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Cloud impacts on photochemistry: building a climatology of photolysis rates from the Atmospheric Tomography mission

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Abstract. Measurements from actinic flux spectroradiometers on board the NASA DC-8 during the Atmospheric Tomography (ATom) mission provide an extensive set of statistics on how clouds alter photolysis rates ($J$ values) throughout the remote Pacific and Atlantic Ocean basins. $J$ values control tropospheric ozone and methane abundances, and thus clouds have been included for more than three decades in tropospheric chemistry modeling. ATom made four profiling circumnavigations of the troposphere capturing each of the seasons during 2016–2018. This work examines $J$ values from the Pacific Ocean flights of the first deployment, but publishes the complete Atom-1 data set (29 July to 23 August 2016). We compare the observed $J$ values (every 3 s along flight track) with those calculated by nine global chemistry–climate/transport models (globally gridded, hourly, for a mid-August day). To compare these disparate data sets, we build a commensurate statistical picture of the impact of clouds on $J$ values using the ratio of $J$-cloudy (standard, sometimes cloudy conditions) to $J$-clear (artificially cleared of clouds). The range of modeled cloud effects is inconsistently large but they fall into two distinct classes: (1) models with large cloud effects showing mostly enhanced $J$ values aloft and or diminished at the surface and (2) models with small effects having nearly clear-sky $J$ values much of the time. The ATom-1 measurements generally favor large cloud effects but are not precise or robust enough to point out the best cloud-modeling approach. The models here have resolutions of 50–200 km and thus reduce the occurrence of clear sky when averaging over grid cells. In situ measurements also average scattered sunlight over a mixed cloud field, but only out to scales of tens of kilometers. A primary uncertainty remains in the role of clouds in chemistry, in particular, how models average over cloud fields, and how such averages can simulate measurements.
1 Introduction

Clouds visibly redistribute sunlight within the atmosphere, thus altering the photolytic rates that drive atmospheric chemistry \( (J_\text{values}) \), as well as the photosynthesis rates on the land and in the ocean. These \( J_\text{values} \) drive the destruction of air pollutants and short-lived greenhouse gases. The NASA Atmospheric Tomography Mission (ATom, 2017; Wofsy et al., 2018), in its charge to measure the chemical radiative transfer over the remote ocean basins, has measured \( J_\text{values} \) while profiling the troposphere \((0–12 \text{ km})\). These measurements reveal a statistical pattern of \( J_\text{values} \) over different geographic, altitude and cloud regimes, which directly challenges current atmospheric chemistry models and provides a new standard test of cloud effects. The observations quantify how clouds alter photochemistry and are compared here with parallel analyses from nine global atmospheric chemistry models.

Since the early models of atmospheric chemistry, the scientific community has tried various approximations and fixes for “those pesky clouds”. Overhead clouds can shadow the sun, resulting in diminished \( J_\text{values} \) beneath and within the lower parts of thick clouds. Cloud scattering results in enhanced \( J_\text{values} \) above and within the tops of clouds. For ideal clouds – uniform layers from horizon to horizon – the models have developed a variety of methods to approximate the 1-D radiative transfer and calculate the \( J_\text{values} \) relative to a “clear sky” (Logan et al., 1981; Chang et al., 1987; Madronich, 1987; Wild et al., 2000; Lefer et al., 2003; Williams et al., 2006; Palancar et al., 2011; Ryu et al., 2017). More realistic treatment of clouds is important in global chemistry as well as air pollution (Kim et al., 2015). For the most part, these chemistry models are provided with the cloud properties and fractional coverage for each grid cell in a column, make some assumptions about the overlap of cloud layers, and then solve the 1-D plane-parallel radiative transfer equation at varying levels of accuracy. In a 3-D world, however, adjacent clouds can block the sun or scatter light even when there are clear skies overhead. Also in a 3-D world, a sunlit adjacent cloud that is not overhead can increase \( J_\text{values} \). It is nigh impossible to specify the 3-D cloud fields at \( \sim 1 \text{ km} \) scale along the ATom flight paths or any similar mission; 3-D high-resolution cloud fields on this are available, but these are limited to CloudSat–CALIPSO afternoon overpasses extended to 200 km wide swaths (see Barker et al., 2011; Miller et al., 2014). Anyway, none of the standard global models can deal with such a 3-D radiative transfer problem. So accepting the model limitations and the inability to match individual measurements, we use the statistics of observed \( J_\text{values} \), enhanced or diminished relative to a clear sky, and ask if the models’ many approximations for the radiative transfer in cloud fields can yield those same net results.

This paper presents the statistical distribution of the measured \( J_\text{values} \) using the CAFS instrument (Charged-coupled device Actinic Flux Spectroradiometer; Shetter and Müller, 1999; Petropavlovskikh et al., 2007) from the first ATom deployment (29 July through 23 August 2016). We build up statistics of the observed \( J_\text{values} \) relative to that calculated for a clear sky under similar conditions. The chemical reactivity of the troposphere (see Prather et al., 2017) is generally proportional to these changes in \( J_\text{values} \), and thus modeling of the variability of clouds is critical for modeling the lifetime of \( \text{CH}_4 \) and the cycling of tropospheric \( \text{O}_3 \).

So what makes this model vs. measurement comparison different? The atmospheric chemistry community has a history of such comparisons, including photolysis rates, dating to the early ozone depletion assessments (Nack and Green, 1974; M&M, 1993) and continuing to recent multi-model projects (Olson et al., 1997; Crawford et al., 2003; PhotoComp, 2010). These comparisons have been limited mostly to simplistic atmospheric conditions and measurements made under clear skies or with uniform 1-D cloud layers for which an accurate solution can usually be calculated. We introduce here the ability to use CAFS “all-sky” measurements made under a semi-objective sampling strategy (i.e., ATom’s tomographic profiling makes pre-planned slices through the troposphere, limited by available airports, diverting only for dangerous weather). We test the collective treatment of clouds and radiation in models insofar as they can match the observed statistics of \( J_\text{values} \). This approach gets to the core of atmospheric photochemistry by combining the range of assumptions and parameterizations for clouds in the models including, among others, cloud optical depths (CODs) and scattering phase functions, two-stream or multi-stream radiative transfer, cloud overlap, or even just parametric correction factors.

Section 2 describes the measurements and models, including how the observational statistics are compiled, what protocol the models used, and how they included cloud fields. This section also documents the differences in mean \( J_\text{values} \), which can be as large as 30% in either “cloudy” or clear conditions. Section 3 introduces the statistical distributions based on the ratio of full sky including clouds to clear sky. Section 4 examines modeling errors and improvements related to this comparison. In the concluding section, Sect. 5, we discuss the current range in modeling cloud effects and how new observational constraints can be developed and used to build better models.

