Atmospheric Carbon Dioxide Variability in the Community Earth System Model: Evaluation and Transient Dynamics during the Twentieth and Twenty-First Centuries

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ABSTRACT

Changes in atmospheric CO2 variability during the twenty-first century may provide insight about ecosystem responses to climate change and have implications for the design of carbon monitoring programs. This paper describes changes in the three-dimensional structure of atmospheric CO2 for several representative concentration pathways (RCPs 4.5 and 8.5) using the Community Earth System Model–Biogeochemistry (CESM1-BGC). CO2 simulated for the historical period was first compared to surface, aircraft, and column observations. In a second step, the evolution of spatial and temporal gradients during the twenty-first century was examined. The mean annual cycle in atmospheric CO2 was underestimated for the historical period throughout the Northern Hemisphere, suggesting that the growing season net flux in the Community Land Model (the land component of CESM) was too weak. Consistent with weak summer drawdown in Northern Hemisphere high latitudes, simulated CO2 showed correspondingly weak north–south and vertical gradients during the summer. In the simulations of the twenty-first century, CESM predicted increases in the mean annual cycle of atmospheric CO2 and larger horizontal gradients. Not only did the mean north–south gradient increase due to fossil fuel emissions, but east–west contrasts in CO2 also strengthened because of changing patterns in fossil fuel emissions and terrestrial carbon exchange. In the RCP8.5 simulation, where CO2 increased to 1150 ppm by 2100, the CESM predicted increases in interannual variability in the Northern Hemisphere midlatitudes of up to 60% relative to present variability for time series filtered with a 2–10-yr bandpass. Such an increase in variability may impact detection of changing surface fluxes from atmospheric observations.

1. Introduction

Carbon dioxide is a dominant anthropogenic greenhouse gas in the earth’s atmosphere. Its atmospheric concentration has risen from 280 ppm preindustrially to 390 ppm in 2011 because of fossil fuel combustion and land use change (R. F. Keeling et al. 1996; Prentice et al. 2000). The extent to which CO2 levels in the atmosphere continue increasing will be the primary determinant of future climate change (Solomon et al. 2007). The trajectory of atmospheric CO2 depends not only on human-mediated fluxes, such as those from fossil fuel combustion or land use change, but also on the response of the natural carbon cycle to changing atmospheric composition and climate (Fung et al. 2005). Currently, 55% of CO2 emitted by human activities is taken up by the ocean or terrestrial ecosystems, with 45% remaining airborne (Le Quéré et al. 2009). The degree to which the efficiency of these sinks will change with climate is an
open question, with implications for the rate of atmospheric accumulation. Therefore, prediction of future climate is closely linked to prediction of carbon cycle dynamics (Friedlingstein et al. 2006).

Numerous studies have documented the responses of terrestrial and oceanic fluxes of carbon to changes in the physical climate system (e.g., Sarmiento and Gruber 2002). Carbon–climate interactions operate on multiple time scales; for instance, drought and high temperature stress can reduce carbon uptake by terrestrial ecosystems over continental-scale regions (Ciais et al. 2005; Phillips et al. 2009) and can trigger increases in biomass burning (e.g., Golding and Betts 2008; Chen et al. 2011) on interannual time scales. Observations of atmospheric CO₂ indicate that the atmospheric growth rate varies substantially from year to year owing to interannual climate modes, such as the El Niño–Southern Oscillation, and their impact on ocean and terrestrial biospheric fluxes (Bacastow 1976; Rayner et al. 1999; Jones et al. 2001; Schwalm et al. 2011). On decadal time scales, increases in westerly winds over the Southern Ocean may have weakened the net ocean uptake of CO₂ over the last three decades (Le Quéré et al. 2007; Lovenduski et al. 2008). In terrestrial ecosystems, lengthening of the growing season in high northern latitudes has increased carbon dioxide uptake in spring and early summer (C. D. Keeling et al. 1996; Randerson et al. 1999), but also carbon release in the early fall (Piao et al. 2008). The carbon losses from these ecosystems are likely caused by increases in soil temperature that may contribute to permafrost thaw and degradation (e.g., Schuur et al. 2009; Koven et al. 2011).

Global climate models have developed over time to include more Earth system processes that feedback onto the climate. With the development of Earth system models (ESMs), interactions between the carbon cycle and the climate system are represented through multiple pathways, enabling the study of climate–carbon feedbacks (e.g., Thornton et al. 2009; Arora and Montenegro 2011). These models allow us to estimate likely CO₂ levels that correspond to particular fossil fuel emission pathways and to better represent uncertainty in climate projections resulting from carbon cycle processes. Important challenges in this regard are evaluating ESM performance and understanding how biases in model structure influence future projections.

Here, we developed several new approaches for evaluating the three-dimensional structure of CO₂ in earth system models. Atmospheric CO₂ is an integrator of surface fluxes, and its variations on seasonal and longer time scales reflect the terrestrial and oceanic processes that control carbon–climate feedbacks (Randerson et al. 2009; Cadule et al. 2010). Although CO₂ perturbations in the atmosphere persist long enough to be well mixed in terms of radiative forcing, small horizontal and vertical gradients in atmospheric CO₂ allow for the diagnosis of regional to continental-scale sources and sinks (e.g., Gurney et al. 2002). Carbon dioxide is emitted and removed largely at the surface, but simulation of atmospheric mixing near the surface is challenging because of difficulties in representing boundary layer and convective processes and the impacts of topographic variations in atmospheric models. As a consequence, simulation of vertical and horizontal mixing of CO₂ varies considerably among different models, and this has important implications for our understanding of CO₂, including the magnitude of sources and sinks (Gurney et al. 2002; Stephens et al. 2007; Yang et al. 2007).

In this study, we analyzed the three-dimensional structure of atmospheric CO₂ in one example of such coupled models, the Community Earth System Model (CESM). In the version of CESM used here (CESM1-BGC), atmospheric CO₂ is represented as a three-dimensional atmospheric tracer, which, in the simulations described here, influenced the atmospheric radiation budget and thus top-of-the-atmospheric energy fluxes (Lindsay et al. 2012, manuscript submitted to J. Climate). Variability of CO₂ in the model is controlled by the geographic and seasonal distribution of natural and anthropogenic surface fluxes as well as by different atmospheric transport processes such as advection, convection, and eddy mixing.

We evaluated the performance of the CESM against several types of atmospheric CO₂ data measured from different platforms, each of which has a different measurement footprint and is therefore impacted by fluxes and transport differently. We used the following datasets: surface flask and in situ data from the NOAA Global Monitoring Division (GMD) (Conway et al. 1994) and the Scripps Institution of Oceanography (SIO) (Keeling et al. 2001), column-average mole fraction observations from the Total Carbon Column Observing Network (TCCON) (Wunch et al. 2011), and aircraft data obtained from both the High-Performance Instrumented Airborne Platform for Environmental Research (HIAPER) Pole-to-Pole Observations (HIPPO) campaigns (Wofsy et al. 2011) and GMD and Department of Energy (DOE) long-term monitoring stations. In our analysis, we examined seasonal cycles in CO₂, vertical and horizontal gradients, and interannual variability, along with multidecadal trends in these datasets. These comparisons were used to infer how accurately the CESM reproduces spatial and temporal variations in fluxes, which in turn provides a means to gauge the realism of future projections by the model for representative concentration pathways (RCPs) (Moss et al. 2010).
In the second part of this paper, we analyzed simulations of twenty-first-century CO2 based on the RCP scenarios developed for phase 5 of the Coupled Model Intercomparison Project (CMIP5). This analysis provides us with insight about the response of carbon fluxes to increased anthropogenic forcing and how variations in atmospheric CO2 may change in time with possible consequences for monitoring strategies. Because of interest in an international climate treaty to monitor and verify national fossil fuel emissions (National Research Council 2010), the carbon monitoring community will require observations of atmospheric CO2 that can differentiate among trajectories of CO2 emissions at subcontinental spatial scales. When considering the information that would be gained by additional observations, it is important to consider the changing background variability—such as seasonal variations and variations caused by transport acting on larger spatial gradients—in the future and how this variability influences our ability to use the network for change detection and carbon flux attribution.

This paper is structured as follows: in section 2, we describe the CESM components relevant for simulating atmospheric CO2 and the atmospheric CO2 data used for comparison with the model. In section 3, we provide results from the comparison of simulated CO2 from the CESM with these datasets. In section 4, we show trends in simulated CO2 for two RCP scenarios. In section 5, we discuss the implications of our comparisons for model development, carbon cycle science, and future atmospheric monitoring strategies.

2. Methods

a. Community Earth System Model

We analyzed simulations from CESM1-BGC, a fully coupled general circulation model with atmosphere, ocean, land, and sea ice components. CESM was developed from model components in the Community Climate System Model version 4 (CCSM4) (Gent et al. 2011). For the purposes of this study, the most significant change with CESM was that the carbon cycle was completely prognostic: feedbacks between the simulated climate and terrestrial and oceanic fluxes of CO2 determined the atmospheric concentration used in the radiative scheme. More details on the biogeochemistry coupling are found in Lindsay et al. (2012, manuscript submitted to J. Climate).

The atmospheric component of CESM is the Community Atmospheric Model (CAM), version 4 (Neale et al. 2013). CAM 4 used a finite volume dynamical core and has an improved parameterization of deep convection relative to its predecessors (Gent et al. 2011). Interannual variability due to El Niño was better captured than in previous versions (Deser et al. 2012), and variables such as land surface temperature and precipitation that affect terrestrial carbon cycling also showed less bias (P. J. Lawrence et al. 2012). For the simulations analyzed here, CAM 4 was run at 0.9° latitude × 1.25° longitude horizontal resolution and had 26 vertical layers arranged in a hybrid pressure sigma coordinate. The total CO2 dry air mole fraction (parts per million) and the dry air mole fraction of the components of CO2 owing to fossil fuel fluxes, land exchange, and ocean exchange were each tracked separately in the model. We primarily analyzed monthly mean CESM output, but used daily mean CO2 for the comparisons with atmospheric aircraft data.

