present climate. High clouds efficiently
56 reflect sunlight, but also strongly reduce the amount of emitted infrared radiation. This compensation makes
57 the surface temperature somewhat less sensitive to changes in high cloud amount than to changes in low

1 cloud amount. In a future climate, warmed by increasing greenhouse gases, most IPCC-assessed climate
2 models simulate a decrease in low-cloud amount, which would increase the warming, but the extent of this
3 decrease is quite model-dependent.

4
5 Although surface temperature is relatively insensitive to changes in high cloud amount, changes in the
6 altitude of a given amount of high clouds can more strongly affect surface temperature. Higher cloud tops
7 increase the greenhouse effect, while reflecting a similar amount of solar radiation. There is strong evidence
8 that high clouds rise further in a warmer climate. This acts as an important positive feedback by preventing
9 some of the additional infrared radiative energy emitted by the surface from leaving the climate system.

10
11 There are other ways clouds may change in a warmer climate. More clouds may be made of liquid drops, and
12 fewer made of ice crystals, changing the overall amount of light reflected. Changing amounts of aerosol
13 particles could cause changes in cloud reflectivity, independent of any caused by greenhouse gases (see FAQ
14 7.2). Changes in wind patterns and storm tracks could also affect the regional and seasonal distribution of
15 cloudiness and precipitation. Some studies suggest that the signal of one such expected trend—a poleward
16 migration of the clouds associated with mid-latitude storm tracks—is already detectable in the observational
17 record. However, because clouds are quite variable from year to year, there is as yet no broadly accepted
18 way to infer global cloud feedback more generally from observations.

19
20 **[END FAQ 7.1 HERE]**

21
22 **[START FAQ 7.2 HERE]**

23 24 **FAQ 7.2: How do Aerosols Affect Climate and Climate Change?**

25
26 *Atmospheric aerosols are small particles suspended in the atmosphere. They come from natural and*
27 *anthropogenic sources, and can affect the climate in multiple and equivocal ways through their interactions*
28 *with radiation and clouds. Overall, it seems very likely that anthropogenic variations in atmospheric*
29 *aerosols have exerted a cooling influence on the Earth, which has diminished anthropogenic greenhouse*
30 *warming.*

31
32 Atmospheric aerosols have a typical lifetime of one day to two weeks in the troposphere, and about one year
33 in the stratosphere. They vary greatly in size, chemical composition and shape. Some aerosols, such as dust
34 and sea spray, are mostly or entirely of natural origin, while other aerosols, such as sulphates and smoke,
35 come from both natural and anthropogenic sources.

36
37 Aerosols affect climate in many ways. First, they scatter and absorb sunlight, which modifies the Earth's
38 radiative balance (see FAQ.7.2, Figure 1). Aerosol scattering generally makes the planet more reflective, and
39 tends to cool the climate, while aerosol absorption has opposite effects, and tends to warm the climate
40 system. The balance between cooling and warming depends on aerosol properties and environmental
41 conditions. Many observational studies have quantified local radiative effects from anthropogenic and
42 natural aerosols, but predicting their global impact requires satellite data and models. Most studies agree that
43 the overall radiative effect from anthropogenic aerosols is to cool the planet. One of the remaining
44 uncertainties, however, comes from soot, an absorbing aerosol which is more difficult to measure than
45 scattering aerosols, and induces a specific cloud response.

46
47 **[INSERT FAQ 7.2, FIGURE 1 HERE]**

48 **FAQ 7.2, Figure 1:** Overview of aerosol-radiation interactions and their impact on climate. The left panels show the
49 instantaneous radiative effects of aerosols, while the right panels show their overall impact after the climate system has
50 responded to their radiative effects.

51
52 Aerosols also serve as condensation and freezing sites, where cloud droplets and ice particles form (see
53 FAQ.7.2, Figure 2). You might assume that more cloud condensation nuclei would increase the amount of
54 low clouds, but cloud formation is largely limited by dynamical processes, so that the net effect on clouds of
55 more aerosols is quite subtle, and remains uncertain. A robust result is that more aerosols generally produce
56 liquid clouds, which are brighter because they have more numerous, smaller cloud droplets. There are many
57 other pathways for aerosol-cloud interactions, particularly in ice—or mixed liquid and ice—clouds, where

1 phase changes are sensitive to aerosols. Quantifying the overall impact of aerosols on cloud amounts and
2 properties is understandably difficult. Available studies, based on climate models and satellite observations,
3 generally indicate that the net effect of anthropogenic aerosols on cloud is to cool the climate system.
4

5 **[INSERT FAQ 7.2, FIGURE 2 HERE]**

6 **FAQ 7.2, Figure 2:** Overview of aerosol-cloud interactions and their impact on climate. The left and right panels
7 represent a clean and a polluted low-level cloud, respectively.
8

9 Since aerosols are distributed unevenly in the atmosphere, they can heat and cool the climate system in
10 patterns that can drive small changes in the weather. These effects are complex, and hard to simulate with
11 current models, but several studies suggest significant effects on precipitation in certain regions.
12

13 Because of their short lifetime, the abundance of aerosols—and their climate effects—have varied over time,
14 in rough concert with anthropogenic emissions of aerosols and their gaseous precursors. Since anthropogenic
15 emissions have increased substantially over the industrial period, this has very likely counteracted some of
16 the warming that would otherwise have occurred from increased concentrations of long-lived greenhouse
17 gases. Aerosols from volcanic eruptions, such as those of El Chichón and Pinatubo, have also caused
18 sporadic cooling periods.
19

20 Over the last two decades, anthropogenic aerosol emissions have decreased in industrialised countries, but
21 increased in developing countries. It is therefore difficult to assess whether the global impact of aerosols has
22 been to cool or warm the planet over this recent period. It is very likely, however, that emissions of
23 anthropogenic aerosols will ultimately decrease, augmenting greenhouse-gas induced warming.
24

25 **[END FAQ 7.2 HERE]**

26
27 **[START FAQ 7.3 HERE]**

29 **FAQ 7.3: Could Geoengineering Counteract Climate Change and What Side Effects Might Occur?**

30
31 *Geoengineering—also called climate engineering—is defined as a deliberate large-scale manipulation of the*
32 *climate system to counteract the consequences of increasing greenhouse gas emissions. Two distinct*
33 *categories of geoengineering methods are usually considered: Solar Radiation Management, also called*
34 *Sunlight Reflection Methods (SRM, assessed in Chapter 7) and Carbon Dioxide Removal (CDR, assessed in*
35 *Chapter 6). Each operates on different physical principles, and acts in different ways. While both methods*
36 *are likely to counter some effects of greenhouse gases, they also carry risks and side effects.*
37

38 ***Carbon Dioxide Removal (CDR) Methods***

39
40 By definition, CDR methods are designed to remove CO₂ from the atmosphere, often through a manipulation
41 of natural carbon cycle processes. The carbon withdrawn from the atmosphere would then be stored in land,
42 ocean or geological reservoirs, where it would have little or no impact on the Earth's energy budget. Some
43 proposed CDR methods would rely on biological processes, such as afforestation/reforestation, carbon
44 sequestration in soils through biochar, bioenergy associated with carbon capture and storage (BECCS) and
45 ocean fertilisation. Others would rely on geochemical processes, such as accelerated weathering of silicate
46 and carbonate rocks—on land or in the ocean—to remove CO₂ from the atmosphere, or directly capture it in
47 industrial facilities (see FAQ.7.3, Figure 1). The captured CO₂ would then be stored in organic form in land
48 reservoirs, and in inorganic form in ocean and geological reservoirs.
49

50 **[INSERT FAQ 7.3, FIGURE 1 HERE]**

51 **FAQ 7.3, Figure 1:** Overview of some carbon dioxide removal methods. (a) nutrients are added to the ocean, which
52 increases oceanic productivity in the surface ocean and transports a fraction of the resulting biogenic carbon downward,
53 (b) alkalinity from solid minerals is added to the ocean, which causes more atmospheric CO₂ to dissolve in the ocean,
54 (c) the weathering rate of silicate rocks is increased, and the dissolved carbonate minerals are transported to the ocean,
55 (d) alkalinity from mined silicate rocks is extracted, then combined with atmospheric CO₂ to produce solid carbonate
56 minerals, (e) atmospheric CO₂ is captured chemically, and stored either underground or in the ocean, (f) biomass is
57 burned at an electric power plant with carbon capture, and the captured CO₂ is stored either underground or in the ocean
58 and (g) CO₂ is captured through afforestation and reforestation to be stored in land ecosystems.

1
2 In general, insofar as CDR methods are effective at removing CO₂ from the atmosphere and keeping the
3 removed carbon away from the atmosphere, they are believed to pose a low risk of unintended climatic side
4 effects. If deployed, they would counter the root cause of CO₂-induced climate change by reducing
5 atmospheric carbon dioxide concentrations, thus reducing the associated radiative forcing and ocean
6 acidification.

7
8 One main issue relates to the capacity and permanence of storage reservoirs. Permanent carbon storage by
9 CDR would decrease climate warming in the long term. However, non-permanent storage strategies, which
10 would eventually see the CO₂ returned to the atmosphere, would only reduce climate change for some
11 limited time. Geological reservoirs could permanently store several thousand PgC, while oceans may also be
12 able to store a further few thousand PgC. In contrast, the estimated capacity of the terrestrial biosphere is
13 only 150 to 200 PgC, which represents only about 10% of the existing living biomass and soil carbon pools.

14
15 It is well known that the land and ocean carbon pools act as a buffer for CO₂ emitted to, and removed from,
16 the atmosphere. At present, about 50% of the CO₂ emitted into the atmosphere from fossil fuel burning is
17 taken up by the land and ocean carbon pools. Should CDR be deployed, the uptake would be reduced or even
18 reverse into a carbon source over a period of several decades. Therefore, to offset past anthropogenic CO₂
19 emissions, CDR techniques would need to remove not just the CO₂ that accumulated in the atmosphere, but
20 also the anthropogenic carbon previously taken up by terrestrial and oceanic sinks.

21
22 Any given biological or chemical weathering CDR method is estimated to be able to remove a maximum of
23 100 PgC (equivalent to about 50 parts per million by volume [ppm] of atmospheric CO₂) over a century.
24 However, that maximum potential cannot typically be achieved, due to constraints such as competing
25 demands for land. Furthermore, buffering could reduce the effective potential to about 25 ppm.

26
27 The atmospheric concentration of CO₂ has risen by more than 75 ppm in the last 50 years. Assuming a
28 maximum CDR sequestration rate of 100 PgC per century as discussed above, any single biological or
29 chemical weathering CDR method would take about three centuries to remove the CO₂ emitted in the last 50
30 years, making it difficult—even for a suite of biological methods—to mitigate climate change rapidly. Direct
31 air capture methods could in principle operate much more rapidly, but are limited by cost, energy use and
32 environmental constraints.

33
34 CDR could have climatic and environmental side effects. For instance, it would tend to counter the general
35 increase in plant productivity expected in a world with higher CO₂ concentrations. A large-scale increase in
36 vegetation coverage, for instance through afforestation or energy crops, could alter surface characteristics,
37 such as surface reflectivity and turbulent fluxes. Many modelling studies have shown that afforestation in
38 seasonally snow-covered boreal regions could accelerate global warming, whereas afforestation in the
39 tropics may be more effective at slowing global warming.

40
41 Ocean-based CDR methods, which rely on enhanced biological production, could have many
42 biogeochemical side effects, most of them being largely unknown, given the complexity of marine
43 ecosystem communities. Macronutrients such as nitrogen and phosphorus would decrease in regions where
44 marine productivity is artificially enhanced, likely leading to a shortage of nutrients downstream and a
45 decrease in productivity. Ocean-based biological CDR methods might also have impact on ocean oxygen
46 levels, and stimulate increased production of DMS, isoprene, and non-CO₂ greenhouse gases such as N₂O
47 and CH₄. Finally acidification of the deep ocean might occur, caused by the respiration associated with the
48 organic matter that is exported downwards.

49 50 ***Solar Radiation Management (SRM) Methods***

51
52 The globally-averaged temperature of the planet is controlled by the amount of sunlight absorbed by the
53 Earth's atmosphere and surface, and by the existence of a greenhouse effect, by which gases and clouds
54 make the atmosphere opaque in the infrared spectrum where terrestrial radiation is emitted. If less incoming
55 sunlight reaches the surface, because of an increase in the reflectivity of the planet, or if energy is emitted to
56 space from a lower altitude because of a less opaque atmosphere in the infrared, the average global surface
57 temperature will be less.

1
2 Suggested geoengineering methods which manage the Earth’s radiative budget are based on this fundamental
3 physical principle (see FAQ7.3, Figure 2). These methods propose to either reduce sunlight reaching the
4 Earth—using mirrors in space—or increase the reflectivity of the planet by making the atmosphere, clouds or
5 the surface more reflective. Another technique proposes to suppress cirrus clouds, as these absorb some of
6 the terrestrial radiation emitted by the Earth.

7
8 **[INSERT FAQ 7.3, FIGURE 2 HERE]**

9 **FAQ 7.3, Figure 2:** Overview of some proposed solar radiation management (SRM) schemes to reflect sunlight to
10 space that would otherwise be absorbed. Illustrated methods, counter-clockwise from upper left, are (a) reflectors in
11 space, (b) aerosols in the stratosphere, (c) enhanced reflectivity of marine clouds, (d) making the ocean surface more
12 reflective, (e) growing more reflective crops, and (f) whitening of roofs and other built structures.

13
14 Basic physics tells us that if any of these methods produce a net flux change of energy from Earth to space,
15 the planet will cool. The picture is complicated, however, because of the many complex physical processes
16 which govern the interactions between the flow of energy, the atmospheric circulation, weather and the
17 resulting climate.

18
19 While the average surface temperature of the planet responds to the energy budget in a rather straightforward
20 way, the temperature at any given location and time is influenced by many other factors. The amount of
21 cooling from SRM will not in general equal the amount of warming caused by greenhouse gases at each
22 location and time. For example, SRM will only change heating rates during the daytime, whereas changes in
23 greenhouse gases affect heating rates during both day and night. This inexact compensation will have some
24 influence on the diurnal cycle of surface temperature, even if the average surface temperature is unchanged.
25 There may be other important changes: a uniform stratospheric aerosol loading might offset global mean
26 CO₂-induced warming, but some regions will cool less than others and polar regions could be left with some
27 residual warming.

28
29 If deployed, SRM could theoretically counteract anthropogenic climate change rapidly, cooling the Earth to
30 pre-industrial levels within one or two decades. We know this from climate models but also from the
31 observed rapid cooling of about 0.5 K that followed the eruption of Mt Pinatubo in 1991.

32
33 Climate consists of many factors besides surface temperature. Consequences for other climate features, such
34 as precipitation, soil moisture, river flow, snowpack and sea ice, may also be important. Both models and
35 theory show that compensating an increased greenhouse effect with SRM will not counter the globally-
36 averaged surface temperature and rainfall equally, and there could be large regional changes. Such imprecise
37 compensation in regional and global climate patterns makes it very unlikely that solar radiation management
38 will produce a future climate that is “just like” the one we experience today, or have experienced in the past.
39 However, climate models indicate that a geoengineered climate with SRM and high atmospheric CO₂ levels
40 could be much closer to 20th century climate than a future climate with elevated CO₂ concentrations and no
41 SRM.

