

Response of the global surface ozone distribution to Northern Hemisphere sea surface temperature changes: implications for long-range transport

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Abstract. The response of surface ozone (O_3) concentrations to basin-scale warming and cooling of Northern Hemisphere oceans is investigated using the Community Earth System Model (CESM). Idealized, spatially uniform sea surface temperature (SST) anomalies of $\pm 1 \,^{\circ}$ C are individually superimposed onto the North Pacific, North Atlantic, and North Indian oceans. Our simulations suggest large seasonal and regional variability in surface O₃ in response to SST anomalies, especially in the boreal summer. The responses of surface O₃ associated with basin-scale SST warming and cooling have similar magnitude but are opposite in sign. Increasing the SST by 1 °C in one of the oceans generally decreases the surface O_3 concentrations from 1 to 5 ppbv. With fixed emissions, SST increases in a specific ocean basin in the Northern Hemisphere tend to increase the summertime surface O₃ concentrations over upwind regions, accompanied by a widespread reduction over downwind continents. We implement the integrated process rate (IPR) analysis in CESM and find that meteorological O₃ transport in response to SST changes is the key process causing surface O₃ perturbations in most cases. During the boreal summer, basin-scale SST warming facilitates the vertical transport of O₃ to the surface over upwind regions while significantly reducing the vertical transport over downwind continents. This process, as confirmed by tagged CO-like tracers, indicates a considerable suppression of intercontinental O₃ transport due to increased tropospheric stability at lower midlatitudes induced by SST changes. Conversely, the responses of chemical O₃

production to regional SST warming can exert positive effects on surface O_3 levels over highly polluted continents, except South Asia, where intensified cloud loading in response to North Indian SST warming depresses both the surface air temperature and solar radiation, and thus photochemical O_3 production. Our findings indicate a robust linkage between basin-scale SST variability and continental surface O_3 pollution, which should be considered in regional air quality management.

1 Introduction

High ground-level ozone (O₃) concentrations adversely impact human health by inducing respiratory diseases and threaten food security by lowering crop yields (Brown and Bowman, 2013; WHO, 2013; Chuwah et al., 2015). Considering the ecotoxicity of O₃, understanding the physical and chemical mechanisms that control atmospheric O₃ concentrations is of great importance. Surface O₃ is produced in the atmosphere via photochemical processing of multiple precursors including volatile organic compounds (VOCs), carbon monoxide (CO) and nitrogen oxides (NO, NO₂). These precursors originate from both natural and anthropogenic sources (Vingarzan, 2004; Simon et al., 2014; Jiang et al., 2016). In addition to local production, transport of O₃ and its precursors from upwind regions and the upper atmosphere can also influence surface O₃ abundance. Stratospheric intrusion events, which lead to vertical downmixing of ozonerich air, can significantly elevate surface O_3 during spring (Grewe, 2006; Lin et al., 2012b; Zhang et al., 2014). The long-range transport of O_3 and its precursors has been extensively studied, and their intercontinental impacts have been evaluated using measurements and model simulations (Parrish et al., 1993; Fehsenfeld et al., 1996; Wild and Akimoto, 2001; Creilson et al., 2003; Simmonds et al., 2004; Fiore et al., 2009; Brown-Steiner and Hess, 2011; Lin et al., 2012a, 2014).

Both photochemistry and dynamic transport collectively affect surface O₃ levels. Important meteorological factors that can impact both photochemistry and transport include atmospheric circulations, solar radiation, air temperature, and relative humidity. Atmospheric circulation considerably determines the timescale and pathway of O₃ transport (Bronnimann et al., 2000; Auvray and Bey, 2005; Hess and Mahowald, 2009). The efficiency of O₃ transport varies concurrently with atmospheric circulations on different scales. Knowland et al. (2015) demonstrated the important role of midlatitude storms in redistributing O₃ concentrations during springtime. The North Atlantic Oscillation (NAO) significantly affects surface and tropospheric O₃ concentrations over most of Europe by influencing the intercontinental transport of air masses (Creilson et al., 2003; Christoudias et al., 2012; Pausata et al., 2012). Lamarque and Hess (2004) indicated that the Arctic Oscillation (AO) can modulate springtime tropospheric O₃ burdens over North America. The shift in the position of the jet stream associated with climate change was found to strongly affect summertime surface O₃ variability over eastern North America (Barnes and Fiore, 2013). Increases in solar radiation and air temperature can increase the rate of the chemical production of O₃ and modulate the biogenic emissions of O₃ precursors (Guenther, 1993; Sillman and Samson, 1995; Peñuelas and Llusià, 2001), especially over highly polluted regions (Ordónez et al., 2005; Rasmussen et al., 2012; Pusede et al., 2015). Increases in humidity can enhance the chemical destruction of O₃ and shorten its atmospheric lifetime (Johnson et al., 1999; Camalier et al., 2007). Therefore, changes in meteorological conditions on various spatial and temporal scales play key roles in determining the surface O₃ distribution. Understanding the mechanisms and feedbacks of the interactions between O₃ and climate has received increasing attention and will be essential for future surface O3 mitigation (Jacob and Winner, 2009; Doherty et al., 2013).

Sea surface temperature (SST) is an important indicator that characterizes the state of the climate system. Its variations strongly perturb the mass and energy exchange between the ocean and atmosphere (Small et al., 2008; Gulev et al., 2013), which influence atmospheric circulation, atmospheric temperature, and specific humidity (Sutton and Hodson, 2005; Frankignoul and Sennéchael, 2007; Li et al., 2008) from regional to global scales (Glantz et al., 1991; Wang et al., 2000; Goswami et al., 2006). It also affects cloud formation and consequently influences incoming solar radiation (Deser et al., 1993; Fallmann et al., 2017). Numerous studies have shown that SST changes over different oceans and at different latitudes lead to significantly different meteorological and climate responses (Webster, 1981; Lau and Nath, 1994; Lau, 1997; Sutton and Hodson, 2007; Sabeerali et al., 2012; Ueda et al., 2015). Details on the SST–climate relationships over individual oceanic regions are summarized in Kushnir et al. (2002).