The Community Land Model (CLM) version 4 was the basis for terrestrial ecosystem fluxes of CO2 (D. M. Lawrence et al. 2012). Like previous versions of the land model, CLM4 simulated fluxes of energy and water between the atmosphere and land surface. This newest version included a carbon–nitrogen (C–N) cycle model that is prognostic in carbon and nitrogen fluxes and in vegetation phenology (Thornton et al. 2007). In the experiments that we analyzed, plant functional types were prescribed, and land-use change fluxes were simulated following the approach described by P. J. Lawrence et al. (2012). The imposed land-use change fluxes are in good agreement with Global Carbon Project (GCP) estimates (Canadell et al. 2007) between 1970 and 1999. For this time period, the GCP estimated a 1.5 ± 0.5 Pg C yr\(^{-1}\) land use source of CO2 to the atmosphere, while the fluxes in CESM are slightly lower at 1.4 Pg C yr\(^{-1}\). Global soil carbon from CLM4 was 514 Pg C, which is approximately a factor of 2 lower than the 1255 Pg C estimate from the Harmonized World Soil Database (HWSD) (Todd-Brown et al. 2013). At high northern latitudes, the low bias in CLM4 was even greater: by a factor of 5 compared to HWSD and by over a factor of 8 compared to the Northern Circumpolar Soil Carbon Database. Global net primary production levels were approximately 52 Pg C yr\(^{-1}\), comparable to remote sensing-derived estimates (Todd-Brown et al. 2013). At high northern latitudes, gross primary production (GPP) from CLM4 was approximately the same as observations from FLUXNET, although in the tropics there was a significant positive bias (Bonan et al. 2011). The combined soil carbon and productivity estimates from CLM indicated that soil organic turnover times in the model were likely too small, with rapid carbon cycling contributing to the low bias in soil carbon stocks. We also note that the CESM runs analyzed here did not include dynamic vegetation changes, which may have
limited the magnitude of simulated changes in regional carbon fluxes during the twenty-first century simulations.

The ocean model influenced atmospheric CO₂ in CESM by physical uptake and transport of CO₂ in overturning water masses and through ocean biogeochemistry. The ocean physical model is described in detail in Danabasoglu et al. (2011). Briefly, the ocean model was run at 1° horizontal resolution with 60 vertical layers. Ocean dynamics were based on the Parallel Ocean Program, version 2 (Smith et al. 2010). The ocean biogeochemical model was the Biogeochemical Element Cycle (BEC) model developed for the CCSM (Moore et al. 2004; Moore and Braucher 2008; Doney et al. 2009). There were few modifications to the model for the CESM implementation (for details see Lindsay et al. 2012, manuscript submitted to J. Climate; Moore et al. 2012, manuscript submitted to J. Climate). Air–sea CO₂ exchange during the twentieth century is described by Long et al. (2013).

We analyzed output from a CESM simulation that followed the historical CMIP5 protocol. The simulation began in 1850 and ended in 2005. Anthropogenic fossil fuel emissions, based on national emission estimates from the Carbon Dioxide Information Analysis Center (CDIAC) (Boden et al. 2009), were prescribed with a 1° spatial resolution and monthly time step (Andres et al. 2012, manuscript submitted to J. Climate). Transient aerosol emissions also were specified, following time series developed by Lamarque et al. (2010). As described above, exchange of CO₂ between the land and atmosphere and ocean and atmosphere was fully coupled within the model. As a result, atmospheric CO₂ responded prognostically to fossil and land use change fluxes.

We also analyzed variations in twenty-first century CO₂ from two CESM simulations that followed different RCP scenarios (Moss et al. 2010; van Vuuren et al. 2011). The RCP4.5 scenario corresponds to a nominal anthropogenic radiative forcing of 4.5 W m⁻² at 2100 and assumes stabilization of emissions by the middle of the twenty-first century. To achieve emissions reductions and radiative forcing stabilization of 4.5 W m⁻², the scenario assumes international participation in emissions mitigation and a common global price for emissions of CO₂ and other greenhouse gases that cover all sectors of the economy (Thomson et al. 2011). The RCP8.5 scenario corresponds to nominal anthropogenic radiative forcing of 8.5 W m⁻² at 2100 and assumes steady population growth with continuing high energy demand and little investment in energy technology, leading to reliance on local and high-carbon-content fuels (Riahi et al. 2011). These simulations began in the year 2005, with initial conditions provided by the historical simulation described above, and were run to 2100. Fossil fuel emission trajectories consistent with the nominal radiative forcing for these two scenarios were developed using two different integrated assessment models (IAMs) (van Vuuren et al. 2011). The regionally aggregated emissions generated by the IAMs were regridded at monthly resolution by T. Nozawa from the Asian–Pacific Integrated Model (AIM) modeling team at the National Institute for Environmental Studies in Japan by scaling the spatial and seasonal patterns for year 2005 fluxes (e.g., Andres et al. 2011). This regridding process accounted for CO₂ emissions from international ships. As such, the spatial and seasonal distribution of emissions may differ from emissions used by other modeling centers for the same CMIP5 experiments. And the seasonal dynamics and finescale spatial structure represent only one possible scenario of fossil fuel emission patterns consistent with the RCP time series. Because the interactive carbon fluxes and aerosol parameterizations in CESM differed from those in the IAMs used to devise the emissions pathways, the resulting radiative forcing differed from the nominal RCP value at 2100.

b. Carbon dioxide observations

We used several data sources to evaluate simulated atmospheric CO₂, including surface, column, and aircraft measurements (Fig. 1). The surface, column, and aircraft observations spanned all latitudes, but several geographic areas were under represented (e.g., Africa and the Amazon). For ease in comparing the observations with the model, we binned the data into the eight latitude zones shown by alternating color bands in Fig. 1 and listed in Table 1.

For surface CO₂ data, we compared CESM to NOAA GMD flask and Scripps in situ observations. The NOAA data record consisted of weekly flasks sampled in a global cooperative network (Fig. 1, Table 1). These data
are compatible to within 0.2 ppm of the WMO CO$_2$ scale (WMO 2011), with a precision of ±0.1 ppm (1σ) (Conway et al. 1994). We selected locations that had near-continuous data coverage from the mid-1990s (or earlier) to the present. We also used monthly mean atmospheric CO$_2$ obtained in situ at the Mauna Loa (MLO) and South Pole (SPO) Observatories by the Scripps Institution of Oceanography (SIO) (Keeling et al. 2001) to compare with the atmospheric growth rate in the model from 1958 to 2010.

For each surface observation site, we sampled CESM at the gridbox containing the observatory. We sampled the model at the nearest altitude (in units of meters above sea level) to that at which observations were obtained. For one site—Baring Head, New Zealand, where observations were only made when the wind was onshore and CO$_2$ stable—we chose a CESM gridbox 10° south of the site to reflect marine background conditions. Similar approaches have been used in earlier atmospheric transport and inversion studies to account for the sampling biases introduced by coarse-resolution global model grids (e.g., Law et al. 2010). We did not use modified sampling locations for other sites because patterns in the shape and phasing of the mean annual cycle qualitatively between CESM and the observations. We note that in our comparisons of NOAA GMD and TCCON observations, we used 24-h mean mole fractions from the model, whereas many of the observations were sampled only during the day. To avoid potential biases from this temporal sampling mismatch, we did not attempt to systematically analyze CO$_2$ variability on the planetary boundary layer over land.

We compared total column CO$_2$ in CESM to data obtained by TCCON observatories (Washenfelder et al. 2006; Deutscher et al. 2010; Messerschmidt et al. 2011; Wunch et al. 2011). Total column CO$_2$ is an observation of the vertically integrated mass of CO$_2$ above a sampling location, normalized by the dry air column to yield a vertically integrated mole fraction. As such, simulated total column CO$_2$ is less sensitive to errors in the model’s vertical mixing parameterization than surface mole fraction (Keppel-Aleks et al. 2011). Variations in total column CO$_2$ integrate fluxes at large scales, so the observations provide little information about local fluxes and, instead, provide strong constraints on hemispheric-scale fluxes (Keppel-Aleks et al. 2012).

TCCON data were obtained using ground-based Fourier transform spectrometry. The data have been calibrated against aircraft profiles and are compatible with respect to the WMO CO$_2$ scale and precise to better than 0.8 ppm (1-σ precision) (Wunch et al. 2010). Only seven TCCON sites had at least one full annual cycle of observations, although several more high northern latitude stations had obtained data during summer, when solar zenith angles permitted retrievals (Table 2). We leveraged sites without a full annual data cycle to constrain the north–south CO$_2$ mass gradient during seasons when observations were available. We sampled CESM at the TCCON locations and integrated the simulated CO$_2$ mole fractions to determine the total column average mole fraction, applying TCCON averaging kernels as described in Keppel-Aleks et al. (2012).

We compared free tropospheric CESM CO$_2$ with aircraft measurements from two sources: the HIPPO campaigns (Wofsy et al. 2011) and biweekly NOAA GMD vertical profiles. Like surface CO$_2$, observations of free tropospheric CO$_2$ measured at a single level in the atmosphere are sensitive to the rate of vertical mixing in the model. Taken as a set, however, the relative agreement between observed and simulated CO$_2$ at several levels have the potential to allow diagnosis of the extent to which differences are due to surface fluxes or atmospheric physics (Stephens et al. 2007; Yang et al. 2007).

We compared free tropospheric CO$_2$ observed during the HIPPO campaign with CESM CO$_2$. HIPPO consisted of five campaigns, with flights occurring during all seasons between 2009 and 2011 (Wofsy et al. 2011). The aircraft obtained a series of vertical profiles from the surface to >14 km while flying between 87°N and 67°S. Flights occurred primarily over the Pacific, although individual campaigns included excursions over, for instance, Australia or the continental United States (Fig. 1). The CO$_2$ mole fraction was measured using several instruments aboard the NCAR Gulfstream V aircraft: two LI-COR nondispersive infrared gas analyzers, a quantum cascade laser system (QCLS), and two flask samplers. In this study, we used the composite recommended CO$_2$$_x$ variable, which is almost exclusively from the QCLS instrument, and has precision of 0.02 ppm and compatibility with respect to the World Meteorological Organization CO$_2$ scale of 0.2 ppm.

