Effect of volcanic emissions on clouds during the 2008 and 2018 Kilauea degassing events

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Abstract. Volcanic eruptions in otherwise clean environments are “natural experiments” wherein the effects of aerosol emissions on clouds and climate can be partitioned from meteorological variability and anthropogenic activities. In this work, we combined satellite retrievals, reanalysis products, and atmospheric modeling to analyze the mechanisms of aerosol–cloud interactions during two degassing events at the Kilauea volcano in 2008 and 2018. The eruptive nature of the 2008 and 2018 degassing events was distinct from long-term volcanic activity for Kilauea. Although previous studies assessed the modulation of cloud properties from the 2008 event, this is the first time such an analysis has been reported for the 2018 event and that multiple degassing events have been analyzed and compared at this location. Both events resulted in significant changes in cloud effective radius and cloud droplet number concentration that were decoupled from local meteorology and in line with an enhanced cloud albedo. However, it is likely that the effects of volcanic emissions on liquid water path and cloud fraction were largely offset by meteorological variability. Comparison of cloud anomalies between the two events suggested a threshold response of aerosol–cloud interactions to overcome meteorological effects, largely controlled by aerosol loading. In both events, the ingestion of aerosols within convective parcels enhanced the detrainment of condensate in the upper troposphere, resulting in deeper clouds than observed under pristine conditions. Accounting for ice nucleation on ash particles led to enhanced ice crystal concentrations at cirrus levels and a slight decrease in ice water content, improving the correlation of the model results with the satellite retrievals. Overall, aerosol loading, plume characteristics, and meteorology contributed to changes in cloud properties during the Kilauea degassing events.

1 Introduction

Aerosol emissions influence Earth’s climate both directly and indirectly. The direct effect involves scattering and absorption of thermal and solar radiation by atmospheric aerosols, while indirect effects involve alteration of the microphysical properties and the global distribution of clouds (Boucher et al., 2013; Twomey, 1977). Both liquid and ice clouds are susceptible to aerosol emissions that can alter their microphysical (i.e., particle size distribution and albedo) and macrophysical properties (liquid and ice water content, cloud lifetime, and cloud fraction) (Lohmann and Feichter, 2005; Boucher et al., 2013; Seinfeld et al., 2016). These effects, collectively known as aerosol indirect effects (AIEs), may offset a significant fraction of the warming induced by greenhouse gas emissions, yet their magnitudes are poorly constrained (Boucher et al., 2013). Additionally, cloud formation is a complex and nuanced physical process occurring on scales far smaller than those resolved by climate models, and the precise feedback mechanisms influencing AIEs across various timescales are not fully understood (Boucher et al., 2013; Klein et al., 2013; Malavelle et al., 2017; Yuan et al., 2011).
The presence of cloud condensation nuclei (CCN; typically sulfates, organics, and nitrate particles) in the atmosphere indirectly impacts Earth’s net radiative balance by increasing the number of cloud droplets, hence altering the scattering and absorption of incoming solar radiation. This effect, historically referred as the first AIE or “Twomey” effect (Twomey, 1977), represents a change in Earth’s albedo and results in net radiative cooling. Smaller droplets are also less efficient at coalescing into rain-bearing clouds. Non-precipitating clouds have a longer lifetime and thereby provide extended coverage, which is known as the second AIE (Albrecht, 1989). Aerosols can also act as ice nucleation particles (INPs; typically dust, soot, and organics), modifying cloud properties at low temperature (Lohmann and Feichter, 2005). Besides these effects, CCN and INP ascend within convective parcels and determine, to a large extent, the onset of precipitation, modifying the release of latent heat within convective clouds (Koren et al., 2005).

Satellite datasets facilitate global observational monitoring of meteorology, ambient aerosol concentrations and distributions, and cloud properties. However, inferring aerosol–cloud interactions (ACIs) from satellite retrievals is difficult due to the concurrent influence of meteorological effects (i.e., variation in temperature, water vapor, and winds from large-scale forcing) on clouds (Lohmann and Feichter, 2005; Gryspeerdt et al., 2017). From a modeling perspective, the relevant scale for ACIs (i.e., tens to hundreds of meters) is typically unresolved in atmospheric general circulation models (AGCMs; Boucher et al., 2013) for which grid resolution is coarser (generally thousands of meters). The parameterization of ACIs in AGCMs is largely dependent on theory, with many assumptions involved, and model outputs are difficult to validate directly against satellite retrievals due to differences in resolution.

Volcanic degassing events in otherwise “clean” environments, where anthropogenic aerosol emissions are minimal, have been used as “natural” experiments to identify characteristics of ACIs in satellite retrievals and to evaluate AIEs in GCMs. This is well-exemplified at the Kilauea volcano (Fig. 1) in the Hawaiian Islands (Eguchi et al., 2011; Yuan et al., 2011; Beirle et al., 2014; Malavelle et al., 2017). Not only are local anthropogenic emissions low in the local environment surrounding Kilauea, but because the volcano is situated on an island approximately 1500 km from the nearest major anthropogenic emissions source (US west coast), there is a low likelihood of multiple emission sources of similar magnitude; therefore, the climatic effects of volcanic emissions may be evaluated against relatively pristine background conditions. Kilauea has had long-term monitoring of emissions, seismic activity, and eruptive behavior over the historical record. Being a low-altitude volcano, Kilauea also provides a unique opportunity to study ACIs for liquid and ice cloud phases since the injection height of the emissions is mainly controlled by the strength of the eruption.

In this work, we assessed the effects of SO$_2$ and ash aerosol emissions on liquid and ice cloud formation during two volcanic degassing events from Kilauea in June 2008 and May 2018 (Fig. 1). In both of these events, atmospheric aerosol loading increased by several orders of magnitude over a few days and remained high for at least 2 to 3 months. The volcanic and seismic activity during peak emissions in 2008 and 2018 has been well-characterized (Nadeau et al., 2015; Neal et al., 2019; Elias and Sutton, 2012; Elias et al., 2020; Wilson et al., 2008; Orr and Patrick, 2009; Orr et al., 2013; Patrick et al., 2019; Kern et al., 2020), revealing that emission rates of SO$_2$ during the 2018 event were conservatively at least $\geq 5\times$ higher than 2008 peak degassing emissions (Kern et al., 2020; Elias et al., 2020). Analysis of the satellite record has shown significant anomalies in aerosol loading and liquid cloud properties during the 2008 event resulting from volcanic emissions (Eguchi et al., 2011; Yuan et al., 2011; Beirle et al., 2014), in agreement with modeling studies (Malavelle et al., 2017). Tang et al. (2020) showed significant effects of the 2018 Kilauea emissions on air quality. However, analyses of ice clouds and of the impacts of the 2018 event on cloud properties and evolution have not yet been reported. This work constitutes a substantial contribution towards identifying ACI signatures for liquid and ice macrophysical and microphysical processes as well as their sensitivity to aerosol loadings, CCN, and INPs.

This work is organized as follows. Section 1.1 provides an overview of each degassing event. Satellite datasets used in the analysis are described in Sect. 2, and the modeling approach is described in Sect. 3. Section 4 discusses the results of our analysis, with an emphasis on liquid and ice cloud interactions. Finally, concluding remarks are given in Sect. 5.

### 1.1 Kilauea degassing events

Kilauea is an active basaltic shield volcano located on the Island of Hawai‘i. Volcanic plumes on Kilauea are generally low-altitude (< 10 km) because the gentle slopes of the volcano provide little protection from strong trade winds and because of the shallow depth of the summit crater (< 100 m) (Elias et al., 2018). Since March 2008, volcanic activity on Kilauea has been marked by summit degassing and flank eruptions with increasing intensity and frequency (Nadeau et al., 2015). Degassing events were likely triggered by rockfalls related to vent widening and/or seismic activity, which then disturbed the lava lake surface, beneath which a layer of gas had accumulated (Orr and Patrick, 2009; Orr et al., 2013). Degassing events produced variable volumes of tephra (ash), and some larger explosive events scattered lithic material (rock and ash) within a radius as large as 50 ha surrounding the vent and produced an ash-rich plume (Elias and Sutton, 2012; Elias et al., 2020; Nadeau et al., 2015). The summit of Kilauea has been in an eruptive state since 2008, and the degassing events of 2008 and 2018 represent brief periods of increased volcanic activity and SO$_2$ emissions, resulting in
Figure 1. © Google Earth imagery of the Kilauea summit crater before and after each degassing event. Clockwise from top left: pre-2008 degassing, post-2008 degassing, during 2018 peak degassing, and post-2018 degassing. The image from 5 June 2011 shows the summit vent, which was not present prior to the 2008 event, passively degassing during an eruptive lull. The image from 30 May 2018 was taken on a cloudy day during the 2018 event, several days after peak observed SO$_2$ emissions (> 100 kt d$^{-1}$) and plume height (< 8 km) (Kern et al., 2020). The red boxes approximately outline the summit crater, while the X symbols indicate the degassing vent.

an optically denser plume relative to passive degassing. Some studies have noted residual effects of the plume downwind of Kilauea long after violent eruptions ceased (Businger et al., 2015; Pattantyus et al., 2018).

