




# THE CONVECTIVE TRANSPORT OF ACTIVE SPECIES IN THE TROPICS (CONTRAST) EXPERIMENT

L. L. PAN, E. L. ATLAS, R. J. SALAWITCH, S. B. HONOMICHL, J. F. BRESCH, W. J. RANDEL, E. C. APEL,  
R. S. HORN BROOK, A. J. WEINHEIMER, D. C. ANDERSON, S. J. ANDREWS, S. BAIDAR, S. P. BEATON,  
T. L. CAMPOS, L. J. CARPENTER, D. CHEN, B. DIX, V. DONETS, S. R. HALL, T. F. HANISCO, C. R. HOMEYER,  
L. G. HUEY, J. B. JENSEN, L. KASER, D. E. KINNISON, T. K. KOENIG, J.-F. LAMARQUE, C. LIU, J. LUO, Z. J. LUO,  
D. D. MONTZKA, J. M. NICELY, R. B. PIERCE, D. D. RIEMER, T. ROBINSON, P. ROMASHKIN, A. SAIZ-LOPEZ,  
S. SCHAUFFLER, O. SHIEH, M. H. STELL, K. ULLMANN, G. VAUGHAN, R. VOLKAMER, AND G. WOLFE





Deep convection over the tropical western Pacific viewed at ~40,000 ft above sea level from the NSF/NCAR Gulfstream V research aircraft during the CONTRAST experiment. Photo credit: Rebecca Hornbrook.

**Airborne observations over the tropical western Pacific warm pool characterize the role of tropical convection in linking oceanic processes to ozone chemistry in the upper troposphere and lower stratosphere.**

**T**he tropical western Pacific (TWP) warm pool is a uniquely important region for Earth's climate system. The warm pool, often defined by sea surface temperature (SST) in excess of 28°C (Wyrski 1989), is the largest source of latent heat release and water vapor into the atmosphere, and the center of action for El Niño–Southern Oscillation (ENSO) (Webster and Lukas 1992). Fueled by the warm sea surface, the region has intense and massive deep convection and precipitation (Liu and Zipser 2015). At the upper troposphere/lower stratosphere (UTLS) level, strong upwelling during the Northern Hemisphere (NH) winter season couples to the deep convection to make the TWP region the largest source for tropospheric air entering the stratosphere (Newell and Gould-Stewart 1981; Fueglistaler et al. 2004; Krüger et al. 2008; Bergman et al. 2012). This is also the region of coldest temperatures in the tropical tropopause layer (TTL) during NH winter (Fueglistaler et al. 2009; Randel and Jensen 2013); therefore, it plays a critical role in controlling the amount of water vapor that enters the lower stratosphere (Newell and Gould-Stewart 1981; Holton and Gettelman 2001; Schoeberl and Dessler 2011). The effect of dehydration in the region is reflected in satellite cloud observations, which show that the region has the largest fraction of high ►

clouds (17–18-km level) (e.g., Yang et al. 2010) and significant occurrence of clouds above the tropopause (Pan and Munchak 2011).

The unique thermal and dynamical behaviors of this region also create a special chemical environment. A number of field studies and satellite data have identified extremely low  $O_3$  in the region, near or below the detection limit of typical ozonesondes. This is especially true at the TTL level where measurements of  $O_3$  below 20 ppbv suggest a major contribution of convectively lifted air from the oceanic boundary layer (e.g., Kley et al. 1996; Crawford et al. 1997; Thompson et al. 2011; Rex et al. 2014). This low-ozone environment may have an impact on the abundance of hydroxyl radical (OH) in the troposphere, which is a significant agent for self-cleaning of the atmosphere. Deep convection connects emissions from oceanic biological processes to the UTLS. Thus, the UTLS over the oceanic warm pool region is expected to have elevated concentrations of organic halogen species such as dibromomethane ( $CH_2Br_2$ ), bromoform ( $CHBr_3$ ), and methyl iodide ( $CH_3I$ ). These compounds are significant contributors to the input of reactive bromine and iodine into the mid- to upper tropical troposphere where they, along with their inorganic breakdown products, represent an important component of the natural background ozone budget (Saiz-Lopez et al. 2012; Wang et al. 2015). These short-lived halogens and their inorganic oxidation products may have additional impacts on stratospheric chemistry (Salawitch et al. 2005; Fernandez et al. 2014; Saiz-Lopez et al. 2015). Our measurements in the TWP were designed to address current uncertainties in both the halogen-mediated loss of ozone in the UTLS as well as the budget and partitioning of reactive halogen inputs to the stratosphere.

Recognizing the importance of the TWP region, a number of large-scale field experiments have been conducted to investigate a range of atmospheric dynamics and chemistry questions. Ocean–atmosphere coupling was the focus of Tropical Ocean and Global Atmosphere Coupled Ocean–Atmosphere Response Experiment (TOGA COARE) (1988), which deployed large arrays of soundings and collected measurements from multiple aircraft (Webster and Lukas 1992). The NASA Global Tropospheric Experiment (GTE) program conducted a series of airborne studies using the DC-8 aircraft and obtained detailed chemical information over the tropical Pacific, including the warm pool region (e.g., Hoell et al. 1999) up to 12-km altitude. The Biomass Burning and Lightning Experiment (BIBLE) campaign, using a Gulfstream II research aircraft, focused on the impact of lightning and biomass burning emissions of aerosols and trace gases on ozone chemistry (Kondo et al. 2002). Despite these studies, the impact of convection on chemistry in the warm pool region has not previously been well characterized, especially at the TTL level (13–17 km). Figure 1 shows the climatological location of the warm pool in the NH winter season and locations of ozonesonde launch sites, which indicate the lack of ozone measurements in the core of the warm pool. The lack of adequate observations of trace-gas composition, including ozone and reactive halocarbons, in the warm pool region leaves a significant gap in our understanding of the chemistry within the TTL and the transport into stratosphere over the TWP.

A broad range of scientific objectives motivated the coordination of three airborne experiments for research over the TWP warm pool. The objectives required aircraft with the appropriate instrument payloads that could cover altitudes from the marine

**AFFILIATIONS:** PAN, HONOMICHL, BRESCH, RANDEL, APEL, HORN BROOK, WEINHEIMER, BEATON, CAMPOS, HALL, JENSEN, KASER, KINNISON, LAMARQUE, J. LUO,\* MONTZKA, ROMASHKIN, SCHAUFFLER, AND ULLMANN—National Center for Atmospheric Research, Boulder, Colorado; ATLAS, DONETS, AND RIEMER—University of Miami, Coral Gables, Florida; SALAWITCH, ANDERSON, AND NICELY—University of Maryland, College Park, College Park, Maryland; ANDREWS AND CARPENTER—University of York, York, United Kingdom; BAIDAR, DIX, KOENIG, AND VOLKAMER—University of Colorado Boulder, Boulder, Colorado; CHEN AND HUEY—Georgia Institute of Technology, Atlanta, Georgia; HANISCO—NASA Goddard Space Flight Center, Greenbelt, Maryland; HOMEYER—University of Oklahoma, Norman, Oklahoma; LIU—Texas A&M University—Corpus Christi, Corpus Christi, Texas; Z. J. LUO—City College of New York, New York, New York; PIERCE—NOAA Satellite and Information Service (NESDIS) Center for Satellite Applications and Research

(STAR), Madison, Wisconsin; ROBINSON AND SHIEH—University of Hawai'i at Mānoa, Honolulu, Hawaii; SAIZ-LOPEZ—Institute of Physical Chemistry Rocasolano, CSIC, Madrid, Spain; STELL—National Center for Atmospheric Research, Boulder, and Metropolitan State University, Denver, Colorado; VAUGHAN—University of Manchester, Manchester, United Kingdom; WOLFE—NASA Goddard Space Flight Center, Greenbelt, and University of Maryland, Baltimore County, Baltimore, Maryland

\***CURRENT AFFILIATION:** Lanzhou University, Lanzhou, China  
**CORRESPONDING AUTHOR E-MAIL:** Laura L. Pan, liwen@ucar.edu

*The abstract for this article can be found in this issue, following the table of contents.*

DOI:10.1175/BAMS-D-14-00272.1

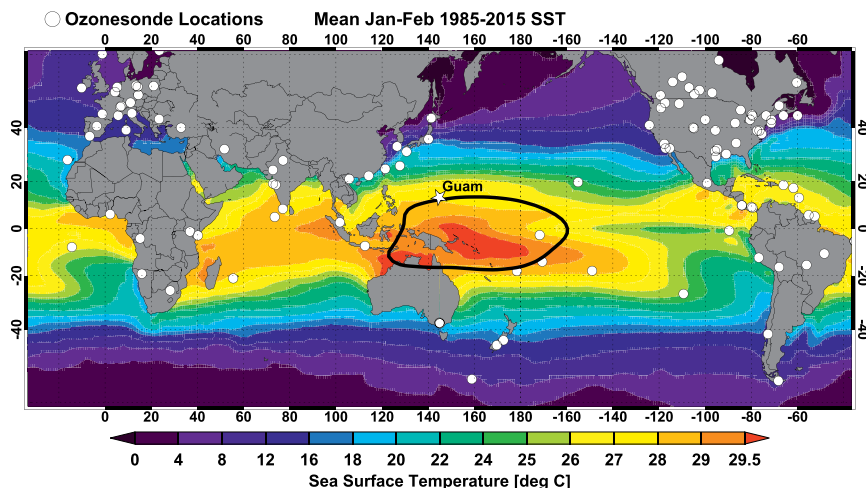
In final form 22 February 2016  
©2017 American Meteorological Society



boundary layer (MBL) to the lower stratosphere (LS). The major objectives addressed by the three aircraft complement include better understanding the role of deep convection in coupling the MBL to UTLS chemical composition, characterization of the processes controlling water vapor and short-lived chemical species transported into the stratosphere, definition of the abundance and partitioning of halogen species and their impact on UTLS ozone, and evaluation of the chemical coupling of ocean and atmospheric oxidation capacity in the warm pool region. The experiments were all based in Guam (13.5°N, 144.8°E) with research flights conducted during January and February 2014. The Airborne Tropical Tropopause Experiment (ATTREX) used the high-altitude NASA Global Hawk (GH) to sample chemical and microphysical parameters at altitudes between ~14 and 18 km [see ATTREX overview by Jensen et al. (2017), this issue]. The Coordinated Airborne Studies in the Tropics (CAST) experiment used the U.K. FAAM BAe 146 research aircraft to measure a large suite of chemical tracers from the MBL up to ~7-km altitude [see CAST overview by Harris et al. (2017), this issue]. The CONTRAST experiment used the National Science Foundation/National Center for Atmospheric Research (NSF/NCAR) Gulfstream V (GV) research aircraft to sample a wide variety of chemical species from the MBL to ~15 km. The concept of the coordinated campaigns is illustrated in Fig. 2.

This article provides an overview for the CONTRAST experiment, including discussions of scientific synergy with the other two coordinated experiments. Within the context of the coordinated campaigns, the main scientific objectives of the CONTRAST experiment were as follows:

- characterizing the influence of deep convection on the chemical composition and the photochemical budget of O<sub>3</sub> at the level of convective outflow over the western Pacific,
- evaluating the budget of organic and inorganic bromine and iodine in the TTL, and



**FIG. 1. Thirty-year climatology of Jan–Feb SST highlighting the TWP warm pool.** The SST, shown by the color shaded contours, is from NOAA Optimum Interpolation data version 2 (Reynolds et al. 2002). The 30-yr mean 193-K temperature contour at the 100-hPa pressure level (black contour) is shown to indicate the coldest region of TTL. Also shown are locations of ozonesonde measurements (white circles) obtained from WOUDC (<http://woudc.org/home.php>), which include the SHADOZ stations.

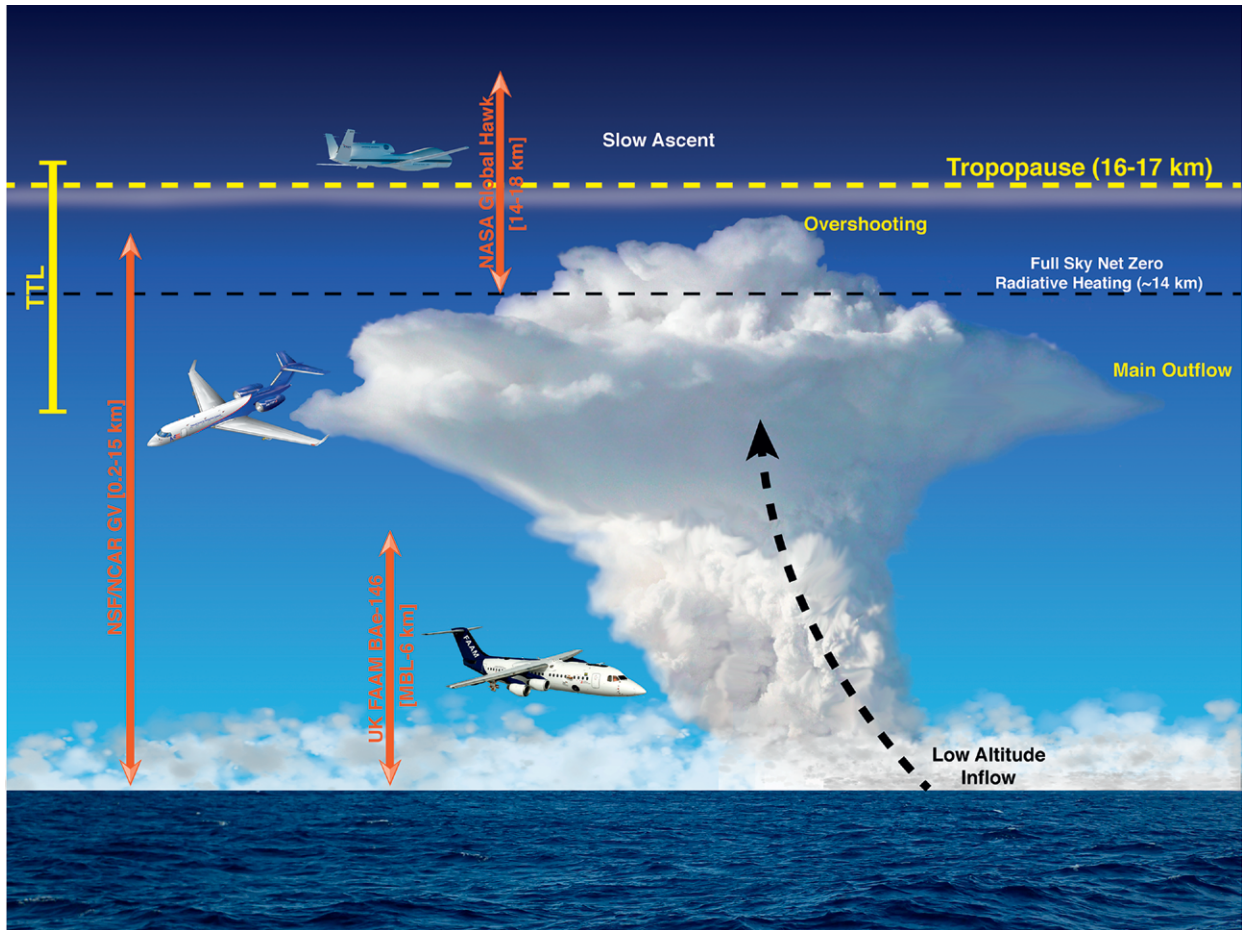
- investigating transport pathways from the oceanic surface to the tropopause using coordinated flights with CAST BAe-146 and ATTREX GH.