For HIPPO, we separated each of the five campaigns into a northbound and southbound leg and binned the data from each leg into the eight north–south regions and averaged the data vertically in four layers of equal depth between the surface and 10 km. We sampled daily mean CESM at every location where a HIPPO observation was available for five consecutive model years and averaged the model estimates using the same latitude–altitude bins.

The second set of aircraft observations consisted of biweekly to monthly profiles sampled by the NOAA Global Monitoring Division and the U.S. Department of Energy (DOE) at 19 aircraft sites in the Northern Hemisphere, primarily over North America (Fig. 1). Measurements were made between the surface and
TABLE 1. Surface flask observations within the eight regional latitude zones used for evaluating the CESM.

<table>
<thead>
<tr>
<th>Region</th>
<th>Station</th>
<th>Acronym</th>
<th>Lat (°N)</th>
<th>Lon (°E)</th>
<th>Operational dates</th>
</tr>
</thead>
<tbody>
<tr>
<td>60°–90°N</td>
<td>Alert, Alaska</td>
<td>ALT</td>
<td>82.5</td>
<td>−62.5</td>
<td>1985–2010</td>
</tr>
<tr>
<td></td>
<td>Barrow, Alaska</td>
<td>BRW</td>
<td>71.3</td>
<td>−156.6</td>
<td>1971–2010</td>
</tr>
<tr>
<td></td>
<td>Storhofdl, Iceland</td>
<td>ICE</td>
<td>63.4</td>
<td>−20.3</td>
<td>1992–2010</td>
</tr>
<tr>
<td></td>
<td>Summit, Greenland</td>
<td>SUM</td>
<td>72.6</td>
<td>−38.4</td>
<td>1997–2010</td>
</tr>
<tr>
<td></td>
<td>Ny-Alesund, Svalbard</td>
<td>ZEP</td>
<td>78.9</td>
<td>11.9</td>
<td>1994–2010</td>
</tr>
<tr>
<td>40°–60°N</td>
<td>Baltic Sea, Poland</td>
<td>BAL</td>
<td>55.4</td>
<td>17.2</td>
<td>1992–2010</td>
</tr>
<tr>
<td></td>
<td>Cold Bay, Alaska</td>
<td>CBA</td>
<td>55.2</td>
<td>−162.7</td>
<td>1978–2010</td>
</tr>
<tr>
<td></td>
<td>Sary Taukum, Kazakhstan</td>
<td>KZD</td>
<td>44.1</td>
<td>76.9</td>
<td>1997–2009</td>
</tr>
<tr>
<td></td>
<td>Plateu Assy, Kazakhstan</td>
<td>KZM</td>
<td>43.3</td>
<td>77.9</td>
<td>1997–2009</td>
</tr>
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<td></td>
<td>Mace Head, Ireland</td>
<td>MHD</td>
<td>53.3</td>
<td>−9.9</td>
<td>1991–2010</td>
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<td></td>
<td>Niwot Ridge, Colorado</td>
<td>NWR</td>
<td>40.1</td>
<td>−105.6</td>
<td>1968–2010</td>
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<tr>
<td></td>
<td>Shemya, Alaska</td>
<td>SHM</td>
<td>52.7</td>
<td>174.1</td>
<td>1985–2010</td>
</tr>
<tr>
<td></td>
<td>Trinidad Head, California</td>
<td>THD</td>
<td>41.1</td>
<td>−124.2</td>
<td>2002–2010</td>
</tr>
<tr>
<td></td>
<td>Ulaan Uul, Mongolia</td>
<td>UUM</td>
<td>44.5</td>
<td>111.1</td>
<td>1992–2010</td>
</tr>
<tr>
<td>15°–40°N</td>
<td>Assekrem, Algeria</td>
<td>ASK</td>
<td>23.2</td>
<td>5.4</td>
<td>1995–2010</td>
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<tr>
<td></td>
<td>Terceira Islands, Azores</td>
<td>AZR</td>
<td>38.8</td>
<td>−27.4</td>
<td>1979–2010</td>
</tr>
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<td></td>
<td>Tudor Hill, Bermuda</td>
<td>BMW</td>
<td>32.3</td>
<td>−64.9</td>
<td>1989–2010</td>
</tr>
<tr>
<td></td>
<td>Izana, Tenerife</td>
<td>IZO</td>
<td>28.3</td>
<td>−16.5</td>
<td>1991–2010</td>
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<tr>
<td></td>
<td>Key Biscayne, Florida</td>
<td>KEY</td>
<td>25.7</td>
<td>−80.2</td>
<td>1972–2010</td>
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<td></td>
<td>Midway Island</td>
<td>MID</td>
<td>28.2</td>
<td>−177.4</td>
<td>1985–2010</td>
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<td></td>
<td>Mauna Loa, Hawaii</td>
<td>MLO</td>
<td>19.5</td>
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</tr>
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<td></td>
<td>Pacific Ocean 20°N</td>
<td>POCN20</td>
<td>20.0</td>
<td>−141.0</td>
<td>1986–2009</td>
</tr>
<tr>
<td></td>
<td>Pacific Ocean 25°N</td>
<td>POCN25</td>
<td>25.0</td>
<td>−139.0</td>
<td>1986–2009</td>
</tr>
<tr>
<td></td>
<td>Point Arena, California</td>
<td>PTA</td>
<td>38.9</td>
<td>−123.7</td>
<td>1999–2010</td>
</tr>
<tr>
<td></td>
<td>Lamont, Oklahoma</td>
<td>SGP</td>
<td>36.8</td>
<td>−97.5</td>
<td>2002–2010</td>
</tr>
<tr>
<td></td>
<td>Tae-Ahn peninsula, South Korea</td>
<td>TAP</td>
<td>36.7</td>
<td>126.1</td>
<td>1990–2010</td>
</tr>
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<td></td>
<td>Wendover, Utah</td>
<td>UTA</td>
<td>39.9</td>
<td>−113.7</td>
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<td></td>
<td>Negev Desert, Israel</td>
<td>WIS</td>
<td>30.9</td>
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<td>1995–2010</td>
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<td></td>
<td>Mt. Waliguan, China</td>
<td>WLG</td>
<td>36.3</td>
<td>100.9</td>
<td>1990–2010</td>
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<tr>
<td>0°–15°N</td>
<td>Mariana Islands, Guam</td>
<td>GMI</td>
<td>13.4</td>
<td>144.7</td>
<td>1979–2010</td>
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<td></td>
<td>Pacific Ocean 0°</td>
<td>POC000</td>
<td>0.0</td>
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<tr>
<td></td>
<td>Pacific Ocean 5°N</td>
<td>POCN05</td>
<td>5.0</td>
<td>−151.0</td>
<td>1987–2009</td>
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<td></td>
<td>Pacific Ocean 10°N</td>
<td>POCN10</td>
<td>10.0</td>
<td>−149.0</td>
<td>1987–2009</td>
</tr>
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<td>Pacific Ocean 15°N</td>
<td>POCN15</td>
<td>15.0</td>
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<td>1986–2009</td>
</tr>
<tr>
<td></td>
<td>Ragged Point, Barbados</td>
<td>RPB</td>
<td>13.2</td>
<td>−59.4</td>
<td>1987–2010</td>
</tr>
<tr>
<td>0°–15°S</td>
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<td>ASC</td>
<td>−8.0</td>
<td>−14.4</td>
<td>1979–2010</td>
</tr>
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<tr>
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<td></td>
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<td></td>
<td>Pacific Ocean 35°S</td>
<td>POCN35</td>
<td>−35.0</td>
<td>−180.0</td>
<td>1987–2009</td>
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</table>
8 km via automated portable flask packages aboard small aircraft. Table 3 lists the sites and the temporal coverage of the data. Like the surface flasks, the aircraft flasks were analyzed at NOAA within one week of sampling (Crevoisier et al. 2010; Biraud et al. 2013; http://www.esrl.noaa.gov/gmd/ccgg/aircraft/index.html).

c. Time series analysis

To compare CESM with observations of the annual cycle, we detrended both the observations and CESM using an iterative approach. At each observing site and altitude bin, a second-order polynomial was fit to the CO₂ time series and then removed, and a mean annual cycle was determined from the thus detrended data by averaging all January data points, all February data points, etc. The mean annual cycle was subtracted from the raw time series, and we again fit a second-order polynomial to the deseasonalized time series. This second polynomial was subtracted from the raw data: we refer to the result as the “detrended time series.” This iterative approach is similar to the one described by Cleveland et al. (1983) and used in several carbon cycle studies (e.g., Conway et al. 1994; Matsueda et al. 2002).

For comparing seasonal cycles between CESM data and observations, we computed the mean annual cycle, as described above, from a multiyear, detrended time series. For CESM and NOAA GMD flask data, we determined the mean annual cycle for 1995–2005; for TCCON and NOAA aircraft data, we used the available data and assumed that trends in the mean annual cycle after 2005 were small. To analyze trends in the seasonal cycle shape and amplitude, we first computed an “interannual” time series by removing the mean annual cycle from the detrended time series and smoothing the resulting time series with a 15-month boxcar filter. This interannual time series was subtracted from the deseasonalized time series to yield high-frequency residuals, and slopes were fitted through all January residuals, all February residuals, etc., to determine the trend for each month (ppm yr⁻¹) (cf. Randerson et al. 1999).

When analyzing interannual variability (IAV), we used a 2–10-yr bandpass filter to remove seasonal variations at high frequency and the secular increase in CO₂ at low frequency. The first and last five years of the combined historical and RCP time series (1850–2100) were removed to avoid edge effects due to the implementation of the bandpass filter. Then we calculated the standard deviation within 30-yr blocks to estimate IAV. Although there were some quantitative differences in the IAV depending on filter cutoffs, the temporal and spatial patterns in IAV were insensitive to the choice of cutoff.

To quantify north–south gradients, we subtracted the southernmost deseasonalized CO₂ mole fraction observation available to recenter the CESM output and the observations. This was done because the background CO₂ in the model was higher than in observations. For surface data, we subtracted the deseasonalized CO₂ mole fraction at SPO. For column data, we subtracted the deseasonalized column mole fraction at Lauder, New Zealand. For HIPPO aircraft profiles, we used the mole fraction at the southernmost surface bin (either 60°–90°S or 40°–60°S, depending on the campaign) to recenter the data for each northbound or southbound leg.