From March to August 2008, summit activity was characterized by degassing bursts from lava in the enlarging vent cavity and small explosive events. During degassing events, the estimated SO$_2$ plume height ranged from 1200 to 2500 m above sea level and SO$_2$ emissions exceeded 10 000 t d$^{-1}$ (Elias et al., 2020). Observational and modeling studies revealed a significant departure of cloud droplet size and optical thickness from their climatological values during the 2008 event, consistent with increased concentrations of atmospheric aerosols within the SO$_2$ plume from the Kilauea summit crater (Eguchi et al., 2011; Yuan et al., 2011; Beirle et al., 2014; Mace and Abernathy, 2016; Malavelle et al., 2017). Figure 1 (top row) shows images before and after the 2008 degassing event. After the 2008 event, the vent can be seen in the SE corner of the summit crater passively degassing from the lava lake during an eruptive pause.

From May to July 2018, summit eruptions and effusive activity in the eastern rift zone (ERZ) were common. Summit activity ejected lithic material and ash ≈ 2000 to 8100 m above the summit vent and SO$_2$ concentrations were elevated (Neal et al., 2019; Kern et al., 2020). ERZ eruptive events (lava fountains < 80 m) were associated with SO$_2$ emissions of ≥ 100 000 t d$^{-1}$ (Neal et al., 2019; Kern et al., 2020; Elias et al., 2018). Multiple cycles of ongoing short pulses of eruptive degassing activity at the vent and longer periods of effusive volcanism in the ERZ caused ongoing aerosol emissions (Patrick et al., 2019). While in 2008 most degassing occurred at the summit vent, in 2018 degassing largely occurred in the ERZ and drained lava beneath the summit vent, causing massive deformation of the summit crater, as shown in Fig. 1 (lower left). Additionally, the ocean entry of 2018 ERZ eruptions caused large clouds of vaporized HCl and water vapor to ascend with the plume (Kern et al., 2020). Overall, the height and geologic composition of the plume in 2018 were similar to more violent, siliciclastic eruption types (i.e., Mt. St. Helens; Mastin et al., 2009); therefore, the 2018 event represented a distinct departure from long-term (decadal) trends. Figure 1 (bottom row) shows the summit crater before, during, and after peak eruptive activity in 2018 and indicates optically denser clouds within the aerosol plume with respect to surrounding clouds during peak degassing (Fig. 1, lower right).

2 Data

Data from NASA’s MODIS instrument and from the Cloud–Aerosol Lidar and Infrared Pathfinder Satellite Observation (CALIPSO) retrievals were used to assess both horizontal and vertical cloud modification, respectively, as done in previous studies for liquid clouds (Yuan et al., 2011; Malavelle et al., 2017; Eguchi et al., 2011; Beirle
et al., 2014; Mace and Abernathy, 2016). The MODIS Aqua Aerosol Cloud Monthly Collection 6 L3 Global 1°
datasets (MYD08_M3) were acquired from the Level-1 and
Atmosphere Archive & Distribution System (LAADS) Dis-
tributed Active Archive Center (DAAC) (https://ladsweb.
nascom.nasa.gov/, last access: 15 June 2020). Our analysis
used MODIS effective radius (Reff; liquid, ice), cloud optical
density (COD; liquid, ice), cloud water path (liquid – LWP;
ice – IWP, and total – TWP), aerosol optical depth (AOD),
and cloud fraction (CF). All variables are defined in Table 1.
The MODIS cloud droplet number concentration (CDNC)
climatology (2003–2015) was obtained from Bennart and
Rausch (2017). For 2018, CDNCs were calculated using the
method of Bennart (2007). Grid box mean LWP and
IWP were calculated by scaling the MODIS product us-
ing the retrieved liquid and ice cloud fractions, respectively.
No significant differences were found by scaling using ei-
ther daily data (MYD08_D3) aggregated into monthly means
(Malavelle et al., 2017) or monthly products (MYD08_M3);
therefore, monthly MODIS products were used in the analy-
sis. MODIS anomalies were calculated as the 3-month aver-
age during peak degassing minus the long-term mean (2003–
2015, excluding 2008). Missing values in MODIS data, pri-
marily found in CDNC and ice products, were smoothed us-
ing cubic spline nearest-neighbor interpolation and a Gaus-
sian filter. Uncertainty of MODIS retrievals is discussed in
Hubanks et al. (2015), Bennartz (2007), and Bennartz and
Rausch (2017) (the latter is for CDNC only).
Vertical cloud fraction profiles from the GCM-Oriented
Cloud CALIPSO Product (CALIPSO-GOCCP) were ob-
tained for comparison with model results. The CALIPSO-
GOCCP dataset is developed with CALIPSO L1 data at
full horizontal resolution (330 m) and vertical resolution
typical for most GCMs (40 levels; Δz = 480 m) (Chepfer
et al., 2010). Instantaneous profiles of the lidar-scattering
ratio are computed and used to infer the vertical and hor-
izontal distributions of cloud fraction. The seasonal mean
uncertainty (2006–2008) for GOCCP cloud fraction is ≤
0.05 above the boundary layer (Chepfer et al., 2010). The
dataset may be used for comparison either directly to AGCM
output or to AGCM and lidar simulator output, and it is
comparable to other standard cloud fraction climatologies
(Chepfer et al., 2010). Comparisons between MODIS Col-
lection 6 and CALIPSO show agreement between MODIS
cloud-phase partitioning and CALIPSO column-wise cloud
profiles (Marchant et al., 2016). GOCCP was used primarily
to assess anomalies on the vertical structure of cloud fraction
and phase partitioning during both events.
Volcanic SO2 emissions were constrained by observa-
tions from the Ozone Monitoring Instrument (OMI) on board
NASA’s EOS/Aura spacecraft (Carn et al., 2015). For Ki-
laua, this dataset only provides “constant” annual SO2 emis-
sion rates. For the 2008 event, we replaced this dataset with
daily varying emissions (Carn et al., 2017; Yang, 2017).
Daily emissions for 2018 were obtained from Li et al. (2020).
Missing values were replaced with Ozone Mapping and
Profiling Suite data (https://so2.gsfc.nasa.gov/, last access:
15 September 2020) whenever possible; otherwise, the near-
est real data point was used. SO2 (cm−2) vertical column
density data were converted to emission rates (kg SO2 s−1)
following the approach of Beirle et al. (2014) (see their
Fig. 6). For uncertainty quantification of OMI retrievals, see

3 Methods
To help explain the observed changes in cloud properties
during the 2008 and 2018 Kilauea degassing events and to what
these changes were sensitive, we undertook a set of AGCM
numerical experiments (Table 2) constrained by observed
SO2 emissions (Carn et al., 2015) and by MERRA-2 (Gelaro
et al., 2017). Our objective was to generate a close repre-
sentation of the clouds formed during each event to under-
stand how aerosol emissions impacted cloud microphysics
and evolution.