The Intergovernmental Panel on Climate Change Fifth Assessment Report (IPCC, 2013) provides strong evidences in Chapter 2 that global SSTs are generally increasing due to the impacts of anthropogenic forcings on global climate change. In addition, regional SST exhibits natural periodic or irregular oscillations with timescales ranging from months to decades. The El Niño-Southern Oscillation (ENSO) is the most influential natural SST variability that originates in the tropical Pacific and has worldwide climate impacts (Philander, 1983; Wang et al., 2012). The Pacific Decadal Oscillation (PDO), defined by ocean temperature anomalies in the northeast and tropical Pacific Ocean, is another long-lived, El Niño-like pattern that persists for several decades (Mantua and Hare, 2002). Over the Indian Ocean, SST anomalies feature a seesaw structure between the western and eastern equatorial regions, known as the Indian Ocean Dipole (IOD) mode (Saji et al., 1999). The North Atlantic Ocean exhibits various modes of low-frequency SST variability (Kushnir, 1994; Wu and Liu, 2005; Fan and Schneider, 2012; Taboada and Anadon, 2012). The mechanisms responsible for SST variability include ocean circulation variability, wind stress, and ocean-atmosphere feedbacks (Frankignoul, 1985; Deser et al., 2010). Aerosols and greenhouse gases (GHGs) emitted from anthropogenic and natural sources also contribute to regional SST variability through modulation of the solar radiation received by the ocean surface (Rotstayn and Lohmann, 2002; Wu and Kinter, 2011; Hsieh et al., 2013; Ding et al., 2014; Meehl et al., 2015).

Considering the distinct roles of regional SST variability in modulating regional climate systems, the impact of regional SST changes on the surface O₃ distribution needs to be explored. Lin et al. (2015) found that more frequent deep stratospheric intrusions appear over the western US during strong La Niña springs because of the meandering of the polar jet towards this region. This process can remarkably increase surface O3 levels in the western US. The La Niña-like decadal cooling of the eastern equatorial Pacific Ocean in the 2000s weakened the long-range transport of O3-rich air from Eurasia towards Hawaii during spring (Lin et al., 2014). Liu et al. (2005) revealed that El Niño winters are associated with stronger transpacific pollutant transport, which also has implications for the long-range transport of O₃. Except for the ENSO impacts, very few studies to date have directly addressed the linkage between SST and O₃. Therefore, a comprehensive understanding of the response of surface O₃ to

SST changes in individual ocean basins is lacking and necessary.

To fill this gap, this study focuses on examining the sensitivity of O_3 evolution over four polluted continental regions in the Northern Hemisphere (i.e., North America (NA, 15– 55° N, 60–125° W), Europe (EU, 25–65° N, 10° W–50° E), East Asia (EA, 15–50° N, 95–160° E) and South Asia (SA, 5–35° N, 50–95° E), defined in Fiore et al., 2009) with respect to nearby basin-scale SST changes. We describe the design of numerical experiments and model configuration in Sect. 2. Surface O₃ responses to regional SST changes are given in Sect. 3. Relevant mechanisms governing the SST–O₃ relationships are discussed in Sect. 4. The impact of basinscale SST changes on intercontinental transport of O₃ is described in Sect. 5. Conclusions are drawn in Sect. 6.

2 Methodology

2.1 Model description and configuration

The Community Earth System Model (CESM, v1.2.2) developed by the National Center for Atmospheric Research (NCAR) is used in this study, configured with the Community Atmosphere Model version 5.0 (CAM5) and the Community Land Model version 4.0 (CLM4). The ocean and sea ice components are prescribed with climatological SST and sea ice distributions. Moist turbulence is parameterized following the Bretherton and Park (2009) scheme. Shallow convection is parameterized using the Park and Bretherton (2009) scheme. The parameterization of deep convection is based on Zhang and McFarlane (1995) with modifications following Richter and Rasch (2008), Raymond and Blyth (1986), and Raymond and Blyth (1992). The cloud microphysical parameterization is following a two-moment scheme described in Morrison and Gettelman (2008) and Gettelman et al. (2008). The microphysical effect of aerosols on clouds are simulated following Ghan et al. (2012). The parameterization of cloud macrophysics follows Conley et al. (2012).

The chemistry coupled in the CAM5 (i.e., CAM5-chem) is primarily based on the Model for O_3 and Related chemical Tracers, version 4 (MOZART-4), which resolves 85 gasphase species and 196 gas-phase reactions (Emmons et al., 2010; Lamarque et al., 2012). A three-mode (i.e., Aitkin, accumulation, and course) aerosol scheme for black carbon (BC), primary organic matter (POM), second organic aerosol (SOA), sea salt, dust, and sulfate was used in our simulations following Liu et al. (2012). The lightning parameterization is modified according to Price et al. (1997) and tropospheric photolysis rates are calculated interactively following Tie et al. (2005). Gaseous dry deposition is calculated using the resistance-based parameterization of Wesely (1989), Walmsley and Wesely (1996), and Wesely and Hicks (2000). The parameterizations of in-cloud scavenging and below8773 bed in detail by

cloud washout for soluble species are described in detail by Giorgi and Chameides (1985) and Brasseur et al. (1998), respectively. Anthropogenic emissions of chemical species are from the IPCC AR5 emission datasets (Lamarque et al., 2010), whose injection heights and particle size distributions follow the AEROCOM protocols (Dentener et al., 2006). The emissions of natural aerosols and precursor gases are prescribed from the MOZART-2 (Horowitz et al., 2003) and MOZART-4 (Emmons et al., 2010) datasets. All emission datasets are available from the CESM data inventory (https:// svn-ccsm-inputdata.cgd.ucar.edu/trunk/inputdata/). The performance of CESM in simulating tropospheric O₃ has been validated by comparing with ozonesondes and satellite observations (Tilmes et al., 2015). The deviations between model and observations are within the range of about 25 %. In general, the model can capture the surface ozone distribution and variability well but may overestimate O3 over the eastern US and western Europe in the summer (Tilmes et al., 2015).