This overview presents the design and implementation of the CONTRAST experiment, the background information for using CONTRAST data, and selected scientific and operational highlights. These highlights are chosen to promote new research opportunities that CONTRAST data may enable. The description of the operation and facility may also serve as a useful reference for future field campaign planning. See the appendix for definitions of the various acronyms used in the text.

**METEOROLOGICAL SETTING OF THE EXPERIMENT.** Located at the northern edge of the warm pool near the division of rising and sinking branches of the Hadley cell in the NH winter season, Guam was an ideal base for the campaign. The GV flights from Guam have access to areas of active convection (to the south) and to the subtropical jet stream (to the north). Furthermore, the flight operations and instrument maintenance were less challenging since the island was in its dry season, and excellent logistical support for airborne operations was available on Guam.

Figure 3 shows the distribution of the 12–14-km cloud fraction from *CloudSat* data, together with additional key dynamical elements important to the Guam operational domain. The wind direction and





**FIG. 2. Concept of the coordinated campaigns.** This schematic highlights the three research aircraft and the altitude ranges of their sampling in relation to the dynamical background. The key feature of the background is deep convection that nearly reaches the tropopause and which pumps marine boundary layer air into the TTL (~13–17 km). The level of full-sky net zero radiative heating, estimated to be near 14 km, is typically used to indicate the transition into mean upward motion driven by large-scale dynamical control (Corti et al. 2005).

the geopotential height (GPH) indicate that, over the region of persistent deep convection, the upper-tropospheric flow is dominated by a large-scale anticyclonic pattern that is symmetrically distributed with respect to the convective axis (Dima and Wallace 2007). The GV nominal operational range is represented in Fig. 3 by the circle of 1,500 nautical miles (n mi) radius. An additional circle of 1,000 n mi radius is shown in Fig. 3 to indicate the range when planning extensive loitering was considered in the flight plan. The northern part of the research domain includes an intense jet stream, illustrated by contours of wind speed greater than  $50 \text{ m s}^{-1}$ . The jet core marks the dynamical division of the tropics and extratropics. The increase of potential vorticity (PV) near the jet indicates the rapid change of air mass character from tropospheric to stratospheric at UTLS levels.

Winter 2014 was characterized as ENSO neutral with a Niño index of about  $-0.5$  [National Oceanic

and Atmospheric Administration (NOAA) Climate Prediction Center (CPC); [www.cpc.ncep.noaa.gov/products/analysis\\_monitoring/enso\\_disc\\_mar2014/](http://www.cpc.ncep.noaa.gov/products/analysis_monitoring/enso_disc_mar2014/)]. SST anomalies over the western Pacific were slightly positive ( $+1^\circ\text{C}$ ) between  $0^\circ$  and  $10^\circ\text{N}$  during January and February on the northern side of the climatological warm pool. The equatorial trough was a common feature in low latitudes during the period with a few periods of enhanced low-level convergence over the study region. Two significant Madden–Julian oscillation (MJO) events occurred during the study period. The first, during the second half of January, was accompanied by the formation of tropical storms Lingling and Kajiki near the Philippines. These storms may have helped Guam receive near-record January rainfall. The second MJO event occurred in late February and was accompanied by the formation of Typhoon Faxai, which was located mostly east of Guam ( $145^\circ\text{E}$ ). Repeated cold fronts/shear lines moved

from the north to Guam's latitude during early 2014, which enhanced surface northeasterlies and promoted low-level convergence and precipitation.

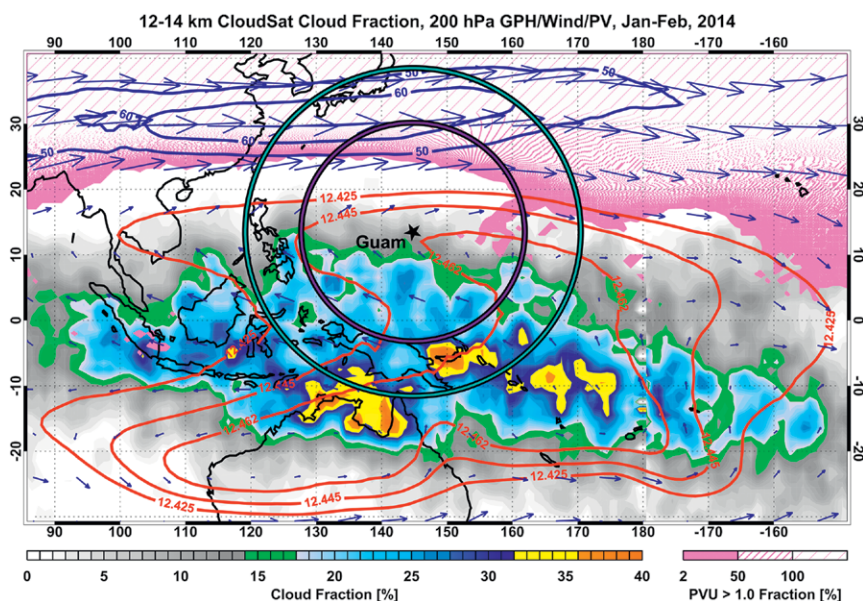
**DESIGN AND OPERATION OF THE EXPERIMENT.** *Platform and payload.* The base in Guam coupled with the range and duration of the NSF/NCAR GV aircraft allowed sampling from MBL to the lower TTL, including the level of the main convective outflow (12–14 km) in the warm pool region. With the CONTRAST payload and aircraft configuration, the GV typically flew at flight altitudes between 13 and 14 km with a flight ceiling near 15 km (~48,000 ft in pressure altitude). Figure 4 and Table 1 summarize the GV payload configuration during the CONTRAST experiment.

To meet the scientific objectives of the CONTRAST mission, the GV payload was designed to characterize the chemical and photochemical environment of TWP at all altitudes, especially the level of convective outflow. The instrument payload had sensors for various trace gases, aerosols, and radiation. The trace-gas measurements were aimed at gases with different sources, atmospheric trends, and chemical lifetimes. These trace-gas properties allowed an examination of long-range and convective transport and chemical reactivity in the CONTRAST region.

To measure organic halogen composition, the payload included a combination of the Advanced Whole Air Sampler (AWAS), a canister sampling system, the trace organic gas analyzer (TOGA), an in situ online gas chromatograph/mass spectrometer (GC/MS), and fast-response instrumentation for CO, CO<sub>2</sub>, and CH<sub>4</sub>. The TOGA and AWAS provide complementary measurements of trace gases. TOGA measured approximately 75 volatile organic compounds (VOCs), including C<sub>3</sub>–C<sub>8</sub> nonmethane hydrocarbons (NMHC),

halocarbons, organic nitrates, and several C<sub>1</sub>–C<sub>4</sub> oxygenated VOC (OVOC) (Apel et al. 2015). The AWAS measured a full suite of NMHC, halocarbons, and organic nitrates at high precision but with lower spatial resolution compared to TOGA. A number of trace gases measured by both systems provided good overlap for data comparison and complementary sampling. In addition, whole air samplers were deployed on all three aircraft [see Jensen et al. (2017) and Harris et al. (2017), this issue] and provide a consistent dataset from the MBL to the LS (see Fig. 9).

Measurements of radiation and relevant reactive gases provided data to define the photochemical environment of the tropical UT, to examine the impact of convective inputs on ozone chemistry, and to evaluate inorganic halogenated products produced from organic precursors. These measurements included actinic flux, ozone, formaldehyde, nitric oxide and nitrogen dioxide (or NO<sub>x</sub>, the sum of the pair), and halogen radical species. The NO<sub>x</sub> measurements also provided information on the



**FIG. 3.** Some key UTLS dynamical elements of the TWP during the CONTRAST campaign period. The figure shows the campaign domain, centered at Guam. The magenta and cyan circles indicate the nominal GV flight range (1,000 and 1,500 n mi radius, respectively). The gray–blue–yellow–orange shading shows the 12–14-km cloud fractions calculated using the *CloudSat* data, indicating the region dominated by frequent deep convection. The pink stipple shading shows the frequency of 200-hPa potential vorticity (PV) > 1 PVU (1 PVU =  $10^{-6} \text{ m}^2 \text{ s}^{-1} \text{ K kg}^{-1}$ ) during the campaign period, based on the GFS analyses. The increase in this frequency of PV > 1 PVU indicates the transition from the tropics to extratropics. The red contours are selected 200-hPa geopotential height (GPH, in km) levels indicating the region under the influence of mean anticyclonic circulation (Dima and Wallace 2007). Blue arrows show the 200-hPa wind field; the 50 and 60  $\text{m s}^{-1}$  wind speed contours are used to indicate the seasonal mean location of the Northern Hemisphere subtropical jet.



TABLE 1. Payload summary.		
Instruments	Measurements	Investigator
Chemistry		
NO <sub>x</sub>	NO, NO <sub>2</sub>	Weinheimer/NCAR Atmospheric Chemistry Observations and Modeling (ACOM) Laboratory
Fast ozone	O <sub>3</sub>	Weinheimer/NCAR ACOM
Vacuum UV (VUV) carbon monoxide	CO	Campos/NCAR ACOM
Picarro	CO <sub>2</sub> , CH <sub>4</sub>	Flocke/NCAR ACOM
TOGA	NMHCs, OVOCs	Apel/NCAR ACOM and Riemer/University of Miami
GT-CIMS	BrO, BrCl, HOBr, ClO	Huey/Georgia Institute of Technology
AMAX	BrO, IO, H <sub>2</sub> CO (remote)	Volkamer/University of Colorado
HAIS Advanced Whole Air Sampler (AWAS)	Trace gases	Atlas/University of Miami
In Situ Airborne Formaldehyde (ISAF)	H <sub>2</sub> CO	Hanisco/NASA Goddard Space Flight Center (GSFC)
Inorganic Br	Br* (Σ BrO + Br)	Atlas/University of Miami and Flocke/NCAR ACOM
Radiation		
HARP	Spectral actinic flux	Hall/NCAR ACOM
State parameters		
State parameters	Lat/lon, P, T, 3D wind	Jensen/NCAR Research Aviation Facility (RAF)
RAF digital video	Forward view	Jensen/NCAR RAF
Microphysics		
Cloud Droplet Probe (CDP) cloud probe	2–50 μm, water droplets, ice crystals	Jensen/NCAR RAF
2D-C precipitation probe	25–1,600 μm, ice, water	Jensen/NCAR RAF
Ultra-High Sensitivity Aerosol Spectrometer (UHSAS) aerosol probe	0.075–1 μm, aerosols	Jensen/NCAR RAF
Water-Based Condensation Nucleus Counter (WCN)	0.01–3 μm, aerosols	Jensen/NCAR RAF
Vertical Cavity Surface-Emitting Laser (VCSEL) hygrometer	Water vapor	Jensen/NCAR RAF

input of lightning-produced nitric oxide in this area of deep convection.

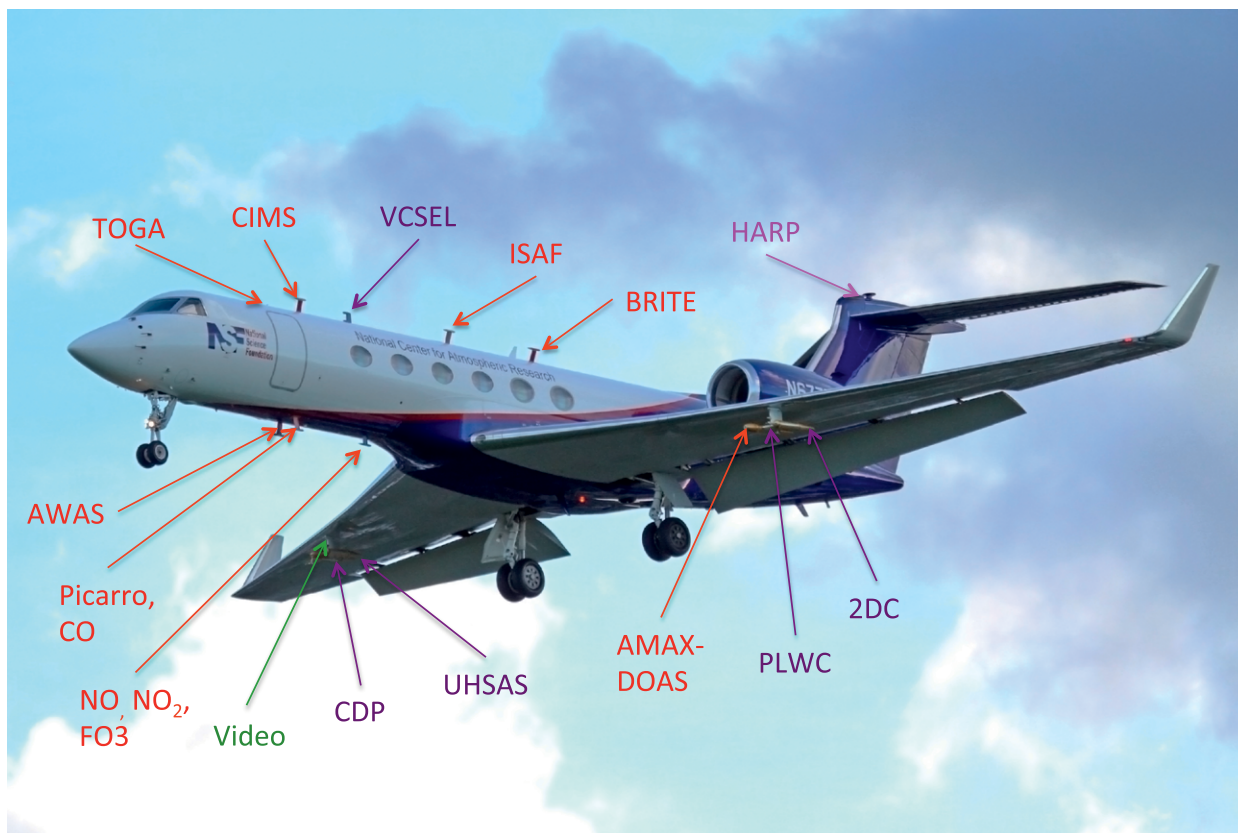
To evaluate the budget and partitioning of bromine in the TTL, organic bromine precursors were measured by AWAS and TOGA. Selected inorganic bromine species were measured in situ by chemical ionization mass spectrometry (CIMS) (Huey 2007), and remote sensing Airborne Multi Axis Differential Optical Absorption Spectroscopy (AMAX-DOAS) (Baidar et al. 2013; Volkamer et al. 2015). In addition, iodine oxide (IO) was measured by AMAX-DOAS (Dix et al. 2013; Volkamer et al. 2015). The AMAX-DOAS and CIMS data also provide a link to comparable measurements on the GH and BAe-146 aircraft. Finally, the payload also included cloud and aerosol measurements for determining aerosol size distributions and for evaluating heterogeneous chemical processing.