3.1 GEOS model description
The NASA Global Earth Observing System (GEOS) ver-
sion 5 was used to analyze and understand the observed mod-
ifications to cloud properties during the 2008 and 2018 Ki-
laua events (Barahona et al., 2014; Molod et al., 2015).
GEOS consists of a set of components that numerically re-
present different aspects of the Earth system (atmosphere,
ocean, land, sea ice, and chemistry), coupled following the
.gov/GEOS_systems/, last access: 10 August 2019). For this
work, the AGCM configuration of GEOS was used. Atmo-
spheric transport of water vapor, condensate and other trac-
ers, and associated land–atmosphere exchanges were com-
puted explicitly, whereas sea ice and sea surface tempera-
ture (SST) were prescribed as time-dependent boundary con-
ditions (Reynolds et al., 2002; Rienecker et al., 2008).
Transport of aerosols and gaseous tracers such as CO was
simulated using the Goddard Chemistry Aerosol and Radi-
ation model (GOCART) (Colarco et al., 2010), which inter-
actively calculates the transport and evolution of dust, black
carbon, organic material, sea salt, and SO2. Dust and sea salt
emissions are prognostic, whereas biomass burning and an-
thropogenic emissions of SO2, black carbon, and organic car-
bon are obtained from the Modern Era Retrospective Reanal-
ysis for Research and Applications Version 2 (MERRA-2)
dataset (Randles et al., 2017). GOCART explicitly calculates the
chemical conversion of sulfate precursors (dimethylsul-
fide, or DMS, and SO2) to sulfate. The aging of carbona-
ceous aerosol is represented by the conversion of hydrophi-
obic to hydrophilic aerosols using an e-folding time of 2 d
(Chin et al., 2009). Using the evolving meteorological fields
from GEOS, for each time step GOCART simulates the ad-
Table 1. Variable definitions and acronyms.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACRI_SNOW</td>
<td>Ice crystal accretion by snow tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>ACRL_(RAIN,SNOW)</td>
<td>Accretion of liquid by rain and/or snow tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>AOD</td>
<td>Aerosol optical depth</td>
<td>–</td>
</tr>
<tr>
<td>AUT</td>
<td>Liquid autoconversion tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>AUTICE</td>
<td>Ice autoconversion tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>CCN</td>
<td>Cloud condensation nuclei</td>
<td>m(^{-3})</td>
</tr>
<tr>
<td>CDNC</td>
<td>Cloud droplet number concentration</td>
<td>m(^{-3})</td>
</tr>
<tr>
<td>CF</td>
<td>Cloud fraction</td>
<td>–</td>
</tr>
<tr>
<td>CNV_DQLDT</td>
<td>Total detrained condensate tendency</td>
<td>mg kg(^{-1}) s(^{-1})</td>
</tr>
<tr>
<td>COD</td>
<td>Cloud optical depth</td>
<td>–</td>
</tr>
<tr>
<td>COND</td>
<td>Liquid condensation tendency</td>
<td>kg(^{-1}) kg(^{-1})</td>
</tr>
<tr>
<td>DEP</td>
<td>Ice crystal growth tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>DCNVl</td>
<td>Ice convective detrainment tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>EVAP</td>
<td>Droplet evaporation tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>HM</td>
<td>Ice splintering tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>ICENC</td>
<td>Ice nucleation tendency</td>
<td>cm(^{-3}) s(^{-1})</td>
</tr>
<tr>
<td>ICNC</td>
<td>Ice crystal number concentration</td>
<td>L(^{-1})</td>
</tr>
<tr>
<td>IWP</td>
<td>Ice water path</td>
<td>g m(^{2})</td>
</tr>
<tr>
<td>LWP</td>
<td>Liquid water path</td>
<td>g m(^{2})</td>
</tr>
<tr>
<td>MELT</td>
<td>Ice melt and/or snowmelt tendency</td>
<td>kg(^{-1}) kg(^{-1})</td>
</tr>
<tr>
<td>NHET_IMM</td>
<td>Ice nucleation by immersion freezing</td>
<td>cm(^{-3})</td>
</tr>
<tr>
<td>SDM</td>
<td>Ice sedimentation tendency</td>
<td>kg(^{-1}) kg(^{-1}) s(^{-1})</td>
</tr>
<tr>
<td>SCF</td>
<td>Supercooled cloud fraction</td>
<td>–</td>
</tr>
<tr>
<td>(R_{\text{eff}})</td>
<td>Effective radius</td>
<td>(\mu m)</td>
</tr>
<tr>
<td>TWP</td>
<td>Total water path</td>
<td>g m(^{2})</td>
</tr>
<tr>
<td>(Q_{\text{liq}})</td>
<td>Liquid mixing ratio</td>
<td>mg kg(^{-1})</td>
</tr>
<tr>
<td>(Q_{\text{ice}})</td>
<td>Ice mixing ratio</td>
<td>mg kg(^{-1})</td>
</tr>
<tr>
<td>WBF</td>
<td>Wegener–Bergeron–Findeisen process (ice)</td>
<td>kg(^{-1}) kg(^{-1})</td>
</tr>
<tr>
<td>WBFSNOW</td>
<td>Wegener–Bergeron–Findeisen process (snow)</td>
<td>kg(^{-1}) kg(^{-1})</td>
</tr>
</tbody>
</table>

Table 2. Simulation experiments performed.

<table>
<thead>
<tr>
<th>Year</th>
<th>Experiment name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2008</td>
<td>2008_1x</td>
<td>Control simulation</td>
</tr>
<tr>
<td></td>
<td>2008_0x</td>
<td>No emissions from Kilauea</td>
</tr>
<tr>
<td></td>
<td>2008_5x</td>
<td>5-fold increase in Kilauea emissions</td>
</tr>
<tr>
<td></td>
<td>2008_PH2km</td>
<td>Volcanic plume height increased to 2 km</td>
</tr>
<tr>
<td>2018</td>
<td>2018_1x</td>
<td>Control simulation</td>
</tr>
<tr>
<td></td>
<td>2018_0x</td>
<td>No emissions from Kilauea</td>
</tr>
<tr>
<td></td>
<td>2018_PH4km</td>
<td>Volcanic plume height increased to 4 km</td>
</tr>
<tr>
<td></td>
<td>2018_PH4km_ash</td>
<td>2018_PH4km plus ice nucleation on ash particles</td>
</tr>
<tr>
<td>2000–2017</td>
<td>GEOSSCLIM</td>
<td>GEOS climatology excepting 2008</td>
</tr>
</tbody>
</table>

Convective transport, and the wet and dry deposition of aerosol tracers. The calculation of AOD is a function of aerosol size distribution, refractive indices, and hygroscopic growth. Each aerosol type is assumed to be externally mixed. Size distributions are prescribed for different types using five bins for dust and sea salt and single log-normal modes for other aerosol components (Colarco et al., 2010; Chin et al., 2009). This approach was also employed to
estimate the aerosol number concentration used in the calculation of aerosol–cloud interactions (Barahona et al., 2014).

Cloud microphysics in GEOS are described using a two-moment scheme wherein the mixing ratio and number concentration of cloud droplets and ice crystals are prognostic variables (Barahona et al., 2014; Morrison and Gettelman, 2008). The two-moment microphysical model links aerosol emissions to cloud properties and predicts the mixing ratio, number concentration, and effective radius of cloud liquid and ice, rain, and snow for stratiform clouds (i.e., cirrus, stratocumulus; Morrison and Gettelman, 2008) and convective clouds (Barahona et al., 2014). Cloud droplet activation is parameterized using the approach of Abdul-Razzak and Ghan (2000). Ice crystal nucleation is described using a physically based analytical approach (Barahona and Nenes, 2009) that includes homogeneous and heterogeneous ice nucleation and their competition. Heterogeneous ice nucleation in the immersion and deposition modes follows Ullrich et al. (2017). Vertical velocity fluctuations were constrained by non-hydrostatic, high-resolution global simulations (Barahona et al., 2017). GEOS has been shown to reproduce the global distribution of clouds, radiation, and precipitation in agreement with satellite retrievals and in situ observations (Barahona et al., 2014).

3.2 Description of simulation experiments

We performed several global integrations of GEOS to best isolate the effects of volcanic aerosol emissions on cloud development. Each simulation was initialized on 1 January of each year and run at a nominal horizontal resolution of 0.5° with 72 vertical levels. The time step was set to 450 s to resolve the large-scale transport of aerosol and condensate. Cloud microphysics were subcycled twice each time step to account for unresolved, fast microphysical processes such as CCN activation (Morrison and Gettelman, 2008). For each event, control runs (identified as 1×, Table 2) used the default model and emissions as described in Sect. 2. A second set of simulations was performed for each event, removing the emissions from Kilauea (0×) to represent background conditions unaffected by the degassing events. Sensitivity studies were also performed. A 5-fold actual emissions run (2008_5×) was performed to compare the 2008 and 2018 events (the 2018 event emitted ≥5 times the aerosol load of the 2008 event) to assess the similarities between the events with respect to increased aerosol loading. Simulations to test the effect of SO2 injection height (i.e., distribution of emissions constrained by plume height) on cloud microphysics were also performed (2008_PH2km and 2018_PH4km). Finally, another numerical experiment was carried out for the year 2018 (2018_PH4km_ash), wherein the effects of ash as active INP were investigated, as described in Appendix A. Besides these experiments, long-term model integration (2000–2017) was performed to represent the climatology of the model (GEOSCLIM). The simulation experiments are summarized in Table 2.