42
43 SRM techniques will have other side effects: radiation management by stratospheric sulphate aerosols can
44 induce stratospheric ozone depletion, especially while chlorine from CFC emissions still resides in the
45 atmosphere, possibly increasing dangerous ultraviolet radiation reaching the surface. It can also alter the
46 ratio of direct to diffuse sunlight reaching the surface, which is known to affect terrestrial ecosystems.
47 Finally, SRM will not mitigate the ocean acidification associated with increasing atmospheric CO₂
48 concentrations.

49
50 Without CDR, high CO₂ concentrations from anthropogenic emissions will persist in the atmosphere for as
51 long as a thousand years, and SRM would have to be maintained for a similar period to remain effective at
52 counteracting global warming. Stopping SRM while CO₂ concentrations were still high would lead to a very
53 rapid warming over one or two decades, severely stressing ecosystem and human adaptation.

54
55 As greenhouse gas concentrations increase, the scale of any SRM deployment would have to increase too, in
56 order to keep pace with the greenhouse effect. This would exacerbate the residual climate change and the
57 risk of abrupt termination. SRM could conceivably help avoid transitions across climate thresholds or tipping

1 points that might, at some point, be unavoidable otherwise. Nevertheless, its side effects make it a high-risk
2 strategy for staving off climate change.

3

4 **[END FAQ 7.3 HERE]**

5

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1 **Tables**

2

3 **Table 7.2:** Global and regional anthropogenic and natural emissions important for atmospheric aerosols. For the anthropogenic components the maximum and minimum values from
 4 available inventories are presented according to Granier et al. (2011). Units are Tg yr⁻¹ except for BVOCs in TgC yr⁻¹ and DMS in TgS yr⁻¹. Dust and sea-spray estimates span the
 5 range in the historical CMIP5 simulations. SOA range is taken from Spracklen et al. (2011). BVOC range from Arneeth et al. (2008).

| Year 2000 Emissions Tg yr ⁻¹ | Dust | | Anthropogenic NMVOCs | | Anthropogenic BC | | Anthropogenic OA | | Anthropogenic SO ₂ | | Anthropogenic NH ₃ | | Source | Natural Global | |
|---|-------------------|------|-------------------------|--------|---------------------|------|---------------------|-------|----------------------------------|--------|----------------------------------|-------|-----------------------------|----------------|------|
| | MIN | MAX | MIN | MAX | MIN | MAX | MIN | MAX | MIN | MAX | MIN | MAX | | MIN | MAX |
| Total | 744 | 8061 | 121.00 | 139.50 | 4.60 | 5.60 | 6.40 | 12.70 | 102.00 | 145.00 | 37.50 | 38.90 | Sea Spray | 1441 | 6786 |
| Western Europe | 0.1 | 71 | 9.20 | 14.30 | 0.32 | 0.38 | 0.32 | 0.40 | 6.10 | 14.10 | 3.40 | 4.50 | PBAP Including spores 28 | 50 | 1000 |
| Central Europe | All Europe | | 2.30 | 3.50 | 0.11 | 0.21 | 0.25 | 0.39 | 4.60 | 10.00 | 1.10 | 1.20 | Dimethylsulphide | 10 | 40 |
| USA | 0.2 | 372 | 13.00 | 17.50 | 0.27 | 0.40 | 0.36 | 0.51 | 13.50 | 17.80 | 3.30 | 4.40 | Monoterpenes | 32 | 121 |
| Canada | All N. America | | 1.50 | 3.40 | 0.04 | 0.04 | 0.03 | 0.06 | 2.20 | 2.90 | 0.51 | 0.60 | Isoprene | 412 | 601 |
| Central America | | | 2.90 | 4.10 | 0.11 | 0.11 | 0.17 | 0.35 | 3.70 | 4.10 | 1.10 | 1.10 | SOA | 50 | 380 |
| South America | 4 | 380 | 8.40 | 12.90 | 0.20 | 0.33 | 0.32 | 0.83 | 3.80 | 8.80 | 3.40 | 3.50 | | | |
| Africa | 519 | 3807 | 10.80 | 14.50 | 0.46 | 0.62 | 1.05 | 1.91 | 5.30 | 8.80 | 2.30 | 2.40 | | | |
| China | 23 | 386 | 11.50 | 24.50 | 0.71 | 1.41 | 1.10 | 3.80 | 19.20 | 21.10 | 8.90 | 13.60 | | | |
| India | 20 | 347 | 7.30 | 10.80 | 0.45 | 0.84 | 1.00 | 3.27 | 4.00 | 7.90 | 3.70 | 8.50 | | | |
| Oceania | 0 | 7.6 | 0.00 | 1.50 | 0.03 | 0.04 | 0.04 | 0.08 | 2.40 | 2.70 | 0.72 | 0.72 | | | |

6

1 **Table 7.3:** Key aerosol properties of the main aerosol species in the troposphere. Brown carbon is a particular type of OA but is treated here as an additional component because it is
 2 light absorbing. The estimate of aerosol burdens and lifetimes in the troposphere are based on the AeroCom models, except for primary biological aerosol particles (PBAP), which
 3 are treated by analogy to other coarse mode aerosol types.

| Aerosol Species | Global Burden | Mass Size Distribution | Sources | Sinks | Lifetime | Key Climate Relevant Properties |
|-----------------|----------------------------|---|--|---|-----------------------------------|---|
| Black carbon | | Freshly emitted: 0–80 nm Aged: accumulation mode | Combustion of fossil fuels, biofuels and biomass | Wet deposition Dry deposition | 1 week to 10 days | Large mass absorption efficiency in the visible |
| Brown carbon | | Freshly emitted: 100–400 nm Aged: accumulation mode | Combustion of biofuels and biomass | Wet deposition Dry deposition | 1 week | Medium mass absorption efficiency in the visible. Light scattering. |
| Organic aerosol | | POA: Aitken mode SOA: nuclei mode Aged OA : accumulation mode Biogenic POA : coarse mode | Combustion of fossil fuel, biofuel and biomass. Continental and marine ecosystems. Some anthropogenic non-combustion activities. | Wet deposition Dry deposition | 1 week | Light scattering. Lens effect when deposited on black or brown carbon. CCN active (depending on aging time and mechanism). IN active (biogenic POA) |
| Sulphate | | Secondary: Nuclei, Aitken, and accumulation mode Primary: coarse mode | Primary: marine and volcanic emissions. Secondary: oxidation of SO ₂ from natural and anthropogenic sources | Wet deposition Dry deposition | 1 week | Light scattering. Very hygroscopic. Lens effect when deposited on black or brown carbon. CCN active. |
| Nitrate | | Accumulation and coarse modes | Oxidation of NO _x | Wet deposition Dry deposition | 1 week | Light scattering. CCN active. |
| Dust | (sensitive to size cutoff) | Coarse and super-coarse modes, with a small accumulation mode | Wind erosion, soil resuspension. Some agricultural practices and industrial activities (cement) | Sedimentation Dry deposition Wet deposition | 1 day to 1 week depending on size | IN active, light scattering and absorption, greenhouse effect. |
| Sea-salt | (sensitive to size cutoff) | Coarse mode and accumulation mode | Wave breaking. Wind erosion. | Sedimentation Wet deposition Dry deposition | 1 day to 1 week depending on size | Light scattering. Very hygroscopic. CCN active. |
| PBAP | | Mostly coarse mode | Terrestrial and oceanic ecosystems | Sedimentation Wet deposition Dry deposition | 1 day to 1 week depending on size | IN active. |

4

1 **Table 7.4:** List of references for each category of estimates displayed in Figure 7.19.

| Estimate | Acronym | References |
|--|------------------|---|
| RFari from CMIP5/ACCMIP models | CMIP5/ ACCMIP | [PLACEHOLDER FOR FINAL DRAFT] |
| RFaci published prior to TAR | TAR | (Boucher and Lohmann, 1995; Chuang et al., 1997; Feichter et al., 1997; Jones et al., 1994; Kaufman and Chou, 1993; Kiehl et al., 2000; Lohmann and Feichter, 1997; Lohmann et al., 2000; Rotstayn, 1999), |
| RFaci published between TAR and AR4 | AR4 | (Chen and Penner, 2005; Chuang et al., 2002; Ghan et al., 2001; Hansen et al., 2005; Jones et al., 2001; Kristjansson, 2002; Ming et al., 2005; Penner et al., 2006; Quaas and Boucher, 2005; Quaas et al., 2004; Rotstayn and Liu, 2003; Rotstayn and Penner, 2001; Suzuki et al., 2004; Takemura et al., 2005; Williams et al., 2001) |
| RFaci published since AR4 | AR5 | (Barahona et al., 2011; Bellouin et al., 2011; Haerter et al., 2009; Kvalevåg and Myhre, 2007; Lohmann et al., 2010; Lohmann et al., 2007; Penner et al., 2011; Rotstayn and Liu, 2009; Schmidt et al., 2012a; Storelvmo, 2011; Storelvmo et al., 2009; Wang and Penner, 2009; Zhou et al., 2012) |
| RFaci using or constrained by satellite observations | SAT | (Bellouin et al., 2012; Dufresne et al., 2005; Lebsock et al., 2008; Quaas and Boucher, 2005; Quaas et al., 2008; Quaas et al., 2009; Storelvmo et al., 2009) |
| RFaci from inverse studies | INV | (Knutti et al., 2002) |
| AFaci published prior to AR4 | AR4 | (Easter et al., 2004; Ghan et al., 2001; Johns et al., 2006; Jones et al., 2001; Kristjansson, 2002; Kristjansson et al., 2005; Lohmann, 2002b; Lohmann and Feichter, 1997; Lohmann et al., 2000; Menon et al., 2002; Ming et al., 2005; Peng and Lohmann, 2003; Penner et al., 2006; Penner et al., 2003; Quaas et al., 2006; Rotstayn, 1999; Rotstayn and Liu, 2005; Rotstayn and Penner, 2001; Storelvmo et al., 2006; Takemura et al., 2005; Williams et al., 2001) |
| AFaci published since AR4 | AR5 | (Chen et al., 2010; Ghan et al., 2011; Hoose et al., 2009; Kirkevåg et al., 2008; Makkonen et al., 2012; Menon and DelGenio, 2007; Ming et al., 2007b; Penner et al., 2011; Quaas et al., 2009; Rotstayn and Liu, 2009; Storelvmo et al., 2008a) |
| AFari+aci in liquid stratiform clouds published prior to AR4 | AR4 | (Ghan et al., 2012; Lohmann and Feichter, 2001; Lohmann et al., 2007; Posselt and Lohmann, 2008; Posselt and Lohmann, 2009; Quaas et al., 2004; Quaas et al., 2006; Quaas et al., 2009; Rotstayn et al., 2007; Salzmann et al., 2010) |
| AFari+aci in liquid stratiform clouds published since AR4 | AR5 | (Ghan et al., 2012; Lohmann et al., 2007; Makkonen et al., 2012; Posselt and Lohmann, 2008; Posselt and Lohmann, 2009; Quaas et al., 2009; Rotstayn et al., 2007; Salzmann et al., 2010) |
| AFari+aci in liquid and mixed-phase stratiform clouds | +MPC | (Hoose et al., 2010b; Hoose et al., 2008; Jacobson, 2006; Lohmann, 2004; Lohmann and Diehl, 2006; Lohmann and Ferrachat, 2010; Lohmann and Hoose, 2009; Salzmann et al., 2010; Storelvmo et al., 2008a; Storelvmo et al., 2008b; Yun et al., 2012), |
| AFari+aci in stratiform and convective clouds | +CNV | (Koch et al., 2009a; Lohmann, 2008; Menon and DelGenio, 2007; Menon and Rotstayn, 2006; Unger et al., 2009; Wang et al., 2011b) |
| AFari+aci from CMIP5/ACCMIP models | CMIP5/ ACCMIP | [PLACEHOLDER FOR FINAL DRAFT] |
| AFari+aci including satellite observations | SAT | (Bellouin et al., 2012; Lohmann and Lesins, 2002; Quaas et al., 2006; Quaas et al., 2009; Sekiguchi et al., 2003) |
| AFari+aci from inverse studies | INV | (Anderson et al., 2003; Andronova and Schlesinger, 2001; Church et al., 2011; Forest et al., 2006; Forest et al., 2002; Gregory et al., 2002; Hansen et al., 2011; Harvey and Kaufmann, 2002; Huber and Knutti, 2011; Libardoni and Forest, 2011; Murphy et al., 2009; Shindell and Faluvegi, 2009; Stott et al., 2006) |

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1 **Table 7.5:** Estimates of aerosol AF (in $W m^{-2}$) in some of the CMIP5 models. The AF are estimated from fixed-SST
 2 experiments using atmosphere-only version of the models listed. Different models include different aerosol effects.

| Modelling Group | Model Name | AFari+aci from All Anthropogenic Aerosols | AFari+aci from Sulphate Aerosols Only |
|--------------------|------------------------------|---|---------------------------------------|
| CCCma | CanESM2 | -0.87 | -0.90 |
| CSIRO-QCCCE | CSIRO-Mk3-6-0 | -1.41 | -1.10 |
| GFDL | GFDL-AM3 | -1.44 ^a | |
| GISS | GISS-E2-R ^b | -1.10 ^a | |
| GISS | GISS-E2-R-TOMAS ^b | -0.76 ^a | |
| IPSL | IPSL-CM5A-LR | -0.72 | -0.71 |
| LASG-IAP | FGOALS-s2 | -0.38 | |
| MIROC | MIROC-CHEM ^b | -1.24 ^a | |
| MIROC | MIROC5 | -1.28 | -1.05 |
| MOHC | HadGEM2-A | -1.22 | -1.16 |
| MPI | MPI-M | | |
| MRI | MRI-CGM3 | -1.10 | -0.48 |
| NCAR | NCAR-CAM5.1 ^b | -1.09 ^a | |
| NCC | NorESM1-M | -1.00 | |
| Ensemble mean | | -1.05 | |
| Standard deviation | | 0.29 | |

3 Notes:

4 (a) From ACCMIP (Shindell et al., 2012a).

5 (b) These models include the BC on snow effect.