**Research flights.** A total of 16 GV research flights were made during the CONTRAST campaign including 3 transit flights and 13 local flights conducted in Guam. The ground tracks of these flights are shown in Fig. 5. The Guam-based flights sampled latitudes between

20°S and 40°N and altitudes between ~100 m and ~15 km above sea level (see flight tracks in altitude–latitude space in Fig. 11). Extensive vertical sampling of the atmosphere in the study region resulted in more than 100 complete profiles during the campaign. These profiles include the eastern and the central Pacific, but most were conducted in the TWP.

The research flights were planned according to seven scenarios, each designed to meet a set of observational objectives:

- 1) Domain survey (RF01–04, RF07, RF14): this type of flight was designed to map out the background distributions and gradients of compounds of different lifetimes and source/sink distribution. Gradients include those from the eastern to western Pacific, as well as those within the Guam domain. This type of flight typically covered the largest sampling range, including excursions out of the nominal domain.
- 2) Fresh convective outflow (RF05, RF09–12, RF14): this type of flight was designed to sample the outflow of ongoing convection and to contrast



**FIG. 4. CONTRAST GV payload configuration.** See Table I for instrument details. The cabin instruments are shown by their inlet locations (with exception of TOGA, its inlet was behind the fuselage in this takeoff photo). The colors are used to indicate the instrument type: chemistry (red), microphysics (purple), radiation (magenta), and the digital camera (green).

the enhancement of short-lived species within outflow to the upper-tropospheric background, which is influenced by more aged convection as well as long-range transport.

- 3) Dawn/dusk (RF08, RF13): this type of flight was designed to investigate photochemical evolution and halogen partitioning that occurs during the transition from daylight to darkness and vice versa. The strategy was to sample the same air mass through the period of solar zenith angle change. To accomplish this, the center of anticyclonic flow was targeted as a relatively stagnant air mass for chemical characterization and photochemical evolution.
- 4) Stratospheric survey (RF06, RF15): this type of flight was designed to sample the northern part of the domain to contrast the chemical composition of the tropical UT with that of the extratropical LS.
- 5) Dynamical boundaries and structures (RF06, RF10, RF14): these flights were special surveys conducted in response to the presence of dynamical boundaries in the flight domain, such as a shear line and a crossing of the ITCZ. The goal was to characterize the role of these dynamical

structures as physical boundaries to chemical composition.

- 6) Ozonesonde collocation (RF09, RF11, RF12, RF14): these flights involved a segment of sampling over Guam or Manus Island (2°S) when there was a coincident ozonesonde launch from the ATTREX or CAST team. The comparison over Manus was particularly important, since ozonesonde calibration issues are challenging in the convectively influenced TTL (Newton et al. 2016).
- 7) BAe-146 or Global Hawk coordination (RF08, RF11, RF12): this type of flight involved two or three aircraft flying on the same day, sampling a geographic region close to the same time. The coordinated flights between the GV and the BAe146 were typically designed to have a segment of “repeated track,” in which the same region and altitude was sampled by the two aircraft sequentially with a short time separation to allow for instrument comparisons.

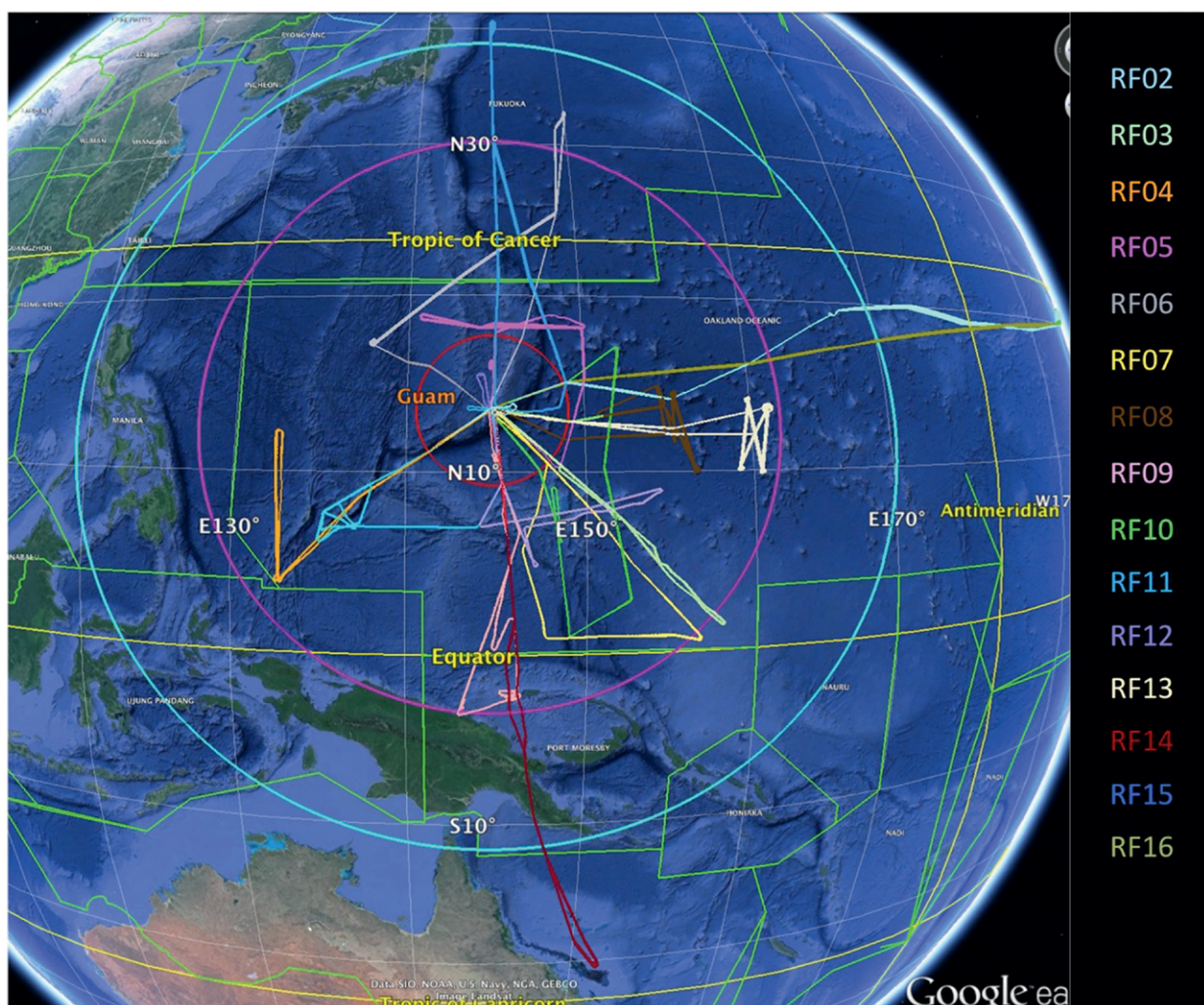
Table 2 provides a brief summary of the research flights with relevant flight scenarios noted by



numbers. As shown in the table, often more than one scenario was involved for a given flight. The table also shows the flight information regions (FIRs) within which each research flight was conducted. The GV operated in the Oakland Oceanic FIR (United States) for the majority of the flight hours. Flight operations in other FIRs [Fukuoka FIR (Japan) and Port Moresby FIR (Papua New Guinea)] were more difficult owing to numerous issues. An example was discussed in the section on “GV–ozonesonde coordinated flights.”

**Forecasting tools and platform.** CONTRAST flight planning and forecasting employed a suite of models that covered the range of dynamical and chemical processes relevant to the science goals. National

Centers for Environmental Prediction (NCEP)’s Global Forecast System (GFS) and NCAR’s Weather Research and Forecasting (WRF) Model aided the meteorological forecasts for CONTRAST. The 0.5° GFS pressure-level data files were used to produce 5-day forecast plots at 6-h increments and were made available in the project’s field catalog (described below). The Advanced Research version of WRF (ARW; Skamarock et al. 2008) was run in real time at the NCAR/Wyoming Supercomputing Center to provide a more detailed 72-h forecast of moist convection, clouds, and winds over the study region. The ARW was configured in a 15-km, 40-level grid covering much of the western Pacific with a 3-km nested grid.



**FIG. 5.** CONTRAST flight operation domain and the GV flight tracks (RF02–RF16). The operational domain is marked by the magenta and cyan circles, which indicate the nominal GV flight range with or without extensive profiling (1,000 and 1,500 n mi radius, respectively). The red ring close to Guam indicates the region covered by the Guam ATC radar. The green lines mark the boundaries of FIRs. As shown, most of the GV flights were operated within the Oakland Oceanic FIR (United States). Three additional FIRs flown were Fukuoka (Japan), Port Moresby (Papua New Guinea), and Brisbane (Australia).

Chemical forecasting was an important part of flight planning. CAM-chem, a global chemistry-climate model (Lamarque et al. 2012) run in the specified dynamics mode (i.e., nudged by observed winds, referred to herein as CAM-chem-SD), was run operationally during the CONTRAST field phase to provide 72-h chemical forecasts for mission planning. The model was configured in  $1^\circ \times 1^\circ$  horizontal resolution with 56 levels from the surface to 2 hPa and used in its specified-dynamics configuration, driven with National Aeronautics and Space Administration (NASA) GMAO/GEOS-5 meteorological fields. The model chemistry includes a detailed representation of tropospheric and stratospheric chemistry (~180 species; ~500 chemical reactions), including very-short-lived (VSL) halogens [details are found in Fernandez et al. (2014) and Saiz-Lopez et al. (2014)]. The model routinely forecast distributions of trace-

gas species including ozone, hydroxyl radical (OH), nitrogen oxides ( $\text{NO}_x = \text{NO} + \text{NO}_2$ ), carbon monoxide (CO), bromine oxide (BrO), iodine oxide (IO), and VSL organic halogens; for example, bromoform ( $\text{CHBr}_3$ ), dibromomethane ( $\text{CH}_2\text{Br}_2$ ), and methyl iodide ( $\text{CH}_3\text{I}$ ). This combination of species assisted the science team in flight planning that could, for example, delineate aged versus more recently polluted air masses, identify areas of stratospheric intrusions, or target air that had been recently lofted by convection.

In addition to CAM-chem-SD, the Real-Time Air Quality Modeling System (RAQMS) (Pierce et al. 2007) and the Monitoring Atmospheric Composition and Climate (MACC) (Flemming et al. 2009), both of which assimilate global satellite data, provided complementary chemical forecast information. In particular, RAQMS provided a number of additional

<b>Flight</b>	<b>Date</b>	<b>Flight type, scenario no.</b>	<b>Flight region</b>
RF01	11 Jan 2014	Research transit, 1	Colorado to Hawaii
RF02	13 Jan 2014	Research transit, 1	Hawaii to Guam
RF03	17 Jan 2014	Domain survey, 1	Guam to SE (Chuuk)
RF04	19 Jan 2014	Domain survey, 1	Guam to SW (Palau)
RF05	22 Jan 2014	Convective outflow, 2	North of Guam
RF06	24 Jan 2014	Jet crossing/pre–postfrontal contrast, 4, 5	North and northwest of Guam, entering Fukuoka FIR (JPN)
RF07	29 Jan 2014	SH convective outflow survey, 1, 4	Guam to equatorial region
RF08	1 Feb 2014	Photochemical evolution–sunset, 3, 7	Guam to east (the stagnation point in the anticyclone)
RF09	4 Feb 2014	Equatorial crossing and Manus ozonesonde, 2, 6	Guam to Port Moresby FIR (PNG)
RF10	8 Feb 2014	Subtropical jet pollution and ITCZ survey, 2, 5	Guam to near equator crossing ITCZ near $6^\circ\text{N}$
RF11	12 Feb 2014	Convective outflow and coordinate flight, 2, 6, 7	Guam to south and southwest
RF12	17 Feb 2014	Convective outflow and coordinated flight, 2, 6, 7	Guam to south and southeast
RF13	19 Feb 2014	Photochemical evolution–sunrise, 3	Guam to east (the stagnation point of the anticyclone)
RF14	22 Feb 2014	Equator crossing, SH TTL survey, and Manus ozonesonde, 1, 2, 6	Guam to Port Moresby FIR (PNG) and Brisbane FIR (AUS)
RF15	24 Feb 2014	Jet crossing and lower-stratosphere survey, 4	Guam to $40^\circ\text{N}$ Fukuoka FIR (JPN)
RF16	28 Feb 2014	Transit flight with limited research, 1	Guam to Honolulu

short-lived MBL tracers, such as dimethyl sulfide (DMS) and methyl ethyl ketone (MEK) during the campaign. The MACC forecast assimilated comprehensive global observations of chemical composition combined with the ECMWF meteorological forecasting system. MACC provided plots for the operation domain of  $\text{O}_3$ , CO,  $\text{CH}_4$ ,  $\text{NO}_x$ , black carbon, HCHO, OH, NO,  $\text{NO}_2$ , OH, and  $\text{HO}_2$  and provided comparisons to the CAM-chem-SD forecasts of chemical fields.