To account for model drift all simulations were run in “replay” mode, wherein pre-computed analysis increments from MERRA-2 were applied to nudge the model state (i.e., horizontal winds and temperature) to the reanalysis every 6 h. The replay technique is more stable and has lower numerical drift than regular nudging (Takacs et al., 2018). Because the two-moment cloud scheme used in this work differed from the single-moment scheme used in MERRA-2, water vapor was not replayed and instead left to evolve with the model physics. Aerosol concentrations are indirectly constrained by the reanalysis since their transport and evolution, as well as the emission of dust and sea salt, depend on the model state (i.e., winds and temperature). The emission of sulfate precursors (SO2) is constrained using satellite retrievals as described in Sect. 3.1. However, aerosol concentrations were not directly nudged to the MERRA-2 product, even though the aerosol increments are also available (Randles et al., 2017). Doing so would have limited the response of clouds to aerosol (via aerosol activation) and vice versa: the response of aerosols to cloud formation and precipitation (via scavenging). Running in replay mode ensured that the effects of meteorological variability were reduced and that our simulations reproduced the assimilated atmospheric state as closely as possible. Replaying temperature (T) also minimized the role of direct and semi-direct effects in modifying cloud properties. Hence, our analysis focuses on the evolution of cloud microphysical properties.

The altitude of the Kilauea summit crater is ≈1200 m, while the top of the well-mixed boundary layer is around 2 km. For this reason, we allow the degassing volcano emissions to be distributed within 1 km above the summit crater and constrained by the boundary layer for all 1× and 0× experiments. For alternate experiments, we set an explicit volcanic aerosol injection height approximating the mean maximum plume height observed during each event (2 km in 2008; Elias and Sutton, 2012; Elias et al., 2020; Eguchi et al., 2011) (4 km in 2018; Neal et al., 2019). In this work, we assumed that the source elevation of emissions (ERZ vs. summit) was irrelevant during peak emissions but that the injection height of aerosols directly into the troposphere at different altitudes (below or above boundary layer processes) influenced cloud microphysics and macrophysical characteristics for liquid and ice clouds. We also assumed the primary aerosol to be SO2 with some percentage of ash, although it is likely that sea salt contributed to ACIs in liquid clouds, and this is recommended for inclusion in future parameterizations.

3.3 Analysis method

MODIS retrievals and the 1× GEOS experiments were compared against the long-term climatology excluding 2008 during peak degassing periods for each event. The 1× simula-
tions were also compared against the $0\times$ results for each year ($1\times-0\times$). In this way, we assessed the effects of volcanic aerosols on cloud formation separated from natural meteorological variability. This also allowed for the assessment of whether passive degassing effects present in $\text{GEOS}_{\text{CLIM}}$ were significant contributors to ACIs as opposed to active degassing events in 2008 and 2018. The $\text{GEOS}_{\text{CLIM}}$ run captured meteorological variability, while the $0\times$ scenario represents an alternative reality without volcanic emissions and with the same meteorological state as the $1\times$ runs. Correlation between observations and the $1\times-0\times$ anomaly would indicate that volcanic emissions forced ACIs that were decoupled from meteorology (i.e., the observed effects would not have been present without elevated emissions). Conversely, correlation with the $\text{GEOS}_{\text{CLIM}}$ anomalies would indicate that the observed anomaly cannot be partitioned from meteorological variability in the regional climate.

Anomalies for each degassing event were calculated as the seasonal mean during peak eruptive periods with long-term seasonal averages removed. We focused on the boreal summer (June–July–August, JJA) for the 2008 degassing event and the transition from the boreal spring to summer (May–June–July, MJJ) for 2018 when emissions from both events were highest and active eruptive events projecting ash and lithics into the volcanic plume occurred. To assess anomalies at the source relative to areas outside the plume, normalized zonal mean anomalies were calculated as

$$v_{\text{lat, norm}}(i) = \frac{v_{\text{lat}}(i)}{\sqrt{\sum_{i=1}^{N} v_{\text{lat}}(i)^2}}.$$  

(1)

where $N$ is the number of latitudes in the domain at $5^\circ$ intervals ($10-25^\circ$ N), $v_{\text{lat}}(i)$ is the seasonal latitudinal average (JJA 2008 or MJJ 2018) at latitude $i$, and $v_{\text{lat, norm}}(i)$ is the seasonal latitudinal anomaly normalized by the latitudinal mean (JJA or MJJ averaged over the climatology) at latitude $i$. To emphasize the location of the plume, we focused on a domain that included only areas west of the source ($25-155^\circ$ NE, $10-180^\circ$ SW).

4 Results and discussion

The goal of this work was to investigate the role of microphysical processes in ACIs during the Kilauea degassing events. To that end, we first show the reliability of our simulations by comparing satellite retrievals and GEOS control simulations (2008_1$\times$ and 2018_1$\times$) and sensitivity experiments (2008_5$\times$, 2008_PH2km, 2018_PH4km, and 2018_PH4km_ash). We then look for common features in both the 2008 and 2018 Kilauea degassing events and interpret their differences. Finally, Sect. 4.4 details the specific microphysical processes involved in cloud modification by the Kilauea emissions. Our results suggest that it is likely that the effects of volcanic emissions on cloud microphysics are specific to cloud phase. When possible, we divide the discussion between effects on ice and liquid clouds.

Anomalies for AOD from MODIS during peak degassing periods are shown in Fig. 2 for JJA 2008 (top left) and MJJ 2018 (bottom left). Figure 2 also shows modeled AOD anomalies for simulations using actual emissions (2008_1$\times$, 2018_1$\times$) calculated against the simulated climatology ($\text{GEOS}_{\text{CLIM}}$) and with zero-emissions scenarios (2008_0$\times$, 2018_0$\times$). Elevated aerosol loadings were apparent beginning at Kilauea (19.4°N, 155.2°W) and extended in a near-Gaussian plume westward across the domain shown in Fig. 2. AOD anomalies outside the plume domain were negligible, supporting the assumption that Kilauea is located in an otherwise “clean” environment relatively untouche by anthropogenic aerosol emissions from North America or East Asia. During both events, the elevated AOD anomalies observed by MODIS extended westward from Kilauea in the path of the Pacific easterly trade winds (Yuan et al., 2011). This is well-reproduced in the GEOS results. AOD anomalies appear weaker compared to $\text{GEOS}_{\text{CLIM}}$ than against $0\times$, particularly near the Kilauea crater. This is due to the fact that there is always some passive degassing near the source, which would be represented in the climatology but not in the $0\times$ experiment. Passive degassing is shown in Fig. 1, where a thin plume coming from the summit crater was evident during an eruptive pause in 2011, a year without a major volcanic event at the site.

Figure 3 shows MODIS retrievals and GEOS simulations of CF anomalies during the two events. Peak anomalies during the 2018 event (Fig. 3; bottom panels) were approximately 2–3$\times$ greater than during the 2008 event (Fig. 3; top panels). In both cases, the $1\times-\text{GEOS}_{\text{CLIM}}$ difference (middle panels) reproduced the spatial distribution of the CF anomaly from the MODIS retrieval (within the plume domain as well as zonal means), whereas the $1\times-0\times$ differences tended to overestimate the anomaly in 2008 and underestimate it in 2018. In 2008, the normalized $1\times-\text{GEOS}_{\text{CLIM}}$ anomaly correlated well with MODIS (Table 4), whereas there was essentially no correlation between the $1\times-0\times$ anomaly and the retrieval, indicating that the observed CF anomaly was mainly driven by meteorologic variability, likely differences in SST (Takahashi and Watanabe, 2016; Boo et al., 2015). Thus, ACIs likely had only a minor effect on CF for the two degassing events. It is possible that the aerosol layer may have locally modified SST, hence indirectly affecting CF. Elucidating this requires coupled ocean–atmosphere simulations and is suggested for future research. The discrepancies between the simulated and MODIS anomalies were relatively larger for CF than for AOD (Figs. 2 and 3). The AOD anomalies are primarily a function of the aerosol load and largely determined by the volcanic events. On the other hand, CF is influenced by many factors including convection, SST, El Niño–Southern Oscillation (ENSO) state, cloud microphysics, and winds, and it is much more sensitive to nat-

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ural variability. Satellite retrievals are also influenced by empirical definitions of cloudy and non-cloudy regions, adding uncertainty (Pincus et al., 2012). Given this, it is remarkable that the CF anomaly against the climatology is in relatively good agreement with the satellite retrieval, demonstrating the skill of GEOS in reproducing clouds during volcanic events.

