Heterogeneous impacts of fire-sourced ozone (O₃) pollution on global

2	crop yields in the future climate scenarios
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10	Abstract
11	Wildfire smoke often aggravates the ozone (O ₃) pollution and negatively affect crop yields. To date,
12	the global impact of fire-sourced O ₃ exposure on crop yields still remained unknown. To address
13	this issue, a multi-stage model was developed to quantify the global wildfire-induced ambient O ₃
14	concentrations in the future scenarios. The results suggested that the relationship between observed
15	K ⁺ and levoglucosan levels with simulated fire-sourced maximum daily average 8-hour (MDA8) O ₃
16	concentration reached 0.67 and 0.73, respectively, indicating the robustness of fire-sourced O ₃
17	estimate. In both of historical and future scenarios, Sub-Sahara Africa (SS: 14.9 ± 8.4 (historical)
18	and 18.3 ± 9.6 (mean of the future scenarios) $\mu g/m^3$) and South America (SA: 4.0 ± 2.5 and 4.7 ± 1.0
19	3.2 µg/m³) showed the highest fire-sourced MDA8 O ₃ concentrations among all of the regions.
20	However, the crop production losses (CPL) caused by O ₃ exposure reached the highest values in
21	China due to very high total crop yields and relatively high wildfire-induced MDA8 O ₃ levels.
22	Moreover, CPL in China was sensitive to emission scenario, indicating the effective emission
23	control could largely decrease fire-sourced O ₃ damage to crop. In contrast, both of SS and SA even
24	showed the higher CPL in low-carbon scenario (SSP1-2.6), suggesting more stringent control
25	measures are required to offset the wildfire contribution. Our findings call for attention on the threat
26	to future global food security from the absence of pollution mitigation and the persistence of global
27	warming.
28	Keywords: MDA8 O ₃ , wildfire, crop yield, Sub-Sahara Africa, China

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Along with the warming climate, large-scale wildfire events have experienced dramatic increases in frequency and intensity in the past decades, and the wildfire seasons have been significantly prolonged in many regions such as the western part of the United States and Australia (Jones et al. 2022, Richardson et al. 2022, Wang et al. 2022). Wildfire often released a large number of gaseous precursors such as carbon monoxide (CO), nitrogen dioxides (NO_x), and volatile organic compounds (VOC) (Anderson et al. 2024, Xu et al. 2022), which could significantly enhance the ozone (O₃) levels through photochemical reactions (Jaffe et al. 2013). Recent studies have revealed that wildfire contributed to 3.6% of ambient all-source O₃ level globally (Xu et al. 2023). The aggravation of O₃ pollution not only poses detrimental effects on human health (Liu et al. 2018), but also reduced the crop yields because the excessive O₃ exposure could affect plant photosynthesis via stomatal uptake (Karmakar et al. 2022, Zhao et al. 2020). Thus, quantifying the negative impacts of fire-sourced O₃ pollution on crop yields was beneficial to propose optimal strategy to ensure agricultural production.

Notably, warming climate in the future not only would increase wildfire burned areas, but also intensified the severity of fire weather (Richardson et al. 2022, Wasserman and Mueller 2023). Moreover, wildfire and heatwave have generated the positive feedback and the mechanism would be further enhanced in the future (Senande-Rivera et al. 2022, Zhao et al. 2024). Meanwhile, the ambient O₃ concentration was very sensitive to air temperature, and the continuous increase of air temperature inevitably aggravate wildfire-related O₃ pollution in the future (Bloomer et al. 2009, Li et al. 2024a, Selin et al. 2009). Therefore, it is necessary to analyze the spatiotemporal characteristics of global wildfire-induced O₃ concentrations especially in the future scenarios, which was favorable to accurately identify the hotspots for wildfire-induced O₃ pollution and to propose effective control measures targeting different future scenarios.