2 Measuring and modeling \( J_\text{values} \) under realistic cloudy skies

Here, we focus on two \( J_\text{values} \): \( \text{O}_3 + \text{hv} \rightarrow \text{O}_2 + \text{O}(^1\text{D}) \) designated \( J-\text{O1D} \) and \( \text{NO}_2 + \text{hv} \rightarrow \text{NO} + \text{O}(^3\text{P}) \) designated \( J-\text{NO2} \). These \( J_\text{values} \) are the most important in driving the reactive chemistry of the lower atmosphere, and each emphasizes a different wavelength region with response to atmospheric conditions. \( J-\text{O1D} \) is driven by short wave-
lengths (<320 nm), where O₃ absorption and Rayleigh scattering control the radiation, whereas J-NO2 is generated by longer wavelengths, where O₃ absorption is not important and Rayleigh scattering is 2 times smaller. Thus, J-NO2 is the more sensitive J to clouds. Both CAFS and the models accept the same spectral data – cross sections and quantum yields from recent assessments (Atkinson et al., 2004; Burkholder et al., 2015) – but the implementation (solar spectrum, Rayleigh scattering, wavelength integration, temperature interpolation) may be different.

Spectrally resolved CAFS measurements of actinic flux (280–650 nm) are used to calculate in situ clear-sky J values denoted J-clear. For J-clear, CAFS uses the Tropospheric Ultraviolet and Visible (TUV) radiative transfer model (Madronich and Flocke, 1999). The model is run with an eight-stream discrete ordinate radiative transfer method with a pseudo-spherical modification to generate actinic fluxes with a 1 nm wavelength grid from 292–700 nm. The calculation is run with no clouds and no aerosols and a fixed surface albedo of 0.06, and applies ozone columns from the satellite Ozone Monitoring Instrument (Levett et al., 2006; Veefkind et al., 2006). CAFS and TUV spectra are processed using the same photolysis frequency code to ensure that the same quantum yield, absorption cross section, and temperature and pressure dependence relationships are applied to the measured and modeled spectra. The strong connection between measurement and model has been established in past campaigns (Shetter et al., 2002, 2003; Hofzumahaus et al., 2004). The ATom CAFS data set used here is a value-added product beyond the standard ATom output (Wofsy et al., 2018), and it is archived with this paper. Only data from the first deployment, ATom-1, were available when this paper was being prepared.

Global chemistry models cannot be used productively in comparisons with individual CAFS observations as noted above, but a statistical comparison of the ratio J-cloudy to J-clear is a useful climatological test. It is difficult for the models to simulate CAFS global data unless there is a very careful sampling strategy to match albedos from the ATom flights over land and cryosphere. Thus, we focus on the two oceanic blocks in the Pacific for which we have a large number of measurements with high sun in ATom-1. See also discussion of ocean surface albedo variations in Sect. 4. The CAFS statistics are derived from the ATom-1 deployment and selected for two remote geographic blocks in order to compare with the models: (block 1) tropical Pacific, 20° S–20° N × 160–240° E, and (block 2) North Pacific, 20–50° N × 170–225° E.

The models here include the six original ones used in the ATom reactivity studies (Prather et al., 2017) plus three additional European global chemistry models. They are described more fully in Table 1, and are briefly designated as GEOS-Chem (GC), GFDL AM3 (GFDL), GISS Model 2E1 (GISS), GSF6 GM1 (GMI), ECMWF IFS (IFS), MOCAGE (MOCA), CESM (NCAR), UCI CTM (UCI), and UM-UKA (UKCA). Additional model information and contacts are given in the Supplement Table S1. All models submitted global 4-D fields (latitude by longitude by pressure for 24 h) for a day in mid-August using their standard treatment of clouds (designated all sky or cloudy here) and then a parallel simulation with clouds and aerosols removed (designated clear sky or clear). In addition to its correlated cloud-overlap model with multiple quadrature column atmospheres to calculate an average J value, UCI also contributed a model version using the B-averaging of cloud fractions (CFs) (Briegleb, 1992) used by most models, designated UCIb (see Prather, 2015). Several models ran the clear-sky case without clouds but with their background aerosols. Globally, aerosols have a notable impact on photolysis and chemistry (Bian et al., 2003; Martin et al., 2003), but over the middle of the Pacific Ocean (this analysis), the UCI model with and without aerosols shows mean differences of order ±1/2 %.

Modeling the effect of clouds on J values began with early tropospheric chemistry modeling. One approach was to perform a more accurate calculation of generic cloud layers offline and then apply correction factors to the clear-sky Js computed in-line: e.g., an increase above the cloud deck and a decrease below (Chang et al., 1987). Another approach used a climatology of overlapping cloud decks to define a set of opaque, fully reflecting surfaces at different levels; e.g., the Js would be averaged over these sub-grid column atmospheres (Logan et al., 1981; Spivakovsky et al., 2000). As 3-D tropospheric chemistry models appeared, the need for computationally efficient J-value codes led to some models ignoring clouds and others estimating cloud layers and applying correction factors to clear-sky Js. With the release of Fast-J (Wild et al., 2000), some 3-D models started using a J-value code that directly simulated cloud and aerosol scattering properties with few approximations. The next complexity, based on general circulation modeling, included fractional cloud cover within a grid cell and thus partial overlap of clouds in each column (Morcrette and Fouquart, 1986; Briegleb, 1992; Hogan and Ilingworth, 2000). This approach later moved on to chemistry models (Feng et al., 2004; Liu et al., 2006; Neu et al., 2007). Monte Carlo solutions for the numerous independent column atmospheres generated by cloud overlap were developed for solar heating (MCICA; Pincus et al., 2003), but random, irreproducible noise was not acceptable in deterministic chemistry transport models. The Cloud-J approach (Prather, 2015) developed a scale-independent 1-D method for cloud overlap based on vertical decorrelation lengths (Barker, 2008a, b). The chemistry models here use a variety of these methods, which range from lookup tables with correction factors, to Fast-J single column, to cloud overlap treatments with Cloud-J: see Table 1. Currently these models do not attempt to define 3-D cloud struc-
treats within a grid square, the approach needed to match individual CAFS \( J \)s.

The CAFS data were collected from ATom-1 during its 10 research flights from 29 July to 23 August 2016. It was not possible for the models to simulate each flight path, including clouds and local solar zenith angles (SZAs) for each measurement. Not all of the models could run with 2016 meteorology, and thus we asked for a day in mid-August and treat that (rightly or wrongly) as typical of the cloud statistics during ATom-1. The meteorological dates are listed in Table 1. This simplification made it possible to attract large participation, but of course it limits the ability to claim that the model statistics are a robust climatology. ATom flights are mostly in daylight and hence a large proportion of CAFS measurements occur at high sun, \( \cos(SZA) > 0.6 \), with more than half at \( \cos(SZA) > 0.8 \) (Supplement Fig. S1). The models report hourly \( J\text{-O1D} \) and \( J\text{-NO2} \) globally over 24 h, and thus all have a similar distribution of \( \cos(SZA) \) but with a
greater proportion at cos(SZA) < 0.4 than the CAFS data. We restrict these comparisons to high sun, cos(SZA) > 0.8, to reduce 3-D effects that are not modeled here, which leaves 11,504 (block 1) and 4,867 (block 2) measurements for the CAFS/TUV 3 s averages. For the models, the number of hourly samples with cos(SZA) > 0.8 is about 18% for both blocks and thus the number of samples depends on model resolution (e.g., 240,000 for UCI and 1,400,000 for NCAR in block 1). Although the analysis here is limited to the Pacific blocks, the global model data are archived with this paper.