To examine the cloud vertical structure during the two degassing events, GEOS results for the 2008_1× and 2018_1× simulations were compared against CALIPSO-GOCCP, as shown in Figs. 4 and 5, respectively. During both events, predominant cloud layers were found at 900 and 200 hPa, corresponding to low-level trade cumulus and in situ formation of thin cirrus clouds, respectively. The cloud vertical structure during both events was similar, indicating a strong meteorological control. This was well-captured by the GEOS simulations; the model, however, tended to overestimate high-level clouds and underestimate low-level clouds, particularly during the 2018 event (Fig. 5). This may result from replaying temperature to the reanalysis (Sect. 3.2) but letting water vapor evolve freely during the simulations. Such a configuration tends to overestimate relative humidity, particularly at cirrus levels, because the two-moment microphysics allows for supersaturation but MERRA-2 does not, and \( T \) is less constrained by observations (Gelaro et al., 2017). The discrepancy may also be exacerbated by artifacts in the retrieval. GOCCP tends to underestimate the presence of thin cirrus clouds and may overestimate low-level clouds in the presence of high aerosol loading (Chepfer et al., 2010). GEOS was, however, able to simulate an anomalous increase in CF above 800 hPa between 20 and 25° N in 2008 as seen in the retrieval. Similarly, there was a strong increase in CF in 2018 across the domain, although GEOS predicted a much larger anomaly effect on cirrus clouds. Again, the discrepancy may be the result of a lack of sensitivity to thin cirrus clouds in GOCCP.

Another interesting feature of Figs. 4 and 5 is the presence of mid-level altocumulus clouds, likely maintaining some supercooled water. In GOCCP, the SCF (i.e., the fraction of condensate remaining as liquid at \( T < 273 \) K) decreased sharply at around 550 hPa in both years but surprisingly remained significant (> 0.1) up to 200 hPa where homogeneous ice nucleation glaciated the remaining water. This was particularly true in 2018 when GOCCP showed enhanced SCF relative to the climatology. GEOS showed similar be-
behavior whereby the supercooled layer (0 < SCF < 1) was deeper in 2018 than in 2008, particularly near the summit crater (≈ 25° N). The supercooled layer was almost 100 hPa higher in GEOS than in GOCCP, indicating differences in ice and liquid partitioning between the model and the retrieval; this may be explained by the different definitions of SCF in each case. In GEOS, SCF is calculated on a mass basis, whereas in GOCCP it corresponds to the frequency of pixels that are classified as ice. The positive anomaly in SCF may suggest a deepening of clouds in the presence of aerosol and is analyzed in Sect. 4.4

4.1 Liquid clouds

Figures 6 and 7 show anomalies in $R_{\text{eff}}$, CDNC, COD, and LWP for MODIS and the GEOS control simulations (Table 2) during the 2008 and 2018 degassing events, respectively. Domain mean anomalies are summarized in Table 3.
Figure 6. Anomaly in liquid cloud properties during JJA 2008 for (from top) $R_{\text{eff}}$ (µm, shown as ER), COD, CDNC (m$^{-3}$), and LWP (g m$^{-2}$). Anomalies are shown (from left) for MODIS observations and for GEOS simulations calculated against GEOS$_{\text{CLIM}}$ and the zero-emissions scenario. The rightmost column shows normalized zonal mean anomalies for MODIS (blue) and GEOS against GEOS$_{\text{CLIM}}$ (green) and zero emissions (black).

Figure 7. Like Fig. 6 but for MJJ 2018.
Table 3. Anomalies for liquid cloud properties, AOD, and CF. Values in parentheses are the $p$ values associated with a one-sided Student’s $t$ test against the long-term mean.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Exp vs. climatology (excluding 2008)</th>
<th>Exp vs. 0×</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$R_{\text{eff}}$ (µm)</td>
<td>COD (–)</td>
</tr>
<tr>
<td>2008_1×</td>
<td>–0.58 (0.08)</td>
<td>4.63 (0.12)</td>
</tr>
<tr>
<td>2008_5×</td>
<td>–1.31 (0.01)</td>
<td>10.70 (0.02)</td>
</tr>
<tr>
<td>2008_PH2km</td>
<td>–0.58 (0.08)</td>
<td>4.63 (0.12)</td>
</tr>
<tr>
<td>MODIS (2008)</td>
<td>–0.75 (0.02)</td>
<td>0.46 (0.18)</td>
</tr>
<tr>
<td>2018_1×</td>
<td>–1.06 (0.15)</td>
<td>7.69 (0.13)</td>
</tr>
<tr>
<td>MODIS (2018)</td>
<td>–0.60 (0.14)</td>
<td>0.69 (0.24)</td>
</tr>
</tbody>
</table>

Table 4. $R^2$ values for the comparison of GEOS normalized zonal anomalies for liquid clouds against MODIS.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>MODIS anomaly vs. GEOS 1×-clim</th>
<th>MODIS anomaly vs. GEOS 1×-0×</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$R_{\text{eff}}$ (µm)</td>
<td>COD (–)</td>
</tr>
<tr>
<td>2008_1×</td>
<td>0.49 (0.70)</td>
<td>0.30 (0.57)</td>
</tr>
<tr>
<td>2008_PH2km</td>
<td>0.49 (0.70)</td>
<td>0.30 (0.57)</td>
</tr>
<tr>
<td>2018_1×</td>
<td>0.01 (0.31)</td>
<td>0.67 (0.58)</td>
</tr>
</tbody>
</table>

Table 4 shows the coefficient of determination ($R^2$) between the MODIS and the GEOS normalized zonal mean anomalies. During both the 2008 and the 2018 events, GEOS was able to capture the geographical distribution of the anomalies apparent in the MODIS retrievals for liquid clouds. Student’s $t$ tests indicated that the anomalies were statistically significant, typically to a 70%–80% level ($p < 0.3$, except for AOD and CF in 2008, which were significant only to 60%) for $R_{\text{eff}}$ and CDNC more so than for other variables (Table 3). For the 2008_1× experiment, the magnitude of the $R_{\text{eff}}$ and CDNC domain-averaged anomalies (–0.58 µm and 16.04 cm$^{-3}$, respectively) was in close agreement with MODIS (–0.75 µm and 21.53 cm$^{-3}$, respectively) and well-correlated with the satellite retrievals (Table 4). These values are also close to the anomalies calculated against the 2008_0× simulation, suggesting a microphysical control on $R_{\text{eff}}$ and CDNC. On the other hand, although the COD and LWP normalized anomalies are well-correlated between GEOS and MODIS, the model overestimated their absolute value. Similarly, in 2018 the GEOS and MODIS anomalies in $R_{\text{eff}}$ (–1.06 µm vs. –0.60 µm) and CDNC (26.65 cm$^{-3}$ vs. 19.43 cm$^{-3}$) were in better agreement than for COD (7.69 vs. 6.69) and LWP (15.30 g m$^{-2}$ vs. 5.09 g m$^{-2}$). Normalized zonal mean anomalies were, however, well-correlated ($R^2 > 0.5$) for CDNC and AOD (Table 4). The discrepancies in LWP and COD absolute anomalies were likely due to variation in SSTs inside the domain and the lack of a cloud albedo–SST feedback in our simulations.

SO$_2$ emissions at the Mauna Loa Observatory (19.5° N, 155.6° W) peaked in JJA 2008 relative to the 1995–2008 seasonal mean, with prevailing La Niña conditions (Potter et al., 2013). Thus, it is likely that the degassing event contributed to lower SSTs, hence lowering surface evaporation rates and LWP. The negative anomaly in LWP in the southernmost part of the domain evident in 2008 is also missing in the 2018 event due to neutral ENSO conditions in the latter (NOAA, 2020), indicating that ENSO exerted a strong meteorological control that could have drowned out ACI signatures in the former. Another reason behind the larger aerosol effects on COD and LWP simulated by GEOS than observed by MODIS may lie in differences in phase partitioning (i.e., liquid vs. ice) (Marchant et al., 2016). To help identify thin cirrus clouds, measured top-of-the-atmosphere (TOA) reflectance at 1.38 µm is used to partition high-altitude cirrus clouds from underlying liquid clouds. This method is strongly influenced by the relative humidity of the atmospheric column. Therefore, areas with low column water va-
por amount may have more clouds partitioned to the ice phase than is realistic. This is important because 2008 was a La Niña year, and the relative humidity in the atmospheric column was below climatological values, so LWP anomalies may appear low due to enhanced partitioning to the ice phase in the MODIS cloud-phase classification algorithm. Even in 2018, most of the total water path (TWP) anomaly resulted from an increase in the ice water path, indicating strong partitioning to the ice phase in the MODIS retrieval (Table 4).