Chapter 7: Clouds and Aerosols

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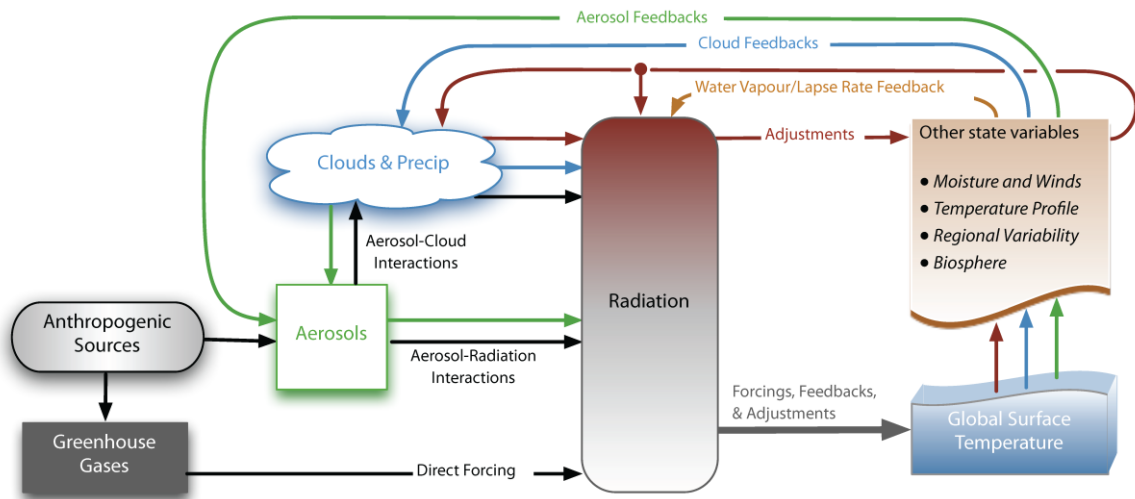
Review Editors: Sandro Fuzzi (Italy), Joyce Penner (USA), Venkatachalam Ramaswamy (USA), Claudia Stubenrauch (France)

Date of Draft: 5 October 2012

Notes: TSU Compiled Version

1 **Figures**

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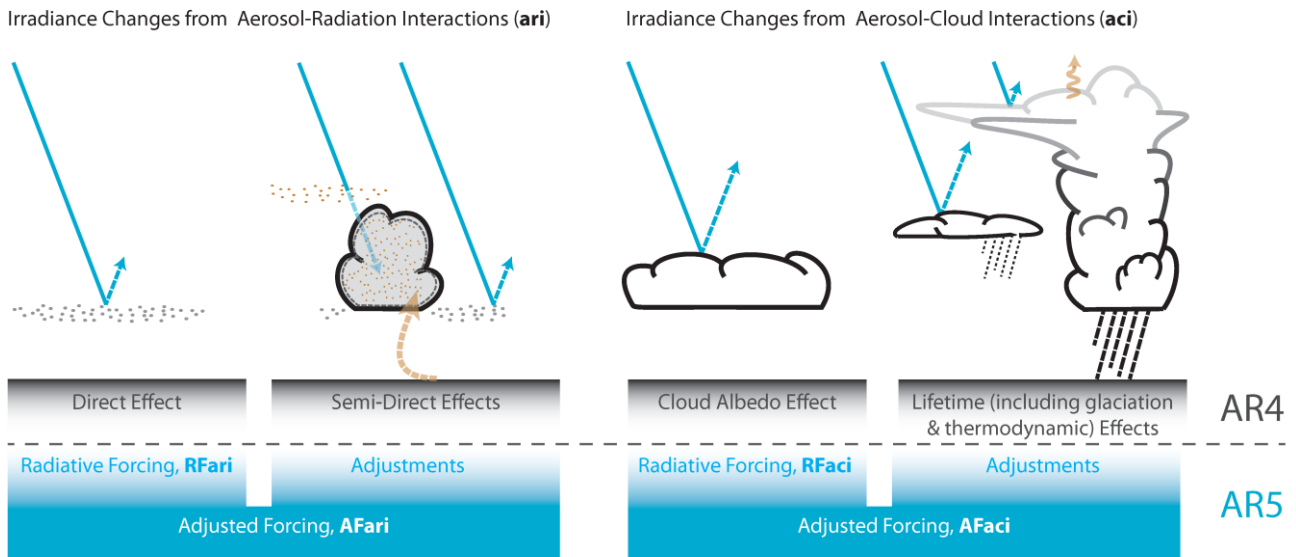
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5 **Figure 7.1:** Overview of feedback and forcing pathways involving clouds and aerosols. Forcing mechanisms are
 6 represented by black arrows; forcing agents are boxes with grey shadows, rapid forcing adjustments (also called rapid
 7 responses) are shown with brown arrows and feedbacks are other-coloured arrows. See text for further discussion.

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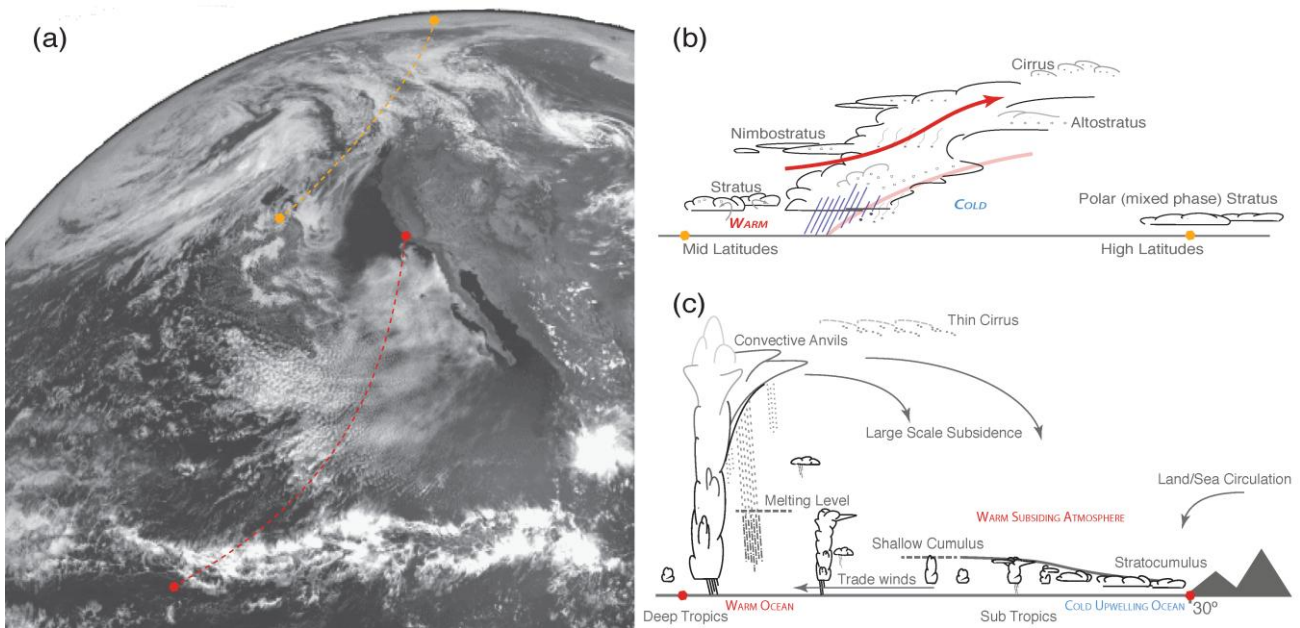
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Figure 7.2: Schematic of the new terminology used in this assessment report for aerosol-radiation and aerosol-cloud interactions and how they relate to the terminology used in AR4. The radiative forcing from aerosol-radiation interactions (abbreviated **RFari**) encompasses radiative effects from anthropogenic aerosols before any adjustment takes place, and corresponds to what is usually referred to as the aerosol direct effect. Rapid adjustments induced by aerosol radiative effects on the surface energy budget, the atmospheric profile and cloudiness contribute to the adjusted forcing from aerosol-radiation interactions (abbreviated **AFari**). They include what has earlier been referred to as the semi-direct effect. The radiative forcing from aerosol-cloud interactions (abbreviated **RFaci**) refers to the instantaneous effect on cloud albedo due to changing concentrations of cloud condensation and ice nuclei. All subsequent changes to the cloud lifetime and thermodynamics are rapid adjustments, which contribute to the adjusted forcing from aerosol-cloud interactions (abbreviated **AFaci**).

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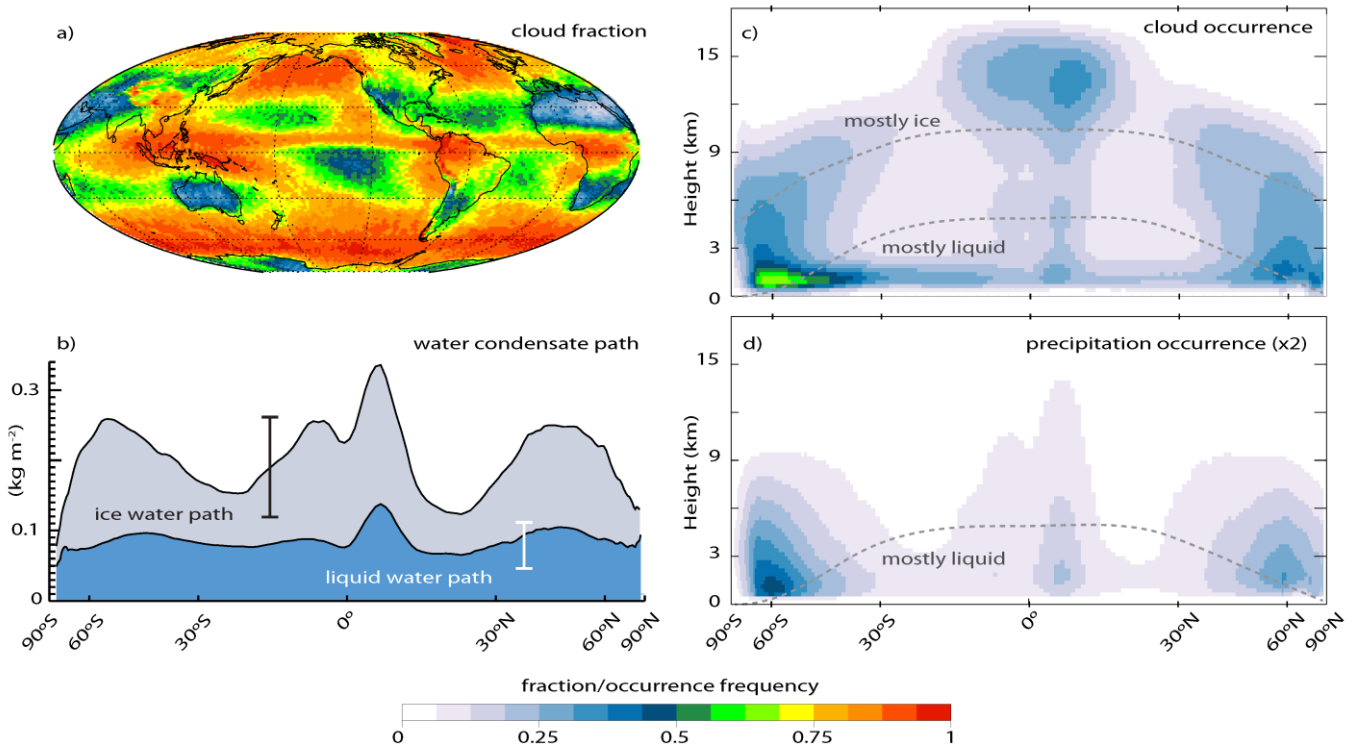
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Figure 7.3: Diverse cloud regimes reflect diverse meteorology. (a) A visible-wavelength geostationary satellite image shows (from top to bottom) expanses and long arcs of cloud associated with extratropical cyclones, subtropical coastal stratocumulus near Baja California breaking up into shallow cumulus clouds in the central Pacific, and mesoscale convective systems outlining the Pacific intertropical convergence zone or ITCZ. (b) A schematic section through a typical warm front of an extratropical cyclone (see orange dots in panel (a)) showing multiple layers of upper-tropospheric ice (cirrus) and mid-tropospheric water (altostratus) cloud upwind of the frontal zone, an extensive region of nimbostratus associated with frontal uplift and turbulence-driven boundary layer cloud in the warm sector. (c) A schematic section along the low-level trade wind flow from a subtropical west coast of a continent to the ITCZ (see red dots in panel (a)), showing typical low-latitude cloud types, shallow stratocumulus above the cool waters of the oceanic upwelling zone near the coast and trapped under a strong subsidence inversion, and shallow cumulus over warmer waters further offshore transitioning to precipitating cumulonimbus cloud systems with extensive cirrus anvils associated with rising air motions in the ITCZ.

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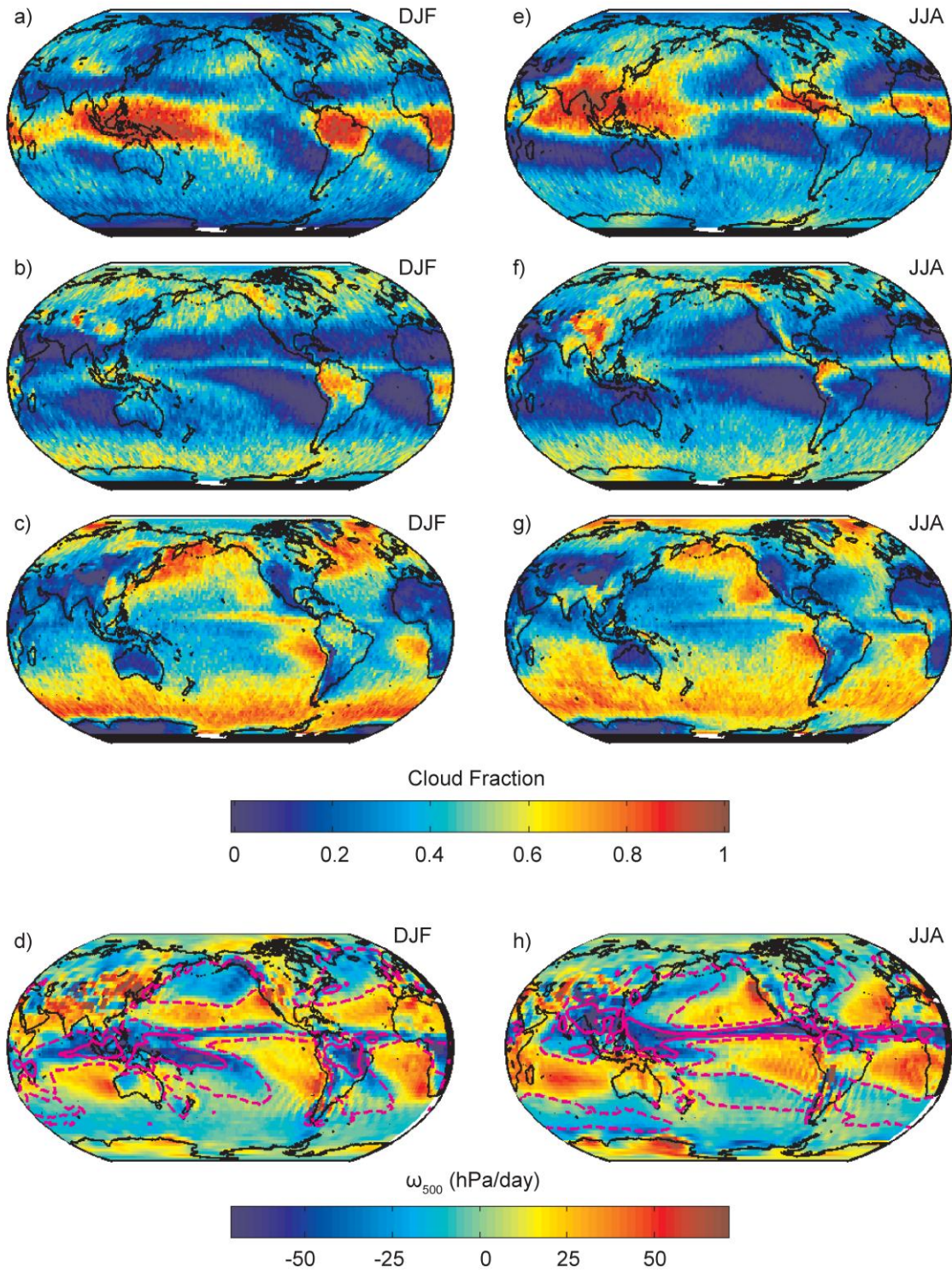
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Figure 7.4: (a) Annual mean cloud fractional occurrence (CloudSat/CALIPSO 2B-GEOPROF-LIDAR dataset for 2006–2011); (b) annual zonal mean liquid water path (blue shading, ocean only, O’Dell et al. (2008) microwave radiometer dataset for 1988–2005; the 90% uncertainty range, assessed to be 70–150% of the plotted value, is schematically indicated by the white error bar) and ice water path (grey shading, from CloudSat 2C-ICE dataset for 2006–2011; the 90% uncertainty range, assessed to be 50–200% of the plotted value, is schematically indicated by the black error bar). (c-d) latitude-height sections of annual zonal mean cloud (including precipitation falling from cloud) occurrence and precipitation (attenuation-corrected radar reflectivity >0 dBZ) occurrence; the latter has been doubled to make use of a common colour scale (2B-GEOPROF-LIDAR dataset). The dashed curves show the annual-mean 0°C and –38°C isotherms.