A quick look at J-cloudy (all sky) profiles of J-O1D and J-NO2 for CAFS (Fig. S2) shows a basic pattern also seen in models. Both Js are larger in the upper troposphere where the direct sunlight is more intense, but in the North Pacific, larger cloud cover and more scattered light almost reverses this pattern with enhanced Js at lower altitudes (>600 hPa). Comparing the variances of J-cloudy (CAFS) and J-clear (TUV) for the tropical Pacific, the J-NO2 variability is driven almost entirely by clouds as expected, while the J-O1D variability is driven firstly by O3 column and sun angle (both CAFS and TUV), while there are clearly cloud contributions (CAFS only) at lower altitudes (>700 hPa).

Figure 1 shows a full comparison of the CAFS J profiles with the 10 model results in four panels (2 Js x 2 geographic blocks). The CAFS Js fit within the range of models; their shape is matched by most models, but the model spread of order 20–30% is hardly encouraging. Differences in these average profiles can have many causes: temperature and O3 profiles, spectral data for both J-O1D and J-NO2, ways of integrating over wavelength, surface albedo conditions, treatment of Rayleigh scattering, basic radiative transfer methods, SZA, and, of course, clouds. In typical comparisons we try to control these differences by specifying as many conditions as possible, but here we want to compare the “natural” Js used in their full-scale simulations (e.g., Lamarque et al., 2013) and thus leave each model to its native atmospheres, spectral data, algorithms, and approximations.

The models show a much tighter match in J profiles under clear-sky conditions (Fig. 2). Typically, eight of the models fall within 10% of their collective mean profile. Some models are obviously different in J-clear (GISS and MOCA for J-O1D, MOCA for J-NO2), and these differences carry through to J-cloudy (Fig. 1). A most important factor in J-O1D is the O3 column, and Fig. S3 shows the modeled O3 columns for August compared with 8 years of OMI observations. MOCA, NCAR, and IFS have low tropical O3 columns, <250 DU vs. observed ∼265 DU, which could lead to higher J-O1D, but this effect is seen only in MOCA.
UKCA has higher tropical columns, > 300 DU, which might explain why their \( J \)-O1D lies in the lower range of the models. Some results, like MOCA’s \( J \)-NO2 and GISS’s \( J \)-O1D point to differences in the implementation of spectral data (e.g., wavelength integration, solar spectrum, temperature interpolation). We expect GC and GMI to be alike since they both use MERRA-2 cloud fields and Fast-J with Briegleb averaging: indeed, they match well except for \( J \)-O1D in the tropical Pacific, yet, they report similar tropical ozone columns.

The ratio of \( J \)-cloudy to \( J \)-clear, shown in Fig. 3, cancels out many of the model differences in Figs. 1 and 2, and as expected GC and GMI are nearly identical. The cloudy : clear ratio identifies new patterns in model differences, whereby some models have ratios close to 1 throughout the troposphere (especially in the Tropics) while others have ratios > 1.1 at altitudes above 800 hPa and < 0.9 below 900 hPa. In a recent model intercomparison with specified chemical abundances (Prather et al., 2018), we found that the tropospheric photochemistry of \( \text{O}_3 \) and \( \text{CH}_4 \) responded almost linearly to cloudy–clear changes in \( J \) values (see Fig. S4; data not shown in Prather et al., 2018). Thus the differing impact of clouds on \( J \) values seen in Fig. 3 will have a correspondingly large impact on global tropospheric chemistry (see Liu et al., 2009).

## 3 The statistical distribution of \( J \)-cloudy to \( J \)-clear

The average ratio of \( J \)-cloudy to \( J \)-clear (Fig. 3) provides only a single measure of the impact of clouds. The CAFS data provide a more acute measure by sampling the range of cloud effects (enhance or diminish \( J \) values) and their frequency of occurrence. A quick look at this range in CAFS data is shown in Fig. S5 with the probability of occurrence of the cloudy : clear ratio defined as \( \ln(\text{J-cloudy}/\text{J-clear}) \) and designated \( r\ln J \). Each curve is normalized to unit area, with the y axis being probability per 0.01 bin (~ 1%) in \( r\ln J \).

We expect that enhancements in \( J \)s occur above clouds and diminishments below, and this is shown in Fig. S5. In the marine boundary layer (900 hPa–surface), there are a greater number of \( r\ln J < -0.10 \), with fewer \( r\ln J > 0.00 \). In the mid-troposphere layer (300–900 hPa), there are frequent occurrences of \( r\ln J > +0.10 \), particularly in the North Pacific where lower level clouds are more extensive than in the Tropics. Likewise, there are times when \( r\ln J < 0.00 \) in the mid-
troposphere when clouds lie overhead. In upper tropospheric layers (100–300 hPa), most of the optically thick clouds are below and $\ln J > 0.00$ is dominant. Thus, our analysis here breaks the atmosphere into these three layers. All figures in this section will be displayed as two-block by three-layer panels with part a ($J$-O1D) and part b ($J$-NO2).

### 3.1 Modeling the distribution of $J$ values

The probability distribution of $\ln J$ for $J$-O1D (Fig. 4a) and $J$-NO2 (Fig. 4b) shows highly varied patterns across the models, but with some consistency. The models and CAFS are not exactly a match, but again, there are some encouraging patterns. The peak $\ln J$ distributions for CAFS will be broadened because the real variation in ocean surface albedo is not simulated in TUV (see discussion in Sect. 4), but this is expected to be of order $\pm 2\%$, and so the overall width (10 to 20%) reflects cloud variability. The modeled and measured distributions are asymmetric and skewed toward $\ln J > 0$ in the free troposphere (100–900 hPa), and toward $\ln J < 0$ in the boundary layer. This pattern is expected since $J_s$ are enhanced above clouds (100–900 hPa) and diminished below. There will also be some occurrences of $\ln J < 0$ in the free troposphere when thick clouds are overhead, but none of the models come close to the CAFS frequency of these occurrences. In general and as expected, this brightening above and dimming below is more evident for $J$-NO2 than for $J$-O1D. Another feature that is somewhat consistent across models and observations is that the wings of the $\ln J$ distribution are wider in the North Pacific than the Tropics. In the boundary layer, observations and models show the reverse with greater cloud effects (dimming, $\ln J < 0$) in the Tropics. Although all the models show this shift from $\ln J > 0$ to $\ln J < 0$ in the peak of their distributions, only a few (GISS, MOCA, NCAR, UCI) have broad wings of large cloud shielding ($\ln J < -0.1$). These models calculate such broad wings for both the Tropics and North Pacific, whereas CAFS only shows this in the Tropics.