Overall, more robust correlations ($R^2 > 0.5$) resulted when the anomalies were calculated against the GEOS$_{\text{CLIM}}$ than against the no-emissions scenarios (Table 4). This indicated that while both GEOS climatology and $0 \times$ cases included aerosol loading consistent with MODIS retrievals (Fig. 2, Table 4), correlations with observed data for liquid clouds were sensitive to meteorological effects captured by the MODIS and GEOS climatologies. MODIS TWP anomalies (for which TWP is the sum of LWP and IWP) in 2018 were about $20 \times$ higher than those reported for the 2008 event (Table 3), suggesting that ACIs for 2018 were not limited to liquid clouds.

We found that effects on liquid clouds were more statistically significant in 2018 than in 2008 for both observed (MODIS) and simulated (GEOS) cloud properties. In 2018, the magnitude and significance level of the simulated anomalies against GEOS$_{\text{CLIM}}$ were close to those calculated against $2018_0 \times$, indicating that increased aerosol loading, regardless of injection height, resulted in the heightened development of liquid cloud droplets and that these effects could be attributed to volcanic degassing as opposed to regional meteorology. The results of the $2008_5 \times$ run were in general highly statistically significant ($p < 0.05$) and agreed in magnitude with the $2018_1 \times$ experiment. The similarities in the magnitudes of simulated anomalies for $2008_5 \times$ and $2018_1 \times$ suggested that increased aerosol loadings would have been sufficient to overcome meteorological effects, which dampened the 2008 JJA anomalies with respect to long-term behavior. This and the similarities in spatial patterns (i.e., anomalies largely constrained by and maximized within the plume domain) for cloud anomalies in JJA 2008 (Fig. 6) and MJJ 2018 (Fig. 7) suggested a threshold response to overcome meteorological effects that was largely controlled by emissions.

### 4.2 Ice clouds

The volcanic plume height during peak degassing periods varied significantly between 2008 and 2018; therefore, we expected the strength of ACIs for ice clouds to be partially dependent on the availability of INPs. It is likely that the plume introduced SO$_2$ and ash particles to the upper troposphere – directly in 2018 and via convection in 2008. The effects of INPs on ice cloud development strongly depend on the microphysical processes dominating cloud evolution as well as the efficiency with which the new particles can promote ice crystal formation (Barahona et al., 2010). Additionally, the presence of aerosols in the upper troposphere was not apparent in a Gaussian-like plume emanating from the Kilauea volcano, as was the case for liquid clouds; therefore, the identification of ACIs for ice clouds was less straightforward.

Figure 8 shows that during the 2008 event, the aerosol plume did not significantly alter the properties of ice clouds. The $2008_1 \times$–GEOS$_{\text{CLIM}}$ anomalies in ice clouds appeared spatially consistent with MODIS anomalies for COD and IWP, whereas $2008_1 \times$–$0 \times$ showed little to no variability in the domain. As shown in Table 6, there was high correlation between MODIS and the $2008_1 \times$–GEOS$_{\text{CLIM}}$ normalized zonal mean anomalies for COD ($R^2 = 0.84$) and IWP ($R^2 = 0.93$). These correlations substantially decreased when considering comparisons between MODIS and the $2008_1 \times$–$2008_0 \times$ difference. For COD in 2008, the latter seemed to retain some correlation ($R^2 = 0.29$) and may indicate some level of microphysical control on the observed anomalies. This suggested that most of the observed ice cloud anomaly in 2008 resulted from meteorological and SST variability. Prevailing La Niña conditions in 2008 likely reduced relative humidity within the atmospheric column, particularly in the upper troposphere, thereby limiting the supply of water vapor necessary for ice crystal nucleation and growth. Anomalous conditions in ice clouds with respect to climatological and background conditions, however, were not statistically significant (Table 5).

Similar to the 2008 event, there seemed to be a strong meteorological component during MJJ 2018 degassing because the spatial correlation of the $2008_1 \times$–GEOS$_{\text{CLIM}}$ results appeared consistent with MODIS anomalies (Fig. 9). Additionally, correlations between MODIS and GEOS$_{\text{CLIM}}$ zonal mean anomalies were higher ($R^2 \geq 0.2$) than for the MODIS vs. $2018_1 \times$–$2018_0 \times$ correlations, with the exception of IWP ($R^2 = 0.32$) (Table 6). Unlike the 2008 event, the $2018_1 \times$–$2018_0 \times$ anomalies showed variability within the domain that was of similar magnitude to and appeared spatially consistent with GEOS$_{\text{CLIM}}$ anomalies (Fig. 9), indicating that not all ACIs in ice clouds could be attributed to meteorology alone. Figure 9 shows similar spatial distributions for observed and simulated anomalies within the plume domain, in particular between 175 and 165° W. Whereas for $R_{\text{eff}}$ and COD anomalous cloud properties appeared to be related to meteorological variability, IWP anomalies appeared to have a strong microphysical component. The MODIS and GEOS$_{\text{CLIM}}$ anomalies for IWP in 2018 were of the same order of magnitude at 20.59 and 30.41 g m$^{-2}$, respectively (Table 5), whereas the GEOS IWP $2008_1 \times$–$2008_0 \times$ difference was $\approx 8.03$ g m$^{-2}$. Although the strength of the observed effect was better represented by the GEOS$_{\text{CLIM}}$ IWP anomaly, the IWP correlation for MODIS vs. $2008_1 \times$–$2008_0 \times$ was slightly higher than against the climatology ($R^2 = 0.32$ vs. $R^2 = 0.23$); we found this difference to
be significant at the 95% confidence level. This strongly suggested a significant microphysical control on the IWP anomaly during the 2018 event.

4.3 Role of injection height and ash content

Domain-averaged CDNC varied widely between the different experiments except for the cases with elevated plume heights (2008\_PH2km and 2018\_PH4km), which essentially overlapped with the 2008\_1\times and 2018\_1\times experiments, respectively (Figs. 10 and 11). Despite this, increasing the plume height yielded sizable differences in some of the anomalies, particularly for ice clouds (Table 5) and for the 2018 event, although the effect was small. Similarities between MODIS and GEOS results for the 2008\_5\times and 2018\_1\times experiments on liquid clouds (Table 3) indicated that increased aerosol loadings during the 2008 event mimicked effects on liquid cloud formation during the 2018 event, most notably for CDNC and AOD. This showed that liquid cloud sensitivity to the first AIE was dominated by aerosol loadings as opposed to plume morphology during the 2008 event. Increasing plume height in concert with aerosol loadings exceeding 20 kt SO\textsubscript{2} d\textsuperscript{-1} (2018\_PH4km) reduced the anomaly in IWP and COD.
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Figure 10. Zonal mean vertical profiles for cloud fraction (CF), liquid ($Q_{\text{liq}}$) and ice ($Q_{\text{ice}}$) mixing ratios, total detrained condensate tendency (CNV_DQLD), cloud droplet (CDNC), and ice crystal (ICNC) number concentration for the 2008 JJA season. Notice that MERRA-2 and GOCCP data are included for CF only.

Figure 11. Equivalent to Fig. 10 but for the 2018 MJJ season.

by less than 5% relative to the 2018_1× experiment (Table 6). Similarly, accounting for ash INP (2018_PH4km_ash) slightly increased the COD and IWP anomalies (Table 5), although it had opposite effects on the ice crystal mixing ratio, $Q_{\text{ice}}$, and ICNC, with the former lower and the latter higher than in the 2018_PH4km experiment (Fig. 5). This is further analyzed in Sect. 4.4. The correlations between GEOS anomalies and MODIS for $R_{\text{eff}}$ and COD in ice clouds improved slightly with the inclusion of ash INP (Table 6).