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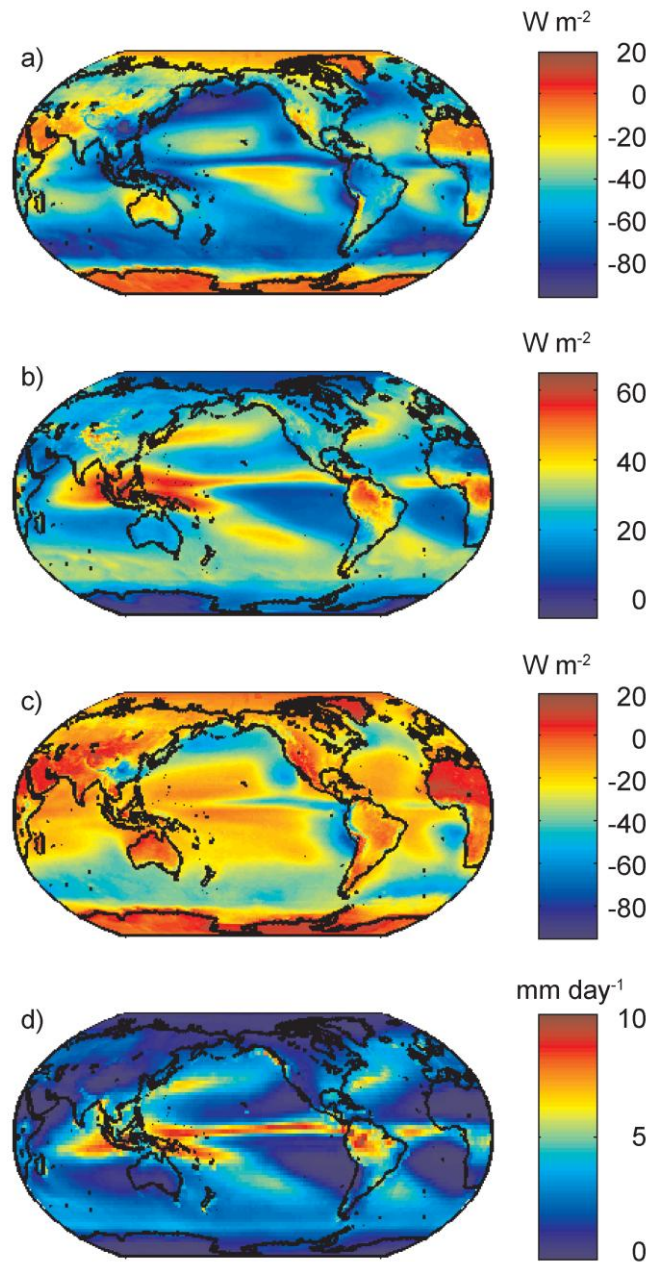
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Figure 7.5: (a-d) DJF mean high, middle and low cloud cover from CloudSat/CALIPSO GEOPROF dataset (2006–2011), and 500 hPa vertical pressure velocity (colours) and GPCP precipitation (magenta contours at 3 mm day⁻¹ in dash and 7 mm day⁻¹ in solid); (e-h) same as (a-d), except for JJA. For low clouds, the CALIPSO-only GOCCP dataset is used at locations where it indicates as larger fractional cloud cover, because the GEOPROF dataset removes some clouds with tops at altitudes below 750 m. Low cloud amounts are likely underrepresented in regions of high cloud (Chepfer et al., 2008), although not as severely as with earlier satellite technologies.

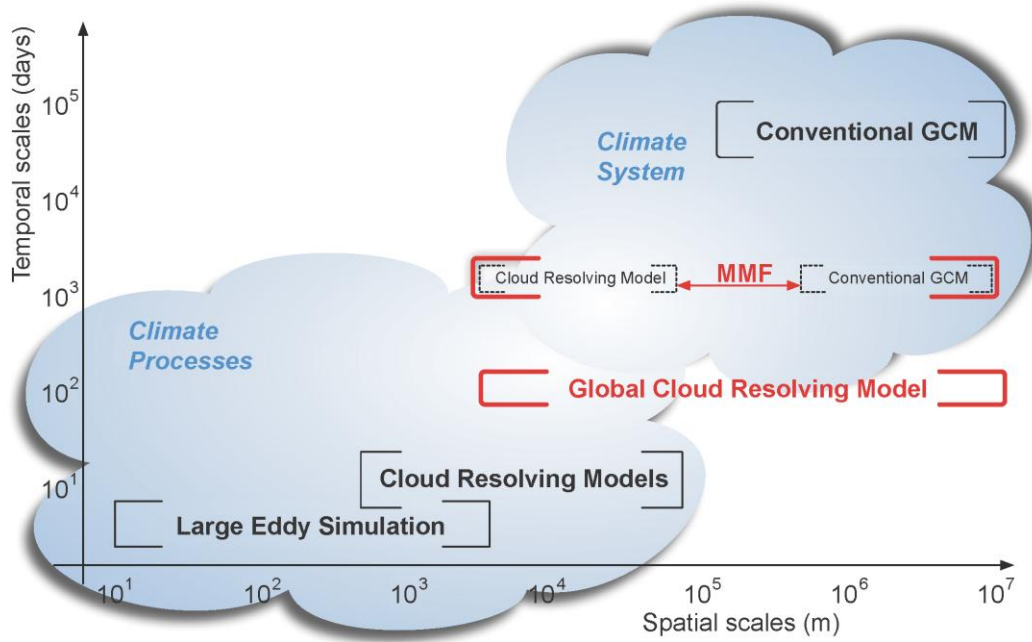
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Figure 7.6: Distribution of annual-mean top of atmosphere (a) SWCRE, (b) LWCRE, (c) net CRE (2001–2010 average from CERES-EBAF) and (d) precipitation (1981–2000 average from GPCP).

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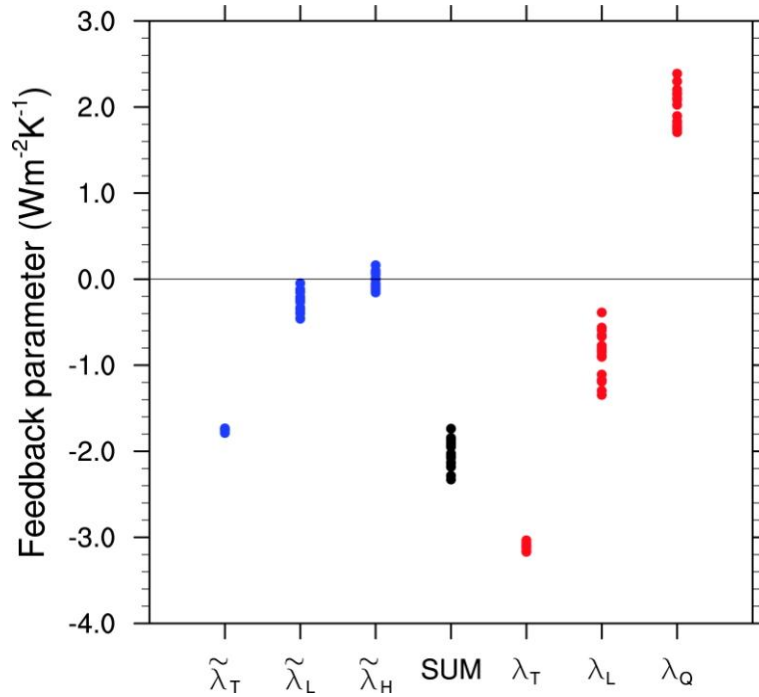
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Figure 7.7: Model and simulation strategy for representing the climate system and climate processes at different space and time scales. Models are usually defined based on the range of spatial scales they represent, shown by square brackets. The temporal scales that can be represented for a given model class can vary; for instance climate models can be run for a few time steps, or can simulate millennia. The figure indicates the typical timescales for which a given model is used. Computational power prevents one model from covering all time and space scales. Since the AR4 the development of global cloud resolving models, and hybrid approaches such as the multi-scale modelling framework, have helped fill the gap between climate system and climate process models.

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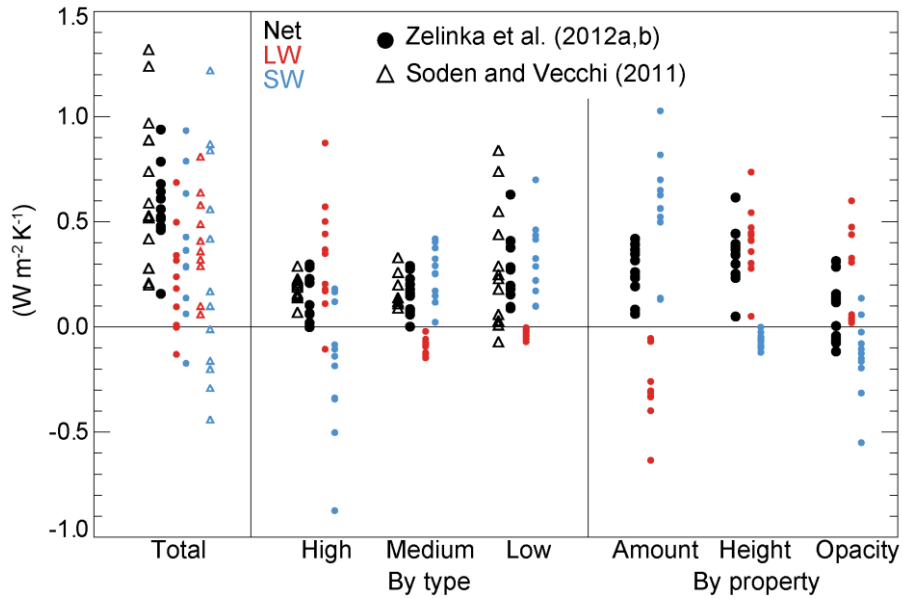
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Figure 7.8: Clear-sky feedback parameters as predicted by [CMIP3] GCMs. Black points in the centre show the total radiative response including the Planck response, with the Planck response and individual feedbacks from water vapour and lapse rate shown to the right in red. On the left are the equivalent three parameters calculated in an alternative, relative humidity-based framework. In this framework the Planck stabilization and each of the two feedbacks are all weaker and more consistent among the models. [PLACEHOLDER FOR FINAL DRAFT: Current working draft of figure taken from Held and Shell (2012); additional data from CMIP5 TBA].

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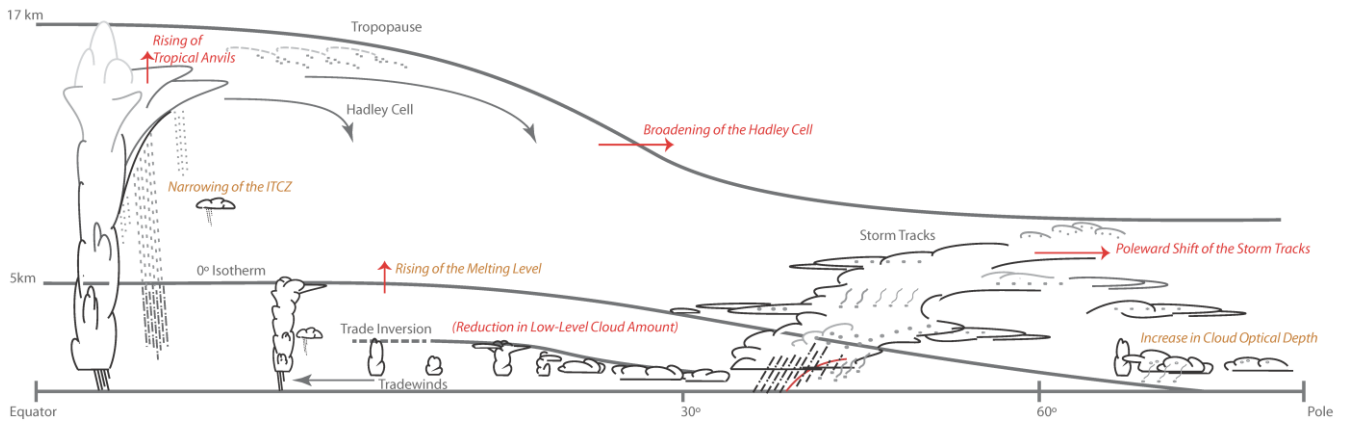
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Figure 7.9: Cloud feedback parameters as predicted by GCMs. Total feedback shown at left by black symbols, broken out into infrared and visible components in red and blue, respectively (Zelinka et al., 2012a; Zelinka et al., 2012b). Centre panel shows components attributable to clouds in different height ranges (see text); values reported by Soden and Vecchi (2011) do not conform exactly to this definition but are shown for comparison, with their “mixed” category assigned “medium”. Right panel shows components attributable to different cloud property changes (not available from all studies).