These diagnostics also identify some CAFS anomalies that have no physical basis in the current models. For example, low-level (surface to 900 hPa) observed cloud enhancements ($\ln J > 0.025$) observed in the tropical Pacific (particularly $J$-O1D) do not appear in the models. Similarly cloud diminishments in the upper troposphere do not appear with the modeled cirrus. A more thorough analysis of the CAFS $\ln J$ with the added deployments should examine if these differences are due to 3-D radiative transfer effects, ocean albedo variations, or missing cloud types in the models. Perhaps, we will identify sun–cloud geometry conditions simply not possible in the 1-D cloud models.
Figure 5a (J-O1D) and b (J-NO2) provide another view of the rlnJ statistics. Figure 5 has the same six blocks as Fig. 4, but plots five quantities schematically in a single line describing each model’s statistics: percent occurrence of diminishment (rlnJ < −0.025), average rlnJ diminishment, percent occurrence of nearly clear sky, percent occurrence of enhancement (rlnJ > +0.025), and average rlnJ enhancement. The horizontal line has a total length of 100% with the thin line (clear sky) centered on 0; see the extended legend in Fig. 5b.

Overall, four models (GC, GFDL, GMI, UKCA) have unusually narrow peak distributions of rlnJ ∼ 0, indicating lesser cloud effects on the Js. The other six models (GISS, IFS, MOCA, NCAR, UCI, UCIb) show a much greater range in Js, with a larger fraction perturbed by clouds (enhanced or diminished by more than 10%). The CAFS observations generally support this latter group. There are individual model anomalies that may point to unusual features: MOCA alone has a peak frequency of enhanced Js at rlnJ ∼ 0.05 in the free troposphere; three models (NCAR, UCI, UCIb) show the largest extended frequency of |rlnJ| > 0.10 in the middle troposphere; UKCA is consistently the most clear-sky model. These model differences are not simply related to the model cloud fields; see discussion of Fig. S6 below.

Immediately above and below extensive thick cloud decks the dimming/brightening of Js exceeds the plotted range of rlnJ of ±0.3 (a factor of 1.35). Most such cloud decks occur around 900 hPa and so the largest brightening occurs in the 100–900 hPa levels, and the greatest dimming at >900 hPa. The top two rows in Fig. 4 give the fraction of samples for which rlnJ > 0.3 on the right side of each plot; the bottom row gives, on the left side, the fraction for which rlnJ < −0.3. The categorization of models and measurements is not simple as many models have shifting magnitudes of this large-scale brightening or dimming. A few models consistently lack these large changes in Js (GC, GMI), and a few always have them (NCAR, UCIb). Large CAFS values are clearly evident in both Js for only two of the six cases: > 900 hPa in the tropical Pacific (13–15% of all Js) and 300–900 hPa in the North Pacific (8–15%). For these cases the CAFS ex-
Figure 4. (a) Probability distribution of the natural log of the ratio of cloudy–clear $J$-O1D values ($r_{lnJ}$) from 10 models and from CAFS during ATom-1. The columns correspond to the two geographic blocks (tropical Pacific, 20° S–20° N × 160–240° E, and North Pacific, 20–50° N × 170–225° E). The rows are the three pressure layers (100–300, 300–900, 900–surface hPa). All histograms sum to 1, but for many models the peak values about $r_{lnJ} = 0$, corresponding to cloud-free skies, are truncated. Where a significant fraction of events does not fit within the ±0.3 range – on the high side for 100–900 hPa and low side for 900–surface hPa – the column of numbers, placed on the appropriate side and color coded to the legend, gives the fraction of occurrences outside the range. (b) Probability distribution of the natural log of the ratio of cloudy–clear $J$-NO2 values ($r_{lnJ}$) from 10 models and from CAFS during ATom-1. In general, $J$-NO2 is more responsive to clouds than $J$-O1D is.

treme fractions are consistent with at least four of the models. Any possible CAFS bias in $r_{lnJ}$ due to TUV modeling (±0.05) is unlikely to affect these results. These extreme fractions, however, are likely sensitive to any sampling bias of flight path with respect to thick cloud decks, and this needs to be assessed with model sampling that matches the ATom-1 profiles of that period.

The large geographic blocks were chosen to acquire representative sampling from the models that would be repeatable over time. It was not possible to acquire month-long or multi-year diagnostics from all the models, and so with the available model results (24 h from a day in mid-August) we subsample the broad tropical Pacific block (20° S–20° N × 160–240° E) into a west (160–200° E), east (200–240° E), and dateline (175–185° E) block. The results of these subsampled statistics are shown for $J$-NO2 in Fig. S7a (100–300 hPa) and b (surface–900 hPa) using the same format as Fig. 5. Each of the 18 panels in Fig. 7a, b represents a single model and a single pressure level for which there are seven bars (sampled distributions of $r_{lnJ}$). The top four bars show the full tropical Pacific (as in Fig. 5b) and then the three subsampled regions. The next three bars are the previously sampled statistics (North Pacific, global 50° S–50° N, and tropical Pacific CAFS). The CAFS bars are the same in each nine-panel plot. Six models show no difference across the subsampling (NCAR, GFDL, IFS, UCI, UKCA, MOCA), while three models using MERRA cloud fields in one way or another (GC, GMI, GISS) show weaker cloud effects in the east
Figure 5.

half of the region. For the most part, the subsampled regions have similar statistics that are distinct from the CAFS observations. Thus our model statistics over a large block may be a representative climatology of cloud effects on \( J \) values. Further sampling tests are recommended for follow-on work, especially to determine if the distinct east–west differences for the three MERRA cloud models are a standard feature.