4.4 Microphysical controls on ACIs

Figures 10 and 11 summarize the experiments performed in this work. Also shown are the CF values from MERRA-2 and GOCCP. In all runs and for both events, the vertical profile of CF was remarkably similar, with two predominant cloud layers associated with shallow cumulus and warm cirrus–altocumulus. This vertical structure was in agreement with GOCCP data (also shown in Figs. 4 and 5). Compared to GOCCP, GEOS tended to overestimate high-level CF and un-
Table 5. Anomalies for ice cloud properties. Values in parentheses are the \( p \) values associated with a one-sided Student’s \( t \) test against the long-term mean.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Exp vs. climatology</th>
<th>Exp vs. 0×</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( R_{\text{eff}} ) (µm)</td>
<td>COD (–)</td>
</tr>
<tr>
<td>2008_1×</td>
<td>0.25 (0.73)</td>
<td>−0.13 (0.19)</td>
</tr>
<tr>
<td>MODIS (2008)</td>
<td>0.53 (0.60)</td>
<td>0.79 (0.40)</td>
</tr>
<tr>
<td>2018_1×</td>
<td>0.95 (0.39)</td>
<td>0.58 (0.21)</td>
</tr>
<tr>
<td>2018_PH4km</td>
<td>1.12 (0.32)</td>
<td>0.58 (0.20)</td>
</tr>
<tr>
<td>2018_PH4km_ash</td>
<td>−4.33 (0.01)</td>
<td>0.63 (0.18)</td>
</tr>
<tr>
<td>MODIS (2018)</td>
<td>0.00 (0.27)</td>
<td>3.03 (0.18)</td>
</tr>
</tbody>
</table>

Table 6. \( R^2 \) values comparing GEOS normalized zonal anomalies for ice clouds against MODIS.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>MODIS anomaly vs. GEOS 1× vs. clim</th>
<th>MODIS anomaly vs. GEOS 1× vs. 0×</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( R_{\text{eff}} ) (µm)</td>
<td>COD (–)</td>
</tr>
<tr>
<td>2008_1×</td>
<td>0.25 (0.84)</td>
<td>0.84 (0.93)</td>
</tr>
<tr>
<td>2018_1×</td>
<td>0.23 (0.15)</td>
<td>0.15 (0.23)</td>
</tr>
<tr>
<td>2018_PH4km</td>
<td>0.25 (0.16)</td>
<td>0.16 (0.20)</td>
</tr>
<tr>
<td>2018_PH4km_ash</td>
<td>0.27 (0.39)</td>
<td>0.39 (0.15)</td>
</tr>
</tbody>
</table>

derestimate it near the surface. This is discussed in Sect. 4; however, here we also note that there is a high uncertainty in the retrieval during these periods. For example, GEOS-simulated high-level clouds were in better agreement with MERRA-2, whereas low-level CF in the latter was much lower than indicated by GOCCP. Examination of the GEOS simulations suggested that CF around 800 hPa was slightly higher for high aerosol loading than for the no-emissions experiments, indicating deepening of clouds from the injection of aerosol. This was supported by an increase in \( Q_{\text{liq}} \) between the 1× and the 0× experiments and runs along an enhanced detrained mass tendency, CNV\_DQLDT, indicating predominant convective effects. In general, the GEOS climatology (CLIM) tended to show lower values of liquid and ice properties than the 0× experiments, with the notable exception of the ice mixing ratio and number concentration in 2008.

These effects can be understood in light of the modification to dominant microphysical tendencies by elevated aerosol emissions. Figures 12 and 13 show the seasonally averaged rates of different cloud microphysical processes that took place during each event. They are distinguished by processes that affect the mass and number concentration of liquid droplets and ice crystals. Both events showed similar features regarding processes affecting liquid condensate and CDNC (Figs. 12a, c and 13a, c). As expected, liquid cloud microphysics were dominated by CCN activation and condensation (i.e., by droplet formation on sulfate particles). Cumulus detrainment (DCNV\_L) is a significant source of condensate but only plays a minor role in determining CDNC. The main sinks of liquid mass and number concentration are droplet autoconversion (AUT) and accretion of cloud droplets by rain (ACRL\_RAIN). The CCN source rate in 2018 (Fig. 13c) was about 3 times that of 2008 (Fig. 12); however, the liquid condensate tendencies shown in panel (a) in both figures were of the same magnitude. Thus, the mechanism for the decrease in \( R_{\text{eff}} \) was an increase in CCN activation and hence CDNC, which did not translate into a proportional increase in liquid mass (i.e., the first AIE). This is revealed in Figs. 10 and 11 as large increases in CDNC, but only slight increases in \( Q_{\text{liq}} \), between the 1× and 0× experiments.

It is not clear why the liquid autoconversion tendencies (AUTs) were almost insensitive to the aerosol load. Most likely, the liquid mass and number concentration sinks were controlled by accretion rather than autoconversion. This is depicted in Fig. 14. Figure 14a and c show that the autoconversion sinks for mass and number were similar across all simulation experiments, even those without volcanic emissions (i.e., 2008_0× and 2018_0×), despite CCN activation tendencies varying by a factor of 4. Our results indicated that ACRL\_RAIN was, however, enhanced for GEOS ex-
Figure 12. Domain-averaged cloud microphysical tendencies for the 2008 JJA season for (a) liquid mass, (b) ice mass, (c) liquid number, and (d) ice number concentration. Shown are Bergeron–Findeisen processes on ice (WBF) and snow (WBFSNOW), melting (MELT), total ice nucleation and immersion freezing only (ICENUC and NHET_IMM, respectively), droplet evaporation (EVAP), convective detrainment of liquid and ice (DCNVL and DCNVI), condensation (COND), liquid and ice autoconversion (AUT and AUTICE, respectively), accretion of liquid by rain and snow (ACRS and ACRL, respectively), CCN activation (CCN), ice sedimentation (SDM), ice crystal growth and accretion by snow (DEP and ACRI_SNOW, respectively), and ice splintering (HM).

Experiments with high aerosol concentrations (Fig. 14c). This suggested an unexpected ACI mechanism. Ingestion of CCN within convective parcels inhibits the formation of precipitation in cumulus clouds; hence, more condensate was detrained to the top. DCNVL was thus enhanced in simulations with high aerosol loading (2018_1 ×, 2008_1 ×, and 2008_5 × in Fig. 14a). This was reflected in the vertical profiles of CNV_DQLDT, Q_liq, and to a lesser extent CF (Figs. 10 and 11). This mechanism created deeper clouds that precipitated from above scavenging the liquid below, which explained the increase in ACRL_RAIN as the CCN activation tendency increased. Because droplets also freeze higher in the convective parcel, it may also lead to convective invigoration, although our setup (i.e., replaying T to the reanalysis) prevented GEOS from explicitly simulating this. These findings are consistent with the deepening of the cloud layers described by Yuan et al. (2011) and the evidence for convective invigoration reported by Mace and Abernathy (2016). They also explain the anomalous enhancement of the mid-level CF and the increased SCF during the two events found in the GOCCP retrievals (Figs. 4 and 5).

Whereas a few processes dominate the microphysics of liquid water, ice microphysics are much more complex. It is clear from Figs. 12 and 13 that ice nucleation (ICENUC), sedimentation (SDM), and growth and sublimation (DEP) were dominant. However, other processes, notably convective detrainment (DCNVI) and the Wegener–Bergeron–Findeisen (WBF) process, still played significant roles in cloud development. Ice autoconversion to snow (AUTICE) was a significant sink for ice mass but not for number concentration. It is likely that the latter was controlled by SDM. This complex microphysical makeup may be one of the reasons why the anomalies for ice clouds were much less evident than for liquid clouds, buffering the former against aerosol perturbations.

It is also interesting that almost every ice tendency was 2–3 times larger in the 2018 (Fig. 13) than in the 2008 event (Fig. 12), while the liquid condensate rates were of the same magnitude across all experiments. The main reason for this was that Q_ice was larger in 2018 than in 2008 (Figs. 10 and 11), likely due to higher SSTs during neutral ENSO conditions relative to La Niña SST cooling in 2008. Warmer SSTs triggered more frequent convection events and enhanced the transport of condensate, water vapor, and frozen condensate to the upper troposphere. Crystals sedimenting from above find a favorable environment to grow, which also enhances splintering processes below, represented by ice splintering (HM) in Fig. 13. Although there was definitely
Figure 13. Like Fig. 12 but for the 2018 MJJ season.