The amplitude of the mean annual cycle was determined by subtracting the minimum monthly mean CO₂ mole fraction from the maximum CO₂ mole fraction from the mean annual cycle. Because we determine the amplitude for both the model and observations from monthly means, damping of the seasonal amplitude due to monthly averaging is common to both data types. The trend in the mean annual cycle was likewise calculated from the month of maximum and minimum CO₂ mole fraction, similar to the methodology in Randerson et al. (1997), as

$$T_{\text{amp}} = 100\% \times \frac{b_{\text{max}} - b_{\text{min}}}{C_{\text{max}} - C_{\text{min}}}$$  \hspace{1cm} (1)

Here $b_{\text{max}}$ and $b_{\text{min}}$ are the slopes of the trends for the months of maximum and minimum CO₂ mole fraction,
and $C_{max}$ and $C_{min}$ are the mean mole fractions for those months. The month of maximum and minimum concentration were not allowed to vary within the decadal periods for which we computed trends in the amplitude.

We diagnosed the phasing of the seasonal cycle using four markers: the seasonal maximum and minimum and the zero crossings between the maximum and the minimum. In the Northern Hemisphere, the first zero crossing occurs in the spring when atmospheric CO2 levels first fall below the annual mean value. The second zero crossing occurs in the fall when atmospheric CO2 increases above its annual mean value after drawdown during the growing season. We determined the day of the year for these four markers by fitting a cubic spline with 1-day time resolution through the mean annual cycle composed of monthly mean data.

### 3. Comparison against contemporary observations

#### a. Atmospheric growth rate

The multidecadal growth rate of atmospheric CO2 in CESM exceeded observations at Mauna Loa during the latter half of the twentieth century (Fig. 2). By 1958, the start of the instrumental record, CESM CO2 was high by 4 ppm in the annual mean (320 ppm compared to 316 ppm in the observations). This difference increased with time such that, by the end of the historical CESM simulation in 2005, the difference was 20 ppm. By integrating the accumulated mass of CO2 for the period from 1958 to 2005, we determined that the airborne fraction of anthropogenic CO2 contributed from fossil fuel emissions and land use change was about 55% in CESM compared to 45% for the observations during this period (Le Quéré et al. 2009), indicating that the combined net

<table>
<thead>
<tr>
<th>Site</th>
<th>Lat (°N)</th>
<th>Lon (°E)</th>
<th>Mean annual cycle (ppm)</th>
<th>TCCON</th>
<th>CESM</th>
<th>Mean annual cycle difference (%)</th>
</tr>
</thead>
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<tr>
<td>Eureka</td>
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<tr>
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<td>0.8</td>
<td>−47</td>
<td></td>
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<tr>
<td>Lauder</td>
<td>−45.0</td>
<td>169.7</td>
<td>0.8</td>
<td>0.6</td>
<td>−25</td>
<td></td>
</tr>
</tbody>
</table>

### 3. Comparison against contemporary observations

#### a. Atmospheric growth rate

The multidecadal growth rate of atmospheric CO2 in CESM exceeded observations at Mauna Loa during the latter half of the twentieth century (Fig. 2). By 1958, the start of the instrumental record, CESM CO2 was high by 4 ppm in the annual mean (320 ppm compared to 316 ppm in the observations). This difference increased with time such that, by the end of the historical CESM simulation in 2005, the difference was 20 ppm. By integrating the accumulated mass of CO2 for the period from 1958 to 2005, we determined that the airborne fraction of anthropogenic CO2 contributed from fossil fuel emissions and land use change was about 55% in CESM compared to 45% for the observations during this period (Le Quéré et al. 2009), indicating that the combined net
The uptake of CO2 by the land biosphere and oceans was too weak in the model. P. J. Lawrence et al. (2012) and Lindsay et al. (2012, manuscript submitted to J. Climate) show that land use change fluxes were in good agreement with the Global Carbon Project estimates for the period from 1970 to 1999 (Canadell et al. 2007). The residual land sink in CESM for this same period, however, was only 1.3 Pg C yr\(^{-1}\), while the observationally based estimate reported by the GCP was 2.0 ± 0.7 Pg C yr\(^{-1}\). (We define the residual land sink here as the flux component that, when added to the net land use change flux, yields the net land flux.) The net air-to-sea flux in CESM also was slightly small (−1.9 versus −2.0 ± 0.4 Pg C yr\(^{-1}\)). However, it is worth noting that the net ocean carbon sink is sensitive to the partial pressure of atmospheric CO2, which was too high in CESM as a consequence of the cumulative impacts of reduced ocean and land carbon uptake. A comparison of the CESM ocean anthropogenic carbon inventory against observations from Sabine et al. (2004) and Khatiwala et al. (2009) provided evidence that cumulative ocean carbon storage was too weak in the model, partly in response to inadequate ocean vertical exchange in the Southern Ocean (Long et al. 2013; F. Hoffman et al. 2013, unpublished manuscript). Weak ventilation of Southern Ocean waters and weak anthropogenic CO2 uptake was a problem in the CCSM 3.1 model as well (Thornton et al. 2009; Wang et al. 2012) and has been observed in other CMIP3 models (Sarmiento et al. 2000).

### b. Seasonal amplitude and phasing

The Mauna Loa time series also indicates that CESM underestimated the amplitude of the mean annual cycle in the Northern Hemisphere (Fig. 2). A more comprehensive comparison using GMD surface flask observations between 1995 and 2005 indicated that the mean annual cycle was underestimated by 25%–40% in different latitude zones in the Northern Hemisphere (Fig. 3, Table 4). There was substantial site-by-site variation in the agreement of the observed and simulated amplitude and phase.

In the Northern Hemisphere, the onset of the growing season in CESM was too early. Both the maximum and first zero crossing (i.e., when CO2 falls below the annual mean value at a given location) occurred too early (Fig. 4). The first zero crossing was early by almost one month throughout the Northern Hemisphere extratropics, while the timing of the maximum was generally too early but varied with latitude. The model did capture the timing of the seasonal minimum in the Northern Hemisphere.

CESM overestimated the mean annual cycle at the surface in the Southern Hemisphere (Fig. 3, Table 4). The mean amplitudes were between 80% and 150% of those determined from flask observations. It is worth noting that the absolute disagreement in amplitude was smaller in the Southern Hemisphere than in the Northern Hemisphere and that the seasonality of Southern Hemisphere CO2 is influenced by Northern Hemisphere terrestrial fluxes (Randerson et al. 1997). In the Southern Hemisphere, the modeled maximum and minimum at surface sites lagged the observations in the tropics and subtropics but were in general agreement south of 40°S (Fig. 4). We repeated this analysis using CESM output at NOAA sampling sites, but at 1 km above the surface (generally within the third model layer) rather than at the surface, to determine whether the biases were caused by rectifier effects in the boundary layer and found similar biases in the mean annual cycle.

The amplitude of the total-column mean annual cycle was also underestimated in CESM (Figs. 5a–d). In the Northern Hemisphere midlatitudes, the seasonal cycle in the column was smaller than observations by 40%–60% (Table 2) and, like the surface data, the seasonal minimum was phase shifted early. Because the seasonal amplitude of column measurements is representative of hemispheric-scale fluxes (Keppel-Aleks et al. 2011), we expected the amplitude at different sites to be coherent and for the total underestimation to reflect the amount by which integrated Northern Hemisphere net ecosystem exchange was underestimated by CLM (Keppel-Aleks et al. 2012). These results indicated that the Northern Hemisphere growing season net flux was too small by around 50% in
CESM and may be related to the underestimation of soil carbon stocks, as discussed in section 5. Low biases in the column-mean annual cycle persisted in the tropics and in Southern Hemisphere midlatitudes, with CESM underestimating the observations by 30% and 40%, respectively (Figs. 5e–g).

c. North–south gradients

The contemporary north–south gradient is determined primarily by the fossil component of CO₂ and annual-mean land and ocean carbon sinks (Tans et al. 1990). In 1850, the north–south gradient in CESM was nearly flat; the mean contrast between Mauna Loa and the South Pole for 1850–60 was 0.05 ± 0.06 ppm standard error (Fig. 6a). The prominent peak in the tropics in the model zonal mean was due to the diurnal covariance of net ecosystem exchange and vertical mixing; at nighttime vertical mixing is suppressed and CO₂ from heterotrophic respiration is trapped within the boundary layer (e.g., Fan et al. 1990; Denning et al. 1995). As a result, the mole fractions in the atmospheric boundary layer above tropical ecosystems were large in the annual mean. If only daytime CO₂ data were averaged, then the peak in tropical latitudes was not observed (Fan et al. 1990). Likewise, the tropical peak was not observed in the zonal mean for the CO₂ total column since this quantity is mostly insensitive to the vertical distribution of CO₂.