Figure 6 shows forecast plots for RF10 as an example, where the CAM-chem-SD forecasted a “CO river” that was primarily a biomass burning plume from Southeast Asia transported by the jet stream to the Guam region, and the WRF model successfully forecast the presence of the ITCZ in the domain. Both the CO river and the ITCZ were successfully

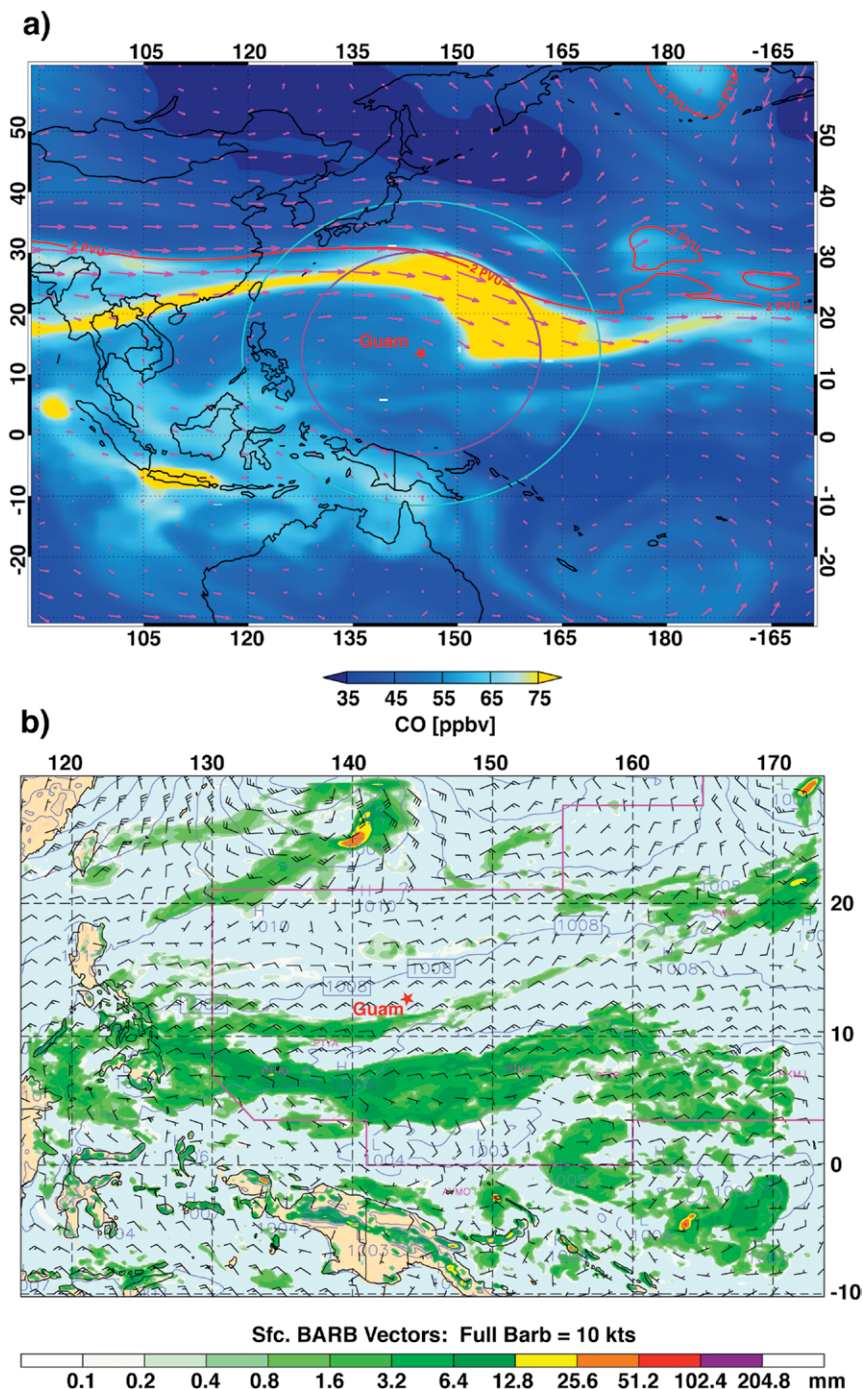


sampled in RF10. The former is further discussed in the section on “Pollution in the tropics and at the TTL level.”

The large suite of forecast materials were integrated and documented on the NCAR EOL field catalog for the CONTRAST project (<http://catalog.eol.ucar.edu/contrast>).

The field catalog provided a virtual operations center that allowed team members to participate in the daily briefings, either in the field or from their home institutions. The catalog also served to document the information flow during the experiment for future reference. In general, the forecast models provided good guidance for flight planning that allowed successful sampling of targeted meteorological and chemical features of interest. The main limitation of the forecast models (including the 3-km ARW simulations) was their difficulty in representing the observed evolution of convective systems over the low-latitude ocean regions. The experience and additional analyses by the forecasters were important in these cases.

**Flight planning and operation.** A daily operations meeting was held in the CONTRAST operations center in Guam on most aircraft maintenance days. The daily meeting was a key step in gathering the science team’s input for flight planning. Typically, different flight scenario options were discussed after the weather briefing and chemical forecast. The priority of different flight options was defined by the campaign science objectives, while meteorological conditions and chemical forecast dictated the practical possibilities of near term flight operations. Planning of the RF10 mission serves



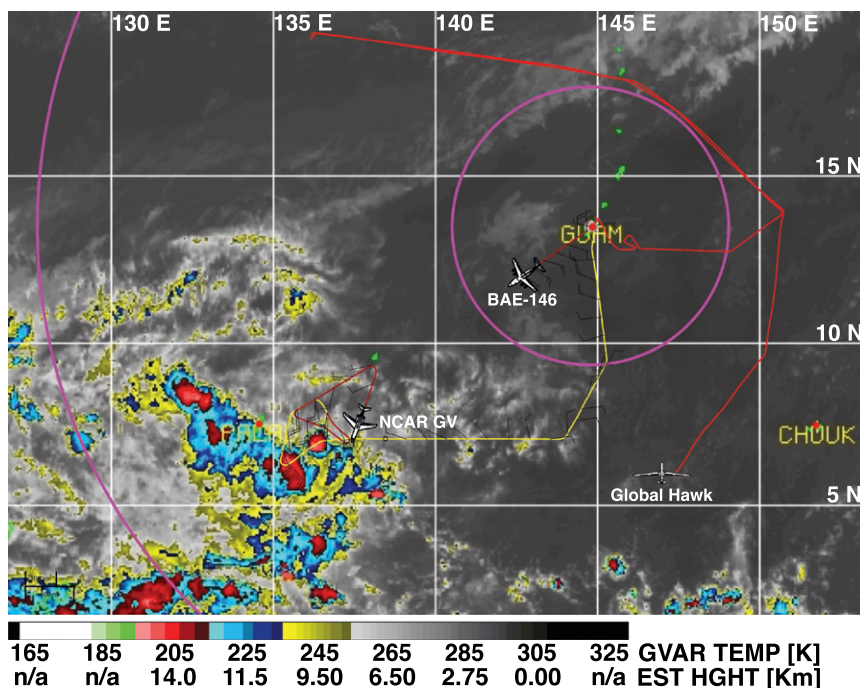
**FIG. 6.** Examples of operational weather and chemical forecast plots for RF10 (flight day 8 Feb 2014). (a) CAM-chem-SD 200-hPa CO (color shade) and wind field (pink arrows) forecast that shows an intense CO plume moving into the Guam domain following the jet stream. Also shown is the 2-PVU contour at 200 hPa, which marks the division of tropical UT and extratropical LS. (b) The WRF forecast of surface wind (wind barb vectors) and precipitating convection (shaded), which predicts an ITCZ near 7°N indicated by the surface wind convergence and the line of convection.

as a good example. After extensive team discussion, both pollution transport (CO river; Fig. 6a) and the impact of dynamical structures on chemical composition (ITCZ; Fig. 6b) were included in the flight plan. Typically, a flight or no-flight decision for the next day was made by the team by the end of the meeting.

Following the daily operations meeting in the morning, the principal investigators (PIs) and the operation center team would proceed with making an initial flight plan that was reviewed by the pilots, who filed the adjusted flight plan with the relevant air traffic control centers (ATCs). The plan was adjusted several hours before flight, if necessary, based on updated meteorological information from models and satellite imagery. The final flight plan was filed two hours before takeoff.

**Nowcasting and real-time decision-making.** Because of rapid development and uncertain movement of convective systems, nowcasting and real-time decision-making were an important part of flight operations. The near-real-time satellite images from MTSAT-2 were essential for providing information on convection. The NCAR EOL field catalog map and mission coordinator map were visible to the flight scientist on board the GV and also to mission support at the operations center. These maps were essential tools for real-time flight monitoring and decision-making. Figure 7 gives an example of the real-time display from the field catalog map, which integrates the real-time aircraft position with the near-real-time operational products. The display shows the real-time position of all three aircraft on 13 February 2014, near 0400 UTC, and the location of the convective systems targeted by the GV. As indicated by the figure, this flight successfully sampled convective outflow from widespread active convection southwest of Guam. An example of trace-gas measurements from this flight will be shown in next section.

The flight on 13 February (RF11) also provided a good example of real-time flight plan adjustment.



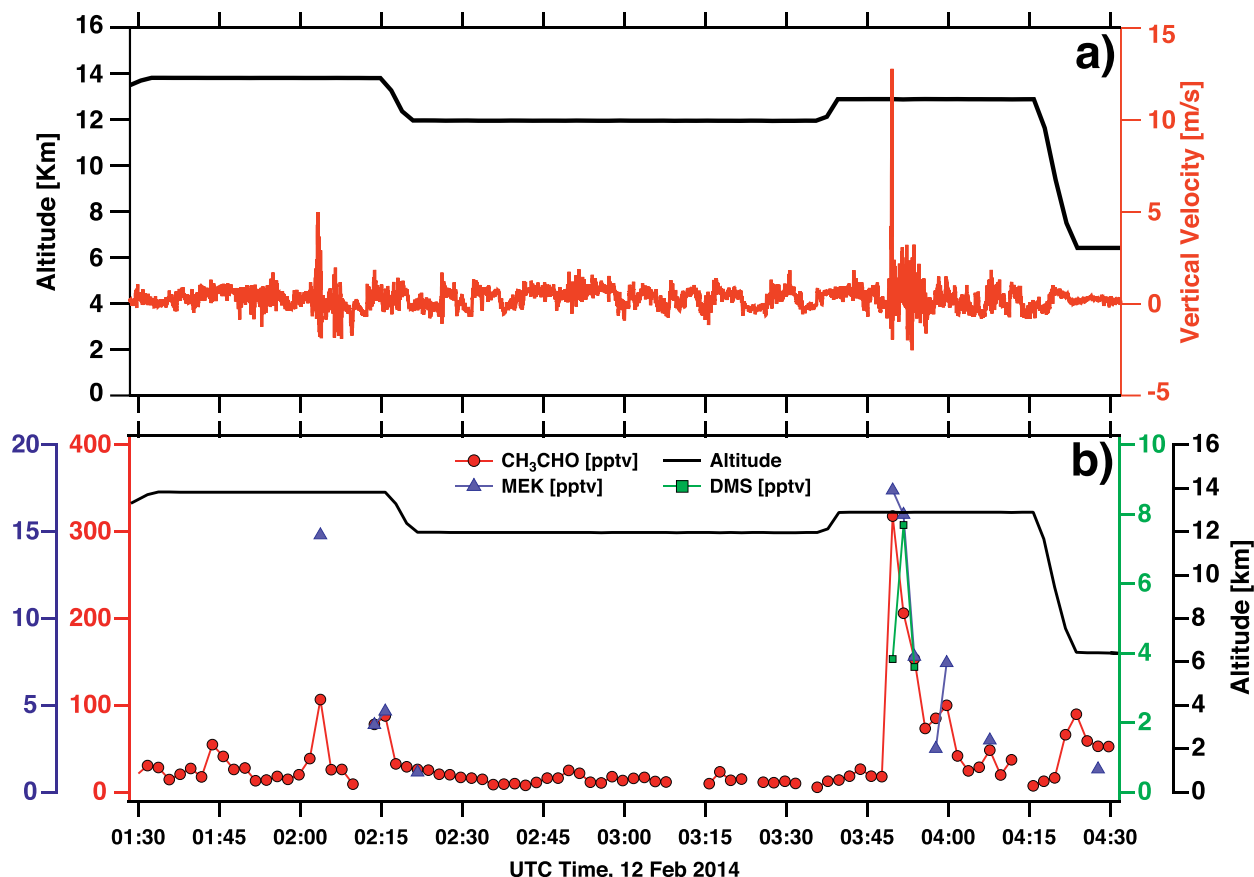
**FIG. 7.** An example of a coordinated flight and the GV targeting active convection. This figure shows the real-time display from the Catalog Map on 13 Feb 2014 near 0400 UTC, around the time the GV sampled the region of convective outflow (see Fig. 8). Active convection was widespread southwest of Guam, as revealed by the MTSAT IR (color shading) and visible (gray) channels. The yellow and red lines show flight tracks for each of the three aircraft (red indicates the track for the last hour).

Initially the flight was planned to pass the convective region at different altitudes to sample the air above and within the convective outflow. The mission scientist requested flight level 43,000 ft to be above the convection when the research aircraft entered the active region for the first pass. The altitude request was based on estimated altitude from the satellite IR image. During this segment, a  $5 \text{ m s}^{-1}$  updraft was sampled (see Fig. 8). The flight level for the second pass was requested at 37,000 ft, but it was found to be too low (visually below the outflow cloud layer). The decision was made to request a third pass at 40,000 ft, which resulted in the sampling of significant boundary layer tracer enhancement associated with a  $13 \text{ m s}^{-1}$  updraft (Fig. 8). The ATC response time over the oceanic region with no radar was at least 15–20 min, which was reflected by the large loop of the GV flight track out of the convective region (Fig. 7).