Figure 14. Comparison of dominant microphysical process rates for (a) liquid mass, (b) ice mass, (c) liquid number, and (d) ice number concentration. Color corresponds to the different experiments listed in Table 2.
a higher rate for ICENUC in 2018 than in 2008 (Fig. 14), the fact that it changed little upon modification of the injection height or by considering ash as INP strongly suggested that changes in the vertical transport of liquid and water vapor, rather than ICENUC, led to the observed anomalies in ice clouds. In fact, accounting for ash INP had a negligible effect on immersion freezing rates (not shown). Including ash INP led to slightly higher ice nucleation rates between 200 and 300 hPa. Because these crystals grow and sediment quickly, they may explain the slightly lower $Q_{\text{ice}}$ values with respect to the 2018 $1 \times$ experiment despite the higher ICNC (Fig. 11).

5 Conclusions

We carried out a comprehensive analysis of the effects of volcanic emissions on cloud formation and microphysical properties during two major events from the Kilauea summit crater, Halemaumau, in the Hawaiian islands, occurring during the late spring and summer of 2008 and 2018. Previous analyses characterized the geologic and atmospheric effects of the 2008 event, but this is the first time that such an analysis has been performed for the 2018 event. We combined satellite data, reanalysis products, and atmospheric modeling simulations to analyze and understand the role of meteorology and aerosol–cloud interactions in cloud evolution.

Several factors contributed to observed and simulated similarities and differences for each event: (1) aerosol loading, (2) plume height and composition, and (3) El Niño–Southern Oscillation conditions. The Kilauea volcano emissions resulted in an aerosol plume that extended westward across the domain driven by easterly trade winds. Aerosol concentrations were anomalously high during both events relative to long-term means and zero-emissions simulations.

The simulated aerosol–cloud interaction signatures common to both events suggested that effects on cloud macrophysical and microphysical properties with respect to the first indirect effect, such as decreased cloud effective radius from increased CCN, were likely decoupled from local meteorological effects. In contrast, changes in liquid water path and cloud lifetime in the presence of enhanced aerosol concentrations were likely dampened by meteorological variability. Thus, even in well-constrained natural experiments such as the ones presented in this paper, caution must be taken when drawing conclusions on the second indirect effect without full consideration of meteorological effects.

The 2018 Kilauea degassing event was stronger and more regionally significant with respect to cloud formation processes. However, when the actual 2008 emissions were increased by 500% (i.e., experiment 2008, $0.5 \times$), computed anomalies in cloud properties were of the same order as for the 2018 event. This and the similarities in the spatial patterns of the anomalies in both events suggest a threshold response of ACIs to overcome meteorological effects, largely controlled by aerosol loading.

For ice clouds, changes in cloud microphysics were significant following the 2018 event, while few, if any, effects were apparent in the 2008 event. Our analysis suggested that effects on ice clouds were largely controlled by aerosol injection height and thus the availability of INPs for ice crystal growth and nucleation. Both satellite and model output showed large positive anomalies for ice water path and cloud optical depth during 2018 across the domain, which indicated that conditions favorable for ice cloud formation present in 2018 were absent in 2008. Ash was present in the volcanic plume for both degassing events, but only the plume in 2018 injected volcanic material to sufficiently high altitudes to potentially impact the formation of ice clouds. Sensitivity experiments using increased emissions, plume height, and introduced ash as an ice nucleation particle (INP) suggested that these changes only slightly amplified anomalies for both liquid and ice clouds.

We performed a detailed analysis of the rates of cloud microphysical processes during the two events. As expected, CCN activation played a major role in determining the cloud droplet number concentration for both cases; however, increased liquid droplet concentration did not lead to a substantial reduction in autoconversion rates. Instead, ingestion of CCN within convective parcels may have caused convective invigoration, enhancing the detrainment of condensate in the free troposphere. As a result, clouds perturbed by aerosols were optically deeper than in the pristine cases. This mechanism also led to enhanced cloud droplet scavenging by accretion.

Microphysical rates in ice clouds were found to be accelerated in 2018 with respect to 2008, likely resulting from higher sea surface temperatures during El Niño conditions in the former (hence more convective inflow to the upper troposphere) and by an increase in convective detrainment in the presence of aerosols. Although the ice nucleation rate in cirrus clouds was substantially higher in 2018 than in 2008, it is likely that it did not control the observed anomaly in ice cloud properties in the former. Accounting for ice formation on ash INP led to slightly enhanced ice nucleation in warm cirrus clouds, thereby facilitating ice sedimentation and decreasing ice water content.

Although the model configuration used in this work presented a clear view of the role of different microphysical processes in determining the evolution of clouds during the 2008 and 2018 events, it precluded a full exploration of the second indirect effect and the feedbacks between aerosol loading and SSTs. The investigation of these effects requires a coupled ocean–atmosphere model and will be the subject of future work. Chemical effects may also lead to complex interactions between the volcanic plume and the formation of clouds. For example, the ocean entry of 2018 ERZ eruptions caused large clouds of vaporized HCl and water vapor to ascend with the plume (Kern et al., 2020). This was a composi-
tional component absent in the 2008 plume that may lead to the injection of CCN to the upper troposphere as well as enhanced convection in the ERZ region. Accounting for these effects requires a detailed parameterization of the chemical evolution of the volcanic plume and is left for future studies.

This work showed that satellite observations provide strong evidence for the effects of aerosols modifying clouds during the Kilauea volcanic events. Model simulations elucidated the underlying microphysical processes involved and provided insight into the role of aerosol–cloud interactions in determining the evolution of clouds. We showed that there were many similarities in cloud anomalies during the 2008 and 2018 degassing events and that the discrepancies are largely attributable to differences in aerosol loading. Our work thus provided an unprecedented view of the mechanisms driving the aerosol indirect effect during volcanic events, helping to advance the understanding of the role of aerosol emissions in climate.
Appendix A: Ice nucleation on ash particles

Sulfates from SO$_2$ oxidation were the primary aerosol associated with volcanic degassing events. There is, however, evidence that ash was co-emitted with SO$_2$ at the Kilauea volcano, which may have an effect on the way the plume interacts with clouds by introducing additional surfaces for ice nucleation (ash is typically considered a poor CCN) (Durant et al., 2008). Ash concentration was likely important during the 2018 event when particulate emissions reached the upper troposphere (Neal et al., 2019; Kern et al., 2020). To assess the impact of ash on ice cloud formation, the ice nucleation parameterization in GEOS was amended as follows.

For mixed-phase clouds, ash was assumed to induce immersion freezing. Maters et al. (2019) reported the active site density, $n_s$, of tephra consisting of glassy lithic fragments and ash particles emitted from Kilauea in May 2018. Only ash particles were found to be active ice nucleation sites. For the ash samples, $n_s$ (cm$^{-2}$) was fit as

$$n_s = \exp(-0.6907T + 176.5992),$$  \hspace{1cm} (A1)

with $240 \leq T \leq 260$ K, where $T$ is temperature. Specific surface area was prescribed as 2.1 m$^2$ g$^{-1}$ (Maters et al., 2019). Ash content in the volcanic plume was assumed to be 0.1%, consistent with literature values (Mastin et al., 2009).

Ash was also assumed to heterogeneously nucleate ice in the deposition mode at low temperature ($T < 236$ K), hence affecting the formation of cirrus clouds. Although the ice nucleation efficiency of ash from the Kilauea summit crater has not been reported, Hoyle et al. (2011) measured a saturation freezing threshold of 110% for ash emissions from the Eyjafjallajökull volcano. This value is used as a first approximation for the Kilauea ash emissions. Ash was then treated as a monodisperse INP (Barahona and Nenes, 2009) and added to the Ullrich et al. (2017) ice nucleation spectrum. The nucleated ice crystal concentration was then calculated using the Barahona and Nenes (2009) parameterization.

Data availability. All data generated in this work are publicly available through the NASA technical reports server (https://trs.nasa.gov, last access: 12 May 2021) (NASA, 2021b) and PubSpace (https://www.nasa.gov/open/researchaccess/pubspace, last access: 12 May 2021) (NASA, 2021c).

Author contributions. DB conceived and supervised this work. KHB prepared the GEOS simulations and the analysis of the satellite retrievals. HB and TY contributed daily emissions files for the volcanic events. SCJ contributed as a scientific consultant and editor of the paper.

Competing interests. The authors declare that they have no conflict of interest.

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