There was only a gradual change in the north–south gradient between 1850 and the beginning of the atmospheric observational record in the 1950s. During 1950–60, the Mauna Loa–South Pole contrast was 1.2 ± 0.1 ppm (Fig. 6a). The growth of the gradient accelerated between 1950 and 2000 such that between 2000 and 2005, the CO₂ contrast between Mauna Loa and South Pole was 3.1 ± 0.1 ppm (standard error) in CESM, in general agreement with the observed contrast of 2.9 ± 0.1 ppm. This growth in the north–south gradient over

### Table 4. Mean surface seasonal cycle amplitudes in NOAA GMD flask data and CESM for eight north–south bins.

<table>
<thead>
<tr>
<th>Region</th>
<th>GMD amplitude* (ppm)</th>
<th>CESM amplitude** (ppm)</th>
<th>Difference (%)</th>
</tr>
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<tbody>
<tr>
<td>60°–90°N</td>
<td>14.7 ± 1.4</td>
<td>11.3 ± 2.6</td>
<td>−23</td>
</tr>
<tr>
<td>40°–60°N</td>
<td>14.0 ± 2.9</td>
<td>8.2 ± 3.2</td>
<td>−41</td>
</tr>
<tr>
<td>15°–40°N</td>
<td>9.4 ± 1.9</td>
<td>5.5 ± 1.6</td>
<td>−41</td>
</tr>
<tr>
<td>0°–15°N</td>
<td>5.1 ± 2.0</td>
<td>3.2 ± 0.2</td>
<td>−37</td>
</tr>
<tr>
<td>0°–15°S</td>
<td>1.8 ± 0.7</td>
<td>2.2 ± 0.8</td>
<td>22</td>
</tr>
<tr>
<td>15°–40°S</td>
<td>1.1 ± 0.3</td>
<td>0.9 ± 0.4</td>
<td>−18</td>
</tr>
<tr>
<td>40°–60°S</td>
<td>1.2 ± 0.3</td>
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<td>50</td>
</tr>
<tr>
<td>60°–90°S</td>
<td>1.3 ± 0.2</td>
<td>1.9 ± 0.7</td>
<td>46</td>
</tr>
</tbody>
</table>

* Average mean annual cycle amplitude for GMD flask sites located within the latitude band.
** Average mean annual cycle sampled at the same GMD flask sites within CESM.
the historical simulations was due almost entirely to the fact that most fossil fuel emissions occurred in the Northern Hemisphere, therefore driving a large pole-to-pole gradient (Fig. 6b). The ocean and land contributions to the mean north–south gradient in the model were relatively small and changed by only a small amount between 1850 and 2000 (Figs. 6c,d).

As atmospheric CO₂ monitoring began in 1958 at only a few locations and more extensive monitoring began in both hemispheres only in the mid-1970s, we could not directly evaluate long-term changes in the north–south gradient within the model. Previous studies (e.g., Heimann and Keeling 1986; Conway and Tans 1999) have extrapolated the contrast between Mauna Loa and the South Pole to zero emissions to investigate natural CO₂ processes that may have existed before the perturbation from fossil fuel CO₂. Although interpretation of this approach can be challenging, it was instructive to analyze the Mauna Loa–South Pole contrast for CESM as it revealed a significantly different pattern than the observations (Fig. 7). A linear regression through the NOAA

FIG. 4. Phasing of the seasonal cycle in atmospheric CO₂ at NOAA GMD flask sites. Solid lines represent observations and dashed lines represent CESM data. In the Northern Hemisphere, the annual maximum mole fraction (blue curves) occurs too early, particularly in the zone from 60° to 90°N. The first zero crossing (i.e., when CO₂ transitions from above to below the annual mean value) occurs too early by 1 month throughout the Northern Hemisphere extratropics and midlatitudes (red). The annual minimum (green) and second zero crossing (cyan) are better simulated. In the tropics, the summer drawdown in CESM lags the observations. In the Southern Hemisphere, the phasing of the seasonal cycle reverses in the midlatitudes, with the annual maximum occurring after the annual minimum. Zero crossings in the Southern Hemisphere are not plotted because they are not clearly defined.

FIG. 5. Mean annual cycle in column CO₂ sampled at seven TCCON sites with a full year of data compared to CESM sampled at the same locations: (a) Bialystok, (b) Park Falls, (c) Lamont, (d) Pasadena, (e) Darwin, (f) Wollongong, and (g) Lauder. The y axis scale has been changed for the Southern Hemisphere sites [panels (e)–(g)].
GMD data had a slope of 0.56 ± 0.08 ppm (Pg C yr\(^{-1}\)) and an intercept of 1.2 ± 0.4 ppm. A simple interpretation might suggest that, in the absence of fossil fluxes, the Northern Hemisphere would have lower CO\(_2\) than the Southern Hemisphere due to a natural Southern Hemisphere source. The CESM results give a slope of 0.39 ± 0.04 ppm (Pg C yr\(^{-1}\)) and a positive intercept of 0.3 ± 0.2 ppm for the observational period from 1958 and 2005. For the full CESM record (1850–2005), the slope was similar 0.42 ± 0.02 ppm (Pg C yr\(^{-1}\)), and the actual difference prior to fossil fuel emissions was 0.2 ± 0.1 ppm. The positive intercept for the recent data in CESM may reflect a lack of Southern Ocean overturning that would limit ventilation of CO\(_2\)-rich waters in the far south (Long et al. 2013). We hypothesize that models with weak Southern Ocean outgassing also may have lower rates of twentieth-century anthropogenic carbon uptake owing to the role of Southern Ocean overturning on both time scales (e.g., Lovenduski et al. 2008; Khatiwala et al. 2009; Lovenduski and Ito 2009). Thus, the model biases in CO\(_2\) growth (Fig. 2) and emissions versus interhemispheric gradient (Fig. 7) may be related.

As with many of the metrics presented in this paper, the observed CO\(_2\) gradients often are a result of multiple factors and thus are only a partial constraint on a single process. Errors in the magnitude, spatial distribution, and timing of land use change emissions, for example, could also affect the trend in the Mauna Loa–South Pole contrast. The preindustrial north–south gradient inferred from the observations is also probably influenced by land–ocean lateral carbon fluxes (e.g., Finzi et al. 2011) that are accompanied by a more uniform breakdown within the ocean of more refractory terrestrial carbon, although inverse modeling results show that these fluxes are small (Jacobson et al. 2007), and by rates of interhemispheric atmospheric mixing. Diagnosing these contributions would require additional simulations and are beyond the scope of this study.

The north–south CO\(_2\) gradient varies considerably as a function of season (Fig. 8). It is highest in late winter when fossil fuels and ecosystem respiration fluxes are the primary land surface fluxes and reverses during summer when photosynthetic drawdown exceeds ecosystem respiration and fossil emissions (Fig. 3). The meridional gradient at the surface in CESM did reasonably well at representing the annual mean of the observations (Fig. 8a). During February, north of 40°, the model was higher than GMD sites by 0.6 ± 0.6 ppm (standard error across 14 sites, Fig. 8b), while in the summer, the model was 6.0 ± 0.6 ppm (standard error across 14 sites) higher than the Global Monitoring Division observations north of 40° (Fig. 8c). The shallow Northern Hemisphere growing season gradient was consistent with underestimation of Northern Hemisphere CO\(_2\) uptake during summer and a correspondingly weak mean annual cycle (Fig. 3). Although the modeled surface CO\(_2\) mole fraction was high in both winter and summer, the model agreed in the annual mean north of 40°N because CESM had excessively large drawdown in spring that lowered the annual mean value.
The north–south CO₂ gradient was likewise underestimated in the column in CESM during Northern Hemisphere summer (Fig. 9). In the annual mean (Fig. 9a), the model and data generally agreed, while in February the CESM was slightly lower than observations in the Northern Hemisphere. In August the CESM was higher than observations throughout the Northern Hemisphere, with an increasing positive bias in northern regions. The two high-latitude sites, Sodankyla and Eureka, operated only during the summer when simulated column CO₂ was 4 ppm higher than the observations.

A comparison of CESM with HIPPO transects revealed how north–south gradients vary with altitude across seasons. For the surface layer, the north–south gradient was overestimated in the model during January. During March and April (Figs. 10b,c), Northern Hemisphere CESM data were lower than HIPPO data, perhaps due to early spring drawdown in CESM. Although initial drawdown was evident along HIPPO flight tracks in June, in CESM this was significantly more advanced for all latitudes and altitudes in the tropics and Northern Hemisphere (Fig. 10d). From June to July (Fig. 10e) the HIPPO data indicated carbon uptake increased in the northern part of the Northern Hemisphere, whereas the simulations showed no additional drawdown. The weak summer
drawdown in CESM was most apparent in August and September (Figs. 10f,g). The model showed surface CO₂ mole fractions higher by 5 ppm and free tropospheric CO₂ mole fractions higher by 2–5 ppm. In autumn CO₂ was generally too high along HIPPO transects (Figs. 10h,i). The pole-to-pole north–south gradients were smaller aloft than at the surface in both observations and simulations for all months, with the magnitude of the north–south gradient aloft tied to the strength of the gradient at the surface. For all of the comparisons shown in Fig. 10, the CESM estimates were a 5-yr mean sampled along the flight tracks to approximate climatological mean patterns. It is important to note that the October and November HIPPO data described above were obtained during 2009 during the transition to a strong La Niña, but this likely has a small effect on the model–data mismatch compared to the weak seasonal exchange in CESM.

d. Vertical structure in the model

The agreement in vertical CO₂ stratification between CESM and HIPPO observations varied considerably as a function of season, with summer CESM data generally less vertically stratified than the HIPPO free tropospheric CO₂ (Fig. 10). In January (Fig. 10a) CESM CO₂ showed vertical gradients of 4 ppm between the surface and 10 km in the Northern Hemisphere, roughly similar to those in the observations. Coincident with growing season onset in March and April (Figs. 10b,c), the vertical gradients in CESM decreased such that, during the peak of the growing season, vertical gradients were quite small. During June and July (Figs. 10d,e) both datasets showed small vertical gradients of 1–2 ppm between the surface and 10 km. In August (Fig. 10f) HIPPO data increased by greater than 5 ppm between the surface and 10 km in regions poleward of 40°N. In CESM, however, the vertical contrast over this altitude range was only 2 ppm. This model–data mismatch persisted into September.

Analysis of the vertical propagation of the annual cycle in NOAA and DOE southern Great Plains (SGP) aircraft data indicated that annual-mean vertical gradients in CESM were broadly consistent with those observed in the Northern Hemisphere (Fig. 11). The mean annual cycle amplitudes in 1-km altitude bins throughout the lower troposphere were normalized to the amplitude of the mean annual cycle below 1 km in an attempt to account for biases in the magnitude of seasonal surface fluxes in the model (e.g., Yang et al. 2007). Between 20° and 40°N the mean annual cycle amplitude decreased somewhat more rapidly with altitude in CESM than it did in the observations (Fig. 11a), which may suggest weak vertical coupling between the surface layer and the free troposphere (Yang et al. 2007), although the seasonal amplitude aloft also may have been diminished by the propagation of fluxes from northern regions, which are demonstrably too weak in CESM. North of 40°N (Figs. 11b,c) the rate of decrease in the mean amplitude was similar for CESM and the observations, although the site-to-site behavior was quite heterogeneous.