2.2 Numerical experiments

We first conduct a control simulation, hereafter referred to as CTRL, with prescribed climatological monthly SSTs averaged from 1981 to 2010 (see Hurrell et al., 2008). We then conduct six perturbation simulations with monthly SSTs that are uniformly increased or decreased by 1 °C in three ocean basins in the Northern Hemisphere: the North Pacific (15–65° N, 100° E–90° W), North Atlantic (15–65° N, 100° W-20° E), and North Indian oceans (5-30° N, 30-100° E; here 5° N is used to attain a relatively larger domain size). The simulations are denoted as Pacific-W, Atlantic-W, and Indian-W for the three warming cases and Pacific-C, Atlantic-C, and Indian-C for the three cooling cases. We defined the latitudinal and longitudinal ranges of these ocean basins mainly based on their geographical features. The boundaries of the prescribed SST anomalies generally align with the edge of the ocean basins, except along the southern side. In each perturbation simulation, we linearly smooth the southern boundaries of these SST anomalies towards the equator to remove the sharp SST anomaly gradients at the edge, following a previous approach (e.g., Taschetto et al., 2016; Seager and Henderson, 2016). We further conduct two sensitivity tests with 1 °C SST warming and 1 °C SST cooling superimposed onto all three ocean basins (i.e., the North Pacific, North Atlantic, and North Indian Ocean) in the Northern Hemisphere, denoted as All-W and All-C, respectively. Air pollution emissions, including biogenic emissions of VOCs, are fixed to distinguish the impacts of SST variation on O₃ transport and photochemistry. All simulations are run for 21 years with the first year used for model spin-up.

To explore the impacts of SST changes on intercontinental transport, an explicit emission tagging technique is used in our simulations following previous studies (Shindell et al., 2008; Doherty et al., 2013). Artificial CO-like tracers emitted from four continental regions, i.e., North America (NA, 15– 55° N, 60–125° W), Europe (EU, 25–65° N, 10° W–50° E), East Asia (EA, 15–50° N, 95–160° E), and South Asia (SA, 5–35° N, 50–95° E), are tracked individually. These tracers are idealized with a first-order decay lifetime of 50 days, which is similar to O₃ (Doherty et al., 2013) and used to single out changes in O₃ transport induced by SST anomalies.

2.3 Integrated process rate (IPR) analysis

To provide a process-level explanation for the response of surface O₃ to regional SST changes, the integrated process rate (IPR) method is applied. This method calculates the accumulated contributions of individual processes (e.g., chemical production and loss, advection, vertical diffusion, dry deposition) to O_3 predictions during the model simulation and has been widely used for air pollution diagnostics (Li et al., 2012; Zhang and Wu, 2013; Tao et al., 2015). In this study, we added the IPR scheme to the CESM framework to track the contribution of six physicochemical processes (i.e., gas-phase chemistry, CHEM; advection, ADVE; vertical diffusion, VDIF; dry deposition, DRYD; shallow convection, SHAL; and deep convection, DEEP) to O_3 concentrations in every grid box. Wet deposition and aqueous-phase chemistry are ignored here due to the low solubility and negligible chemical production of O₃ in water (Jacob, 1999). Therefore, CHEM represents the net production (production minus loss) rate of O₃ due to gas-phase photochemistry. DRYD represents the dry deposition fluxes of O₃, which are an important sink for O₃. The other IPR terms (i.e., ADVE, VDIF, SHAL, and DEEP) represent contributions from different transport processes. The IPR scheme tracks and archives the O₃ flux in each grid from every process during each model time step. The sum of the O_3 fluxes from these six processes matches the change in the O₃ concentration. The IPR method has been widely used in air quality studies to examine the cause of pollution episodes (Wang et al., 2010; Li et al., 2012). When applied in climate sensitivity analysis (usually measuring the difference between two equilibriums), the net change in all IPRs approaches zero. Typically, the positive changes in IPRs are mainly responsible for the increase in surface O₃, which may further induce O₃ removal to balance this forcing in a new equilibrium. Therefore, here, the IPR analysis is used not to budget the SST-induced O₃ concentration changes but rather to help examine the relative importance of different transport and chemical processes in driving the sensitivity of O3 to SST forcing. Its performance is verified by comparing the predicted hourly O₃ changes with the sum of the individual fluxes from the six processes. As shown in Fig. S1 in the Supplement, the hourly surface O₃ changes are well represented by the sum of these fluxes in the model.

3 Response of surface O₃ concentrations to SST changes

Seasonally and regionally averaged surface O₃ changes in each SST perturbation simulation for the four highly populated continental regions and three ocean basins defined in our study are given in Tables 1 and S1 in the Supplement, respectively. The responses of the surface O₃ concentrations to basin-scale SST changes (i.e., ± 1 °C) are mainly below 3 ppbv in the Northern Hemisphere (Tables 1 and S1), though larger anomalies (i.e., up to 5 ppbv) are also observed over the eastern coast of China, the Indian subcontinent, and certain oceanic areas (Figs. 1 and S2). This SST-O₃ sensitivity is comparable to previous findings. For instance, Bloomer et al. (2009) reported a positive O3-temperature relationship of 2.2–3.2 ppbv $^{\circ}C^{-1}$ across the rural eastern US. Wu et al. (2008) found that summertime surface O_3 may increase by 2-5 ppbv over the northeastern US in the 2050s. Additionally, Fiore et al. (2009) demonstrated an intercontinental decrease in surface O₃ of no more than 1 ppbv in response to 20 % reductions in anthropogenic emissions within a continental region. Our study indicates that basin-scale SST changes alone may exert significant effects on the surface O₃ above a specific ocean basin and its surrounding continents.