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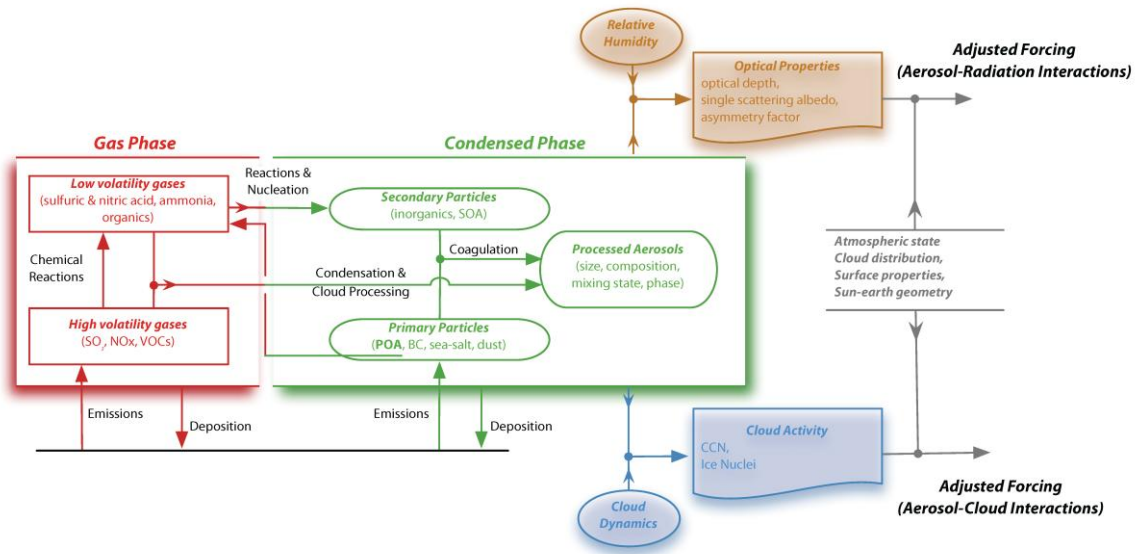
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Figure 7.10: Robust cloud responses to greenhouse warming (those simulated by most models and possessing some kind of independent support or understanding). Key climatological features (tropopause, freezing level, circulations) are shown in grey. Changes anticipated in a warmer climate are shown in red (if contributing positive feedback) or brown (if contributing little or ambiguously to feedback); no robust mechanisms contribute negative feedback. Changes include rising high cloud tops and melting level, and increased polar cloud cover and/or optical thickness (high confidence); broadening of the Hadley Cell and/or poleward migration of storm tracks, and narrowing of the ITCZ (medium confidence); and reduced low-cloud amount and/or optical thickness (low confidence). Confidence assessments are based on degree of GCM consensus, strength of independent lines of evidence from observations or process models, and degree of basic understanding.

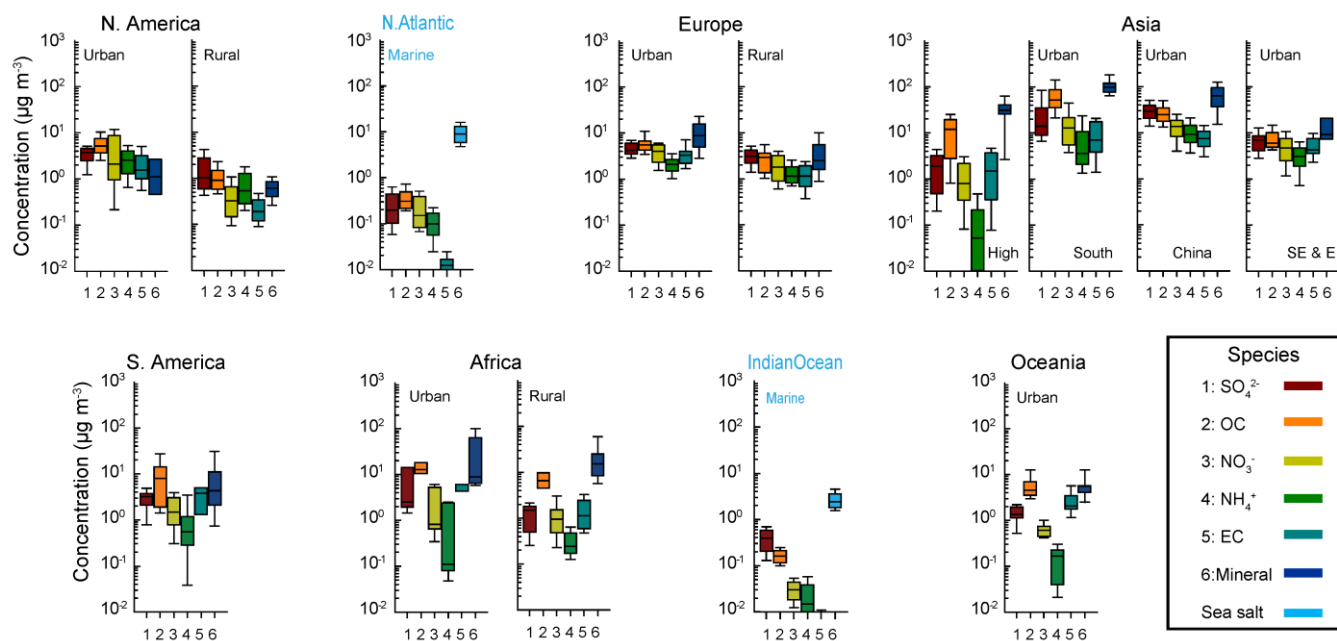
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Figure 7.11: Overview of atmospheric aerosol processes and environmental variables influencing aerosol-radiation and aerosol-cloud interactions. Gas-phase processes and variables are highlighted in red while particulate-phase processes and variables appear in green. Although this figure shows a linear chain of processes from aerosols to forcings, it is increasingly recognized that aerosols and clouds form a coupled system with two-way interactions.

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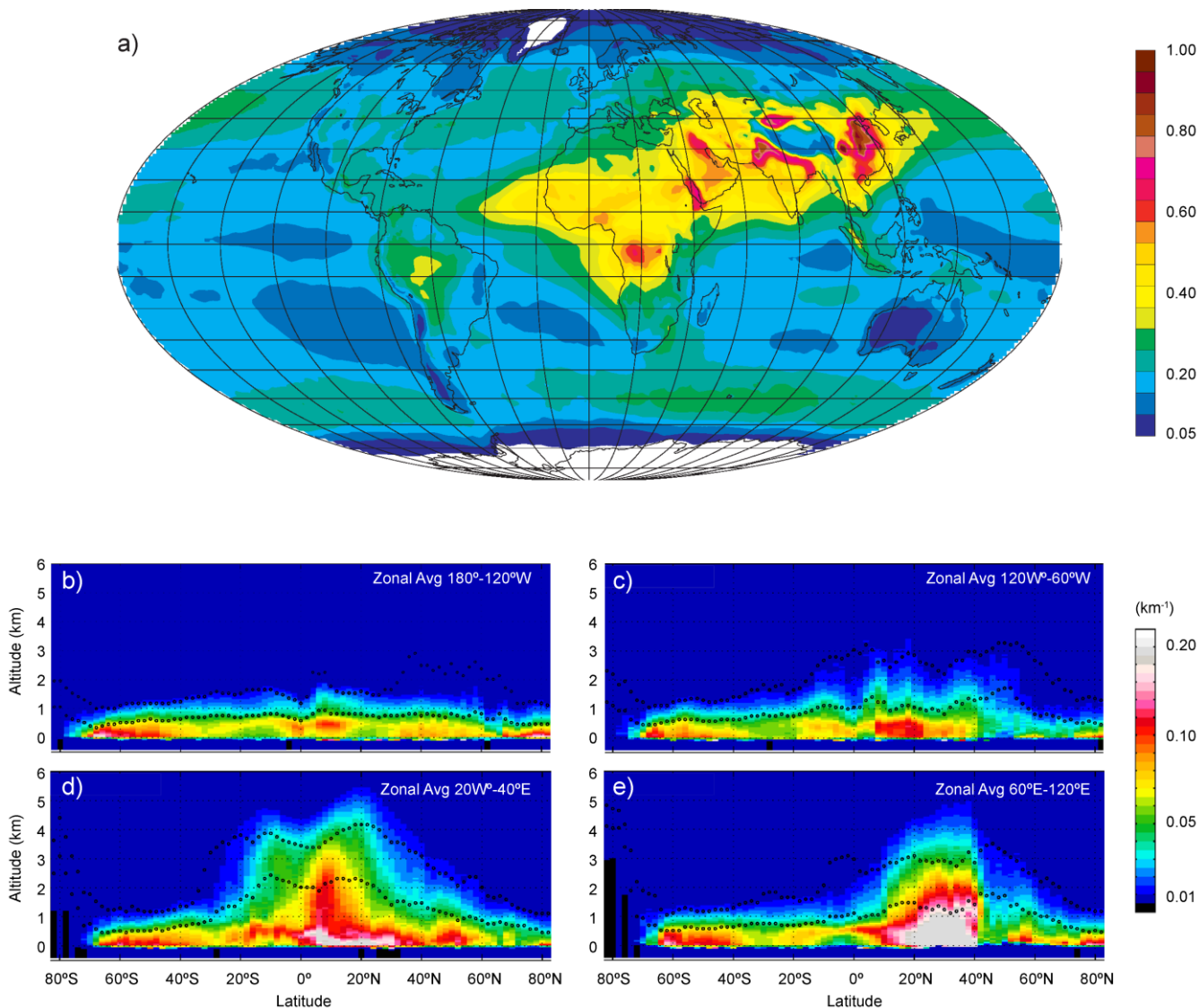
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Figure 7.12: Bar chart plots summarizing the annual, seasonal or monthly mean mass surface concentration ($\mu\text{g m}^{-3}$) of seven major aerosol components for particles with diameter smaller than $10 \mu\text{m}$, from various rural and urban sites in six continental areas of the world with at least an entire year of data, and two marine sites. For each location, the panels represent the median, the 25–75 percentiles (box), and the 10–90 percentiles (whiskers) for each aerosol component. These include: 1) urban North America. (Chow et al., 1993; Ito et al., 2004; Kim et al., 2000; Liu et al., 2005; Malm and Schichtel, 2004; Sawant et al., 2004); rural North America (Chow et al., 1993; Liu et al., 2005; Malm and Schichtel, 2004; Malm et al., 1994); 2) marine northern hemisphere Atlantic Ocean (Ovadnevaite et al., 2011; Rinaldi et al., 2009); 3) urban Europe (Hueglin et al., 2005; Lenschow et al., 2001; Lodhi et al., 2009; Lonati et al., 2005; Perez et al., 2008; Putaud et al., 2004; Querol et al., 2006; Querol et al., 2008; Querol et al., 2001; Querol et al., 2004; Rodriguez et al., 2002; Rodriguez et al., 2004; Roosli et al., 2001; Viana et al., 2006; Viana et al., 2007; Yin and Harrison, 2008); rural Europe (Gullu et al., 2000; Hueglin et al., 2005; Kocak et al., 2007; Putaud et al., 2004; Puxbaum et al., 2004; Querol et al., 2009; Querol et al., 2001; Querol et al., 2004; Rodriguez et al., 2002; Rodriguez et al., 2004; Salvador et al., 2007; Theodosi et al., 2010; Viana et al., 2008; Yin and Harrison, 2008; Yttri, 2007); 4) high Asia, with altitude larger than 1680 m. (Carrico et al., 2003; Decesari et al., 2010; Ming et al., 2007; Qu et al., 2008; Ram et al., 2010; Rastogi and Sarin, 2005; Rengarajan et al., 2007; Shresth et al., 2000; Zhang et al., 2001; Zhang et al., 2012; Zhang et al., 2008); urban South Asia (Chakraborty and Gupta, 2010; Khare and Baruah, 2010; Kumar et al., 2007; Lodhi et al., 2009; Raman et al., 2010; Rastogi and Sarin, 2005; Safai et al., 2010; Stone et al., 2010); urban China (Cheng et al., 2000; Hagler et al., 2006; Oanh et al., 2006; Wang et al., 2003; Wang et al., 2005b; Wang et al., 2006; Xiao and Liu, 2004; Yao et al., 2002; Ye et al., 2003; Zhang et al., 2002; Zhang et al., 2012; Zhang et al., 2011); rural China (Hagler et al., 2006; Hu et al., 2002; Zhang et al., 2012) [PANEL MISSING]; urban China (Cheng et al., 2000; Hagler et al., 2006; Oanh et al., 2006; Wang et al., 2003; Wang et al., 2005b; Wang et al., 2006; Xiao and Liu, 2004; Yao et al., 2002; Ye et al., 2003; Zhang et al., 2002; Zhang et al., 2012; Zhang et al., 2011); South-East and East Asia (Han et al., 2008; Khan et al., 2010; Kim et al., 2007; Lee and Kang, 2001; Oanh et al., 2006); 5) South America (Artaxo et al., 1998; Artaxo et al., 2002; Bourotte et al., 2007; Celis et al., 2004; Fuzzi et al., 2007; Gioda et al., 2011; Mariani and Mello, 2007; Martin et al., 2010; Morales et al., 1998; Souza et al., 2010); 6) urban Africa (Favez et al., 2008; Mkoma, 2008; Mkoma et al., 2009b); rural Africa (Maenhaut et al., 1996; Mkoma, 2008; Mkoma et al., 2009a; Mkoma et al., 2009b; Nyanganyura et al., 2007; Weinstein et al., 2010); 7) marine southern hemisphere Indian Ocean (Rinaldi et al., 2011; Sciare et al., 2009); 8) urban Oceania (Chan et al., 1997; Maenhaut et al., 2000; Radhi et al., 2010; Wang et al., 2005a; Wang and Shooter, 2001).

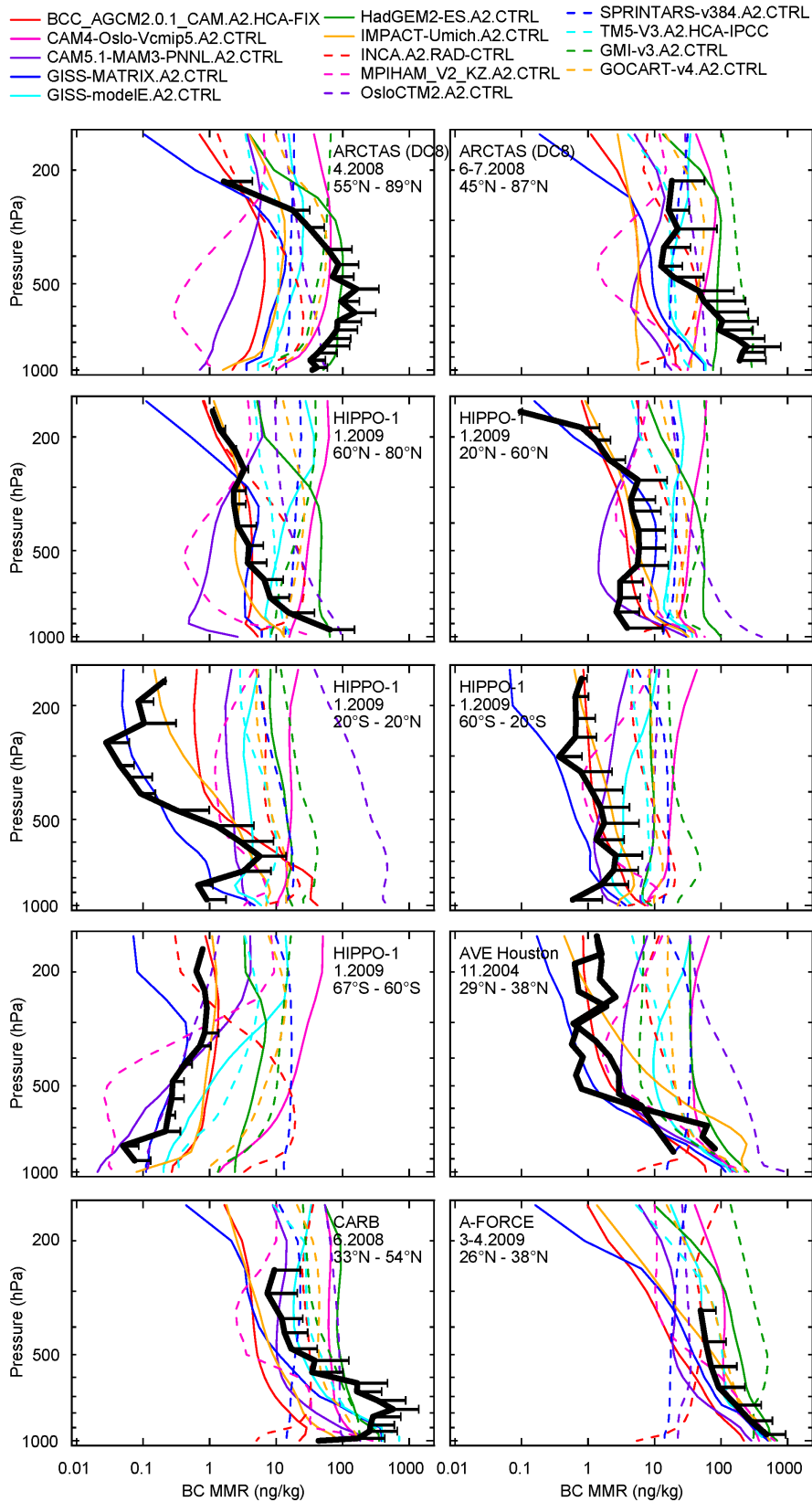
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Figure 7.13: a) spatial distribution of the AOD (unitless) from the MACC model with assimilation of MODIS AOD (Benedetti et al., 2009; Morcrette et al., 2009) for the year 2010; b) to e) latitudinal vertical cross-sections of the aerosol extinction coefficient (km^{-1}) for four longitudinal bands (180°W to 120°W, 120°W to 60°W, 20°W to 40°E, and 60°E to 120°E) from the CALIOP instrument (Winker et al., 2009). The black circles show the extinction scale height for 90% (top) and 63% (bottom) of the average AOD.