A growing body of studies have focused on the wildfire contribution to O₃ pollution. Lee et al. (2024) employed the generalized additive model (GAM) to predict the wildfire-related O₃ concentration in the United States and found wildfire increased maximum daily average 8-hour (MDA8) O₃ concentration across the entire country (Lee and Jaffe 2024). Besides, Xu et al. (2023) have quantified that the wildfire led to average 3.2 μg/m³ increase of O₃ concentration globally using the GEOS-Chem model. Unfortunately, most of the current studies assessed the contribution

of historical wildfire to ambient O₃ level, and the estimates showed large uncertainties associated with the burned areas, fuel consumption, and fuel types. Moreover, most of these studies only focused on the historical estimates, while only two studies explored the wildfire contribution to O₃ pollution in the future scenarios (Yang et al. 2022, Yue et al. 2015). Both of these studies only focused on wildfire in North America, whereas the future wildfire contribution to O₃ pollution in other regions are still unknown. Moreover, their negative impacts on crop yields are also not clear. In fact, the global wheat yield losses reached 0.95% (around 20 t/km²) per ppb O₃ increase (Guarin et al. 2019). Although the current contribution ratio of wildfire to all-source O₃ level is not high, the higher wildfire risk and total crop yields in the future scenarios highlights the seriousness of crop yield losses.

Here, our study developed an ensemble machine-learning model to predict fire-sourced MDA8 O₃ levels under four future scenarios (SSP1-2.6, SSP2-4.5, SSP3-7.0, and SSP5-8.5). Then, the spatiotemporal variations of these concentrations and the key drivers behind them were further revealed. Finally, a crop yield loss assessment framework was applied to quantify the negative impacts (crop yield losses) of wildfire-induced O₃ exposure on global crop yield. The hotspots of crop yield losses in different scenarios should be determined and the appropriate control measures should be proposed to reduce the economic losses.

2. Materials and methods

2.1 Data preparation

Most ground-level MDA8 O₃ observations focused on East Asia, India, Western Europe, and the contiguous United States. Daily MDA8 O₃ data during 2015-2019 over China were collected from the Ministry of Ecology and Environment of China. The observation network comprises of 2,000 monitoring sites distributed across various land-use types (Figure S1). Quality assurance for the ground-level observations in China was performed based on the HJ 630-2011 specifications. The dataset of daily MDA8 O₃ concentrations from 2015 to 2019 in India were collected from the Central Pollution Control Board (CPCB) online database (https://app.cpcbccr.com/ccr/#/caaqmdashboard-all/caaqm-landing). The detailed data quality assurance/control has been introduced by Gurjar et al. (2016). Ground-level observation dataset for member countries of the European Economic Area were collected from the European Environment Agency. The data quality control of

European Environment Agency was explained by Keller et al. (2021). The dataset of daily MDA8 O₃ levels in more than 200 monitoring sites across the United States were downloaded from the website of https://www.epa.gov/ (Figure S1). The quality control of these observations in EPA was carefully introduced by (Lamsal et al. 2015). Observation data in other countries and territories were downloaded from the website of OpenAQ (https://openaq.org/). After the data cleaning and quality control, more than 300,000 daily MDA8 O₃ measurements in 3015 sites were collected to simulate the global O₃ concentrations. For O₃, 1 part per billion (ppb) was approximated as 1.96 μg/m³ based on the standard air pressure and temperature (25.5 °C and 101.325 kPa). The Unite of O₃ was changed into μg/m³ unified.