### 3.2 Analyzing cloud effects

With a graphical synopsis of the rlnJ probability distributions in Fig. 5a, b, some model features become more obvious. We define nearly cloud-free conditions as being within \( \pm 2.5 \% \) of clear-sky \( J \)s, and show the frequency of these with the length of the thin line in the center of the plots. Starting with the \( J \)-O1D in the upper tropical Pacific, we find five models (Group 1: GC, GFDL, GMI, GISS, UKCA) show no effect of clouds more than 50 \% of the time. The other 40–50 \% of the time, they show enhanced \( J \)-O1D, cloud brightening expected from clouds below (thick lines on the right side of the plot). For the other five models (Group 2: IFS, MOCA, NCAR, UCI, UCldb), these clear-sky equivalent \( J \)s occur only 10–20 \% of the time, with cloud brightening enhancements occurring at 80–90 \%. Surprisingly, both model groups show the same average magnitude, +10 \% (Xs on the right side), for their enhanced \( J \)s. Thus Group 2 models will have systematically greater \( J \)-O1D in the upper tropical Pacific than the Group 1 models (i.e., the 10 \% enhancement occurs twice as often). In the North Pacific, this pattern holds although both groups show slightly greater frequency of enhanced \( J \)s, e.g., 50 to 60 \% for Group 1 and 80 to 90 \% for Group 2. For \( J \)-NO2, the results are similar, but with greater average magnitude of enhancement for cloudy skies (20 \% vs. 10 \%) and a slightly greater frequency of occurrence (thick line on the right). For this upper tropospheric layer, none of the models show significant occurrence of diminished \( J \) values from overhead clouds (rlnJ < −0.025) as seen in CAFS for 2–12 \% of the measurements.

In the middle troposphere (300–900 hPa, middle panels), the patterns in clear-sky frequency remain unchanged, but there is a shift to cloud dimming for 5 to 20 \% of the time. This shift to more cloud obscuration is much greater in CAFS than in any model. Group 1 models show consistently more frequent cloud obscuration (10–20 \%) than Group 2 models.
Figure 5. (a) Frequency of occurrence and magnitude of change in J-O1D caused by clouds. The panels and data sources are the same as in Fig. 4ab. The horizontal lines all have length 1 and show the fraction of (i) cloud-diminished Js (thick left segment, cloudy : clear ratio < 0.975), (ii) nearly cloud-free Js (thin central segment, 0.975 < ratio < 1.025), and (iii) cloud-enhanced Js (thick right segment, ratio > 1.025). Each line is plotted with its nearly cloud-free segment centered on 0. The mean magnitude of diminishment/enhancement corresponding to the thick line segments is plotted as an “X” on each line segment, using the x axis [−1, +1] as the natural log of the cloudy : clear ratio. Ratio changes of −20 % and +20 % are shown as dashed vertical grid lines. The Xs are not shown when the frequency of occurrence of either thick segment is < 0.02. For an example of how to read these figures consider the panel in row 3 column 1 (J-O1D, Tr. Pac, 900–srf). The GFDL model has about 22 % cloud-diminished Js (left segment) with an average value of 22 % below clear-sky Js (the X on the left side); most of the remaining, 76 %, are nearly cloud-free (central thin segment). The GISS model has (from left to right) 42 % cloud-diminished Js, 53 % cloud-free Js, and only 5 % cloud-enhanced Js for a total of 100 %; the cloud-diminished Js average about 28 % less than clear-sky Js (the X on the left) while the cloud-enhanced Js average about 22 % greater (the X on the right side). See legend in Fig. 5b. (b) Frequency of occurrence and magnitude of change caused by clouds in J-NO2.
obscuration leads to increased occurrence of \( rlnJ < -0.025 \) compared to the middle troposphere (in both CAFS and models). Even though the frequency changes, the average magnitude of diminishment when there is cloud obscuration (denoted by Xs) does not change much across models in either region. For CAFS in the tropical Pacific, however, the diminishment when there is cloud obscuration is much larger in the boundary layer. These differences could be caused by GMI calculating \( J \) values with a single-column atmosphere (SCA) containing clouds with Briglebe (CF\(^{1/3} \)) averaging and UCI calculating \( J \) values with four quadrature column atmospheres (QCAs); see Table 1. Unfortunately, when UCI mimics the B-averaging (with model UCIb), the differences remain. See further discussion of Fig. S6 in Sect. 4.1.

4 Model difficulties and development

The \( J \)-value statistics here depend on (i) the cloud fields used in the models, (ii) the treatment of cloud overlap statistics, (iii) the radiative transfer methods used, and (iv) the spectral data on sunlight and molecular cross sections. These components are deeply interwoven in each model, and it is nearly impossible to have the models adopt different components except for (iv), for which there has been a long-standing effort at standardization (e.g., the regular IUPAC and JPL reviews of chemical kinetics; Atkinson et al., 2004; Burkholder et al., 2015). These components are briefly noted in Table 1.

4.1 Cloud optical depths and overlap statistics

The models reported their average in-cell cloud optical depth (per 100 hPa) and cloud fraction over the two Pacific blocks in Fig. S6. Averaged cloud optical depths (defined for the visible region 500–600 nm) all tend to peak below 850 hPa in the Tropics and decline with altitude. There is clear evidence of mid-level (400–800 hPa) clouds, but only small COD (total < 0.25) at cruise altitudes (100–300 hPa). The North Pacific block has 2–4 times larger low-altitude COD. The plotted CF is the COD-weighted average over 24 h and all grid cells in the block. Note that for COD the cloud is spread over each model layer, and hence the in-cloud optical depth is estimated by COD/CF. CF is high, 5–15 % below 850 hPa, drops off with altitude as does COD, but peaks at 10–20 % near 200 hPa corresponding to large-scale cirrus. Some of these differences in COD and CF are large enough to explain model differences, but there is no clear pattern between \( J \) values and clouds as noted in Sect. 3.2. A more thorough analysis and comparison of the modeled cloud structures would involve the full climate models and satellite data (Li et al., 2015; Tsushima et al., 2017; Williams and Bodas-Salcedo, 2017), beyond the scope here.

4.2 Sensitivity of \( rlnJ \) to small cloud optical depth

To relate total COD to a shift in \( rlnJ \), the UCI off-line photochemical module Cloud-\( J \) was run for marine stratus (CF = 1) with a range of total CODs from 0.01 to 100. The cloud was located at about 900 hPa and \( rlnJ \) evaluated at 300 hPa. The plot of \( rlnJ \) vs. log COD for a range of SZAs is shown in Fig. S8. A 10 % enhancement (\( rlnJ = +0.10 \)) occurs at COD = 5 for \( J\text{-O1D} \) and COD = 3 for \( J\text{-NO2} \), demonstrating the greater sensitivity of \( J\text{-NO2} \) to clouds. Thus model
average total COD (ranging from 0.8 to 11 in Fig. S6, assuming CF = 1) should produce large shifts in rlnJ. Marine stratus with typical COD ∼ 10 or more would produce rlnJ-O1D of +0.16 and rlnJ-NO2 of +0.30. Thus clear-sky J values (defined here as ±0.025 in rlnJ) require COD < 1 for J-O1D and < 0.3 for J-NO2. A COD ∼ 1 is not that large since these clouds are highly forward scattering and have an isotropic-equivalent optical depth that is 5 times smaller.