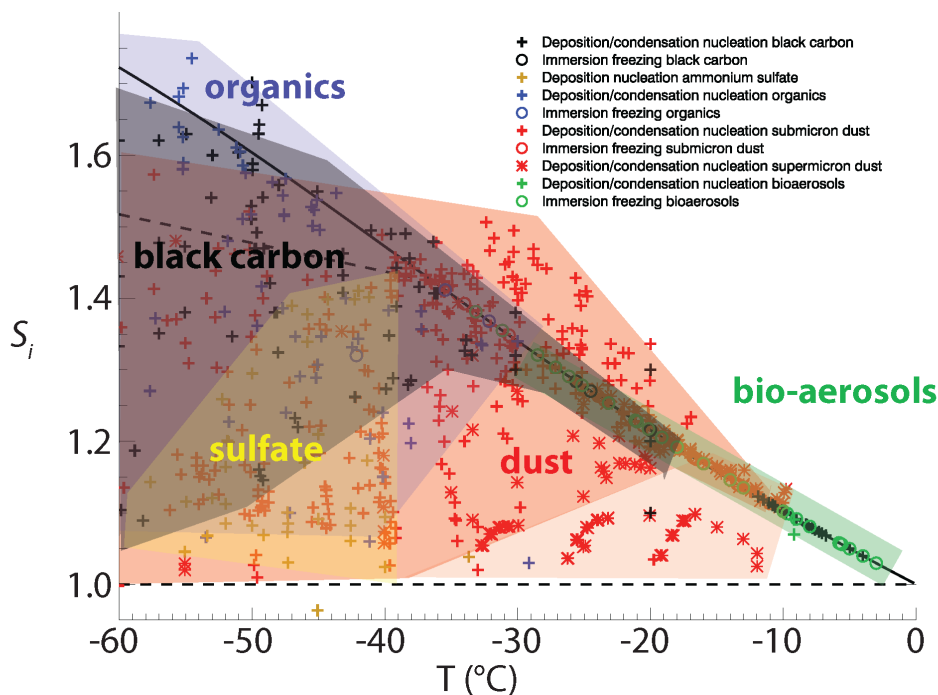
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Figure 7.14: Comparison of profiles of BC mass mixing ratios (ng kg^{-1}) as measured by airborne SP2 instruments during the ARCTAS (Jacob et al., 2010), HIPPO1 (Schwarz et al., 2006; Schwarz et al., 2010) and A-FORCE (Oshima et al., 2012) campaigns and as simulated by a range of AeroCom models. The model values are averages for the month corresponding to each field campaign.

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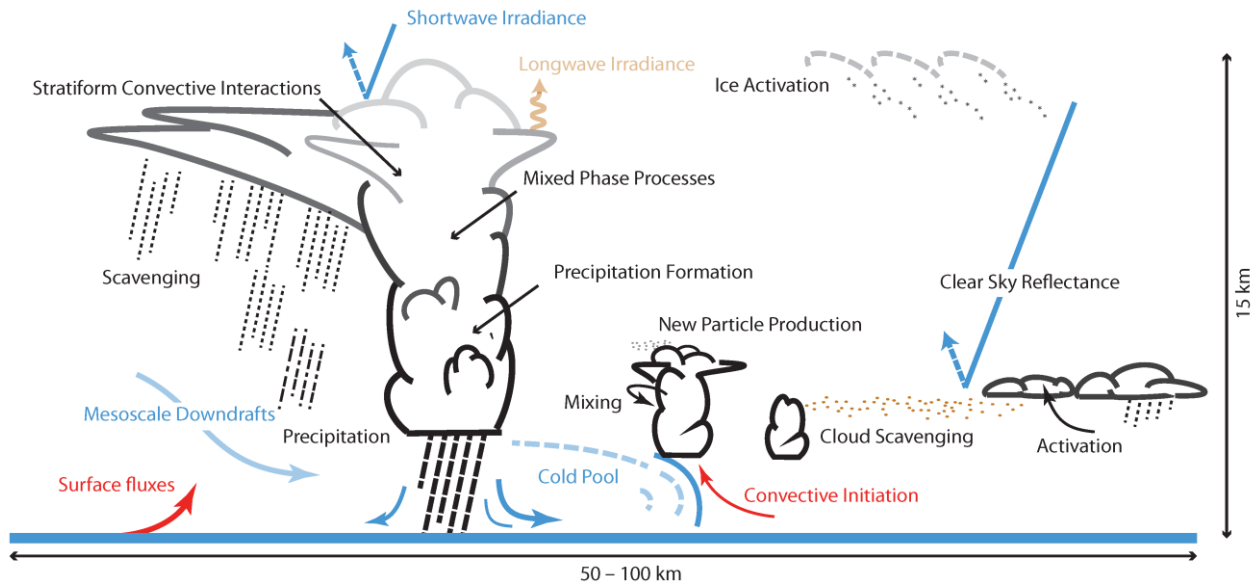
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4 **Figure 7.15:** The onset temperatures and relative humidities for deposition/condensation freezing and immersion
 5 freezing for **bioaerosols** (Ahern et al., 2007; Diehl et al., 2001; Iannone et al., 2011; Kanji et al., 2011; Mohler et al.,
 6 2008; Mortazavi et al., 2008; von Blohn et al., 2005; Yankofsky et al., 1981), **mineral dusts** (Archuleta et al., 2005;
 7 Bundke et al., 2008; Connolly et al., 2009; Cziczo et al., 2009; Field et al., 2006; Kanji and Abbatt, 2006; Kanji et al.,
 8 2011; Knopf and Koop, 2006; Koehler et al., 2010; Kulkarni and Dobbie, 2010; Lüönd et al., 2010; Mohler et al., 2006;
 9 Murray et al., 2011; Niedermeier et al., 2010; Niemand et al., 2012; Roberts and Hallett, 1968; Salam et al., 2006;
 10 Schaller and Fukuta, 1979; Welte et al., 2009; Zimmermann et al., 2008), **organics** (Baustian et al., 2010; Kanji et al.,
 11 2008; Petters et al., 2009; Prenni et al., 2007; Shilling et al., 2006; Wagner et al., 2010; 2011; Wang and Knopf, 2011;
 12 Zobrist et al., 2007), **solid ammonium sulphate** (Abbatt et al., 2006; Baustian et al., 2010; Mangold et al., 2005;
 13 Shilling et al., 2006; Wise et al., 2009; 2010) and **BC (soot)** (Crawford et al., 2011; DeMott, 1990; DeMott et al., 1999;
 14 Diehl and Mitra, 1998; Dymarska et al., 2006; Fornea et al., 2009; Gorbunov et al., 2001; Kanji et al., 2011; Mohler et
 15 al., 2005), from a compilation of experimental data of sub- and super-micron aerosol particles in the literature (for
 16 references see supplementary material). The large range of observed ice nucleation onset conditions is due to different
 17 experimental setups, particle sizes, activated fractions and chemical composition. Only those IN species for which at
 18 least three papers exists are shown. The solid line refers to saturation with respect to liquid water and the dashed line
 19 refers to the homogeneous freezing of solution droplets after Koop et al. (2000). Adapted from Hoose and Möhler
 20 (2012).

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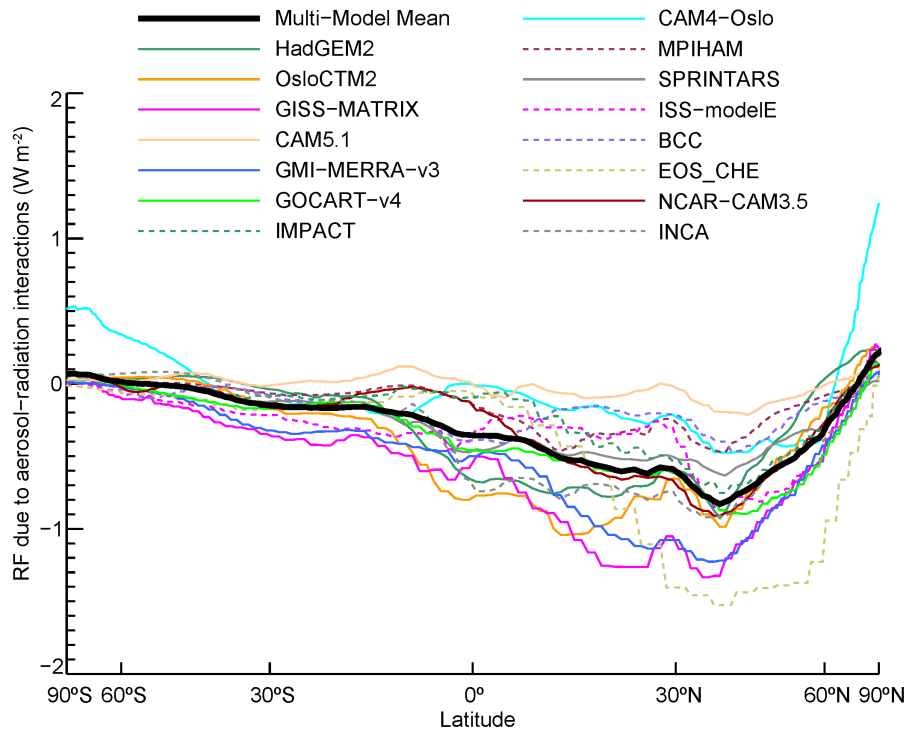
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Figure 7.16: Schematic depicting the myriad aerosol-cloud-precipitation related processes occurring within a typical GCM grid box. The schematic conveys the importance of considering aerosol-cloud-precipitation processes as part of an interactive system encompassing a large range of spatiotemporal scales. Cloud types include low-level stratocumulus and cumulus where research focuses on droplet activation, mixing, cloud scavenging, and new particle formation; ice-phase cirrus clouds where a key issue is homogeneous freezing; and deep convective clouds where some of the key questions relate to aerosol influences on liquid, ice, and liquid-ice pathways for precipitation formation, cold pool formation, and scavenging. These processes influence the short- and longwave forcing of the system and hence climate.

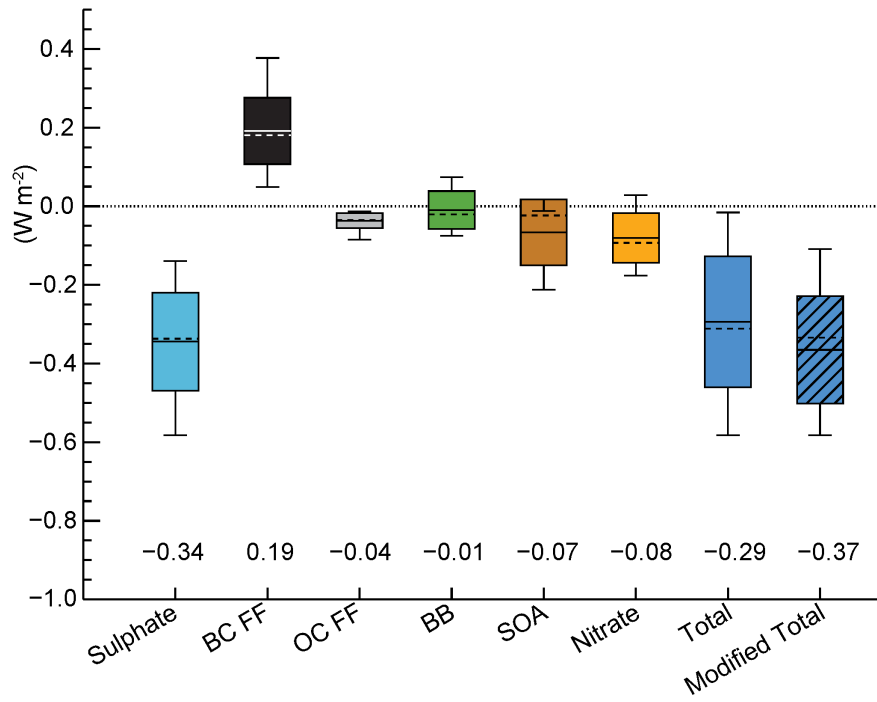
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Figure 7.17: Annual zonal mean RFar (in W m⁻²) due to all anthropogenic aerosols from the different AeroCom II models. No adjustment for missing species in certain models has been applied. The multi-model mean is shown with a black solid line. Adapted from Myhre et al. (2012).