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GEOS-Chem (v13.4.0) model was utilized to estimate atmospheric MDA8 O₃ concentrations during Jan. 1-Dec. 31 during 2015-2019, 2045-2049, and 2095-2099 periods. In our study, the years of 2015-2019 was regarded as the historical period, whereas the years of 2045-2049 and 2095-2099 were regarded as the future period. This model comprises of a complex chemistry mechanism of tropospheric NO_x-VOC-O₃-aerosol (Geddes et al. 2015, Zhao et al. 2017). This model for O₃ estimates during historical period and future scenario were driven by MERRA2 and GCAP2 CMIP6 reanalysis meteorological factors, respectively (Bali et al. 2021, Zhang 2016). The future scenario includes SSP1-2.6 (low-carbon emission scenario), SSP2-4.5 (middle-carbon emission scenario), SSP3-7.0 (traditional energy scenario), and SS5-8.5 (high energy consumption scenario). A global simulation was performed at a spatial resolution of 2 × 2.5° resolution (Bindle et al. 2021, Wainwright et al. 2012). The historical anthropogenic emission inventory during 2015-2019 was downloaded from Community Emissions Data System (CEDS) (Hoesly et al. 2018). The anthropogenic and wildfire emissions during 2045-2049 and 2095-2099 were collected from the website of https://esgf-node.llnl.gov/search/input4mips/. Wildfire emission during 2015-2019 was obtained from GFED (Chen et al. 2023, Pan et al. 2020, Peiro et al. 2022, van Wees et al. 2022). Some other natural emission such as the lightning NO_x emission was collected from http://geoschemdata.wustl.edu/ExtData/HEMCO/OFFLINE LIGHTNING/v2020-03/MERRA2/ (Li et al. 2022, Nault et al. 2017, Verma et al. 2021). The whole simulation processes included four steps. Firstly, we run the GEOS-Chem model with all emissions (including wildfires) to establish reference O₃ concentrations (Baseline simulation). Second, we repeated the simulation while

excluding wildfire emissions with the same meteorological conditions (MERRA2 and GCAP2_CMIP6) and anthropogenic emission inventory (CEDS). Third, we computed the wildfire-induced O_3 by subtracting zero-out results from the baseline. At last, we compare modeled O_3 concentrations with observational data (e.g., ground-based measurements) to assess uncertainty.

Meteorological factors including 2 m dewpoint temperature (D2m), surface pressure (Sp), 2 m temperature (T2m), and total precipitation (Tp), 10 m wind component (U10 and V10) during 2015-2019 were collected from the fifth-generation European Centre for Medium-Range Weather Forecasts Reanalysis (ERA-5). All of these meteorological data showed the same spatial resolution of 0.25°×0.25°. For the estimates in the future scenarios, the CMIP6 dataset in four scenarios (e.g., SSP1-2.6, SSP2-4.5, SSP3-7.0, and SSP5-8.5) were also applied to predict MDA8 O₃ concentrations during 2015-2019, 2045-2049, and 2095-2099. The dataset includes simulated O₃ concentrations, 2-m air temperatures, wind speed at 850 and 500 hPa, total cloud cover, precipitation, relative humidity, and short-wave radiation. The modelled meteorological parameters and chemical compositions derived from multiple earth system models were integrated into the machine-learning model. The detailed models are introduced in our previous studies (Li et al. 2024b). The elevation was collected from ETOPO at a spatial resolution of 1°. Additionally, the land use type data were downloaded from the reference of Liu et al. (2020).

2.2 Model development

A multi-stage model was developed to estimate the global fire-sourced MDA8 O₃ concentrations (Figure S2). In the first stage, the ground-level MDA8 O₃ levels, meteorological factors, land use types, and simulated O₃ levels derived from GEOS-Chem model were integrated into XGBoost model to simulate the full-coverage MDA8 O₃ levels during 2015-2019. In the second stage, the simulated O₃ concentrations and meteorological parameters in four scenarios (SSP1-2.6, SSP2-4.5, SSP3-7.0, and SSP5-8.5) during 2015-2019, 2045-2049, and 2095-2099 were collected from CMIP6 dataset including 16 earth system models. Then, the data in the future scenarios were integrated into the XGBoost model to further calibrate the CMIP6 modeling results based on historical dataset (2015-2019) derived from the first stage model. This stage could obtain the calibrated MDA8 O₃ concentrations in different scenarios during 2015-2019, 2045-2049, and 2095-2099. The detailed equations of XGBoost model are summarized as follows:

$$F^{(t)} = \sum_{i=1}^{n} [l(y_i, y^{\Lambda^{(t-1)}}) + \partial_{y^{(t-1)}} l(y_i, y^{\Lambda^{(t-1)}}) f_t(x_i) + \frac{1}{2} \partial_{y^{(t-1)}}^2 l(y_i, y^{\Lambda^{(t-1)}}) f_t^2(x_i)] + \Omega(f_t)$$
(1)

where $F^{(t)}$ represents the cost function at the t-th period; ∂ denotes the derivative of the function;