## SCIENTIFIC AND OBSERVATIONAL HIGHLIGHTS. Sampling convective outflow.

One of the key observational objectives was sampling air masses in regions of active convective outflow, as well as nearby air masses not influenced by recent convection, to quantify the influence of convection on the composition of



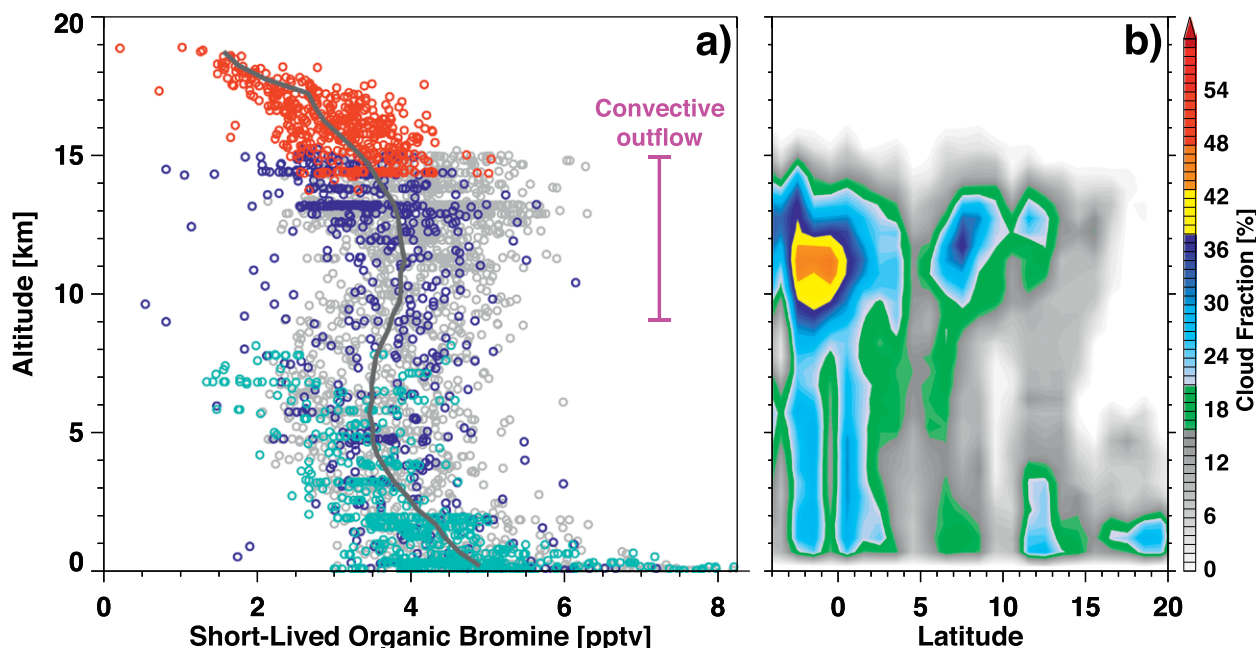


**FIG. 8.** Time series of flight altitudes, vertical velocity, and mixing ratios of selected short-lived compounds during a 3-h segment of RF11 (12 Feb 2014) when the GV was targeting the outflow from active convection (see GV location and the region of active convection during this time period in Fig. 7). The active convection regions are indicated by increases in updraft velocity ( $\sim 5$  and  $13 \text{ m s}^{-1}$ , respectively). For the aircraft location and the convection, see Fig. 7, which shows satellite IR channel near 0400 UTC. As indicated by the track in Fig. 7, the GV repeatedly targeted the convective region at different altitudes. The two time periods of successful sampling of fresh outflow (associated with the updraft velocity  $5$  and  $13 \text{ m s}^{-1}$ ) the GV was at  $43,000 \text{ ft}$  ( $\sim 14 \text{ km}$ ) and  $40,000 \text{ ft}$  ( $\sim 11 \text{ km}$ ), respectively. The location of the updraft was in the vicinity of the aircraft position shown in Fig. 7. The tropopause was at  $\sim 17 \text{ km}$ . The corresponding enhancements of short-lived species in the UT are shown using TOGA observations of acetaldehyde ( $\text{CH}_3\text{CHO}$ ), methyl ethyl ketone [MEK;  $\text{CH}_3\text{C}(\text{O})\text{CH}_2\text{CH}_3$ ], and dimethyl sulfide (DMS;  $\text{CH}_3\text{SCH}_3$ ). DMS and MEK are below the detection limit ( $1 \text{ pptv}$ ) for the majority of the UT time segment with the exception of the updraft region, especially during 0349–0354 UTC.

the TWP troposphere. Targeting active convection is a significant challenge for flight operations in general, but especially over oceanic domains such as the TWP because of the absence of ground-based weather radar coverage. Aided by the real-time information described in section 3, the GV succeeded in targeting active convection during multiple flights in CONTRAST.

Figure 8 shows an example of the impact of active convection on the chemical composition of the UT using data acquired during RF11 conducted on 13 February 2014. The figure shows a 3-h segment of the flight, during which the GV sampled the same convective system (as shown in satellite IR map in Fig. 7) three times at different altitudes. Convectively generated turbulence and the region of strong upward motion

are indicated by the vertical wind velocity. Two time periods of active convection sampled are marked by the  $\sim 5 \text{ m s}^{-1}$  (near 0200 UTC) and the  $\sim 13 \text{ m s}^{-1}$  (between 0349 and 0354 UTC) updraft events. Selected chemical species measured by the TOGA instrument demonstrate the strong impact of convection on the composition of UT air masses, indicated by the contrast of the air masses from inside to outside of convective outflow. The three selected species are all relatively short lived with estimated lifetimes ranging from less than a day for acetaldehyde (Millet et al. 2010) and DMS (Langner and Rodhe 1991) to days for MEK (e.g., Calvert et al. 2011). These species can all be produced by marine biological processes (Carpenter et al. 2010). They were all observed at elevated mixing ratios in the marine



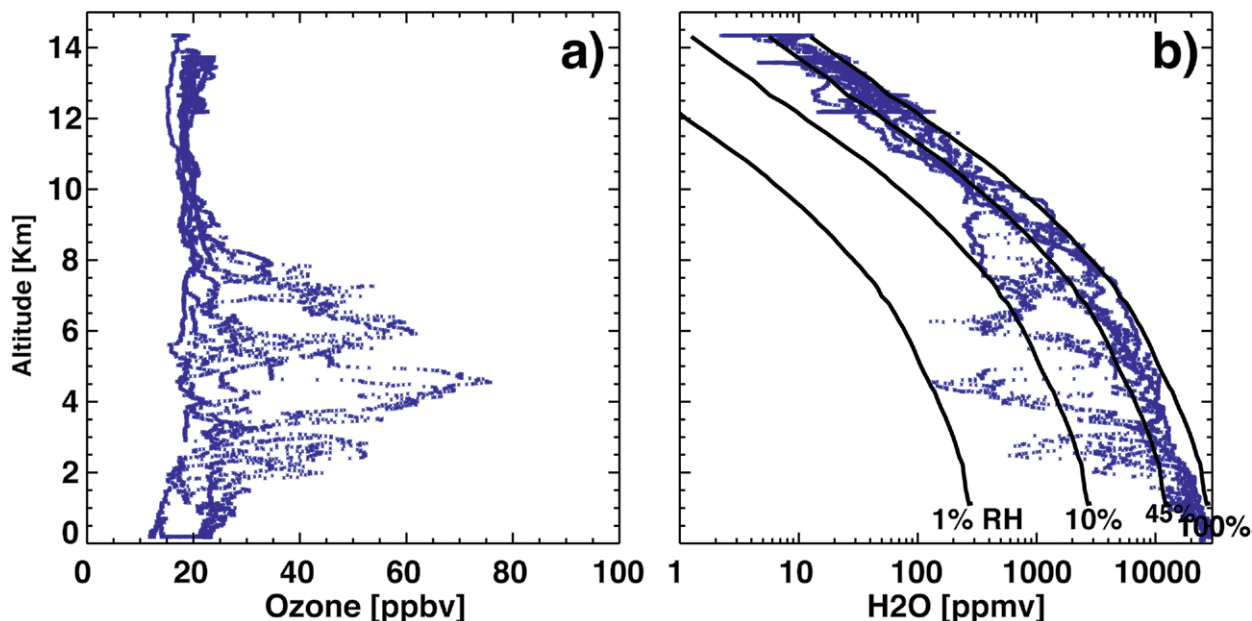
**FIG. 9.** (a) Vertical distribution of the total bromine content from two short-lived organic bromocarbons ( $2 \times \text{CH}_2\text{Br}_2 + 3 \times \text{CHBr}_3$ ) from GV TOGA (gray) and whole air samplers on all three aircraft (cyan: BAe146, blue: GV, red: Global Hawk). The gray line represents the median profile derived using all four datasets. (b) Latitude–height cross section of the mean cloud fraction of the region ( $140^\circ$  and  $150^\circ\text{E}$ ) during the period of the CONTRAST campaign, calculated using CloudSat satellite data.

boundary layer during this flight. At the flight level of  $\sim 12$  km, these species exhibit low mixing ratios characteristic of the background UT outside of the active convection region. The low background levels reflect the rapid decrease of their abundance with altitude. Significant enhancement was observed during both periods of sampling near a convective updraft. In the vicinity of the  $13 \text{ m s}^{-1}$  updraft, the mixing ratios show values only slightly less than in the boundary layer, indicating very little processing or mixing took place during transport of the air masses between the surface and the 12-km level.

**Impact of convection on vertical distribution of VSL halocarbons.** An important objective of CONTRAST was to quantify the abundance of the halogenated VSL compounds throughout tropospheric column of the TWP. Bromine chemistry is important for the photochemistry of  $\text{O}_3$  in both the stratosphere and the tropical troposphere (e.g., Ko et al. 1997; Saiz-Lopez and von Glasow 2012; Salawitch et al. 2005; Frieler et al. 2006; Carpenter and Reimann 2014). Additionally, measurements and modeling suggest that halogen cycles involving both bromine and iodine could contribute to chemical loss of  $\text{O}_3$  in the tropical atmosphere (e.g., Saiz-Lopez et al. 2012, 2015; Dix et al. 2013; Carpenter et al. 2013; Volkamer et al. 2015; Wang et al. 2015), though we focus just on bromine here.

Figure 9a shows the profile in the TWP of the total bromine content of the two most important bromine-bearing VSL species,  $\text{CH}_2\text{Br}_2$  and  $\text{CHBr}_3$ , using data from all three aircraft. Detailed descriptions of the data and the result of intercomparisons are given by Andrews et al. (2016). These bromocarbons are produced by biological processes in the tropical ocean and have lifetimes in the upper troposphere of  $\sim 17$  days ( $\text{CHBr}_3$ ) and  $\sim 150$  days ( $\text{CH}_2\text{Br}_2$ ) (Carpenter and Reimann 2014). The measurements shown in Fig. 9a demonstrate the impact of convection on VSL bromocarbons and highlight the vertical structure of total bromine that can be obtained by synergistic sampling conducted by three aircraft. The profile displays a pronounced reverse “S” shape, with strong decrease above the MBL and enhancement between 10- and 15-km altitude, which is a clear signature of convective uplifting of MBL air. The layer of enhancement is largely consistent with the peak in cloudiness as derived from the CloudSat cloud profiling radar (Fig. 9b) (Stephens et al. 2008). The vertical distribution of the cloud fraction serves as an illustration of the convective outflow layer. Numerous details concerning the outflow (e.g., the level where maximum mass is detrained) can be obtained from analysis of the radar profiles of the anvils, as discussed in Takahashi and Luo (2012). Hence, cloud radar and VSL chemical compound measurements





**FIG. 10. (a) Ozone and (b) water vapor data from RF04, as an example of the ozone profile structure, with layers of ozone enhancement that are coincident with layers of water vapor reduction. Relative humidity (RH) levels corresponding to the measured water vapor and temperature are also shown (black curves).**

offer consistent and complementary views of the deep convective outflow and transport in the TWP.

The amount of total VSL bromine at the tropopause (about 17 km) defines one pathway for transport of bromine from VSL compounds to the stratosphere, termed source gas injection (SGI). The result in Fig. 9a, when supplemented by the bromine content of other VSL species such as  $\text{CHBr}_2\text{Cl}$ ,  $\text{CH}_2\text{BrCl}$ , and  $\text{CHBrCl}_2$  that will soon be available, will represent the first experimental determination of SGI of bromine in the TWP. The other route for stratospheric supply of bromine from VSL compounds, product gas injection (PGI), represents cross-tropopause transport of inorganic species. The value of PGI will be quantified by ATTREX measurements of BrO in the tropical lowermost stratosphere as well as CONTRAST measurements of BrO in the extratropical lowermost stratosphere, obtained on RF15 by the CIMS and DOAS instruments.

**Ozone over the TWP warm pool.** The CONTRAST research flights included significant profile measurements spanning from near the surface to above 14 km. Together with takeoffs and landings there were nearly 100 vertical profiles over the domain. CONTRAST has provided the first extensive in situ measurements of ozone within the UT over the warm pool of the TWP, where routine sampling does not occur and very few prior observations [e.g., Rex et al. (2014) and references therein] are available. Ozone profiles typical of the CONTRAST measurements,

from RF04, are shown in Fig. 10a. Ozone in the UT ( $>9$  km) was near 20 ppbv throughout the experiment, with little variability either in space or time (Fig. 11). Ozone mixing ratios near the surface and in the MBL were typically between 10 and 20 ppbv, as shown in Fig. 10a for RF04, as well as throughout other CONTRAST flights (Fig. 11). Consequently, the persistent upper-tropospheric ozone values near 20 ppbv are consistent with the strong influence of convective outflow of low-level air (Pan et al. 2015).

The airborne measurements during CONTRAST often showed enhanced ozone over the lower-middle troposphere ( $\sim 3$ – $9$  km), with values typically 40–80 ppbv (Fig. 10a). The enhanced ozone occurred in persistent layers and filaments, which were furthermore characterized by extremely dry air (relative humidity  $< 45\%$ ) (Fig. 10b). The strong correlation of enhanced ozone with dry layers is similar to previous observations (e.g., Thouret et al. 2001; Hayashi et al. 2008), although the layers observed in CONTRAST were more persistent. These dry, enhanced ozone layers over  $\sim 3$ – $9$  km (320–340 K in potential temperature coordinates) were a ubiquitous feature observed during CONTRAST. The layers occurred so frequently that the statistical distribution of ozone over this altitude region exhibits a bimodal distribution, with a primary mode near 20 ppbv and an enhanced mode centered on ozone near 60 ppbv (Pan et al. 2015).