As shown in Fig. 1, seasonal changes of up to 5 ppbv in the mean surface O₃ concentration are observed during boreal summers, mainly in coastal regions and remote oceans. Surface O₃ changes in response to positive and negative SST anomalies generally exhibit a consistent spatial pattern but are opposite in sign, suggesting robust relationships between surface O₃ levels and SST anomalies (Fig. 1). An increase in summertime SST over a specific ocean basin tends to increase the surface O₃ concentration over the upwind regions but reduce this concentration over downwind continents. For instance, a 1 °C warming over the North Pacific leads to a widespread decrease of approximately 1 ppbv in surface O_3 over the North Pacific, North America, and the North Atlantic (Table S1) but may enhance the surface O₃ by nearly 3 ppbv over southern China. Similarly, the SST warming over the North Atlantic decreases the surface O₃ levels by 1–2 ppbv over the North Atlantic and Europe but increases $(\sim 1 \text{ ppbv})$ those over North America and the North Pacific. For the North Indian Ocean, positive SST anomalies tend to increase the surface O₃ over the Indian Ocean and Africa but decrease the surface O₃ over South and East Asia (Fig. 1). During the boreal winter, a widespread decrease in surface O₃ associated with the warming of different oceans is observed. Significant changes (e.g., up to 5 ppbv) mainly occur over remote ocean areas. Over populated continents, the response of the surface O₃ to basin-scale SST changes is typically insignificant. Details are shown in Fig. S2.

Our simulations reveal that different oceans can exert distinct region-specific effects on the O_3 distribution. The effects of three individual warming–cooling cases (i.e., Pacific-W, Atlantic-W, and Indian-W and Pacific-C, Atlantic-C, and

Table 1. Seasonally (i.e., DJF: December, January, and February; MAM: March, April, and May; JJA: June, July, and August; and SON: September, October, and November) and regionally averaged (only land grid boxes are included) changes in surface O_3 concentrations (ppbv) for basin-scale SST perturbation cases relative to the control simulation. Positive changes that are significant at the 0.05 level evaluated using the Student's *t* test are marked in bold.

Ozone (ppbv)			DJF	MAM	JJA	SON
North Pacific	+1 °C	North America	-0.27^{*} -0.50*	-0.42^{*} -0.26	-0.92^{*}	-1.03^{*} -0.29
		Earope Fast Asia	-0.88*	-0.71^{*}	0.10	0.17
		South Asia	-1.00^{*}	0.30	0.43	0.43*
	-1 °C	North America	0.00	0.57*	0.55*	0.82*
		Europe	0.19	0.15	-0.47^{*}	0.47^{*}
		East Asia	0.30	-0.17	-0.22	-0.67^{*}
		South Asia	0.04	-0.24	0.03	-0.40
North Atlantic	+1 °C	North America	0.03	0.49	0.50*	0.53*
		Europe	0.30*	0.06	-1.61^{*}	-0.89^{*}
		East Asia	-0.52^{*}	-0.68^{*}	-0.62^{*}	-0.25
		South Asia	-0.20	-1.46*	-1.28^{*}	-0.82^{*}
	$-1 ^{\circ}\mathrm{C}$	North America	-0.07	-0.10	0.10	-0.17
		Europe	0.00	0.00	0.07	0.06
		East Asia	0.16	-0.08	0.80*	-0.60^{*}
		South Asia	-0.20	-0.40	0.30	-0.10
North India	+1 °C	North America	-0.25	-0.04	-0.16	-0.10
		Europe	-0.30	0.08	-0.12	0.19
		East Asia	-0.53^{*}	-0.77^{*}	-0.28	-1.78^{*}
		South Asia	-1.00^{*}	0.14	-1.67*	-2.75^{*}
	$-1 ^{\circ}\mathrm{C}$	North America	0.04	0.17	0.04	0.25
		Europe	0.05	-0.07	-0.13	-0.24
		East Asia	-0.06	0.15	0.55*	0.33
		South Asia	-0.03	0.57	1.70*	1.31*

* Significant at the 0.05 level using the Student's t test and 20 years of model results.

Indian-C) on surface O_3 distributions are further summed up to compare with the combined warming–cooling cases (i.e., ALL-W and ALL-C). The responses of surface O_3 to a hemispheric SST anomaly generally resemble the sum of responses to individual regional SST changes (see Figs. S3 and S4). This indicates that the effect of a generalized SST warming on surface O_3 can be decomposed into individual regional SST forcings. We now analyze the processes that impact the dependence of SST on the O_3 distribution using simulations that increase the SST.

4 Mechanism of SST-induced surface O₃ changes

4.1 Process-level response to SST changes

In this study, IPR analysis is used to evaluate the contribution of different physicochemical processes to O_3 evolution. The SST-induced, process-level O_3 changes are spatially averaged over four populated continental regions (i.e., NA, EU, EA, and SA; Fig. 2) and three ocean basins (i.e., the North Pacific, North Atlantic, and North Indian oceans; Fig. S5). In most cases, VDIF and DRYD are the key processes controlling the O_3 variation. The downward transport of O_3 through diffusion (VDIF) is an important source of surface O_3 , while DRYD acts as a sink. Both processes are simultaneously determined by the strength of turbulence. Here, we define a new term TURB as the sum of DRYD and VDIF, and it can capture the overall effect of turbulence changes on surface O_3 concentrations. In addition, we merge SHAL and DEEP as CONV to represent the total contribution of convective transport to surface O_3 (Figs. 2 and S5). More detailed IPR results are shown in Figs. S6 and S7.