- 148 $\hat{\sigma}_{v^{(t-1)}}^2$ means the second derivative of the function; l refers to the differentiable convex loss function
- that reveals the difference of the predicted O₃ level (y) of the i-th instance at the t-th period and
- the target value (y_i); $f_t(x)$ is the increment; $\Omega(f_t)$ reflects the regularizer. Maximum tree depth
- and learning rate are 20 and 0.1, respectively.
- In the third/final stage, the calibrated MDA8 O₃ concentrations based on previous two-stage
- models were utilized to correct the bias of GEOS-Chem output. Due to the uncertainty of
- 154 GFED/anthropogenic emission inventory and chemical mechanism, the simulated MDA8 O₃
- concentration often largely biased from the ground-level observations. Therefore, it is necessary to
- use the assimilated results to optimize the wildfire-induced concentrations. The detailed equations
- are summarized as follows:

$$O_{3_opt_fire} = O_{3_cal_total} \times (O_{3_chem_fire} / O_{3_chem_total})$$
 (2)

- where $O_{3_opt_fire}$ is optimized wildfire-induced MDA8 O_3 concentration in the final stage.
- 160 $O_{3_cal_total}$ is calibrated total MDA8 O₃ concentration. $O_{3_chem_fire}$ is simulated wildfire-induced
- MDA8 O_3 concentration using GEOS-Chem model. $O_{3_chem_total}$ is simulated total MDA8 O_3
- concentrations using GEOS-Chem model. The ratios of fire-sourced O₃ concentrations and the total
- 163 O₃ concentrations during historical and different climate scenarios were not invariable, which were
- estimated by GEOS-Chem based on different meteorological conditions and emission scenarios.
- All of the independent variables obtained from various sources were resampled to 0.25° grids
- using Kriging interpolation. For the machine-learning model development, it was necessary to
- eliminate some redundant independent variables and then determine the optimal variable group. The
- redundant variables were identified based on the fact that the overall predictive accuracy could
- degrade after the removal of these variables. 10-fold cross-validation method was applied to
- examine the predictive accuracy of XGBoost model.
- 171 The modelling accuracy of wildfire emission to MDA8 O₃ cannot be evaluated directly,

whereas the modelling performance of total MDA8 O₃ concentrations could be assessed. Some typical statistical indices (supporting information) were applied to evaluate the modelling accuracy of this model on the basis of the ground-level observations.

2.3 The crop yield loss estimate

Maize, rice, spring wheat, and winter wheat were major food crops globally, and they were sensitive to O₃ stress. A typical AOT40 exposure index was defined to assess the negative impact of O₃ exposure on crop yields. The AOT40 index was calculated by summing the hourly mean O₃ levels above 40 ppb during the 8 h over the crop growing season.

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$$AOT_{40}(ppbh) = \sum_{i=1}^{n} ([CO_3]_i - 40) [CO_3] \ge 40 \text{ ppb } (3)$$

where [CO₃]_i is the hourly O₃ (ppb), and n denotes the number of hours over the growing season. This growing season was determined by the University of Wisconsin Center for Sustainability and the Global Environment (UW SAGE) global crop calendar containing the planting and harvest dates by crop species and variety (Sacks et al., 2010; Schiferl et al., 2018). To date, some OTC/FACE experiments have been applied to assess the adverse effects of elevated O₃ concentrations on maize, rice, spring wheat, and winter wheat. The relationships between AOT40 and the relative yields (RY) for major crops have also been developed in recent years. The detailed equations are shown in Table S1. The relative yield loss (RYL) of crop is defined as

$$RYL=1-RY(4)$$

The estimated yield and economic losses are not only related to the RYL, while also associated with the grain yield in each grid. The detail equations are shown as follows:

$$CPL_i = RYL_i \times CP_i / (1 - RYL_i)$$
 (5)

where CPL_i is the estimated crop production loss and CP_i is the actual crop production in each grid during the study period.