The mechanisms producing the structure shown in Fig. 10 are a topic of active research. Contributions

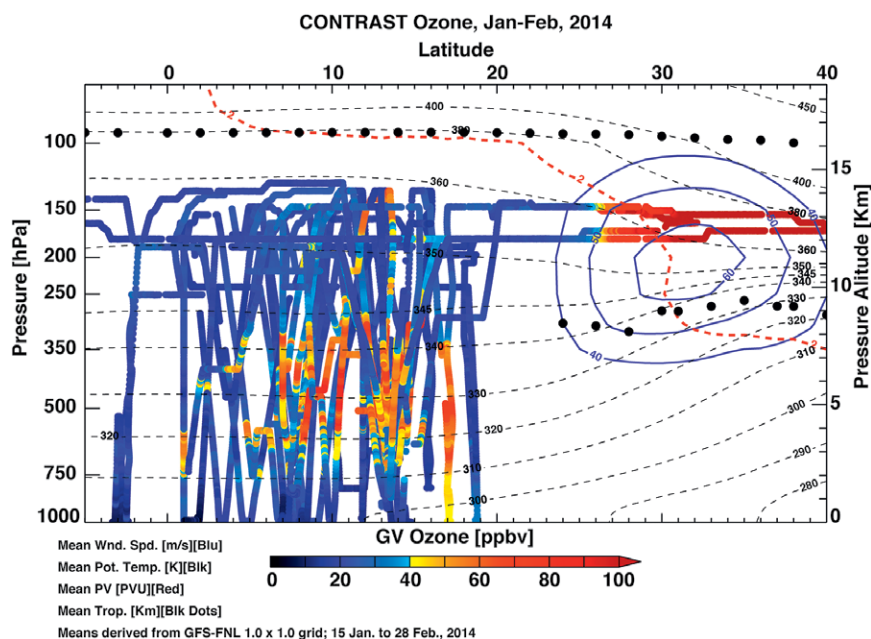
from large-scale dynamics (i.e., transport and mixing) as well as ozone production in plumes of biomass burning (Anderson et al. 2016) are leading candidates under investigation. Figure 11 shows that the occurrence of the ozone layers is mainly in the ~320–340-K isentropic levels. At these levels isentropic mixing could connect the air mass in the tropical low to midtroposphere to the UTLS region in midlatitudes near the subtropical jet. This structure suggests a large-scale dynamical influence, which is consistent with the interpretations of the dry layers by the TOGA COARE community (e.g., Yoneyama and Parsons 1999). On the other hand, tracer measurements show a significant signature of combustion of plant matter that provides compelling chemical evidence that biomass burning also contributes significantly to the enhanced O<sub>3</sub> layers (Anderson et al. 2016). Because the dataset includes a large suite of chemical species that can provide chemical signatures of transport and chemical processing, CONTRAST data can be used to better quantify the contributing mechanisms of these high ozone–low water structures.

**Pollution in the tropics and at the TTL level.** Anthropogenic impact on the remote TWP region was the main focus of several previous experiments (Gregory et al. 1999; Kondo et al. 2002; Jacob et al. 2003). Although not explicitly included in the original objectives of the CONTRAST experiment, the influence of human activity on the chemical composition of the remote TWP warm pool region is directly relevant to ozone budget, oxidizing capacity, and impact of convection for this region of the atmosphere. CONTRAST data have provided the first intensive chemical sampling of the troposphere overlying the TWP warm pool during NH winter. During the experiment, GV flights targeted a number of cases where a significant difference of polluted versus clean air masses was observed. These cases include 1) measurements both north and south of a well-defined ITCZ (RF10); 2)

measurements in front of and behind a shear line, which moved into tropics from the northeast Asian continent (RF06); 3) sampling of a pollution plume brought into the TWP domain by the jet stream and the region's anticyclonic flow (RF10, the CO river); and 4) observations of the polluted TTL over Papua New Guinea (PNG) versus the more pristine TTL over the Coral Sea (RF14).

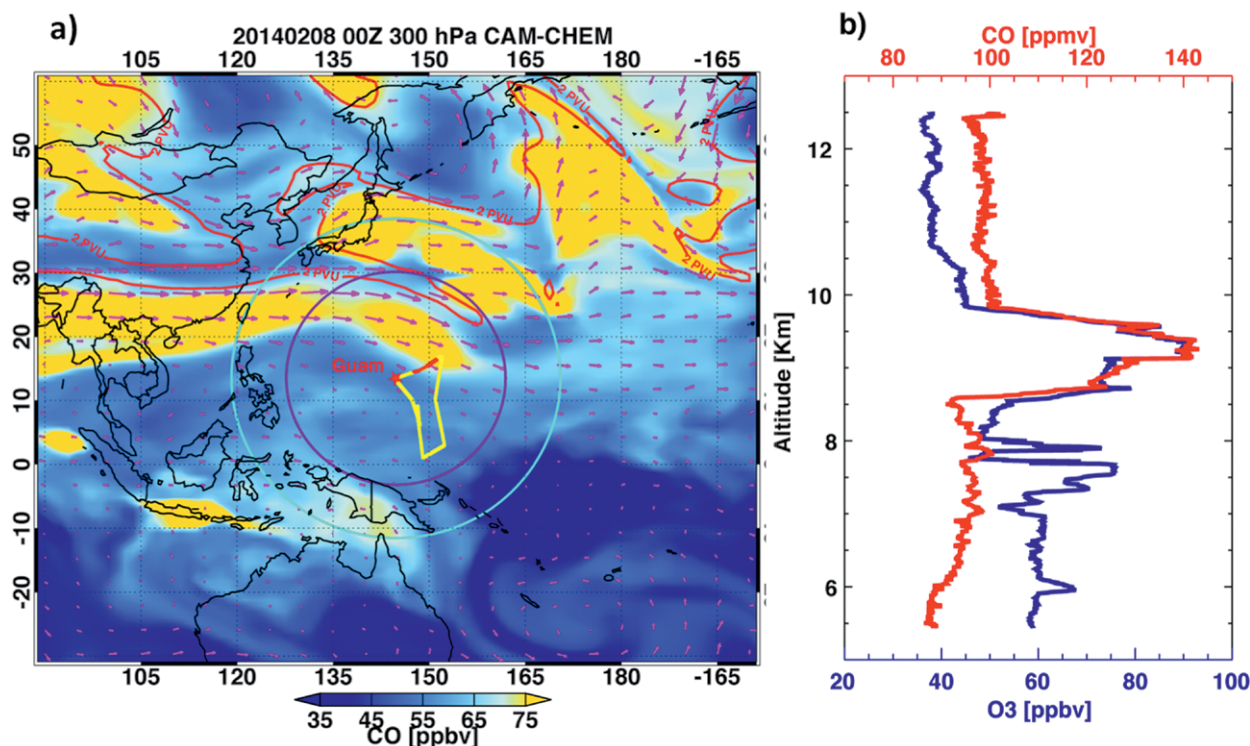
Figure 12 shows the successful sampling of the long-range transport of air pollution on 10 February 2014 (RF10). Based on the chemical forecast by NCAR CAM-chem-SD (see Fig. 6) and MACC CO field (not shown), the flight was designed to target a pollution plume that was flowing into the Guam domain from Southeast Asia and along the equator flank of the jet stream. The flight succeeded in documenting the highest mixing ratio of CO measured over the TWP in the entire campaign (~130 ppbv), providing an interesting case characterized as the CO river. The figure also shows a layer of high ozone well correlated with elevated CO, indicating the associated photochemical production of O<sub>3</sub> from pollutants emitted over the continent. Additional hydrocarbon measurements by TOGA suggest biomass burning may have played a role in the origin of this plume.

The TTL survey flight on 22 February 2014 (RF14) observed stark differences between strongly polluted



**FIG. 11.** The GV flight tracks for all flights colored by the O<sub>3</sub> mixing ratio. Also included are mean large-scale dynamical structures of the campaign period and domain, represented by the tropopause height (black dots), 2-PVU contour (red dash), subtropical jet location (40, 50, 60 m s<sup>-1</sup> horizontal wind, blue contour), and mean isentropes, all given in sectional zonal mean for 130°–160°E longitude range.





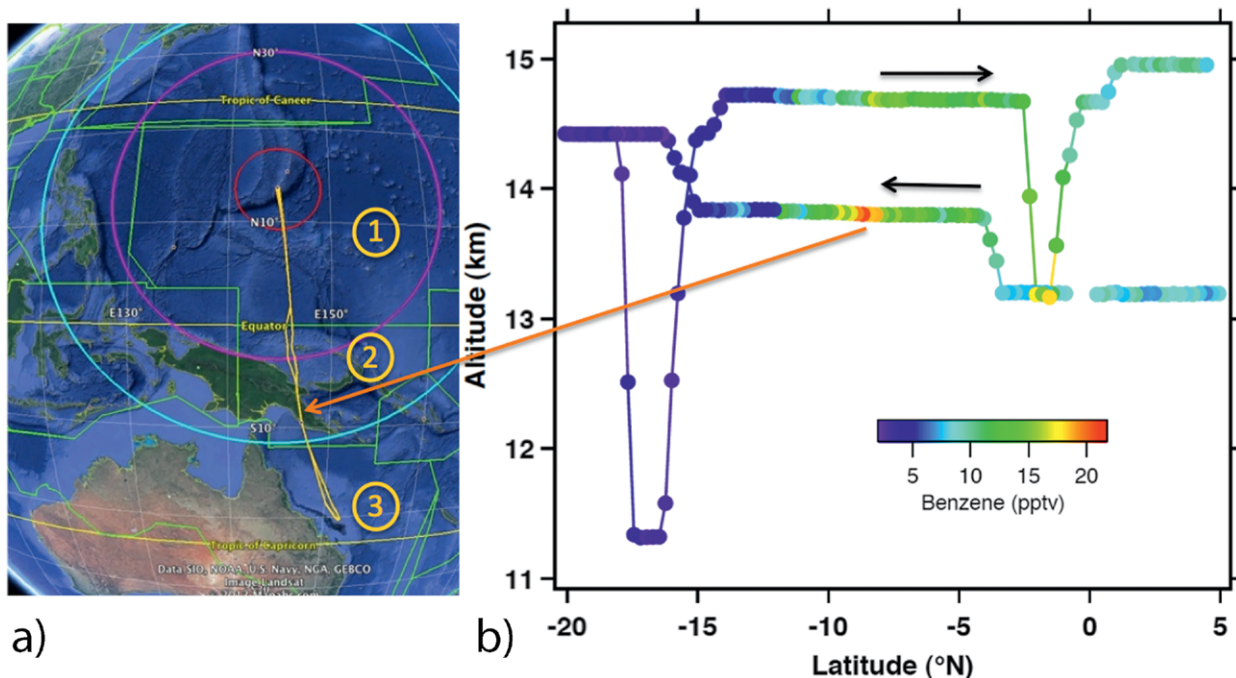
**FIG. 12.** (a) CAM-chem-SD model simulation of CO at the 300-hPa level as well as the aircraft flight track (yellow) and (b) the GV measurements of CO and O<sub>3</sub> in the CO river during RF10 on 8 Feb 2014. The portion of the flight track corresponding to the in situ profile shown in (b) is marked in red on the map in (a). The maximum CO enhancement is shown to be ~8.5-km altitude (~300-hPa pressure level).

and clean regions of the TTL. This 9-h and 30-min flight covered a large-latitude range, from Guam to the Coral Sea (13.5°N–20°S) (Fig. 13a). The flight provided the southernmost set of observations for the entire campaign. Most of the flight was downwind from active convection. While under convective influence, the TTL was moderately polluted for most of the flight, as shown by elevated levels of benzene (Fig. 13b). The highest level of pollution was observed over the island of Papua New Guinea (~8°S), where the GV flew through moderate deep convection and sampled the influence of direct convective pumping of biomass burning on the island. A region of extremely clean air was sampled over the Coral Sea (~18°S), indicated by the mixing ratio of benzene being near the detection limit of the TOGA instrument. This clean air region provided unique measurements to chemically characterize the pristine tropical UT conditions with no immediate pollution source and under calm weather. The contrast of clean versus polluted air in the TTL is important for characterizing the impact of anthropogenic activity, coupled with deep convection around the maritime continents, on the composition of the UT.

*GV–ozonesonde coordinated flights.* A significant fraction of prior ozone profile measurements in the

tropical troposphere have been obtained by ozonesondes. These measurements include data from the Southern Hemisphere Additional Ozonesondes (SHADOZ) network (Thompson et al. 2011), which does not routinely sample the warm pool atmosphere over the TWP (Fig. 1 shows ozonesonde launch locations), as well as ozonesondes launched from several ship-based field experiments in the TWP [Rex et al. (2014) and references therein]. Extremely low-ozone values at the level of convective outflow in UT (i.e., at the base of the TTL) have been reported from these ozonesonde measurements in the TWP (Kley et al. 1996; Rex et al. 2014). The low-ozone environment has been hypothesized to suppress the presence of hydroxyl radicals (OH), which controls the lifetime of a large suite of chemical species (Rex et al. 2014). However, the physical interpretation of these low-ozone cases is challenged by the uncertainty in background current calibration of ozonesondes (Vömel and Diaz 2010).