In the Pacific-W case, a 1 °C SST warming over the North Pacific increases VDIF over eastern China in JJA (Fig. S8), which is insignificant if averaged over the whole East Asian region. Meanwhile, this Pacific warming considerably reduces VDIF over North America (Fig. S6). The corresponding decrease in TURB over North America mainly determines the surface O_3 reduction in JJA and SON, while the reduction in CONV exerts an additional negative impact (Fig. 2). In the Atlantic-W case, increases in VDIF are also



Figure 1. Changes in the summertime (June–August) surface O₃ concentrations (ppbv) in the Northern Hemisphere induced by 1 °C warming (**a**, **b**, **c**) and 1 °C cooling (**d**, **e**, **f**) in the North Pacific Ocean (**a**, **d**), North Atlantic Ocean (**b**, **e**), and North Indian Ocean (**c**, **f**) relative to the CTRL. The four major regions of interest (i.e., NA: $15-55^{\circ}$ N, $60-125^{\circ}$ W; EU: $25-65^{\circ}$ N, 10° W– 50° E; EA: $15-50^{\circ}$ N, $95-160^{\circ}$ E; and SA: $5-35^{\circ}$ N, $50-95^{\circ}$ E) are marked with red polygons. The + symbols denote areas in which results are significant at the 0.05 level, evaluated using the Student's *t* test and 20 years of data.

observed over the upwind regions (i.e., North America) in JJA. However, these increases are accompanied by commensurate decreases in DRYD, resulting in an insignificant overall change in TURB (Fig. 2). Therefore, the increase in CHEM is mainly responsible for the surface O₃ increase over North America in JJA. TURB is more relatively important over Europe (only in JJA and SON), leading to reduced surface O₃ abundance. In the Indian-W case, both CHEM and CONV are reduced over South Asia in JJA, leading to overall reductions in surface O₃ over the Indian subcontinent (Fig. 2). The IPR analysis over the ocean basins shows that the warming of the North Pacific or North Atlantic induces reductions in VDIF and CHEM, which are responsible for the significant decrease in surface O₃ above these regions in JJA (Fig. S7). Conversely, North Indian Ocean warming enhances DEEP and VDIF, leading to a local increase in surface O₃ in JJA.

The IPR analysis indicates that, in general, an SST increase in the North Pacific or North Atlantic is more likely to enhance the vertical diffusion of O_3 over upwind regions (i.e., East Asia or North America, respectively) but suppress

this diffusion over the ocean basin as well as downwind continents in JJA (Fig. S8). These opposite changes in VDIF over upwind and downwind regions lead to distinct surface O_3 responses. Changes in CHEM enhance surface O_3 formation in most cases. An exception is in South Asia, where CHEM and DEEP dominate the reduction in surface O_3 over the region in JJA associated with the North Indian Ocean warming. In the following subsections, the mechanisms of the SST– O_3 relationship for the four polluted continents are further explored. Here we focus on boreal summers since the surface O_3 response to SST changes is more robust during this period than other seasons.

4.2 Response of photochemical O₃ production to SST increases

Changes in the net production rate (i.e., chemical production rate minus loss rate) of O_3 at the surface in JJA associated with basin-scale SST increases are shown in Fig. 3. The peak changes are mainly confined to regions where O_3 precursors are abundant (e.g., South and East Asia and North



Figure 2. Seasonally averaged changes in the IPR contributions (bars, ppbv h^{-1} , left scale) and surface O₃ concentrations (hollow circles, ppbv, right scale) for Pacific-W (**a**, **d**, **g**, **j**), Atlantic-W (**b**, **e**, **h**, **k**), and Indian-W (**c**, **f**, **i**, **l**) relative to the CTRL. Values are regionally averaged over NA (**a**, **b**, **c**), EU (**d**, **e**, **f**), EA (**g**, **h**, **i**), and SA (**j**, **k**, **l**). TURB is defined as the sum of VDIF and DRYD. CONV is the sum of DEEP and SHAL. IPR contributions from the four processes (i.e., TURB, ADVE, CHEM, and CONV) are represented by different colors. A more detailed IPR result is shown in Fig. S6.

America). For example, a warmer North Pacific SST exerts a positive (negative) impact on net O_3 production in the northern (southern) regions of East Asia. Similarly, the warming of the North Atlantic promotes a dipole impact on the surface O_3 production over North America, while the warming of the North Indian Ocean significantly decreases the net O_3 production rate over South Asia.

As emissions are fixed in all simulations, the change in net O₃ production is driven by SST-induced meteorological changes (e.g., air temperature, air humidity, and solar radiation). An increase in SST of 1 °C in any ocean basin leads to a widespread enhancement of the surface air temperature (i.e., the air temperature at 2 m) over most continental areas (Fig. 4). An exception is the North Indian Ocean, where an increase in SST tends to cool the Indian subcontinent by 1– 2 °C. This temperature decrease is not only limited to the surface but also spreads to 600 hPa (Fig. S9). Associated with this temperature decrease is a remarkable reduction in the solar radiation received at the surface (more than $15 \,\mathrm{W}\,\mathrm{m}^{-2}$, Fig. S10). Previous studies have indicated that moist convection is more sensitive to the SST changes in the tropical oceans than in mid- or high-latitude oceans (Lau and Nath, 1994; Lau et al., 1997; Hartmann, 2015). The SST increase over the North Indian Ocean tends to strengthen the moist convection that eventually facilitates cloud formation in the upper troposphere (Roxy et al., 2015; Xi et al., 2015; Chaudhari et al., 2016). The latent heat released from convective activities significantly warms the air temperature over the upper troposphere (Sabeerali et al., 2012; Xi et al., 2015). Meanwhile, the corresponding increase in cloud cover reduces the solar radiation reaching the surface of the Indian subcontinent and thus the air temperature of the lower troposphere in that region. These processes lead to opposite air temperature changes between the upper and lower troposphere over South



Figure 3. Perturbations of the surface net O₃ production rate $(1 \times 10^6 \text{ molecules cm}^{-3} \text{ s}^{-1})$ for (a) Pacific-W, (b) Atlantic-W, and (c) Indian-W relative to the CTRL in the boreal summer. The + symbols denote areas in which the results are significant at the 0.05 level, evaluated using the Student's *t* test and 20 years of data.