The data about actual crop production in each grid were collected from The Agricultural Model Intercomparison and Improvement Project (AgMIP). The average value of simulated crop yields based on four models including DSSAT-Pythia, pDSSAT, LPJ-GUESS, and LPJ-ML were applied to estimate the actual crop production in each grid during 2015-2019, 2045-2049, and 2095-2099. We selected the simulate results of these models because they showed the better accuracy.

3. Results and discussions

3.1 Model evaluation

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Multi-source information data were integrated into the multi-stage model to predict firesourced MDA8 O₃ concentrations globally. At first, the global MDA8 O₃ simulation was evaluated. As illustrated in Figure S3, the 10-fold cross-validation (CV) results suggested that the R² value for MDA8 O₃ estimate reached 0.72. The root mean square error (RMSE) and mean absolute error (MAE) for MDA8 O₃ were 18.1 and 13.2 μg/m³, respectively (Figure S3). The CV R² value in our study reached 0.72, which was higher than that estimated by Liu et al. (2020) (0.64), indicating the satisfied predictive accuracy of O₃ estimates. However, the result was slightly lower than that (R²: 0.80 and 0.81) estimated by Xu et al. (2023) and Delang et al. (2021). It was supposed that the training samples in our study was much less than those used by Xu et al. (2023) (2000-2019 simulation) and Delang et al. (2021) (1990-2019 simulation). It was well known that the predictive accuracy was strongly dependent on the sample size (Li et al. 2020a, Li et al. 2020b). Overall, the predictive performance of ambient O₃ pollution was robust. Although the prediction capability of this model has been well validated, the accuracy for the fire-sourced MDA8 O₃ estimates could not be directly tested. It is well-known that potassium (K⁺) is often considered to be a fingerprint of wildfire, and thus we employ the relationship between ground-level K+ observations and wildfire-induced MDA8 O3 concentrations to examine the modelling accuracy. As shown in Figure S3, the correlation (R value) between observed K⁺ levels and fire-sourced MDA8 O₃ concentrations reached 0.67 (146 training samples), which was above 0.5 (p < 0.01). The results have confirmed that the wildfire-induced O_3 estimate showed the satisfied predictive performance. Although K⁺ has been often applied to reflect the wildfire contribution, the K⁺ could be also derived from anthropogenic emission and dust resuspension. To further validate the modelling performance of wildfire-related MDA8 O₃, the strong fire fingerprint (levoglucosan) was employed to construct the relationship with fire-sourced MDA8 O₃ concentrations. The results suggested that the R value (R = 0.73) was even higher than that between observed K^+ levels and fire-sourced MDA8 O₃ concentrations. Overall, the predictive performance was close to some previous studies (Childs et al. 2022, O'Dell et al. 2019, Xu et al. 2023), and thus we could use the result to further perform the data analysis.