Quantification of O<sub>3</sub> at the level of convective outflow in the UT over the TWP, and assessment of ozonesonde calibration, were motivating factors for the planning of a GV overflight of Manus Island (2°S) within a short window of an ozonesonde launch by the CAST team. Overflights of Manus by the GV



**FIG. 13. Chemical gradients at the TTL level shown by TOGA Benzene data from RF14 flight on 22 Feb 2014. (a) The flight track shows the large-latitude range ( $\sim 15^{\circ}\text{N}$ – $20^{\circ}\text{S}$ ) sampled. The flight was conducted in three FIRs: 1) Oakland Oceanic (United States), 2) Port Moresby (Papua New Guinea), and 3) Brisbane (Australia). (b) Benzene mixing ratio from TOGA highlights the strong contrast of air over PNG ( $\sim 8^{\circ}\text{S}$ ) and the Coral Sea (south of  $10^{\circ}\text{S}$ ). The black arrows above the tracks indicate the outbound/inbound flight directions.**

were carried out on 4 and 22 February 2014 (RF09 and RF14). Flight permission issues and air traffic control challenges posed significant limitations to the flights and prevented the GV from performing a complete profile near the site of the ozonesonde launch. Specifically, RF09 was the first flight operation crossing the equator into the Port Moresby FIR working with PNG ATC. The flight plan started with MBL sampling first once entered Port Moresby FIR, followed by ascending to the flight ceiling near the Manus Island. When GV arrived on station near the Manus Island for collocated profiling with the ozonesonde launch, the PNG ATC kept the plane at flight level 34,000 ft for a commercial flight to pass at 38,000 ft. This altitude hold used up the filed flight pattern, which was planned to loop up to the TTL level. The ATC did not give the team a chance to repeat the flight pattern at higher altitudes once the commercial airliner passed. The lessons learned from RF09 led to a different strategy in planning for RF14, resulting in a successful partial profile at the TTL level with the collocated ozonesonde from Manus Island in this second attempt. Despite the difficulties in operations, the two flights obtained collocated partial profiles of  $\text{O}_3$  that have been used to validate the ozonesonde calibration (Newton et al. 2016).

Figure 14 shows the flight track, ozonesonde profile, and the collocated GV partial  $\text{O}_3$  profile between 13 and 15 km during RF14. The GV data were obtained for the altitude range where the question of whether near-zero ozone in the balloon observations is actually real (Kley et al. 1996) or is due to varying background current (Vömel and Diaz 2010) had yet to be resolved. The GV in situ  $\text{O}_3$  instrument, based on chemiluminescence (Ridley et al. 1992), has a low detection limit of 0.1 ppbv and high vertical resolution of near 7 m during profiling. For the entire campaign, the lowest measured amount of ozone in the upper troposphere was 13 ppbv.

The intercomparison results show a consistent picture. By not exposing the ozonesonde to ozone during preparation, and by assuming a constant background current measured in the laboratory immediately before launch, ozonesonde profiles were measured and agreed with the GV to 3 ppbv between 150 and 200 hPa (Newton et al. 2016). This is within the expected error limits of the sondes. Further, during the entire CAST ozonesonde campaign the minimum repeatable ozone concentration in the TTL from the Manus ozonesondes was 12 ppbv, again very consistent with the GV.

**FORWARD FROM THE FIELD PHASE.** The CONTRAST experiment succeeded in obtaining a large suite of trace-gas measurements, together with



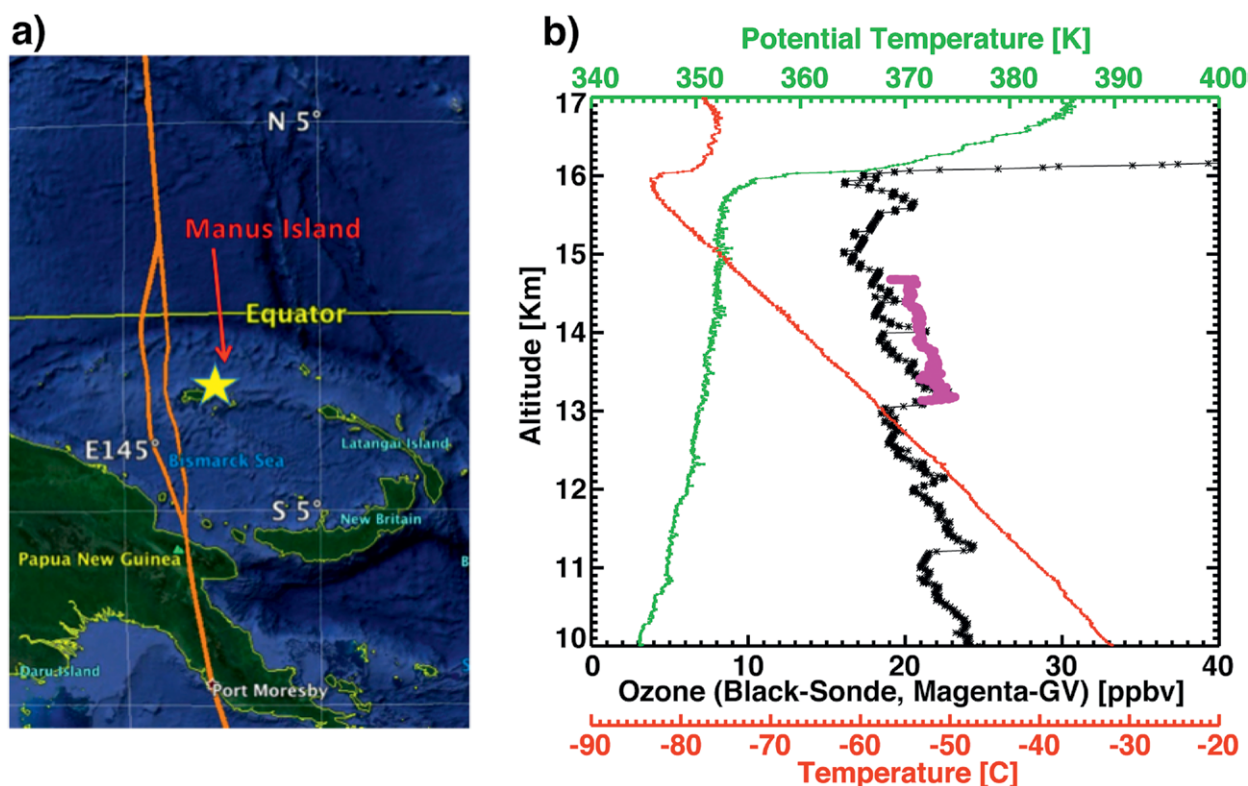
dynamical and microphysical variables, under a wide range of conditions over the TWP. The intensive sampling of actinic flux, ozone, water vapor, and other active species will allow constrained photochemical calculation of hydroxyl radical over the TWP, which will lead to a better understanding of the processes that control atmospheric oxidation capacity in the tropical western Pacific. Oxidation processes in the tropical atmosphere play a major role in the global budget of many trace gases, including greenhouse gases. The region has added significance because the major transport of tropospheric air into the lower stratosphere occurs over the TWP.

The flights successfully sampled air masses in convective updraft and outflow both near and downstream of active convection. The data will enable individual case studies and statistical quantification of the impact of convection and convective transport on the vertical distribution of a wide range of compounds with different photochemical lifetimes.

Within the context of the coordinated experiments, the CONTRAST, ATTREX, and CAST campaigns obtained the vertical distribution of a large number of reactive gases relevant to chemistry–climate

interactions, especially a unique dataset of halogenated VSL species from the oceanic surface to the LS over the TWP. These measurements, along with measurements of inorganic halogen product gases, will add new capability to evaluate the processes that control the reactive halogen chemistry, extending from the MBL, through the TTL, and into the LS. The ocean sources and sinks of these gases, as well as a range of oxygenated organic gases (such as formaldehyde) will be examined from the measurements that were collected during the campaigns. Accurate representation of the impact of deep convection on the chemical environment is a major challenge for CCMs. Observational information from CONTRAST will provide important constraints, especially for the remote TWP troposphere.

Finally, CONTRAST data provided the first intensive in situ observations of ozone in the TTL over the oceanic warm pool. The measurements filled an important data gap by coordinating in situ measurements with collocated ozonesonde profiles, which resulted in a new understanding of ozonesonde uncertainty (Newton et al. 2016) and quantified the low-ozone level over the TWP TTL. The repeated



**FIG. 14.** (a) GV–ozonesonde coordinated flight during RF14, 22 Feb 2014 with Manus Island ozonesonde launch. GV flight track is shown in orange on the map. (b) GV in situ ozone, sampled during a dip from west of Manus Island from 45,000 to 41,000 ft, is shown in magenta, while ozonesonde measurements of ozone (black), temperature (red), and potential temperature (green) are also shown.

vertical profiles of chemical distributions during CONTRAST also led to the discovery of a bimodal ozone distribution in the free troposphere between ~3 and 9 km over the TWP (Pan et al. 2015). The controlling mechanism(s) of the two modes, and how well the modes are represented in global CCMs, are topics of active research (Anderson et al. 2016 Randel et al. 2016). CONTRAST measurements also provided new constraints on the oxidation capacity over the TWP (Nicely et al. 2016) and on the amount and sources of inorganic bromine in the tropical upper troposphere (Chen et al. 2016).

CONTRAST data are publicly available for all researchers and can be obtained at [http://data.eol.ucar.edu/master\\_list/?project=CONTRAST](http://data.eol.ucar.edu/master_list/?project=CONTRAST).

**ACKNOWLEDGMENTS.** Funding for this work was provided by the National Science Foundation (NSF) via its sponsorship of the National Center for Atmospheric Research (NCAR). The CONTRAST experiment was sponsored by the (NSF). We acknowledge the excellent field project support provided by NCAR/EOL during flight operations. We give special thanks to the GV pilots and mission coordinators for their knowledge, skills, and dedication; these were essential to the success of this campaign. The MACC forecasts were funded by the European Union's Seventh Framework Programme (FP7) under Grant Agreement 283576. The views, opinions, and findings contained in this report are those of the author(s) and should not be construed as an official National Oceanic and Atmospheric Administration or U.S. government position, policy, or decision.

## APPENDIX. List of abbreviations.

ATTREX:	Airborne Tropical Tropopause Experiment
BIBLE:	Biomass Burning and Lightning Experiment
CAM-chem:	Community Atmosphere Model with Chemistry
CAST:	Coordinated Airborne Studies in the Tropics
CCM:	Chemistry–Climate Model
CONTRAST:	Convective Transport of Active Species in the Tropics
ECMWF:	European Centre for Medium-Range Weather Forecasts
ENSO:	El Niño–Southern Oscillation
EOL:	Earth Observing Laboratory
FAAM:	Facility for Airborne Atmospheric Measurements
FIR:	Flight information region
GEOS-5:	Goddard Earth Observing System Model, version 5
GFS:	Global Forecast System
GMAO:	Global Modeling and Assimilation Office
GTE:	Global Tropospheric Experiment
HIAPER:	High-Performance Instrumented Airborne Platform for Environmental Research
IGAC:	International Global Atmospheric Chemistry
ITCZ:	Intertropical convergence zone
MACC:	Monitoring Atmospheric Composition and Climate
MJO:	Madden–Julian oscillation
MTSAT:	Multifunctional Transport Satellite
RAQMS:	Real-time Air Quality Modeling System
SHADOZ:	Southern Hemisphere Additional Ozonesondes
SPARC:	Stratospheric Processes and their Role in Climate
TOGA COARE:	Tropical Ocean and Global Atmosphere Coupled Ocean–Atmosphere Response Experiment
TORERO:	Tropical Ocean Troposphere Exchange of Reactive Halogens and Oxygenated Hydrocarbons
WCRP:	World Climate Research Programme
WOUDC:	World Ozone and Ultraviolet Radiation Data Centre
WRF:	Weather Research and Forecasting Model

## REFERENCES

- Anderson, D. C., and Coauthors, 2016: A pervasive role for biomass burning in tropical high ozone/low water structures. *Nat. Commun.*, **7**, 10267, doi:10.1038/ncomms10267.
- Andrews, S. J., and Coauthors, 2016: A comparison of very short-lived halocarbon (VSLS) and DMS aircraft measurements in the Tropical West Pacific from CAST, ATTREX and CONTRAST. *Atmos. Meas. Tech.*, **9**, 5213–5225, doi:10.5194/amt-2016-94.