Figure 4. Changes in the surface air temperature ($^{\circ}$ C) for (a) Pacific-W, (b) Atlantic-W, and (c) Indian-W relative to CTRL in the Northern Hemisphere in the boreal summer. The + symbols denote areas in which the results are significant at the 0.05 level, evaluated using the Student's *t* test and 20 years of data.

Asia in response to the North Indian warming (as shown in Fig. S9), which may further suppress the development of deep convection over the Indian subcontinent.

Previous studies have indicated that air temperature positively affects both O_3 production and destruction rates (Zeng et al., 2008; Pusede et al., 2015). As shown in Fig. S11, changes in the net O_3 production rate are mainly dominated by O_3 production over continents but by O_3 destruction over oceans. An increase in SST leads to a widespread enhancement of the air temperature, resulting in a positive change in the net O_3 production over most continental regions (Fig. 3). However, a warmer SST also increases the air humidity (Fig. S12), which enhances O₃ destruction over most coastal and oceanic areas. In addition, over South Asia, a warming of the North Indian Ocean decreases solar radiation and air temperature and simultaneously increases air humidity, which jointly exert negative effects on O₃ production in that region.

4.3 Response of physical O₃ transport to SST increases

In Sect. 4.1, our IPR analysis highlights multiple physical processes (i.e., vertical diffusion, convection, and advection) that are important in modulating surface O_3 concentrations.



Figure 5. Changes in the surface pressure (color contours, Pa) and 850 hPa wind (arrows, $m s^{-1}$) for (a) Pacific-W, (b) Atlantic-W, and (c) Indian-W relative to the CTRL in the boreal summer.

However, the role and relative importance of each process exhibit large spatial heterogeneity. In this section, we explore the key factors controlling physical O_3 transport in response to basin-scale SST changes.

The changes in the surface pressure and wind pattern induced by a basin-wide SST increase are shown in Fig. 5. Generally, a warming of any ocean basin will lead to a lowpressure anomaly centered to its west at low latitudes, which is caused by SST-induced convective activity. Additionally, the warming of the Indian Ocean induces an anticyclonic anomaly over the subtropical western Pacific, which has been documented in previous studies (Yang et al., 2007; Li et al., 2008). As shown in Fig. 6, the surface pressure reduction induced by SST warming in any ocean basin is closely associated with enhanced upward motions, suggesting a substantial enhancement in deep convection over tropical oceans. Previous studies have identified an SST threshold (approximately 26-28 °C) to generating deep convection (Graham and Barnett, 1987; Johnson and Xie, 2010). Therefore, the sensitivity of deep convection to an SST anomaly is strongly dependent on the distribution of base SST. The enhanced upward motion in response to a uniform increase in basin-scale SST mainly occurs over regions with high climatological SST (Fig. 6). Regions with a low climatological SST have little effect on the vertical movement of air masses.

Strengthened deep convection will trigger large-scale subsidence over nearby regions through the modulation of largescale circulation patterns, which may suppress convective transport (Lau et al., 1997; Roxy et al., 2015; Ueda et al., 2015). This effect is verified by the decreases in upward velocity at 500 hPa. As depicted in Fig. 6, significant decreases in upward velocity occur over regions adjacent to the strengthened deep convection. Similar effects are also observed over higher latitudes or remote oceans (Fig. S13). Meanwhile, the air temperature increase in response to regional SST warming is more significant above the lower troposphere, which leads to a decrease in the vertical temperature gradient (Fig. S9). These factors tend to restrain the vertical exchange of air pollutants at midlatitudes, which facilitates surface O_3 accumulation over polluted continental



Figure 6. Spatial pattern of vertical velocity changes at 500 hPa (color contours, $1 \times 10^{-2} \text{ Pa s}^{-1}$) for (**a**) Pacific-W, (**b**) Atlantic-W, and (**c**) Indian-W relative to the CTRL in the boreal summer. Positive values indicate upward motion. Red polygons denote the regions where the surface pressure responses to SST anomalies are significant (see Fig. 5a–c). The + symbols indicate areas in which the results are significant at the 0.05 level, evaluated using the Student's *t* test and 20 years of data.

regions in JJA but may weaken the intrusion of O_3 from the upper troposphere to the surface in most unpolluted areas. This process helps to explain the widespread decrease in surface O_3 over unpolluted regions associated with an SST increase, as described in Sect. 3, and can be further verified by the widespread reduction in VDIF shown in Fig. S8.

The surface pressure anomalies induced by SST changes can play a dominant role in modulating surface O_3 transport at specific locations. For example, the low-pressure anomaly centered over the subtropical northwestern Pacific in the Pacific-W case causes the convergence of wind in the lower troposphere (Fig. 5a). Consequently, surface O_3 pollution is enhanced in southern China due to an increase in O_3 transport from more polluted northern China (Fig. 7a). The vertical distribution of the corresponding O_3 changes also shows that the increase in O_3 over southern China occurs below 700 hPa, accompanied by noticeable decreases above 700 hPa as well as over nearby northern China (Fig. 7d). The IPR analysis also indicates that the increases in advective transport and downward turbulent transport are mainly responsible for the surface O_3 increase in southern China.

In the Atlantic-W case, the SST warming-induced surface pressure anomalies lead to substantial O_3 redistribution, especially over the North Atlantic Ocean (Fig. 7b). For North America, the changes in horizontal O_3 fluxes have no significant effect on the O_3 concentration increase. In addition, O_3 changes are observed to be larger in the upper troposphere than at the surface (Fig. 7e). As demonstrated in Sect. 4.1, the response of lower-altitude O_3 over North America to the North Atlantic warming is mainly caused by enhanced chemical production rather than physical transport.