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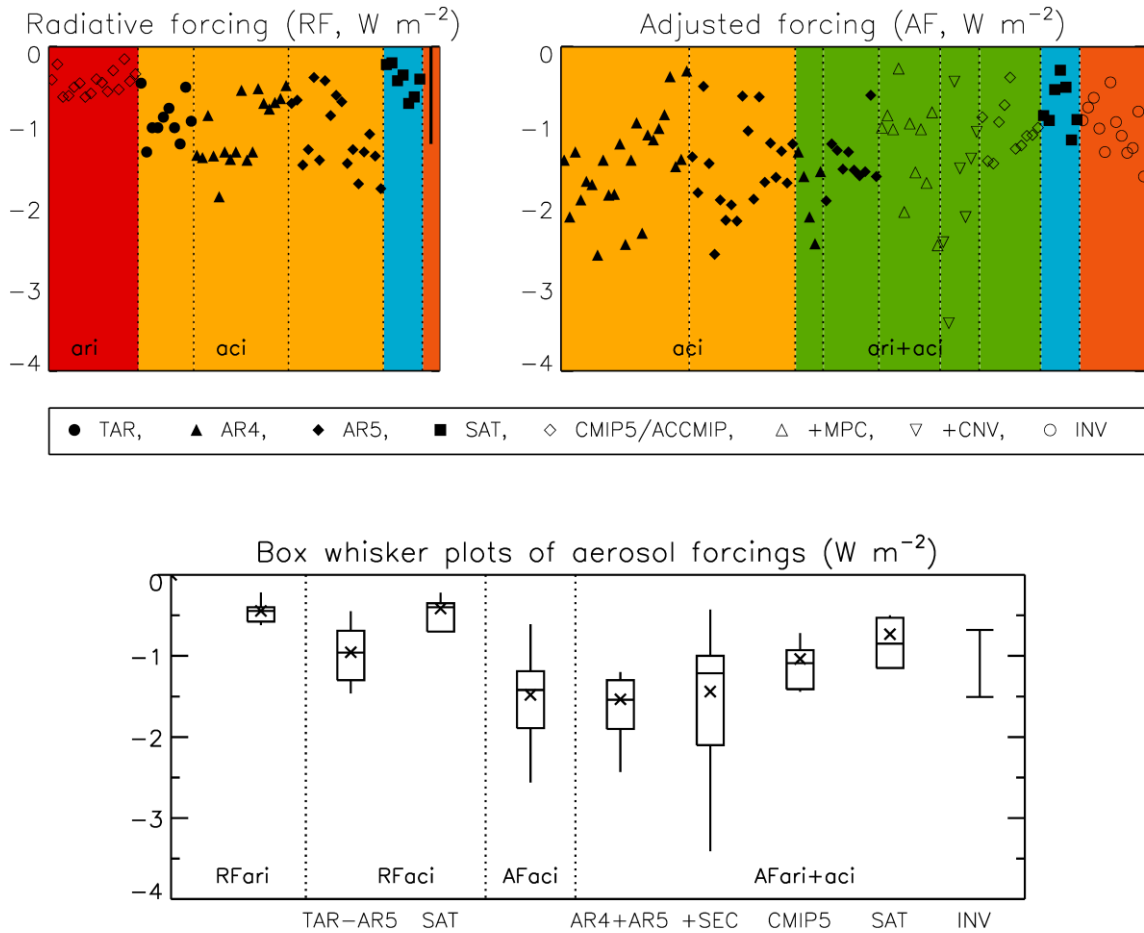
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Figure 7.18: Mean (solid line), median (dashed line), one standard deviation (box) and full (min-max) range (whiskers) for RForc (in $W m^{-2}$) from different aerosol types from AeroCom II models. The forcings are for the 1850 to 2000 period. Adapted from Myhre et al. (2012).

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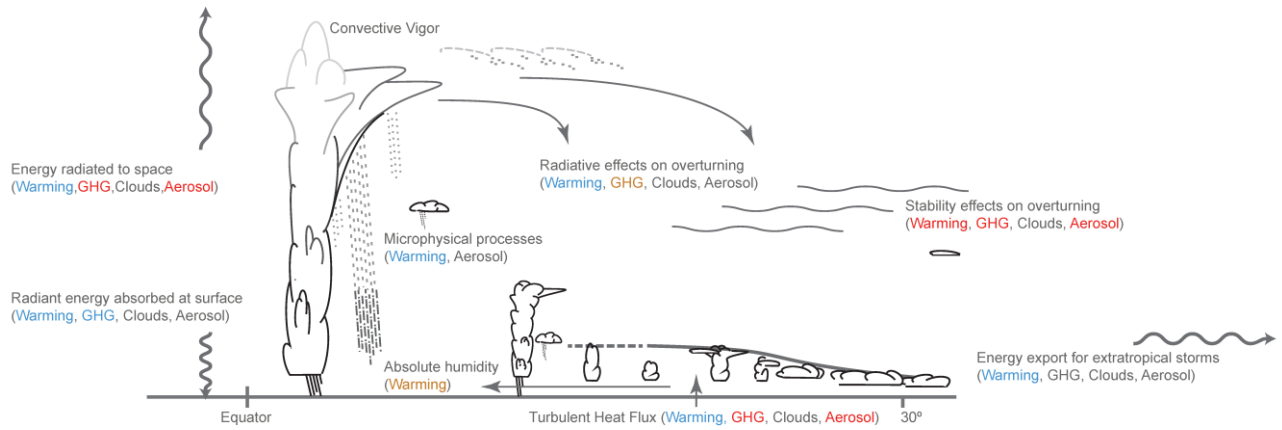
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Figure 7.19: Upper panel: GCM, satellite and inverse estimates of RFari, RFaci, AFaci and AFari+aci. Each symbol represents the best estimate per model and paper as detailed in Table 7.4. The RFaci studies are divided into those from GCMs published prior to TAR, AR4 and AR5, those including satellite data (SAT) and the inverse estimate (INV). AFaci and AFari+aci studies from GCMs on liquid stratiform clouds are also divided into those published prior to AR4 and AR5 and from the CMIP5/ACCMIP models. GCM estimates that include adjustments beyond aci in liquid stratiform clouds are marked +MPC when including aci in mixed-phase clouds and are marked +CNV when including aci in convective clouds. For RFaci from inverse estimates the range instead of the best estimate is given because it is only one study.

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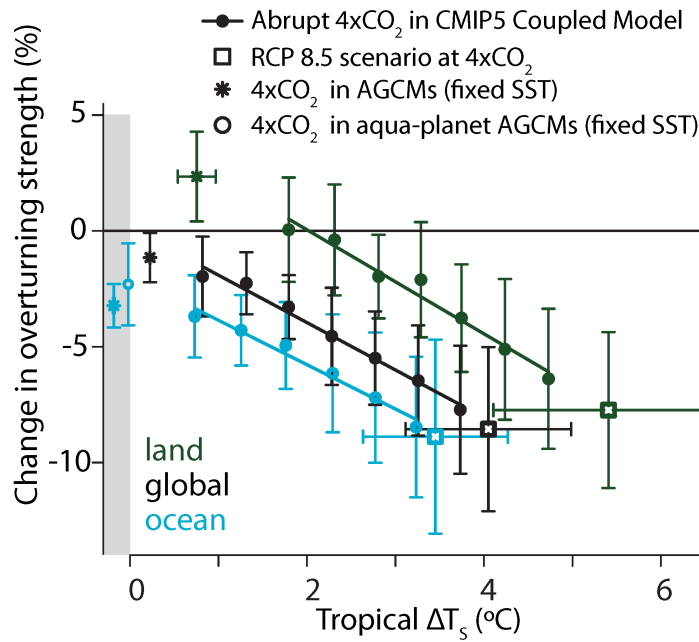
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Figure 7.20: Illustration of major drivers affecting precipitation. Radiative drivers cool the atmosphere, warm the surface, and thereby provide the energy for evaporation and condensation/precipitation. Circulations organize and distribute precipitation. The ability of changes in the position or strength of circulation features to change the distribution of precipitation is referred to as a dynamic effect, the ability of warmer circulations to transport more water vapour and thereby change the amount of precipitation is referred to as a thermodynamic effect. The immediate effect of warming, greenhouse gases, clouds and aerosols on precipitation is indicated by blue or red if their change over the 20th century is thought to have changed a precipitation driver in a way that will increase, respectively decrease, precipitation. Grey indicates that changes are unknown or have multiple effects of different sign.

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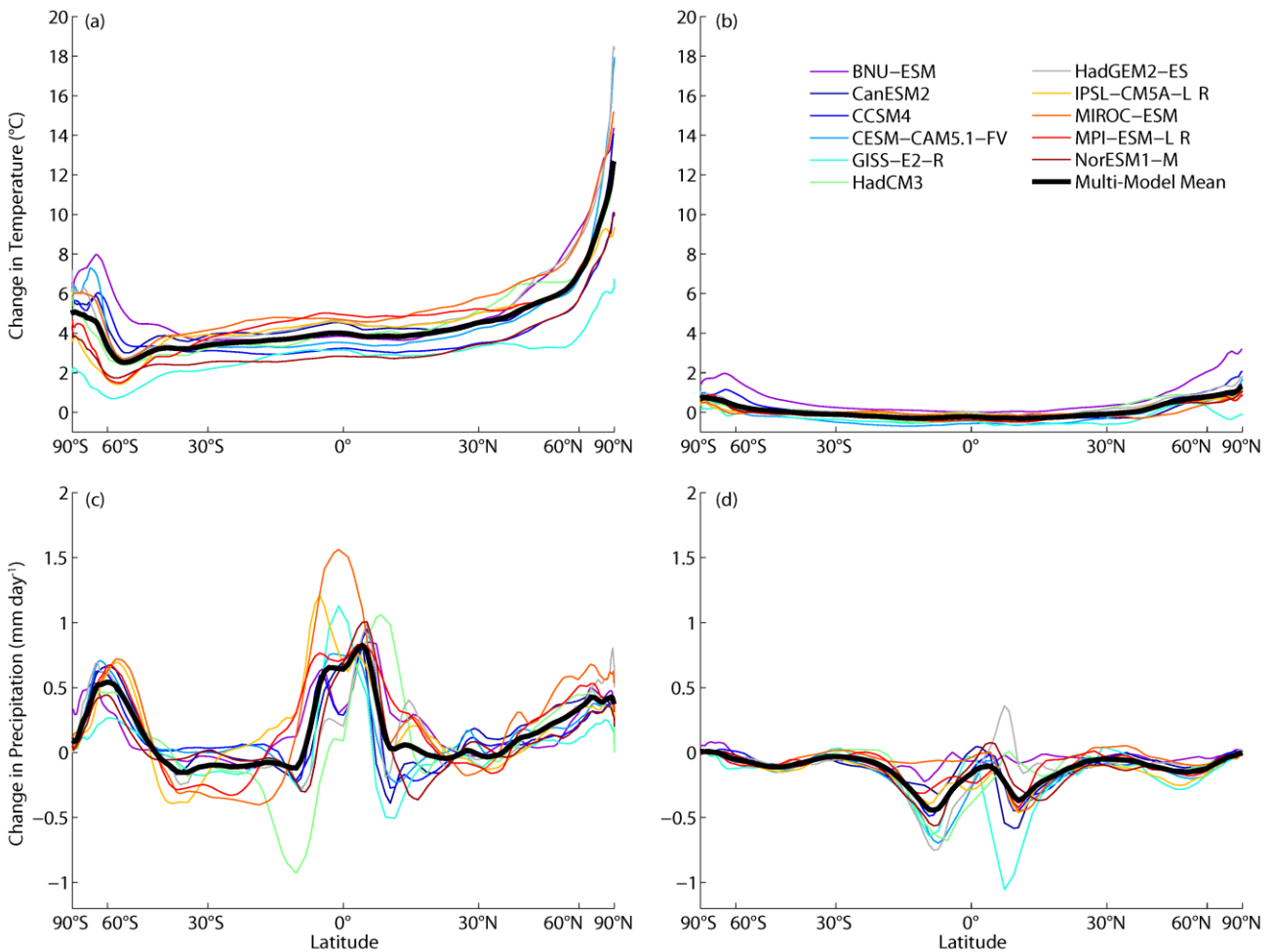
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Figure 7.21: Illustration of the response of the large-scale overturning to increasing CO₂ concentrations (adapted from Bony et al., 2012). Approximately half of the response is evident before any warming is felt, but additional warming continues to slow down the circulation and adds linearly to the rapid adjustment. The rapid adjustment is different over land and ocean, with the increase in CO₂ initially causing an intensification of the circulation over land. The robustness of the result is illustrated by the common behaviour of 15 CMIP5 models, irrespective of the details of their configuration.

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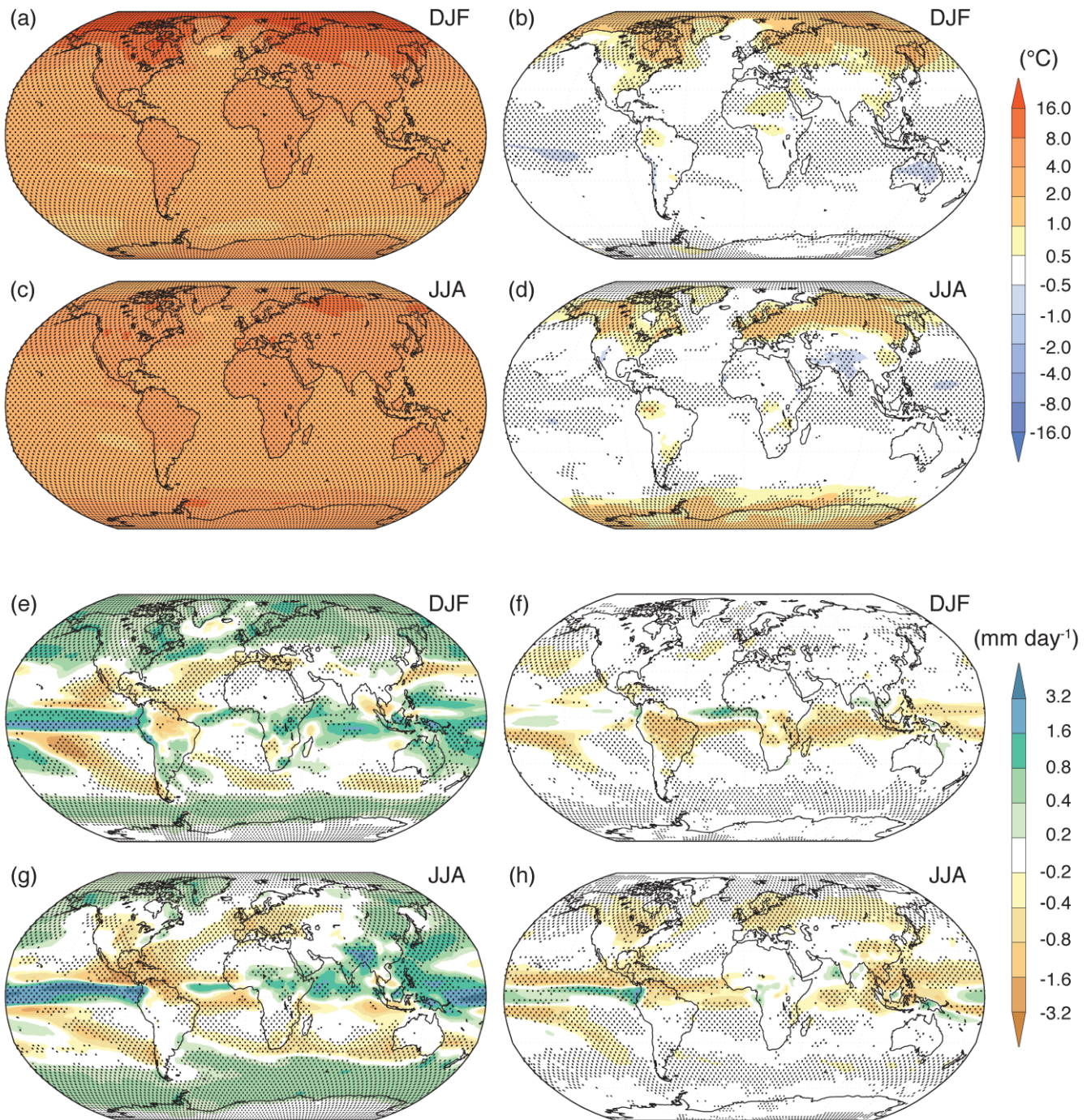
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Figure 7.22: Zonally- and annually-averaged change in surface air temperature ($^{\circ}\text{C}$) for (a) an abrupt $4 \times \text{CO}_2$ experiment and (b) the GeoMIP G1 experiment for 11 coupled atmosphere-ocean general circulation models. (c) and (d) Same as (a) and (b) but for the change in precipitation (mm day^{-1}). In the GeoMIP G1 experiment (Kravitz et al., 2011) an abrupt fourfold increase in CO_2 concentration is balanced by a reduction in total solar irradiance to produce a top of atmosphere flux imbalance of less than $\pm 0.1 \text{ W m}^{-2}$ during the first 10 years of the simulation. All changes are relative to the pre-industrial control experiment and averaged over years 21–50.