4.3 Averaging over clouds affects clear-sky fraction

Comparison of the CAFS-ATom measurements of J values with modeled ones presents a fundamental disconnect, but one that we must work through if we are to test the J values in our chemistry–climate models with measurements. A CAFS observation represents a single point with unique solar zenith and azimuth angles within a unique 3-D distribution of clouds and surface albedos. Ozone column and temperature also control J values but are less discontinuous across flight path and model grids. One can define a column atmosphere (CA) for each J value in terms of the clouds directly above/below and the surface albedo, as would be measured by satellite nadir observations. The CAFS-measured actinic flux includes direct and diffuse light, which depends on all the neighboring CAs out to tens of kilometers. Adjacent clouds can either increase or decrease the scattered sunlight at the measurement site depending on the location of the sun.

By including cloud fractional coverage from the meteorological models, and attempting in various ways to describe cloud overlap, the models here recognize that the atmosphere is not horizontally homogeneous. Yet, for cost effectiveness and non-random J values, most modeling solves the radiative transfer problem for a 1-D plane-parallel atmosphere that is horizontally homogeneous. Most chemistry models adopt a simple averaging procedure to create a single, horizontally homogeneous cloudy atmosphere in each grid cell and then solving for a single J value (see Table 1). The UCI model uses decorrelation lengths to determine cloud overlap, to generate a set of independent column atmospheres (ICAs), to generate four quadrature column atmospheres (QCAs), and to calculate four J values, which are then averaged to get a single J value. In either case, the radiative transfer solution is 1-D and there is one J value per grid cell given to the chemistry module (and analyzed here).

What would the probability distribution rlnJ look like if we used the UCI J values from the QCAs before averaging? For this, we collect the statistics on total COD for the two geographic blocks and compare the sub-grid QCA CODs against averaging approaches in Fig. 6. This histogram (blue dots) is our best estimate of the distribution of total COD from a 1-D nadir perspective of all the sub-grid ICAs. The UCI J values are calculated as the average over four QCAs as representatives of the ICAs. The two SCA models use simple averaging (COD × CF, green dots) and B-averaging (COD × CF²/2, orange dots), which reduces the cloud fraction but assumes maximal cloud overlap. In these cases, a single J-value calculation is made with the SCA. When the clouds are simply averaged over the grid cell (∼ 1° × 1°), the clear-sky occurrence drops to 7% and the deep cumulus disappears. When Briegleb (1992) B-averaging is used, there is only slightly more clear sky (12%). Both averaging methods also reduce the occurrence of thick cumulus. When run at lower resolution, both averages find less clear sky, while the QCA statistics are not affected by resolution in the range 50–200 km. This averaging, of either J values or clouds, explains why most models do not produce a single, sharp peak at rlnJ = 0.

At sufficiently high model resolution, where CF is either 0 or 1, a new problem arises because the radiative transfer problem is now clearly 3-D. The 1-D radiative transfer used here would produce a very sharp clear-sky peak in rlnJ that is not seen in CAFS. The CAFS rlnJ distribution is widened in part by TUV albedo biases, but also because it is effec-
ivamente a weighted average of cloud conditions over tens of kilometers or more and thus has lower frequency of clear sky than the 1-D ICAs do. The Group 1 models with large peak distributions at rlnJ ~ 0 appear to be basically incorrect since averages over these model resolutions (>0.5°) should reduce clear-sky occurrence. There is probably a sweet spot in model resolution at about 20 km where the model statistics, even with 1-D RT, should match the observed statistics.

4.4 Ocean surface albedo

We chose our Pacific blocks for this comparison to avoid large aerosol contributions and to be oceanic to avoid large variations in surface albedo. Nevertheless, the ocean surface albedo (OSA) is variable (Jin et al., 2011), but most of these models, including TUV, assume a uniform low albedo in the range of 0.05 to 0.10. For the modeled ratio J-cloudy to J-clear, using a fixed albedo is not so important since both J values use the same albedo. For the CAFS / TUV ratio, however, it is essential to have the TUV model use the OSA that best corresponds to the sea surface conditions under the CAFS measurement. Work on the CAFS / TUV calibration seeks to achieve this zero bias, and it continues beyond the cutoff date of the ATom-1 data used here. The OSA affects our 2 Js differently: for J-O1D with peak photolysis about 305 nm, the OSA under typical conditions (SZA = 20°, surface wind = 10 m s⁻¹, chlorophyll = 0.05 mg m⁻³) is 0.038, while for J-NO2 with peak photolysis at 380 nm, the OSA is 0.048. OSA depends critically on the incident angle of radiation, increasing from 0.048 at 20° to 0.068 at 50° (380 nm). Rayleigh scattered light has on average larger incident angles than the solar beam for CAFS measurements and is reflected more than the direct beam. Rayleigh scattering is much more important for J-O1D than J-NO2.

The UCI stand-alone photolysis model was rewritten to include a lower boundary albedo that varies with angle of incident radiation, and is now designated Cloud-J version 8 (v8). In Cloud-J there are five incident angles on the lower surface: the direct solar beam and the four fixed-angle downward streams of scattered light. The OSA modules are adapted from the codes of Séférian et al. (2018) based on Jin et al. (2011), but we do not use their approximation for a single “diffuse” radiation since all of Cloud-Js scattered light is resolved by zenith angle. The resulting albedo is a function of wavelength, wind speed, and chlorophyll; it is computed for each SZA and the four fixed scattering angles. As a test of the importance of using a more realistic OSA, we used Cloud-J v8 to compute the ratio of J with our OSA module to J using a constant fixed albedo, with both Js calculated for clear sky. We calculate this rlnJ (log of the ratio of J-clear (OSA) to J-clear (single albedo)) for a range of SZAs, wind speeds, and chlorophyll comparing to fixed albedos of 0.00, 0.06, and 0.10 as shown in Fig. S9. Because the range in conditions, there is no single offset, but we have a probability distribution whose width in rlnJ due to a range of surface albedo has a magnitude that affects the interpretation of measurement–model differences. For high sun (SZA = 0–40°), choosing an optimal fixed albedo of 0.06 results in little mean bias, although individual errors are about ±2 %. This error is based on conditions for cos(SZA) > 0.8. If the fixed albedo differs from this optimum (e.g., 0.00 or 0.10), then bias errors of 2 % to 10 % appear, and the width of the distribution expands greatly. For SZA = 40–80°, the optimum fixed albedo starts showing bias and has a much broader range of errors under different circumstances (wind, chlorophyll, SZA). Thus J values calculated using unphysical, simplistic fixed ocean surface albedos can have errors of order ±10 % depending on ocean surface conditions and the angular distribution of direct and scattered light at the surface. These errors will not directly affect the model results here since the cloudy–clear differences used a self-consistent albedo in each model. Overall, however, there is a need for chemistry models to implement a more physically realistic OSA.