The North Indian SST warming leads to a low-pressure anomaly centered over the Arabian Sea (Fig. 5c). The warming of the North Indian Ocean strengthens the upward motion of air at low latitudes and further induces a convergence of highly polluted air over the Indian Ocean. The effects of this process on O_3 concentrations are observed to be more significant in the upper troposphere (Fig. 7f). According to the IPR analysis, the surface O_3 increase over the Indian Ocean



Figure 7. Top three rows: changes in O₃ concentrations (color contours, ppbv) and horizontal fluxes (arrows, mol cm⁻² s⁻¹) at the surface level for (**a**) Pacific-W, (**b**) Atlantic-W, and (**c**) Indian-W relative to the CTRL in the boreal summer. Bottom row: zonal average of the tropospheric O₃ changes (color contours, ppbv), wind fluxes in CTRL (red arrows, m s⁻¹), and the wind flux perturbation (black arrows, m s⁻¹) in (**d**) Pacific-W, (**e**) Atlantic-W, and (**f**) Indian-W relative to the CTRL in the boreal summer. The red rectangles in (**a**), (**b**), and (**c**) denote the longitudinal range used for the zonal averages in (**d**), (**e**), and (**f**), respectively. The vertical wind velocity is amplified 1000 times to make it comparable to the horizontal wind velocity.

is mainly caused by the enhanced vertical transport of O_3 to the surface through deep convection and vertical diffusion processes (Fig. S7). However, over the nearby Indian subcontinent, the suppressed convection tends to decrease surface O_3 in that region (Fig. 2).

5 Implications for O₃ long-range transport

The findings above indicate that, in general, a basin-scale SST increase in the Northern Hemisphere is more likely to enhance atmospheric stability at midlatitudes, which may suppress air pollutants from lofting to the free troposphere. This process potentially has large effects on O_3 interconti-

nental transport. Following previous work (e.g., Doherty et al., 2013; Fang et al., 2011), we use passive CO-like tracers to demonstrate the potential effect of regional SST changes on long-range O₃ transport. A warming of North Pacific SSTs by 1 °C tends to increase the East Asian CO tracer concentrations by nearly 6% at the surface (Fig. 8b), which is accompanied by a significant reduction (~4%) in eastward transport to North America. Similarly, for the North American tracer, a warming of North Atlantic SSTs by 1 °C increases (~1%) the concentrations over downwind Europe (Fig. 8d). The response of the South Asian CO tracer to North Indian Ocean warming also shows a decreasing tendency over



Figure 8. Panels (**a**, **c**, **e**): difference in the surface concentration (ppbv) of a CO-like tracer emitted from (**a**) East Asia for Pacific-W, (**c**) North America for Atlantic-W, and (**e**) the South Asia for Indian-W relative to the CTRL in the boreal summer. Panels (**b**, **d**, **f**): the percentage changes in the surface concentration of a CO-like tracer emitted from (**b**) East Asia for Pacific-W, (**d**) North America for Atlantic-W, and (**f**) South Asia for Indian-W relative to the CTRL in the boreal summer. Red polygons denote the region where the CO-like tracer is emitted from. The + symbol denotes areas in which the results are significant at the 0.05 level, evaluated using the Student's *t* test and 20 years of data.

downwind regions, but the patterns are more complicated over the source region in this case (Fig. 8e). Because the CO-like tracers added in the simulation have a fixed decay lifetime, their concentration changes are completely caused by the SST-induced transport anomalies. The decrease in CO tracer concentrations over downwind regions suggests that the warming of basin-scale SST tends to suppress the longrange transport of air pollutants. Additionally, in the Pacific-W case, changes in the East Asian CO tracer (Fig. 8a) generally resemble the changes in surface O3 over East Asia (Fig. 7a), indicating the dominant effect of physical transport on the O₃ distribution over East Asia. Regarding the North American CO tracer in response to the North Atlantic warming or the South Asian CO tracer in response to the North Indian Ocean warming, their concentration changes are spatially inconsistent with those of O₃ (see Figs. 7 and 8). This further indicates the distinct roles that different basin-scale SSTs play in nearby air quality.

Further investigations of zonal wind suggest that an increase in SST over different oceans consistently decreases the westerly winds at lower midlatitudes $(25-45^{\circ} \text{ N})$ in the Northern Hemisphere but increases these winds at higher

latitudes (Fig. 9). In general, increases in the geopotential height induced by basin-scale SST warming are more significant at midlatitudes than at other latitudes, which is consistent with the air temperature changes. Consequently, the meridional geopotential height gradient decreases at lower latitudes but increases at higher latitudes, leading to corresponding changes in the westerly winds. The latitude band at 25-45° N covers many polluted regions (i.e., North America and East Asia). A weakened westerly wind may reduce longrange O₃ transport. As demonstrated in Sect. 4.3, the basinscale SST increases also exert negative effects on the upward transport of air masses at midlatitudes. Therefore, the decreases in CO tracer concentrations over downwind regions (Fig. 8a and c) can be explained by both suppressed vertical transport and weakened westerly winds. In the Indian-W case, the SST increase over North India leads to a lowpressure anomaly above the Arabian Sea due to the enhanced deep convection (as discussed in Sect. 4.3). The corresponding anomalous cyclonic circulation may be responsible for the dipole of the South Asian CO tracer changes over the source region depicted in Fig. 8e.



Figure 9. Zonally averaged changes in zonal wind (color contour, m s⁻¹) and geopotential height (contour, m) for (**a**) Pacific-W, (**b**) Atlantic-W, and (**c**) Indian-W relative to the CTRL in the boreal summer. Solid and dashed black lines in the contours indicate positive and negative geopotential height anomalies, respectively (contour interval: 5 m). The + symbol denotes areas in which the zonal wind changes are significant at the 0.05 level, evaluated using the Student's *t* test and 20 years of data.

In addition, we also find a hemispheric-scale decrease in peroxyacetyl nitrate (PAN), a reservoir of O_3 precursors (NO_x and HO_x) that facilitate the long-range transport of O_3 , during the warming of different oceans (Fig. S14). This decrease is likely caused by the increase in the thermal decomposition of PAN in response to the air temperature increase (Jacob and Winner, 2009; Doherty et al., 2013).