3.2 Spatiotemporal trends of fire-sourced O₃ concentrations

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Global variations of fire-sourced MDA8 O₃ concentrations in historical and future scenarios are shown in Figure 1 and 2. From 2015 to 2019, the fire-sourced MDA8 O₃ level was in the order of Sub-Saharan Africa (SS) ($14.9 \pm 8.4 \,\mu\text{g/m}^3$) > South Asia (SA) ($4.0 \pm 2.5 \,\mu\text{g/m}^3$) > China ($1.6 \pm$ $0.7 \mu g/m^3$) > United States (US) $(1.3 \pm 0.9 \mu g/m^3)$ > Europe $(1.2 \pm 0.4 \mu g/m^3)$. In future scenarios, fire-sourced MDA8 O3 levels display marked spatial variability across different Shared Socioeconomic Pathways (SSPs). MDA8 O₃ showed the higher concentrations in some regions such as SS, SA, and US. Among all of the scenarios, fire-sourced O₃ levels displayed the highest concentrations in SS. It was assumed that this region possessed extensive burned area (52%) and higher biomass fuel consumption (5200 g C m⁻²) compared with other regions (van Wees et al. 2022). Following SS, SA also exhibited the higher wildfire-related MDA8 O₃ concentrations. The elevated concentrations of fire-sourced O₃ levels in SA were closely associated with exceptionally high fuel consumption (8600 g C m⁻²) (Chen et al. 2023, van Wees et al. 2022) though the burned areas were not very high among all of the regions. In addition, it should be noted that many previous studies have confirmed US showed the higher wildfire-induced PM_{2.5} or other aerosol components compared with many other regions (e.g., East Asia and South America) (Park et al. 2024, Xu et al. 2023). However, it did not show the higher O₃ concentrations in nearly all of the scenarios in our study. It was assumed that the MDA8 O₃ concentration exhibited significant latitudinal distribution (decreasing with the increase of latitude) globally. Both of China and Europe showed very low burned areas (0.2%) and fuel consumption (950 g C m⁻²), and thus the fire-sourced MDA8 O₃ concentrations were relatively lower compared with SS and SA. Besides, the fire-sourced MDA8 O₃ levels exhibited significant inter-annual trends and large discrepancy between different scenarios. The global average fire-sourced MDA8 O₃ concentrations showed overall increase from 2010s $(1.3 \pm 0.7 \,\mu\text{g/m}^3)$ to 2090s (SSP1-2.6, SSP3-7.0, and SSP5-8.5: 1.9 ± 0.9 , 1.6 ± 0.8 , and $1.4 \pm 0.7 \,\mu\text{g/m}^3$) for nearly all of the scenarios. The global average wildfirerelated MDA8 O₃ concentrations (the average of 2040s and 2090s) followed the order of SSP3-7.0 $(1.6 \pm 0.9 \,\mu\text{g/m}^3) > \text{SSP}5-8.5 \,(1.5 \pm 0.8 \,\mu\text{g/m}^3) > \text{SSP}1-2.6 \,(1.4 \pm 0.8 \,\mu\text{g/m}^3)$. The highest wildfirerelated MDA8 O₃ levels in SSP3-7.0 (air temperature: about 1.8°C higher than SSP1-2.6) and SSP5-8.5 (air temperature: about 2.3°C higher than SSP1-2.6) scenarios were contributed by the increased

fuel consumption and the warmer condition because O₃ level was more sensitive to air temperature increase (Wang et al. 2021, Wu et al. 2021).

Nevertheless, different regions showed distinct long-term trends. Wildfire-related MDA8 O₃ levels in nearly all of the regions in SSP3-7.0 scenario (air temperature: about 1.1°C higher than historical period) showed remarkable increases compared with the historical period because the warmer condition facilitated the rapid increase of O₃ level (Zhao et al. 2020). For low-carbon scenario (SSP1-2.6), the wildfire-related MDA8 O₃ concentrations in China, Europe, and US showed the relatively lower O₃ levels, whereas SA and SS still increased by 40% and 64%, respectively. The results suggested that the low-carbon pathway cannot effectively reduce the wildfire-induced O₃ pollution in both of SA and SS.