Consistent with the HIPPO profiles, CESM CO₂ showed more vertical stratification at the GMD and DOE aircraft sites during winter than summer (Fig. 12). The observed summer profiles varied by 6 ppm from the surface to 10 km, but were mostly uniform in CESM, except for a small enhancement in CESM below 1 km. This surface enhancement was not present at all sampling
locations and may simply be a consequence of the diurnal rectifier. We were not able to test this because CESM data were not archived at a temporal resolution higher than 1 day for most of the atmosphere (for all atmospheric levels above the surface layer).

The somewhat sluggish rate of atmospheric vertical mixing (suggested by Fig. 11a) may explain why the seasonal cycle amplitudes agreed better at the surface in the Northern Hemisphere (Table 4) than in the column (Table 2). The weak vertical gradients during Northern Hemisphere summer in CESM compared to both HIPPO and NOAA aircraft profiles (Figs. 10, 12) may be mostly caused by weak surface fluxes. Similarly, in winter the reasonable agreement between model and observations

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**Fig. 10.** Free tropospheric CO$_2$ measured during the HIPPO campaigns (black) and sampled in CESM at corresponding locations (red) for the data averaged into eight latitude bands and four altitude bins: the surface to 2.5 km (circles), 2.5–5.0 km (squares), 5.0–7.5 km (triangles), and 7.5–10.0 km (diamonds). Each panel represents a different northbound or southbound campaign in (a) January 2009 (southbound), (b) March 2010 (southbound), (c) April 2010 (northbound), (d) June 2011 (southbound), (e) July 2011 (northbound), (f) August 2011 (southbound), (g) September 2011 (northbound), (h) October 2009 (southbound), and (i) November 2009 (northbound).
may result from the combined effects of weak biosphere emissions outside the growing season and weak transport. A more conclusive diagnosis of transport, as discussed in section 5, will require additional simulations with new tracers. We note that, at some locations in the Northern Hemisphere, the surface mole fraction was higher in CESM during winter than in the NOAA and DOE observations (Fig. 8b) or HIPPO observations (Fig. 10a), even though surface fluxes were weaker. Any surface enhancement in CAM (e.g., the increase below 1 km in Fig. 12) also could have contributed to the mean annual cycle at different altitudes by normalizing the mean annual cycle amplitude in each layer by the amplitude for the corresponding data type (observation or model) in the layer centered at 0.5 km, similar to the methodology described by Yang et al. (2007), to account for differences in surface fluxes. The sites are separated by latitude for clarity in plotting.

e. East–west gradients

We present one example of growing east–west gradients created by increasing fossil fuel CO₂ at the NOAA GMD site on the Tae-Ahn peninsula in South Korea, where the air is transported from heavily populated regions in China and industrial centers in South Korea (Turnbull et al. 2011), relative to background stations in the Northern and Southern Hemispheres. These background stations included Mace Head, which samples relatively clean airflow from the Atlantic, Mauna Loa, and Samoa. We detrended each site relative to the South Pole; thus, each site retained the signature of the increasing north–south gradient (Fig. 6). Deseasonalized NOAA GMD data from the early 1990s to 2010 showed that the increase relative to the South Pole on the Tae-Ahn peninsula since 1992 was faster than the increase at...
Mace Head and Mauna Loa by about a factor of 3 (0.17 ppm yr\(^{-1}\) compared to 0.06 ppm yr\(^{-1}\)) (Table 5), and that the trends were correlated with Chinese emissions (Fig. 13a). This pattern across the different stations was consistent with observations that China and Korea emissions increased at a rate that was several fold larger than the hemispheric mean during this interval and that the Tae-Ahn peninsula site was in close proximity to these source regions (Boden et al. 2009; Turnbull et al. 2011). At all three Northern Hemisphere sites, CO\(_2\) increased faster than at Samoa, due at least in part to the fact that the north–south gradient is increasing with time (Fig. 6). CESM output for the same time period revealed a similar trend at Tae-Ahn compared to the observations. The fossil fuel component of CO\(_2\) was primarily responsible for the higher increase at the site relative to the other Northern Hemisphere sites, although we did not carry separate tracers for China and Korea (or other countries) to enable further attribution (Table 5). The growth rate trends were slightly smaller at Mace Head and Mauna Loa than in the observations, but the fossil component of the trend was similar in magnitude to the observed total at each site. Given that CESM was able to capture the faster accumulation rate of CO\(_2\) at the Tae-Ahn peninsula relative to more remote stations (Table 5) and that this gradient is likely to increase many fold in the future with greater energy consumption in China (Figs. 13b,c), there is potential to use the existing monitoring network to constrain regional increases in fossil emissions over time (National Research Council 2010).

**f. Trends in the seasonal cycle**

Analysis of trends in the mean annual cycle of atmospheric CO\(_2\) during the contemporary period may reveal how CO\(_2\) varies in response to contemporary climatic variations and changes in atmospheric composition (Table 6). Previous studies have noted changes in the seasonality of CO\(_2\) (e.g., C. D. Keeling et al. 1996; Randerson et al. 1997; Piao et al. 2008) owing mostly to increased seasonality in land biosphere fluxes. We calculated the trend in the seasonal cycle amplitude using Eq. (1) at 17 NOAA GMD sites that had mostly continuous coverage during 1980–2005 and sampled the model for the corresponding period. The agreement in the calculated trends varied considerably among sites. Both observations and the CESM showed consistent increases of around 0.5% yr\(^{-1}\) to the mean annual cycle amplitude at two high northern latitude sites, Alert and Barrow. In addition we found that in CESM there were negative trends on the shoulders of the growing season at these sites, suggesting lengthening of the growing season that was not evident in the observed trends. At mid and low latitudes of the Northern Hemisphere between 0° and 40°N, the observations generally had smaller positive trends. Within this zone, CESM generally overestimated the observed amplitude increases. Randerson et al. (1997) noted that the role of transport acting on gradients in fossil fuel CO\(_2\) plays a significant role in setting the seasonal cycle at stations near the equator in both Northern and Southern Hemispheres. As a result, increases in the north–south gradient from increasing fossil fuel emissions over this 25-yr analysis...
period likely contributed to some of the average positive trends observed in the NOAA GMD measurements and in CESM.

g. Interannual variability

Interannual variations in CESM were smaller than those in GMD flask data. Table 6 lists the IAV in the CO2 time series for the same 17 sites discussed above from 1980 to 2005. CESM IAV was approximately 50% that of observed CO2 IAV during this time period. The interannual variability (IAV) was somewhat larger in the Northern than in the Southern Hemisphere in the data, whereas the two hemispheres had similar IAV in the model.

Interannual variability in CO2 remained relatively constant during the historical simulation (Figs. 14a,b). We took advantage of the fact that land, ocean, and fossil components of atmospheric CO2 were tagged separately in CESM to explore the sources of interannual variability in each latitude band (Fig. 14). In CESM, land CO2 was the primary source of interannual variability from 1850 to 2005, even in the Southern Hemisphere. The IAV in the fossil component of CO2 grew during the twentieth century in all latitude bands, but the increase was largest in the Northern Hemisphere between 15° and 60°N where most fossil fuel CO2 was emitted. The fossil component of CO2 generally showed larger IAV than the ocean component from 1850 to 2005, except in the tropical bands between 15°S and 15°N.

The sum of the IAV for the fossil, land, and ocean components was greater than the IAV for the total CO2, suggesting that variations in one component were compensated by the other components. An asynchronous phasing of land and ocean fluxes to ENSO variability in the model (Lindsay et al. 2012, manuscript submitted to J. Climate) is probably a driver of some of this compensation, which was also noted in Doney et al. (2006). Assuming that the fossil fuel component of the IAV was well captured by CESM for the historical simulations (i.e., source emissions were approximately correct and atmospheric transport was not a large error source on these longer time scales), these results indicated that interannual variability owing to terrestrial biosphere and ocean fluxes, in particular interannual variability in the land biosphere, was too small in CESM given the underestimate of IAV at GMD sites (Fig. 6). The implementation of nitrogen coupling in the C–N biogeochemistry model may be partly responsible for reducing the magnitude of net primary production (Thornton et al. 2007) and, thus, the contribution of this gross flux component to net ecosystem exchange variations driven by year-to-year changes in climate. Interannual variability in fires from the model was also lower.
than estimates derived from satellite observations of burned area and active fires (van der Werf et al. 2010; Kloster et al. 2010).

4. Evolution of CO$_2$ variability in RCP scenarios

In the representative concentration pathways RCP4.5 scenario, global emissions increased from 8 Pg C yr$^{-1}$ in 2005 to 9.0 Pg C yr$^{-1}$ in 2040 before declining to around 4 Pg C yr$^{-1}$ in 2088 and leveling off (Fig. 15a), resulting in an atmospheric CO$_2$ mole fraction of 610 ppm at Mauna Loa in 2100 (Fig. 15b). In contrast, emissions increased to almost 30 Pg C yr$^{-1}$ by the end of the century in the RCP8.5 scenario, resulting in a mole fraction of 1150 ppm (Figs. 15a,b). Between 2005 and 2100 global mean surface air temperature increased by 1.7 K in RCP4.5 and by 4.1 K in RCP8.5 (Fig. 15c). Increasing fossil fuel emissions and climatic changes led to changing patterns in fluxes with GPP, net primary production (NPP), and heterotrophic respiration increasing during the twenty-first century in both simulations (Figs. 15d,e), albeit at a reduced rate in the RCP4.5 simulation. These changes in ecosystem fluxes, as well as the prescribed reductions in land use change for RCP4.5, led to a net ecosystem carbon balance (NECB) lower than the contemporary value in RCP4.5, while the NECB increased in the RCP8.5 scenario (Fig. 15f). We discuss the effect of these changing fluxes on atmospheric CO$_2$ in the following section.

a. Twenty-first-century seasonal cycles

The mean annual cycle in future atmospheric CO$_2$ (2065–95) increased in both CESM RCP simulations relative to the contemporary period (1975–2005) (Fig. 16). At each of the sites in Fig. 16, the amplitude of the mean annual cycle increased in both scenarios relative to the contemporary period, although the changes were larger in RCP8.5.