Thus, it is reasonable to infer that, in general, the increased thermal decomposition of PAN, the weakened midlatitude westerlies, and the reduced vertical air transport may exert a joint reduction effect on the intercontinental transport of O_3 during basin-scale SST increases.

6 Summary

In this paper, we investigate the responses of surface O_3 to basin-scale SST anomalies in the Northern Hemisphere. The latest version of CESM (version 1.2.2) is used in our simulation, forced with climatological and stationary SST anomalies $(\pm 1 \,^{\circ}C)$ in the North Pacific, North Atlantic, and North Indian oceans, respectively. The responses of surface O₃ associated with these SST changes are evaluated. Results of similar magnitude but opposite sign are observed for the SST warming versus cooling simulations for each ocean basin, suggesting robust connections between the SST anomalies and surface O3 changes. The regionally and seasonally averaged surface O₃ changes over four continental regions (i.e., NA, EU, EA, and SA) produce wide seasonal and regional variability (varying from 1 to 3 ppbv). The warming of the North Pacific leads to nearly 3 ppbv increases in the surface O₃ over southern China in summer, with corresponding decreases over North America (~ 1 ppbv). Similarly, the North Atlantic SST warming elevates the surface O₃ pollution over North America while reducing the surface O_3 (nearly 1–2 ppbv) over Europe. Changes in the North Indian SST exert significant impacts (1–3 ppbv) over South and East Asia during the entire year.

Process analysis indicates that dry deposition and vertical diffusion are two major processes governing the surface O₃ balance. The increase in SST in different ocean basins tends to increase the contributions of vertical diffusion to surface O₃ over upwind regions while greatly restraining those over downwind continents. These processes generally lead to widespread decreases in surface O₃, which are partially offset by increases in air-temperature-dependent chemical production rates. Specifically, the chemical production changes are mainly responsible for the surface O₃ increases over North America in response to the North Atlantic SST warming but exert a negative effect on South Asia in response to the North Indian SST warming. Decreases in the convective transport of O₃ to the surface associated with North Indian warming are significant over South Asia and exert a negative impact on surface O₃ concentrations. Advective transport has a positive effect on surface O₃ in southern China in the Pacific-W case.

We further show that air temperature is an important factor controlling the surface O_3 responses to SST anomalies. Reductions in the surface O_3 chemical production in South Asia associated with North Indian SST warming can be explained by the corresponding SST-induced decreases in ground-level air temperature and solar radiation. Meanwhile, the widespread increase in air temperature associated with basin-scale SST warming is more likely to promote O_3 production over other highly polluted regions.

Conversely, SST increases at low latitudes over different oceans enhance deep convection in summer, which promotes convergence at the surface, as well as upward motions at low latitudes. The corresponding surface pressure anomalies centered over the east coast of East Asia associated with the North Pacific warming and over the Arabian Sea associated with the North Indian warming tend to increase the surface O_3 above through exchanges with the surrounding highly polluted air. The basin-scale SST increases in the Northern Hemisphere reduce the tropospheric temperature gradient at midlatitudes that restrains vertical transport of O_3 over continents and weakens the westerlies at lower midlatitudes. The response of the CO tracer also suggests that these factors may jointly exert a negative effect on the intercontinental transport of O_3 .

This study highlights the sensitivity of O₃ evolution to basin-wide SST changes in the Northern Hemisphere and identifies the key chemical or dynamical factors that control this evolution. Idealized and spatially uniform SST anomalies are used to explore the general mechanisms governing SST-O₃ relationships. We find that the SST changes over tropical regions exert considerable impacts on surface O₃ levels. The increase in tropical SST over different ocean basins enhances deep convection, which further triggers large-scale subsidence over nearby and remote regions. These enhanced convective activities also tend to release more latent heat over the upper troposphere and significantly increase the air temperature there. These processes influence large-scale circulation patterns and lead to opposite surface O₃ responses over upwind and downwind regions related to a specific ocean basin. This finding provides valuable implications for the potential surface O3 change in response to future warming or cooling of individual oceans.

Additionally, the sensitivity tests with 1 °C SST warming and cooling superimposed onto all three ocean basins further show in general that the SST forcing on O₃ distribution is geographically additive. A number of studies have used the decomposed SST anomalies for different regions to identify their relevant roles in a particular climate response (e.g., Sutton and Hodson, 2005; Camargo et al., 2013; Ueda et al., 2015). A linear assumption that the influence of large-scale SST anomaly pattern on the atmosphere can be generally constructed by the linear combination of the influences of individual SST patches has been verified by previous studies, especially for the tropical regions where the signal-to-noise ratio is higher (e.g., Fan et al., 2016; Seager and Henderson, 2016). Therefore, our study also helps to understand the roles different ocean basins in the Northern Hemisphere play in modulating surface O₃ distributions in a global-wide SST warming condition associated with climate change.

Overall, this study may guide the management of regional O_3 pollution by considering the influence of specific SST variability. However, cautions should be taken in interpreting our results in the real world since observed surface O_3 variabilities are induced by various factors including O_3 precursor emissions and atmospheric conditions. Realistic SST anomalies over different oceans are more complicated (usu-

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ally not uniformly distributed) and often intercorrelated with each other (Fan et al., 2016). They may exert joint effects on modulating surface O_3 distributions. To provide more precise understanding about the SST– O_3 relationship over a specific region, additional sensitivity tests regarding smaller patches of SST variability are necessary.

Data availability. The CESM release code is downloadable through the Subversion repository path: https://svn-ccsm-models. cgd.ucar.edu/cesm1/. All input datasets are available from the CESM data inventory (https://svn-ccsm-inputdata.cgd.ucar.edu/trunk/inputdata/). The simulation results are available from the corresponding author, Junfeng Liu (jfliu@pku.edu.cn), upon request.

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

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