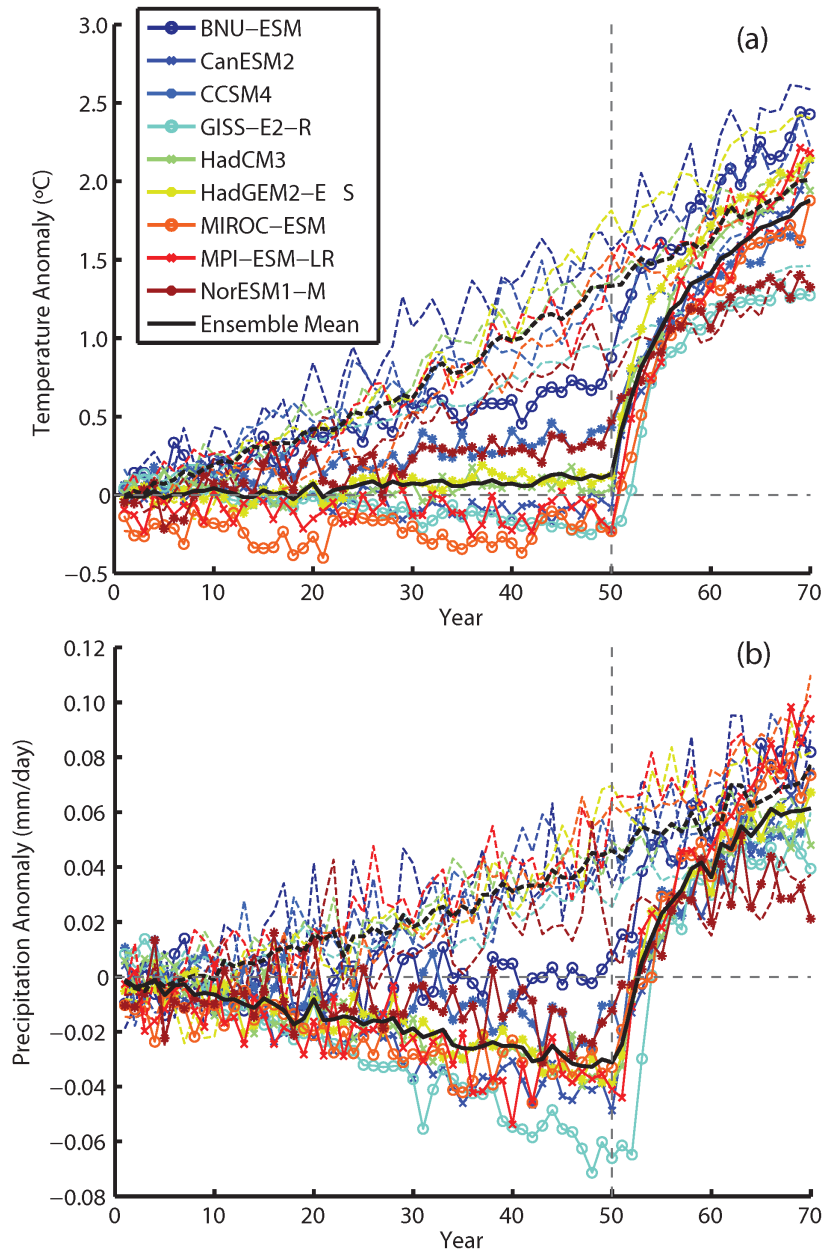
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Figure 7.23: Multi-model mean of the change in surface air temperature (°C) averaged over December, January and February months for (a) the abrupt $4 \times \text{CO}_2$ simulation and (b) the GeoMIP G1 experiment. (c-d) same as (a-b) but for the June, July and August months. (e-h) same as (a-d) but for the change in precipitation (mm day^{-1}). All changes are relative to the pre-industrial control experiment and averaged over years 21-50. Stippling denotes agreement on the sign of the temperature anomaly in at least 75% of the models in panels.

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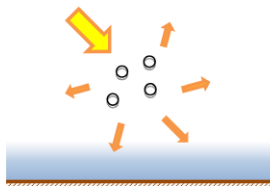
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Figure 7.24: Timeseries of surface temperature ($^{\circ}\text{C}$, left) and precipitation change (mm day^{-1} , right) for GeoMIP experiment G2, relative to each model’s $1 \times \text{CO}_2$ reference simulation. Solid lines are simulations using SRM to balance a $1\% \text{ yr}^{-1}$ increase in CO_2 concentration until year 50 after which SRM is stopped. Dashed lines are for $1\% \text{ CO}_2$ increase simulations with no SRM.

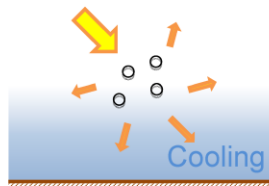
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Aerosol-radiation interactions

Scattering aerosols

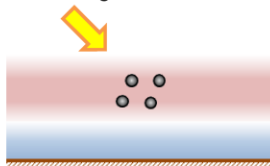


Aerosols scatter solar radiation. Less solar radiation reaches the surface, which leads to a localized cooling.

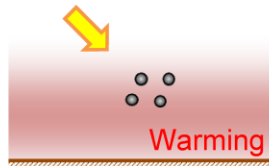


The atmospheric circulation and mixing processes spread the cooling regionally and in the vertical.

Absorbing aerosols



Aerosols absorb solar radiation. This heats the aerosol layer but the surface, which receives less solar radiation, can cool locally.



At the larger scale there is a net warming of the surface and atmosphere because the atmospheric circulation and mixing processes redistributes the thermal energy.

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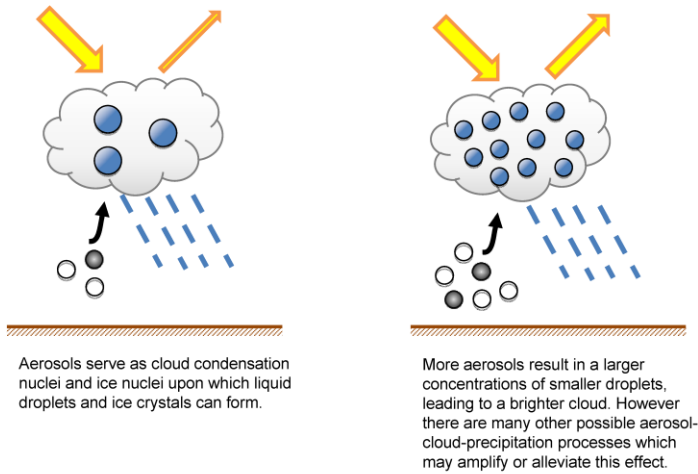
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FAQ 7.2, Figure 1: Overview of aerosol-radiation interactions and their impact on climate. The left panels show the instantaneous radiative effects of aerosols, while the right panels show their overall impact after the climate system has responded to their radiative effects.

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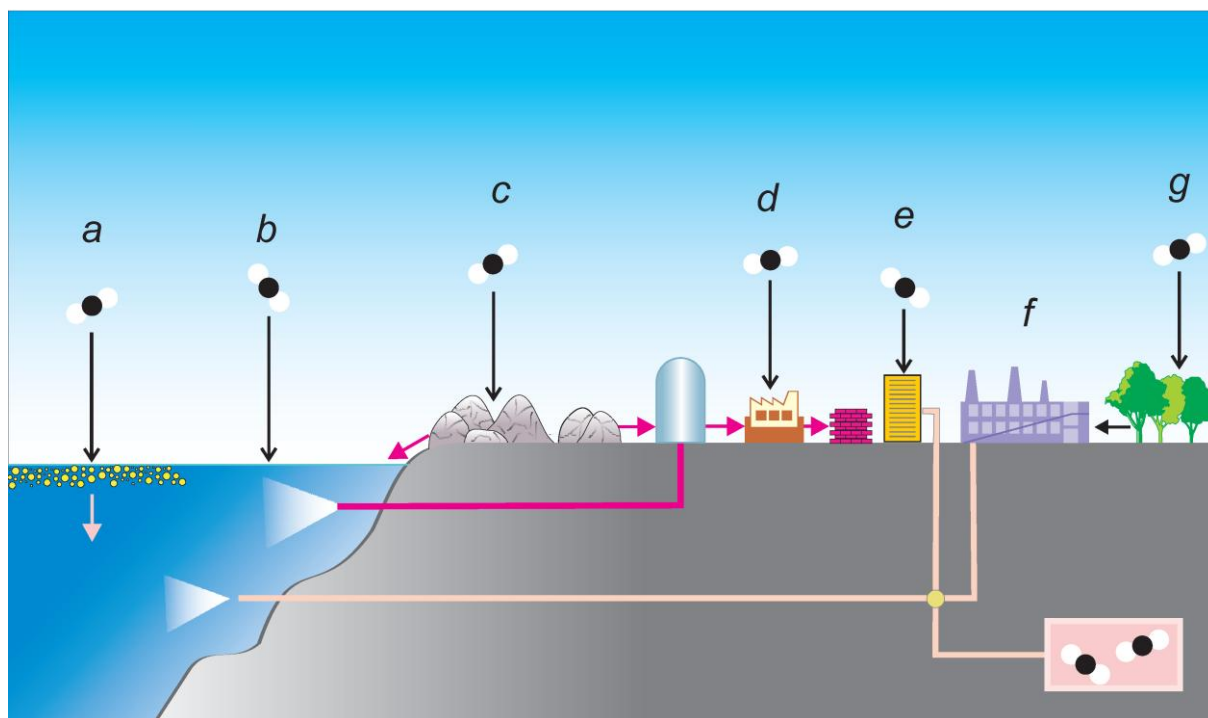
Aerosol-cloud interactions



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FAQ 7.2, Figure 2: Overview of aerosol-cloud interactions and their impact on climate. The left and right panels represent a clean and a polluted low-level cloud, respectively.

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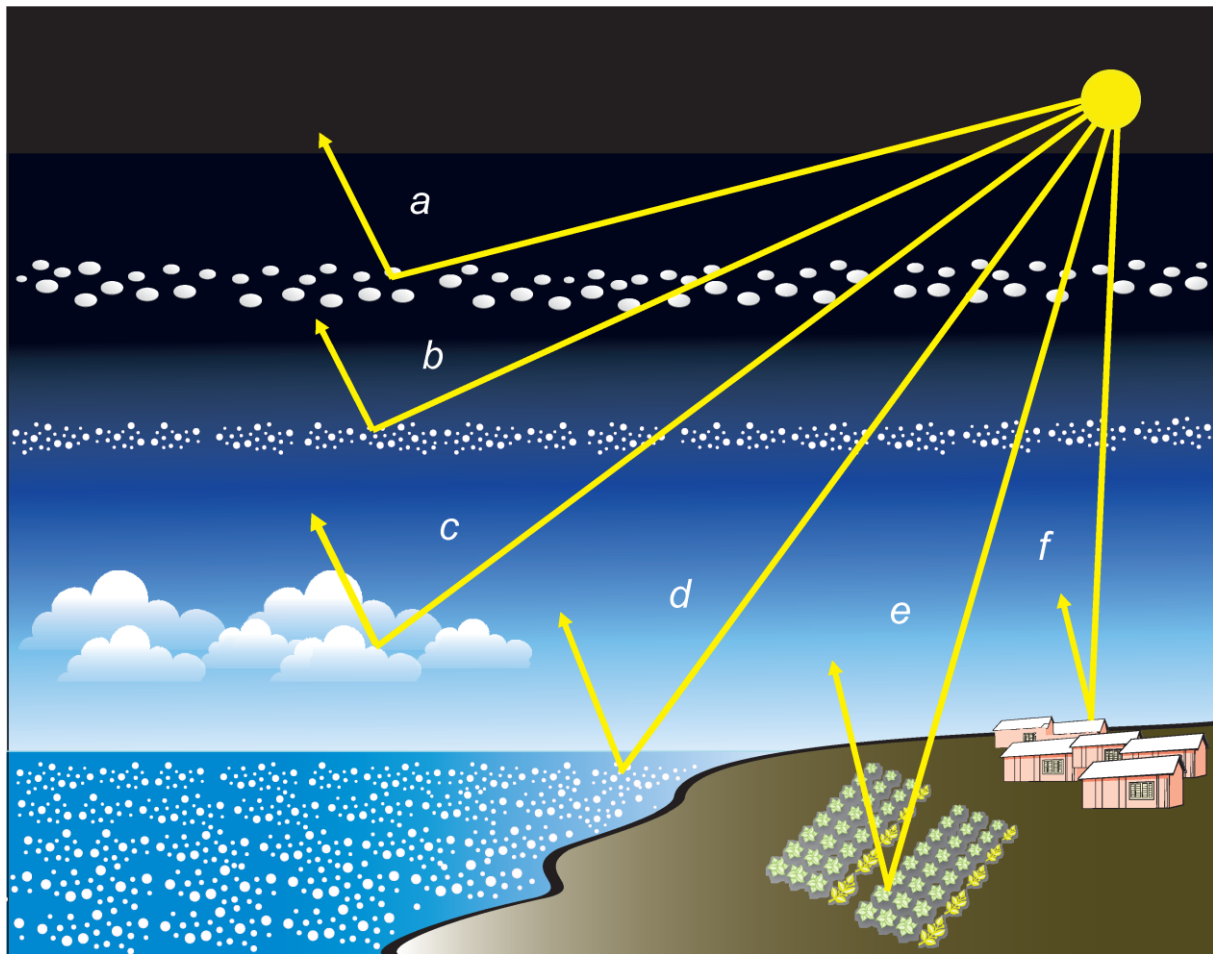
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FAQ 7.3, Figure 1: Overview of some carbon dioxide removal methods. (a) nutrients are added to the ocean, which increases oceanic productivity in the surface ocean and transports a fraction of the resulting biogenic carbon downward, (b) alkalinity from solid minerals is added to the ocean, which causes more atmospheric CO₂ to dissolve in the ocean, (c) the weathering rate of silicate rocks is increased, and the dissolved carbonate minerals are transported to the ocean, (d) alkalinity from mined silicate rocks is extracted, then combined with atmospheric CO₂ to produce solid carbonate minerals, (e) atmospheric CO₂ is captured chemically, and stored either underground or in the ocean, (f) biomass is burned at an electric power plant with carbon capture, and the captured CO₂ is stored either underground or in the ocean and (g) CO₂ is captured through afforestation and reforestation to be stored in land ecosystems.

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FAQ 7.3, Figure 2: Overview of some proposed solar radiation management (SRM) schemes to reflect sunlight to space that would otherwise be absorbed. Illustrated methods, counter-clockwise from upper left, are (a) reflectors in space, (b) aerosols in the stratosphere, (c) enhanced reflectivity of marine clouds, (d) making the ocean surface more reflective, (e) growing more reflective crops, and (f) whitening of roofs and other built structures.