The growth in the mean annual cycle had contributions from fossil, ocean, and land components. At all four sites, the fossil fuel contribution to seasonality was negligible in the historical simulation from 1975 to 2005 and in the RCP4.5 scenario where, from 2065 to 2095, mean fossil fuel emissions were actually lower than contemporary emissions. In contrast, fossil fuel emissions contributed significantly to CO$_2$ seasonality in the RCP8.5 scenario for 2065–95 (Figs. 16e–h). We caution that the seasonality of fossil fuel emissions in the RCP scenarios was derived from late-twentieth-century seasonal patterns scaled by the change in annual mean emissions, which resulted in increasing seasonality of emissions throughout the RCP8.5 scenario. In reality, energy use might show a smaller seasonal cycle in a warmer climate as, for example, winter heating burdens decrease and air conditioning demands increase during summer (Gregg et al. 2009).

The change in the mean annual cycle owing to ocean exchange was larger at high-latitude sites than at low-latitude sites (Figs. 16i–l). The ocean CO$_2$ amplitude at both Barrow and the South Pole increased, and the phasing shifted toward earlier maxima and minima.

The terrestrial mean annual cycle amplitude increased in the Northern Hemisphere in both RCP simulations. The positive high-latitude trend in CESM during 1975–2005 (Table 6) remained steady during the RCP4.5 scenario. The trend in the seasonal amplitude at Barrow

<table>
<thead>
<tr>
<th>Site</th>
<th>Lat</th>
<th>Period</th>
<th>GMD trend (% yr$^{-1}$)</th>
<th>CESM trend (% yr$^{-1}$)</th>
<th>GMD IAV (ppm)</th>
<th>CESM IAV (ppm)</th>
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<tr>
<td>Alert</td>
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<td>1980–2005</td>
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<td>0.5</td>
<td>0.8</td>
<td>0.4</td>
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<tr>
<td>Cold Bay</td>
<td>55.2</td>
<td>1980–2005</td>
<td>0.2</td>
<td>0.7</td>
<td>0.9</td>
<td>0.4</td>
</tr>
<tr>
<td>Shemya</td>
<td>52.7</td>
<td>1985–2005</td>
<td>0.0</td>
<td>0.3</td>
<td>0.7</td>
<td>0.4</td>
</tr>
<tr>
<td>Niwot Ridge</td>
<td>40.1</td>
<td>1980–2005</td>
<td>−0.1</td>
<td>2.0</td>
<td>0.7</td>
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<tr>
<td>Azores</td>
<td>38.8</td>
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<td>0.0</td>
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<tr>
<td>Midway Island</td>
<td>28.2</td>
<td>1985–2005</td>
<td>0.1</td>
<td>0.7</td>
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<td>0.4</td>
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<td>Key Biscayne</td>
<td>25.7</td>
<td>1985–2005</td>
<td>0.2</td>
<td>0.2</td>
<td>0.8</td>
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<td>Mauna Loa</td>
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<td>0.3</td>
<td>0.7</td>
<td>0.3</td>
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<td>Kumukahi</td>
<td>19.5</td>
<td>1980–2005</td>
<td>−0.1</td>
<td>0.3</td>
<td>0.6</td>
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<tr>
<td>Guam</td>
<td>13.4</td>
<td>1980–2005</td>
<td>−0.1</td>
<td>0.2</td>
<td>0.8</td>
<td>0.3</td>
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<td>1.0</td>
<td>2.3</td>
<td>0.9</td>
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<td>Ascension Island</td>
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<td>1980–2005</td>
<td>1.1</td>
<td>−0.5</td>
<td>0.6</td>
<td>0.4</td>
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<td>Christmas Island</td>
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<td>Samoa</td>
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<td>0.3</td>
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<td>Palmer Station</td>
<td>−64.9</td>
<td>1980–2005</td>
<td>−0.7</td>
<td>1.4</td>
<td>0.7</td>
<td>0.3</td>
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<tr>
<td>South Pole</td>
<td>−90.0</td>
<td>1980–2005</td>
<td>−1.0</td>
<td>1.3</td>
<td>0.6</td>
<td>0.3</td>
</tr>
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</table>

Table 6. Amplitude trends in the mean annual cycle and interannual variability (IAV). GMD flask sites were selected that have nearly continuous data coverage from 1980 to 2005 and sampled CESM coincidentally. The amplitude trend is calculated from Eq. (1). Interannual variability is defined as the standard deviation in bandpass-filtered (2–10-yr filter cutoffs) time series.
accelerated in RCP8.5, and the spring drawdown moved earlier in RCP8.5 relative to RCP4.5 (Fig. 16m). These changes in the seasonality in CO2 were due to an increase in net primary production during spring and summer at mid to high northern latitudes and a decrease in the seasonality of heterotrophic respiration. At Mauna Loa, the mean annual cycle amplitude of land CO2 increased by 28% from the contemporary period (1975–2005) to the end of the twenty-first century (2065–95) in the RCP8.5 scenario, in response to both land and fossil fuel influences (Fig. 16n). Gross primary production in CLM increased during summer by about 30%, and the growing season onset occurred earlier in the spring at high northern latitudes. Heterotrophic respiration, which has a strong dependence on temperature, likewise increased in the annual mean, although its seasonality decreased throughout the twenty-first century in the RCP8.5 scenario, probably as a result of proportionally larger temperature increases during winter and spring. In the Southern Hemisphere, trends in the terrestrial contribution to the mean annual cycle were relatively small (Figs. 16o,p).

b. Evolution of the north–south gradient

The north–south gradient in CO2 continued to increase during the twenty-first century while fossil fuel emissions increased. The Mauna Loa–South Pole contrast grew from 3.2 ppm in 2000 to 5.2 ppm in 2050 in RCP4.5 but declined in the second half of the twenty-first century as emissions fell below the contemporary levels (Fig. 6a). In RCP8.5, the CO2 contrast between Mauna Loa and the South Pole grew to 14.2 ppm in 2100 (Fig. 6a).

Changes in atmosphere–ocean exchange also impacted the north–south gradient in the RCP simulations.

Fig. 14. Interannual variability (IAV) in surface CESM CO2 for eight latitude bands in the (left) Northern Hemisphere and (right) Southern Hemisphere: (a),(b) IAV for the total CO2 in CESM, (c),(d) IAV for the fossil CO2 component, (e),(f) IAV for the land component of CO2, and (g),(h) IAV for the ocean component of CO2. The solid symbols show IAV from RCP8.5 and the open symbols show IAV from RCP4.5.
The ocean component of CO$_2$ decreased relative to the global mean between 40° and 70°S (Fig. 6c), consistent with a continued Southern Ocean sink for anthropogenic CO$_2$, enhancing the total north–south gradient. Deep convective mixing and marine export production in the high-latitude North Atlantic were greatly reduced by the end of the century (Moore et al. 2012, manuscript submitted to J. Climate), weakening the ocean uptake of atmospheric CO$_2$ in the Northern Hemisphere. The annual mean gradient owing to land fluxes showed the smallest changes in the twenty-first century (Fig. 6d), with increases smaller than 1 ppm in the Northern Hemisphere midlatitudes at the end of the twenty-first century in RCP8.5.

c. East–west gradients

Monitoring east–west gradients in CO$_2$ within the Northern or Southern Hemisphere may become increasingly important in the future for verification of CO$_2$ emissions agreements. In this section, we discuss changes in both surface and column CO$_2$ in relation to changing surface fluxes between the contemporary period (1975–2005) and the end of the RCP8.5 scenario (2065–95). For contemporary conditions, the most noticeable signal in east–west gradients in surface CO$_2$ was the CO$_2$ enhancement over tropical land relative to the oceans due to increases in photosynthesis and nighttime trapping of respired CO$_2$ in a stable boundary layer (Fig. 17a). Enhancements in surface CO$_2$ also occurred over the industrialized regions of Europe, East Asia, and eastern North America. Notably, column CO$_2$ changes were negligible in the tropics (Fig. 17d), while the east–west gradients caused by fossil fuel emissions in the Northern Hemisphere were still apparent.

East–west gradients at the surface increased by the end of the twenty-first century in the RCP8.5 scenario in northern midlatitudes and in the tropics (Figs. 17b,c). In northern midlatitudes, much of the increase in land–ocean CO$_2$ differences by 2100 appeared to be linked with increases in fossil fuel emissions, as described above in section 3e. In the tropics, in contrast, enhancements in surface CO$_2$ over forests were not matched by column CO$_2$ increases (Figs. 17c,d), suggesting that the increases at the surface were due primarily to the rectifier effect operating on larger gross fluxes (Figs. 15d,e) rather than changing net fluxes (Chapin et al. 2006). We examined GPP, ecosystem respiration, and net land-use change fluxes and found that GPP fluxes increased by about 25% throughout the tropics. The total carbon loss from land use change in CESM decreased by about 40% for the Amazon at the end of the twenty-first century relative to 1975–2005 in RCP8.5, also confirming that the changes in tropical surface mole fractions were due to
rectifier effects. For tropical Africa the integrated carbon loss due to land use change was higher by a factor of 6 compared to the period from 1975 to 2005 but still constituted a small enough source that it did not lead to observable east–west gradients in the column.

Fossil fuel emissions increased significantly in the RCP8.5 scenario and were a primary driver of the changes in the east–west column CO₂ gradients in northern mid-latitudes. The east–west gradients in the column increased considerably from China to the mid Pacific over the twenty-first century. Specifically, column differences between the NOAA Tae-Ahn peninsula site (36°N, 126°E) and the mid-Pacific (36°N, 155°W) increased from 0.5 ppm during 1975–2005 to 2.6 ppm during 2065–95.

We also quantified the trend in the surface enhancement relative to SPO at Tae-Ahn and the three background sites explored in section 3e with respect to fossil fuel emissions from China for the two emission scenarios (Figs. 13b,c). All four sites showed a linear trend with respect to Chinese emissions, even while emissions decreased and the difference in CO₂ at each site relative to the South Pole declined at the end of the RCP4.5 simulation. The linear regression slope of the Tae-Ahn to SPO CO₂ gradient versus Chinese emissions was 5.0 ± 0.5 ppm (Pg C yr⁻¹) higher than the slopes computed from neighboring sites (at similar latitudes) in both simulations. The relative slopes were similar in both RCP4.5 and RCP8.5 because emissions from China accounted for about 25% of global emissions in both scenarios over the twenty-first century, compared to about 20% in 2005. China’s contribution to global emissions increased rapidly at the beginning of the RCP4.5 scenario and peaked.