5 Discussion

The importance of clouds in altering photolysis rates (Js) and thence tropospheric chemistry is undisputed. On a case level it is readily observed, and on a global level, every model that has included cloud scattering finds significant changes in chemical rates and budgets. For example, the inclusion of cloud layers by Spivakovsky et al. (2000) caused a shift in peak OH abundance from the boundary layer to just above it, resulting in a shift to colder temperatures (and lower reaction rates) for the oxidation of CH₄-like gases, even with the same average OH abundance. Other than single-column, idealized, off-line tests of radiative transfer methods, we have few methods to constrain modeled Js under cloudy conditions.

The impact of clouds on Js is large, simply by looking at the profile of mean Js (Figs. 1, 2, and 3), and it greatly complicates the comparison of J values across models. The modeled mean clear-sky profiles, including the CAFS TUV model (Fig. 2), tend to agree within ±10 % with the exception of MOCA, GISS, and sometimes IFS. However, the all-sky (cloudy) profiles (Fig. 1) have a much wider spread, except for J-O1D in the North Pacific for which there is inexplicably a core group of eight models within ±10 %. The observed CAFS cloudy profiles show a distinctly different profile in the North Pacific vs. the tropical Pacific for both J-O1D and J-NO2 (Fig. 1) and especially in the CAFS / TUV derived J-cloudy to J-clear ratios (Fig. 3). Many models follow this typical profile in the tropical Pacific (100–900 hPa; Fig. 3), but some (NCAR, UCI, UCLA) show large shifts to enhanced J values immediately above the marine boundary layer. Whether this fundamental shift in cloud regimes is robust remains uncertain, and may be resolved with the full set
of ATom deployments or with careful model studies; see below.

A more informative diagnostic of cloud effects is the statistical distribution of the individual $J$-cloudy to $J$-clear ratios ($rlnJ$). To model clouds correctly we need to understand how frequently and by how much clouds interfere with $J$ values. The statistical distribution of $rlnJ$ shows distinct patterns and classes of models. In the free troposphere (100–900 hPa), all have a sharp edge at $rlnJ = 0$ with hardly any diminished $J$ values ($rlnJ < 0$), but the small-cloud-effects models (GC, GFDL, GISS, GMI, UKCA) show a dominant peak at $rlnJ \sim 0$ while the large-cloud-effects models (IFS, MOCA, NCAR, UCI, UCib) have extensively enhanced $J$ values with large fractions at +25% or more. In the marine boundary layer (surface–900 hPa), this pattern is reversed with a sharp edge for all models at $rlnJ \sim +0.03$, and the large-cloud-effects models showing more diminished $J$ values. CAFS/TUV has its own pattern, which is very broadened, but generally supports the large-cloud-effect models. If further analysis can tighten the CAFS $rlnJ$ distribution, then we may be able to discriminate among the models in one of the classes. One robust result from the models is that large cloud effects ($|rlnJ| > 0.05$) are always asymmetric, splitting in sign above and below 900 hPa. If the symmetric spread in CAFS $rlnJ$ remains robust with additional deployments and efforts to reduce CAFS-TUV inconsistencies, then we have a clear challenge for the cloud climatologies used in the chemistry models.

More work on the CAFS/TUV data could help better discriminate among the model classes identified here. For one, we need to sample over different seasons and synoptic conditions to build a more robust climatology. Fortunately, the additional three deployments (ATom-2, -3, -4) will provide these. They occur in different seasons, which is a consideration, but the tropical oceanic data can probably be combined. The analysis can be extended to the Atlantic. CAFS or similar measurements from other aircraft missions could also be added. Other tasks involve tightening the spread in $rlnJ$ by looking for potential measurement–model noise (e.g., aircraft–sun orientation, sun–cloud geometries) and by improving the TUV clear-sky modeling (e.g., a more accurate ocean surface albedo derived from observed surface wind, chlorophyll, SZA). An unresolved issue is how to treat aerosols, both in the models and in CAFS. If clear sky includes aerosols then TUV must be able to infer an aerosol profile for all measurements, and likewise the models need to be careful in how they calculate cloudy–clear ratios. Fortunately for most of the oceanic ATom measurements, aerosol optical depths are small, but where they are large (e.g., Saharan dust events) the daily satellite mapping should provide adequate coverage for the TUV modeling. Developing these cloudy–clear $J$-value statistics over land will be more difficult due to the higher inherent variability in albedo and aerosol profiles.

More effort is needed from the modeling community to characterize the key factors driving these model differences in photolysis rates under realistic, cloudy conditions. This might include sensitivity runs that address aerosol and surface albedo impacts for each model. We would also need a better characterization of the cloud distributions used in chemistry models, including comparison with satellite climatologies (Cesana and Waliser, 2016; Ham et al., 2017), to understand how cloud fraction and overlap affects the $J$s used in the photochemical calculation of a column atmosphere. Models can help assess the ATom CAFS statistics for representativeness by checking on flight routing, multiple days, or different years.

The 3-D nature of the radiation field measured by aircraft presents a more fundamental challenge. The observations average over cloud fields out to tens of kilometers, and even if the atmospheric column at the aircraft is clear, neighboring clouds alter $J$ values. The chemical models are coarse resolution compared with the CAFS measurements and average over wider range of cloud fields, almost eliminating the occurrence of clear-sky conditions. Even if they calculate a distribution of $J$ values for the cloud statistics in a column (like Cloud-$J$), they combine these to deliver a single average $J$ value for the chemistry module, again reducing the occurrence of clear-sky $J$ values. Thus, the result above (i.e., that five models have about 50% nearly clear-sky $J$ values down to the surface) is inexplicable unless they have column optical depths of 0.3 or very small effective cloud fractions. At some model resolution, probably of order tens of kilometers, the CAFS measurements may be a statistical representation over that grid, and our comparisons, even with 1-D radiative transfer over a range of ICAs, may be more consistent. At these scales, the problem of a strong zenith angle dependence remains (Tompkins and Giuseppe, 2007). Super-high-resolution models (~1 km) are becoming available in regional or nested-grid models (Kendon et al., 2012; Schwartz, 2014; Berthou et al., 2018), and one might hope that our problem is now solved because each single-column atmosphere (SCA) explicitly resolves cloud overlap, with each grid cell being either cloudy or clear. The calculation of photolysis and solar heating rates is not simplified, however, because now the SCAs interact with neighbors. Calculating the correct rates at any location, or even the average over a region, would require that we calculate the ratio of cloudy to clear over a 20 km domain of cloud-resolved grid cells.
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Supplement of

Cloud impacts on photochemistry: building a climatology of photolysis rates from the Atmospheric Tomography mission

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Contents of this file
   Figures S1 to S9
   Table S1