- Apel, E. C., and Coauthors, 2015: Upper tropospheric ozone production from lightning NO<sub>x</sub>-impacted convection: Smoke ingestion case study from the DC3 campaign. *J. Geophys. Res. Atmos.*, **120**, 2505–2523, doi:10.1002/2014JD022121.
- Baidar, S., H. Oetjen, S. Coburn, B. Dix, I. Ortega, R. Sinreich, and R. Volkamer, 2013: The CU Airborne MAX-DOAS instrument: Vertical profiling of aerosol extinction and trace gases. *Atmos. Meas. Tech.*, **6**, 719–739, doi:10.5194/amt-6-719-2013.
- Bergman, J. W., E. J. Jensen, L. Pfister, and Q. Yang, 2012: Seasonal differences of vertical-transport efficiency in the tropical tropopause layer: On the interplay between tropical deep convection, large-scale vertical ascent, and horizontal circulations. *J. Geophys. Res.*, **117**, D05302, doi:10.1029/2011JD016992.
- Calvert, J., A. Mellouki, and J. Orlando, 2011: *Mechanisms of Atmospheric Oxidation of the Oxygenates*. Oxford University Press, 1634 pp.
- Carpenter, L. J., and Coauthors, 2010: Seasonal characteristics of tropical marine boundary layer air measured at the Cape Verde Atmospheric Observatory. *J. Atmos. Chem.*, **67**, 87–140, doi:10.1007/s10874-011-9206-1.
- , S. M. MacDonald, M. D. Shaw, R. Kumar, R. W. Saunders, R. Parthipan, J. Wilson, and J. M. C. Plane, 2013: Atmospheric iodine levels influenced by sea surface emissions of inorganic iodine. *Nat. Geosci.*, **6**, 108–111, doi:10.1038/ngeo1687.
- , S. Reimann, J. B. Burkholder, C. Clerbaux, B. D. Hall, R. Hossaini, J. C. Laube, and S. A. Yvon-Lewis, 2014: Ozone-depleting substances (ODSs) and other gases of interest to the Montreal Protocol. Scientific assessment of ozone depletion: 2014, Global Ozone Research and Monitoring Project Rep. 55, World Meteorological Organization, 1.1–1.79. [Available online at [www.wmo.int/pages/prog/arep/gaw/ozone\\_2014/documents/Full\\_report\\_2014\\_Ozone\\_Assessment.pdf](http://www.wmo.int/pages/prog/arep/gaw/ozone_2014/documents/Full_report_2014_Ozone_Assessment.pdf).]
- Chen, D., and Coauthors, 2016: Airborne measurements of BrO and the sum of HOBr and Br<sub>2</sub> over the tropical west Pacific from 1 to 15 km during the Convective Transport of Active Species in the Tropics (CONTRAST) experiment. *J. Geophys. Res. Atmos.*, **121**, 12 560–12 578, doi:10.1002/2016JD025561.
- Corti, T., B. P. Luo, T. Peter, H. Voemel, and Q. Fu, 2005: Mean radiative energy balance and vertical mass fluxes in the equatorial upper troposphere and lower stratosphere. *Geophys. Res. Lett.*, **32**, L06802, doi:10.1029/2004GL021889.
- Crawford, J. H., and Coauthors, 1997: Implications of large scale shifts in tropospheric NO<sub>x</sub> levels in the remote tropical Pacific. *J. Geophys. Res.*, **102**, 28 447–28 468, doi:10.1029/97JD00011.
- Dima, I. M., and J. M. Wallace, 2007: Structure of annual-mean equatorial planetary waves. *J. Atmos. Sci.*, **64**, 2862–2880, doi:10.1175/JAS3985.1.
- Dix, B., S. Baidar, J. F. Bresch, S. R. Hall, K. S. Schmidt, S. Wang, and R. Volkamer, 2013: Detection of iodine monoxide in the tropical free troposphere. *Proc. Natl. Acad. Sci. USA*, **110**, 2035–2040, doi:10.1073/pnas.1212386110.
- Fernandez, R. P., R. J. Salawitch, D. E. Kinnison, J.-F. Lamarque, and A. Saiz-Lopez, 2014: Bromine partitioning in the tropical tropopause layer: Implications for stratospheric injection. *Atmos. Chem. Phys.*, **14**, 17 857–17 905, doi:10.5194/acpd-14-17857-2014.
- Flemming, J., A. Inness, H. Flentje, V. Huijnen, P. Moinat, M. G. Schultz, and O. Stein, 2009: Coupling global chemistry transport models to ECMWF's integrated forecast system. *Geosci. Model Dev.*, **2**, 253–265, doi:10.5194/gmd-2-253-2009.
- Frieler, K., and Coauthors, 2006: Toward a better quantitative understanding of polar stratospheric ozone loss. *Geophys. Res. Lett.*, **33**, L10812, doi:10.1029/2005GL025466.
- Fueglistaler, S., H. Wernli, and T. Peter, 2004: Tropical troposphere-to-stratosphere transport inferred from trajectory calculations. *J. Geophys. Res.*, **109**, D03108, doi:10.1029/2003JD004069.
- , A. E. Dessler, T. J. Dunkerton, I. Folkins, Q. Fu, and P. W. Mote, 2009: Tropical tropopause layer. *Rev. Geophys.*, **47**, RG1004, doi:10.1029/2008RG000267.
- Gregory, G. L., and Coauthors, 1999: Chemical characteristics of Pacific tropospheric air in the region of the intertropical convergence zone and South Pacific convergence zone. *J. Geophys. Res.*, **104**, 5677–5696, doi:10.1029/98JD01357.
- Harris, N. R. P., and Coauthors 2017: Coordinated Airborne Studies in the Tropics (CAST). *Bull. Amer. Meteor. Soc.*, **98**, 145–162, doi:10.1175/BAMS-D-14-00290.1.
- Hayashi, H., K. Kita, and S. Taguchi, 2008: Ozone-enhanced layers in the troposphere over the equatorial Pacific Ocean and the influence of transport of midlatitude UT/LS air. *Atmos. Chem. Phys.*, **8**, 2609–2621, doi:10.5194/acp-8-2609-2008.
- Hoell, J. M., and Coauthors, 1999: Pacific Exploratory Mission in the tropical Pacific: PEM-Tropics A, August–September 1996. *J. Geophys. Res.*, **104**, 5567–5583, doi:10.1029/1998JD100074.
- Holton, J. R., and A. Gettelman, 2001: Horizontal transport and the dehydration of the stratosphere. *Geophys. Res. Lett.*, **28**, 2799–2802, doi:10.1029/2001GL013148.
- Huey, L. G., 2007: Measurement of trace atmospheric species by chemical ionization mass spectrometry:

- Speciation of reactive nitrogen and future directions. *Mass Spectrom. Rev.*, **26**, 166–184, doi:10.1002/mas.20118.
- Jacob, D. J., and Coauthors, 2003: The Transport and Chemical Evolution over the Pacific (TRACE-P) aircraft mission: Design, execution, and first results. *J. Geophys. Res.*, **108**, 9000, doi:10.1029/2002JD003276.
- Jensen, E. J., and Coauthors, 2017: The NASA Airborne Tropical Tropopause Experiment (ATTREX): High-altitude aircraft measurements in the tropical western Pacific. *Bull. Amer. Meteor. Soc.*, **98**, 129–143, doi:10.1175/BAMS-D-14-00263.1.
- Kley, D., P. J. Crutzen, H. G. J. Smit, H. Vömel, S. J. Oltmans, H. Grassl, and V. Ramanathan, 1996: Observations of near-zero ozone concentrations over the convective Pacific: Effects on air chemistry. *Science*, **274**, 230–233, doi:10.1126/science.274.5285.230.
- Ko, M. K. W., N.-D. Sze, C. J. Scott, and D. K. Weisenstein, 1997: On the relation between stratospheric chlorine/bromine loading and short-lived tropospheric source gases. *J. Geophys. Res.*, **102**, 25 507–25 517, doi:10.1029/97JD02431.
- Kondo, Y., and Coauthors, 2002: Effects of biomass burning, lightning, and convection on O<sub>3</sub>, CO, and NO<sub>y</sub> over the tropical Pacific and Australia in August–October 1998 and 1999. *J. Geophys. Res.*, **107**, 8402, doi:10.1029/2001JD000820.
- Krüger, K., S. Tegtmeier, and M. Rex, 2008: Long-term climatology of air mass transport through the Tropical Tropopause Layer (TTL) during NH winter. *Atmos. Chem. Phys.*, **8**, 813–823, doi:10.5194/acp-8-813-2008.
- Lamarque, J.-F., and Coauthors, 2012: CAM-chem: Description and evaluation of interactive atmospheric chemistry in the Community Earth System Model. *Geosci. Model Dev.*, **5**, 369–411, doi:10.5194/gmd-5-369-2012.
- Langner, J., and H. Rodhe, 1991: A global three-dimensional model of the tropospheric sulfur cycle. *J. Atmos. Chem.*, **13**, 225–264, doi:10.1007/BF00058134.
- Liu, C., and E. J. Zipser, 2015: The global distribution of largest, deepest, and most intense precipitation systems. *Geophys. Res. Lett.*, **42**, 3591–3595, doi:10.1002/2015GL063776.
- Millet, D. B., and Coauthors, 2010: Global atmospheric budget of acetaldehyde: 3-D model analysis and constraints from in-situ and satellite observations. *Atmos. Chem. Phys.*, **10**, 3405–3425, doi:10.5194/acp-10-3405-2010.
- Newell, R. E., and S. Gould-Stewart, 1981: A stratospheric fountain? *J. Atmos. Sci.*, **38**, 2789–2796, doi:10.1175/1520-0469(1981)038<2789:ASF>2.0.CO;2.
- Newton, R., G. Vaughan, H. M. A. Ricketts, L. L. Pan, A. J. Weinheimer, and C. Chemel, 2016: Ozone profile from the West Pacific Warm Pool: Measurements and validation. *Atmos. Chem. Phys.*, **16**, 619–634, doi:10.5194/acp-16-619-2016.
- Nicely, J. M., and Coauthors, 2016: An observationally constrained evaluation of the oxidative capacity in the tropical western Pacific troposphere. *J. Geophys. Res. Atmos.*, **121**, 7461–7488, doi:10.1002/2016JD025067.
- Pan, L. L., and L. A. Munchak, 2011: Relationship of cloud top to the tropopause and jet structure from CALIPSO data. *J. Geophys. Res.*, **116**, D12201, doi:10.1029/2010JD015462.
- , and Coauthors, 2015: Bimodal distribution of free tropospheric ozone over the tropical western Pacific revealed by airborne observations. *Geophys. Res. Lett.*, **42**, 7844–7851, doi:10.1002/2015GL065562.
- Pierce, R. B., and Coauthors, 2007: Chemical data assimilation estimates of continental U.S. ozone and nitrogen budgets during the Intercontinental Chemical Transport Experiment–North America. *J. Geophys. Res.*, **112**, D12S21, doi:10.1029/2006JD007722.
- Randel, W. J., and E. J. Jensen, 2013: Physical processes in the tropical tropopause layer and their roles in a changing climate. *Nat. Geosci.*, **6**, 169–176, doi:10.1038/ngeo1733.
- Randel, W. J., L. Rivoire, L. L. Pan, and S. B. Honomichl, 2016: Dry layers in the tropical troposphere observed during CONTRAST and global behavior from GFS analyses. *J. Geophys. Res. Atmos.*, **121**, doi:10.1002/2016JD025841.
- Rex, M., and Coauthors, 2014: A tropical West Pacific OH minimum and implications for stratospheric composition. *Atmos. Chem. Phys.*, **14**, 4827–4841, doi:10.5194/acp-14-4827-2014.
- Reynolds, R. W., N. A. Rayner, T. M. Smith, D. C. Stokes, and W. Wan, 2002: An improved in situ and satellite SST analysis for climate. *J. Climate*, **15**, 1609–1625, doi:10.1175/1520-0442(2002)015<1609:AIISAS>2.0.CO;2.
- Ridley, B. A., F. E. Grahek, and J. G. Walega, 1992: A small, high-sensitivity, medium-response ozone detector for measurements from light aircraft. *J. Atmos. Oceanic Technol.*, **9**, 142–148, doi:10.1175/1520-0426(1992)009<0142:ASHSMR>2.0.CO;2.
- Saiz-Lopez, A., and R. von Glasow, 2012: Reactive halogen chemistry in the troposphere. *Chem. Soc. Rev.*, **41**, 6448–6472, doi:10.1039/c2cs35208g.
- , and Coauthors, 2012: Estimating the climate significance of halogen-driven ozone loss in the tropical marine troposphere. *Atmos. Chem. Phys.*, **12**, 3939–3949, doi:10.5194/acp-12-3939-2012.

- , R. P. Fernandez, C. Ordóñez, D. E. Kinnison, J. C. Gómez Martín, J. F. Lamarque, and S. Tilmes, 2014: Iodine chemistry in the troposphere and its effect on ozone. *Atmos. Chem. Phys.*, **14**, 13 119–13 143, doi:10.5194/acp-14-13119-2014.
- , A., and Coauthors, 2015: Injection of iodine to the stratosphere. *Geophys. Res. Lett.*, **42**, 6852–6859, doi:10.1002/2015GL064796.
- Salawitch, R. J., D. K. Weisenstein, L. J. Kovalenko, C. E. Sioris, P. O. Wennberg, K. Chance, M. K. W. Ko, and C. A. McLinden, 2005: Sensitivity of ozone to bromine in the lower stratosphere. *Geophys. Res. Lett.*, **32**, L05811, doi:10.1029/2004GL021504.
- Schoeberl, M. R., and A. E. Dessler, 2011: Dehydration of the stratosphere. *Atmos. Chem. Phys.*, **11**, 8433–8446, doi:10.5194/acp-11-8433-2011.
- Skamarock, W. C., and Coauthors, 2008: A description of the Advanced Research WRF version 3. NCAR Tech. Note NCAR/TN-475+STR, 113 pp., doi:10.5065/D68S4MVH.
- Stephens, G. L., and Coauthors, 2008: CloudSat mission: Performance and early science after the first year of operation. *J. Geophys. Res.*, **113**, D00A18, doi:10.1029/2008JD009982.
- Takahashi, H., and Z. Luo, 2012: Where is the level of neutral buoyancy for deep convection? *Geophys. Res. Lett.*, **39**, L15809, doi:10.1029/2012GL052638.
- Thompson, A. M., S. J. Oltmans, D. W. Tarasick, P. von der Gathen, H. G. J. Smit, and J. C. Witte, 2011: Strategic ozone sounding networks: Review of design and accomplishments. *Atmos. Environ.*, **45**, 2145–2163, doi:10.1016/j.atmosenv.2010.05.002.
- Thouret, V., J. Y. N. Cho, M. J. Evans, R. E. Newell, M. A. Avery, J. D. W. Barrick, G. W. Sachse, and G. L. Gregory, 2001: Tropospheric ozone layers observed during PEM-Tropics B. *J. Geophys. Res.*, **106**, 32 527–32 538, doi:10.1029/2001JD900011.
- Volkamer, R., and Coauthors, 2015: Aircraft measurements of BrO, IO, glyoxal, NO<sub>2</sub>, H<sub>2</sub>O, O<sub>2</sub>–O<sub>2</sub> and aerosol extinction profiles in the tropics: Comparison with aircraft-/ship-based in situ and lidar measurements. *Atmos. Meas. Tech.*, **8**, 2121–2148, doi:10.5194/amt-8-2121-2015.
- Vömel, H., and K. Diaz, 2010: Ozone sonde cell current measurements and implications for observations of near-zero ozone concentrations in the tropical upper troposphere. *Atmos. Meas. Tech.*, **3**, 495–505, doi:10.5194/amt-3-495-2010.
- Wang, S.-Y., and Coauthors, 2015: Active and widespread halogen chemistry in the tropical and subtropical free troposphere. *Proc. Natl. Acad. Sci. USA*, **112**, 9281–9286, doi:10.1073/pnas.1505142112.
- Webster, P. J., and R. Lukas, 1992: TOGA COARE: The Coupled Ocean–Atmosphere Response Experiment. *Bull. Amer. Meteor. Soc.*, **73**, 1377–1416, doi:10.1175/1520-0477(1992)073<1377:TCTCOR>2.CO;2.
- Wyrutki, K., 1989: Some thoughts about the West Pacific Warm Pool. *Proc. Western Pacific Int. Meeting and Workshop on TOGA COARE*, Nouméa, New Caledonia, ORSTOM, 99–109.
- Yang, Q., Q. Fu, and Y. Hu, 2010: Radiative impacts of clouds in the tropical tropopause layer. *J. Geophys. Res.*, **115**, D00H12, doi:10.1029/2009JD012393.
- Yoneyama, K., and D. B. Parsons, 1999: A mechanism for the intrusion of dry air into the tropical western Pacific region. *J. Atmos. Sci.*, **56**, 1524–1546, doi:10.1175/1520-0469(1999)056<1524:APMFTI>2.0.CO;2.