FIG. 16. Simulated mean annual cycle in atmospheric CO₂ at four GMD sites: Barrow (BRW), Mauna Loa (MLO), Samoa (SMO), and the South Pole (SPO) for 1975–2005 (historical simulation) and the later part of the twenty-first century (RCP4.5 and RCP8.5). The seasonal cycle in total CO₂ is separated into its fossil, ocean, and land components. Note the change of scale for the y axis for each site.
at 30% in 2050. During the latter half of the twenty-first
century, Chinese emissions and the Tae-Ahn to SPO CO$_2$
gradient decreased in RCP4.5 but remained on the
same trend line as for the first half of the century, shown
in Fig. 13b.

The interannual variability in the growth rate at Tae-
Ahn also was greater than at the northern midlatitude
background stations, as shown in Fig. 13, likely because
of this station’s proximity to source regions and, thus, for
year to year differences in meteorology to modify large
CO$_2$ spatial gradients (Turnbull et al. 2011). This high
level of variability may complicate the use of atmospheric
observations to infer surface fluxes (National Research
Council 2010). We note that RCP4.5 and RCP8.5 repre-
sented only two possible trajectories for atmospheric
CO$_2$ and that simulated spatial gradients were sensitive
to the prescribed distribution of emissions across dif-
ferent countries and regions. Further examination of the
differences in east–west gradients in atmospheric CO$_2$—
both at the surface and through the free troposphere—
between scenarios with similar emissions but different
geographic distributions is needed to determine at what
point the current observational network can detect
changes in regional fossil fuel emissions and to determine
how best to improve the observational network such that
emissions from China, for example, can be distinguished
from emissions originating from other countries within
East Asia.

d. Interannual variability

In the RCP4.5 scenario, interannual variability (IAV)
did not significantly change during the twenty-first cen-
tury in either the Northern or Southern Hemispheres
(Fig. 14). In contrast, Northern Hemisphere IAV in CO$_2$
increased throughout the RCP8.5 scenario, driven by
increasing variability in fossil fuel and land fluxes. In
general, increasing interannual variability can be caused
by increases in the variability of atmospheric circulation,
increased variability in surface fluxes, or larger CO$_2$ gra-
dients (likely from increases in fossil fuel emissions) in-
teracting with atmospheric mixing processes that may or
may not change significantly with climate warming. In
the case of fossil fuel IAV, it is likely that some of the
increase was caused by transport acting on larger at-
mospheric gradients since the prescribed fossil fuel
emissions were smooth in time. In the case of land CO$_2$,
all three of the mechanisms described above may have
contributed to the positive trend. Globally integrated
terrestrial GPP increased during the twenty-first century
in both scenarios (Fig. 15d), as did the Northern Hemi-
sphere growing season length; in response, periods of
drought and temperature stress induced larger year to
find that ocean anthropogenic carbon inventory in CCSM-3 (the predecessor to CESM1-BGC) is low by 20%.

Weak overturning of water masses in the Southern Ocean likely explain both the small ocean sink in CESM and the mismatch in the inferred preindustrial interhemispheric CO₂ gradient documented here. Long et al. (2013) investigate the uptake of atmospheric CO₂ by the world’s oceans and find that, while fluxes of CO₂ into the Southern Ocean are larger than observed, the inventory of anthropogenic CO₂ is too low due to weak formation rates of Antarctic Intermediate Water and Subantarctic Mode Water. This result is corroborated by Weijer et al. (2012), who show that CFC enhancements do not penetrate below the mixed layer as much as indicated by observations. Weak exchange between the surface and deeper water masses subsequently results in low sequestration of anthropogenic CO₂ and reduced outgassing of natural CO₂ from the deeper water masses. Analysis of changing north–south contrasts in atmospheric CO₂ and Δ¹³C suggest that the stationary gradient in atmospheric CO₂ is determined mostly by ocean fluxes, corroborating our result (Keeling et al. 2011). Long et al. (2013), however, also show that the ocean anthropogenic carbon inventory is low elsewhere, including the North Atlantic, which may be due to bias in the formation of mode and intermediate waters.

Land fluxes in CESM contributed to discrepancies between atmospheric CO₂ in CESM and observations on shorter time scales, with implications for sources and sinks and the sensitivity of these carbon fluxes to climate change. Weak mean annual cycle amplitudes in atmospheric CO₂ throughout the Northern Hemisphere and weak equator-to-pole CO₂ gradients during the Northern Hemisphere growing season provided evidence that the growing season net flux is too small in CESM. This phenomenon was noted in an analysis of TCCON total column data for other land models, and atmospheric tracer simulations with net ecosystem production flux perturbations show that weak growing season fluxes in boreal ecosystems are the cause (Keppel-Aleks et al. 2012). From the mass constraints provided by column observations, we infer that the CESM growing season net flux in the Northern Hemisphere was too small by 40%–60%, with much of this deficiency in boreal or temperate forest ecosystems. Part of this discrepancy could be due to biases in the climate driving the CESM land fluxes: P. J. Lawrence et al. (2012) show that the simulated seasonal cycle in temperature in high-latitude regions generally agrees with observations, although precipitation is slightly too high. Simulations with CLM driven by historical climate patterns are underway as part of the Trendy analysis project and will allow us to
quantify the effects of the CESM climate on the coupled system. Rapid turnover of soil carbon, consistent with simulated soil carbon inventory at high latitudes being only about 10% of observations (Todd-Brown et al. 2013), may also have contributed to the small seasonality since more rapid cycling of summer polar-orbiting NPP inputs within labile litter and soil pools could potentially offset simultaneous carbon gains from photosynthesis. Given that high latitudes are expected to experience amplified warming (e.g., Holland and Bitz 2003), it is crucial that coupled models such as CESM capture the mechanics of land fluxes within these ecosystems, including permafrost hydrology (D. M. Lawrence et al. 2012) and carbon processes (Koven et al. 2011).

The early onset of the Northern Hemisphere growing season potentially has important implications for the climate response of CESM to further warming during spring. P. J. Lawrence et al. (2012) show that seasonal patterns in land surface temperature are well represented in the Northern Hemisphere, suggesting that model deficiencies in the representation of phenology or cold hardening processes in conifers are more likely to be primary drivers than biases in climate. Richardson et al. (2012) found that many biogeochemical models simulate early onset to growing season because phenology is inaccurately parameterized. Since accurate representation of model phenology is crucial for partitioning fluxes of energy and for determining the sensitivity of vegetation to climate stressors (Richardson et al. 2012), we recommend further analysis and refinement of phenology algorithms in CLM.

The diagnostics that we have developed in this paper provide a means to test future generations of coupled carbon–climate models. Because most CO₂ observations, including those with the longest observational records, are made within the atmospheric boundary layer, most model evaluation has been conducted against this type of observation (e.g., Cadule et al. 2010). By extending our analysis to column and free tropospheric observations, we were able to more thoroughly examine the fluxes and atmospheric physics in CESM. We could not positively establish whether a vertical transport bias in CESM contributed to the weak vertical gradients in the model relative to aircraft observations because the mean annual cycle at the surface and at all layers was set by the interplay of fluxes (which were established by the comparison with column observations to be too weak) and transport. We suggest that more careful analysis of the vertical distribution of water vapor, carbon isotopes, and other regional carbon tracers in CAM would be useful for evaluating the vertical mixing parameterization on seasonal time scales, in addition to adding passive tracers such as radon or SF₆ (e.g., Denning et al. 1999).

The CESM simulations analyzed here indicated that patterns of variability in atmospheric CO₂ during the twenty-first century will likely depend on future climate forcing and CO₂ emissions. CESM simulated plausible increases in the mean annual cycle and north–south and east–west gradients during the twenty-first century, but the magnitude of these increases should be considered as having large uncertainties owing to the model’s bias toward low temporal variability in the historical simulation. CESM simulations of the twenty-first century showed the mean annual cycle amplitude increasing in all latitude bands, with the magnitude of the increase affected by climatic changes, emission rates, and atmospheric CO₂ levels. Although these simulations provided evidence that the shape of the mean annual cycle will continue to change, with a trend toward earlier drawdown of CO₂ in the spring, this result must be considered in light of the bias in the timing of the growing season onset that we documented for CESM during the contemporary period.

Although both RCP4.5 and RCP8.5 simulations show increased interannual variability in fossil CO₂ at all latitude bands, interannual variability in land and ocean CO₂ varied considerably as a function of latitude and between the two scenarios. Some of the increased variability may be due to atmospheric dynamics interacting with larger spatial gradients: small interannual shifts in the position of the jet stream, for instance, may change the origin of CO₂ sampled at a given observatory significantly. We propose more in-depth analysis to determine how both atmospheric dynamics and fluxes may change in the future so that monitoring networks can be optimized to discern such variability from interannual trends.

Changes in regional CO₂ patterns over source regions may be one of the fingerprints by which we can potentially monitor fossil fuel emissions (National Research Council 2010). Here we showed that temporal variability in atmospheric CO₂ increased as a function of fossil fuel emissions, with IAV changes in the RCP8.5 scenario outpacing those in the RCP4.5 scenario. As patterns in spatial and interannual variability change, interpretation of atmospheric observations may become more complicated and may require different observational strategies to extract information pertaining to fossil fuel emissions and terrestrial and ocean exchange. We note that the spatial coverage of CO₂ observations is expected to increase markedly in the coming decades as satellite missions using a variety of sensing methods and orbital geometries come online and as routine aircraft sampling of atmospheric CO₂ increases. Monitoring of CO₂ in urban areas also is likely to expand in the coming decades (Duren and Miller 2012). An expanded suite of
atmospheric tracer observations, including, for example, O$_2$:N$_2$ ratios, $^{13}$C and $^{14}$C isotopes, and coemitted species like CO and CH$_4$, will also be crucial for attributing changing patterns in atmospheric CO$_2$ to natural and anthropogenic causes.

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