This Supplementary Material includes figures and tables referenced in the main text and useful for understanding this work, but not essential to its findings.
Figure S1. The probability distribution of solar zenith angles (SZAs) as sampled by ATom with the CAFS instrument (red) and by two models (UCI & NCAR) using regular sampling over the two geographic blocks: (1, left) Tropical Pacific and (2, right) North Pacific. The points represent the relative number of samples in each 0.10 interval of cos(SZA). ATom flights in this region were generally centered about noon, and hence CAFs has a higher proportion of high cos(SZA) values. For most of the analysis here only J's with cos(sza) > 0.8 are used, and with this restriction the number of CAFS measurements is 11,504 (block 1) and 4,867 (block 2). For UCI the number of points is 240,000 (block 1) and 120,000 (block 2); while for the higher resolution NCAR model, the values are 1,400,000 and 700,000, respectively.
Figure S2. Mean CAFs profiles of J-O1D (left) and J-NO2 (right) from ATom-1 made under all-sky conditions for high-sun conditions, cos(sza) > 0.8. The 3 geographic blocks shown here are the standard blocks 1 (blue, Tropical Pacific, 20°S – 20°N x 160°E – 240°E), and 2 (red, North Pacific, 20°N – 50°N x 170°E – 225°E, plus a global block (black, 55°S – 55°N, all longitudes, including the Atlantic). Each 3-sec observation is averaged with equal weight in 100-hPa pressure bins. The standard deviation for block 1 (blue, *) is shown along with that from the corresponding TUV model calculation for clear sky (blue, o). The patterns here are expected: J-O1D is sensitive to O3 absorption and hence has higher upper-tropospheric values in the tropics; J-NO2 is more sensitive to cloud scattering and hence has higher values in the lower troposphere of the North Pacific. The standard deviations show that J-O1D variance is driven by O3 and SZA, and not by clouds; while the J-NO2 shows the opposite.

Figure S3. Total ozone columns (in DU) versus latitude in August for 9 models and 8 years of OMI observations (2010-2017, Levelt et al., 2006; Veefkind et al., 2006). Most curves are the monthly zonal means, but some data are from a single day, or just the ATom flight track for several days (e.g., GFDL, explains the lack of smoothness). Differences here can explain only some of the spread in clear-sky J-O1D values in Figure 2.
Figure S4. Effects of clear sky (blue points), simple averaged clouds (gray points), and fractional cloud overlap (the reference case, 1:1 dashed line) using the UCI CTM. X-axis is the same as the Y-axis, but for the reference model UCI 2016. Direct parcel-by-parcel comparison of modeled 24-hour reactivities (top row: P-O3, L-O3, L-CH4; all ppb/day) and photolysis rates (J-NO2, J-O1D; all /sec) calculated for a data stream of 14,880 simulated air parcels from 60 °S to 60 °N, 0.5 km to 12 km altitude, and along 180 °W, see Prather et al., 2018. Each point is an average of the 5 simulated dates in August (8/01, 8/06, 8/11, 8/16, 8/21). UCI 2016 ref uses full cloud quadrature and the newly implemented decorrelation lengths for cloud overlap (Prather, 2015). From the figure, the different cloud treatments are clearly visible in the shifts in J-NO2 and J-O1D, and they have largest effect on P-O3 as compared with L-O3 and L-CH4. On average, the clear-sky simulation has 3-4% lesser J-values and similar changes in all 3 reactivities. The averaged-cloud method has about 12 % greater J-values (mostly above clouds), increasing reactivities by 5 % (L-O3 and L-CH4) to 10% (P-O3).
Figure S5. Histogram of the natural log of the ratio of cloudy-to-clear J values (J-O1D & J-NO2), designated rlnJ, which is calculated from the 10-sec CAFS data and sorted into 3 pressure bins (blue, 100 – 300; red, 300 – 900; black, 900 – surface hPa). Note that the upper level has a peak near 0.0, corresponding to clear sky; the lowest level extends more to values <0; and that the N. Pacific for <900 hPa has a large number of cloud enhanced J-values (rlnJ >0). The CAFS data were binned at 0.01 rlnJ and smoothed with a 1-2-1 filter six times.
Figure S6. Averaged in-cell cloud optical depth (COD, at ~600 nm and per 100 hPa) and cloud fraction (CF) over the two geographic blocks studied here (Tropical Pacific, 20 °S – 20 °N x 160 °E – 240 °E, and North Pacific, 20 °N – 50 °N x 170 °E – 225 °E). Averages are made over 24 hours for one day in mid-August as used in this paper. For these data (unlike the J-value statistics), all hours are weighted equally independent of solar zenith angle. Column total COD for each region is given in the order/color-indexed numbers on the right of each COD panel.
Figure S7a. The effect of sampling different regions of the tropical Pacific in terms of the frequency of occurrence and magnitude of changes caused by clouds in J-NO2 at 100-300 hPa. See Figure 5. Each of the 9 panels shows a single model with different sampling and also the CAFS observations over the tropical Pacific (bottom black bar and X's). The top purple bar is the entire tropical Pacific as defined in Figure 5 (20S-20N, 160E-240E) and the next three bars (blue, green, yellow) sub-sample this region into West (160E-200E), East (200E-240E) and 180E (175E-185E), respectively. The next two bars are the North Pacific (as in Figure 5) and a global sampling (50S-50N, all longitudes).

Figure S7b. The effect of sampling different regions of the tropical Pacific in terms of the frequency of occurrence and magnitude of changes caused by clouds in J-NO2 at surface-900 hPa. See Figure S7a.
Figure S8. The ratio of J-cloudy/J-clear calculated for J-O1D (left) and J-NO2 (right) at 300 hPa for a marine stratus cloud (CF = 100%) at about 900 hPa. The natural log of the ratio (rlnJ) is calculated for a cloud optical depth (COD) ranging from 0.1 to 100. The calculation uses Cloud-J (Prather, 2015), which does not scale COD, uses the truncated (order 8) phase function, and applies a constant lower boundary albedo = 0.05. The incident SZA ranges from 0º (black) to 60º (red) in 10º intervals (blue). The rlnJ's in this paper are restricted to 0º – 40º. A 5% enhancement (rlnJ = 0.05) occurs at COD = 2 for J-O1D and COD = 1 for J-NO2, demonstrating the greater sensitivity of J-NO2 to clouds.

Figure S9. Probability distribution of the natural log of the ratio of J-values calculated using an interactive ocean surface albedo (OSA) to J-values using a fixed albedo. J-O1D values (left) and J-NO2 values (right) assume clear skies for both OSA and fixed-albedo. This format is similar to Figure 4. The interactive OSA model includes a dependence on wavelength and incident-angle now in Cloud-J v8, see text. The fixed albedo is denoted by 'a = ' in the legend. These distributions show the relative error in J-values calculated in typical models using a fixed albedo instead of the more physically based OSA models (Séférian et al., 2018). For both cases the reflected sunlight is assumed to be isotropic. The OSA depends on surface wind (sampled uniformly from 1 to 21 m/s) and chlorophyll abundance (sampled uniformly in log from 0.01 to 30 mg/m³). The plots are also split between the lowermost troposphere (lower panels, >850 hPa) and free troposphere (upper, 200 – 850 hPa).
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