3.3 The crop yield losses caused by O₃ exposures

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As shown in Figure 3 and 4, the global crop yield losses caused by fire-sourced O₃ exposure have been quantified based on the equations 3-5. During historical period, the global fire-sourced O₃ caused 3.1 (2.4-3.8), 1.7 (1.5-1.9), 24 (21-27), and 43 (39-47) t/km² crop losses for maize, rice, spring wheat, and winter wheat, respectively. Compared with the historical period, CPL values in different future scenarios displayed large discrepancy. In SSP1-2.6 scenario, CPL of maize, rice, spring wheat, and winter wheat associated with fire-sourced O₃ exposure were 1.1 (0.9-1.3), 0.5 (0.4-0.6), 4.6 (4.1-5.4), and 4.6 (3.5-5.2) t/km², respectively (Figure S4-S11). However, CPL for maize (2.1 (1.9-2.3) and 2.4 (2.1-3.0) t/km²), rice (1.1 (0.9-1.3) and 1.3 (1.1-1.5) t/km²), spring wheat (557 (486-628) and 184 (154-218) t/km²), and winter wheat (258 (208-308) and 19 (14-22) t/km²) caused by fire-sourced O₃ exposure experienced dramatic increases in SSP3-7.0 and SSP5-8.5 scenarios (Figure S4-S11). There are two reasons accounting for the fact. First of all, the wildfire-related O₃ exposures showed marked increase in high-emission scenarios (Yang et al. 2022, Yue et al. 2017). Moreover, the crop yields also displayed substantial increases in both of these scenarios because rapid increase of fertilizer consumption (Brunelle et al. 2015, Randive et al. 2021). In addition, CPL caused by fire-sourced O₃ exposure also suffered significant spatial difference. During the historical period, the total CPL for four major foods caused by fire-sourced O₃ exposure in China, Europe, US, SA, and SS were 1451 (1302-1650), 65 (54-82), 61 (48-70), 56 (52-59), and 404 (372-425) t/km², respectively. In the future scenario (SSP1-2.6, SSP3-7.0, and SSP5-8.5), the

total CPL for four major foods caused by fire-sourced O₃ exposure in China, Europe, US, SA, and SS were 23 (19-28) (711 (630-802) and 339 (299-375)), 14 (12-16) (684 (596-768) and 32 (28-34)), 11 (8-12) (19 (17-22) and 21 (18-23)), 14 (12-15) (35 (30-39) and 21 (18-24)), 298 (272-320) (160 (145-179) and 745 (641-840) t/km², respectively. In both of historical and future scenarios, SS, SA, and China showed the higher CPL compared with other regions. The higher CPL in SS and SA might be attributable to the higher fire-sourced O₃ concentrations and crop yields. The higher CPL in China might be associated with exceptionally high crop yields though the wildfire-induced O₃ level was not very high. For most regions, CPL showed the higher values in high-emission scenarios (SSP3-7.0 and SSP5-8.5). Although SS and SA also showed the higher CPL in high-emission scenarios (SSP5-8.5), the CPL values of SS and SA in SSP1-2.6 scenario were still very high. The results suggested that the low-carbon policy still cannot effectively weaken local agricultural damage of fire-sourced O₃ exposure.

3.4 Implications and limitations

Our study developed a multi-stage machine-learning model based on the multi-source information data to predict the fire-sourced MDA8 O₃ concentrations at the global scale. It is the first study to use the ground-level observations as the constraint to improve the O₃ estimates in the future scenarios. The results confirmed that the model showed the better predictive accuracy and transferability.

Our assessment highlighted the severity and scale of the fire-sourced MDA8 O₃ level and a notable increasing trend in the future scenarios. Especially in high-emission scenarios (SSP3-7.0 and SSP5-8.5), the fire-sourced MDA8 O₃ showed the higher concentrations compared with the low-carbon scenario. Therefore, the global mean temperature increase should be limited to 2.0 °C or 1.5 °C above pre-industrial levels. In addition, both of SS and SA showed the highest wildfire-induced MDA8 O₃ concentration compared with other regions, indicating these hotspots should be determined to propose some control measures. For instance, wildfires could be partially controlled via effective evidence-based fire management and appropriate planning (González-Mathiesen and March 2021, Gonzalez-Mathiesen et al. 2021). Some prevention policy should be proposed to reduce agricultural waste incineration and some prescribed fires (Koul et al. 2022, Lange and Gillespie 2023). Some wildlands could be also changed into agricultural or commercial lands to

reduce the occurrence frequency of forest wildfire (Mansoor et al. 2022).

Besides, the impacts of fire-sourced O₃ pollution on crop yields were also quantified. The results confirmed China was faced of serious crop production losses, which was even higher than those in SS and SA because the higher crop production and increasing O₃ pollution risk in the future scenarios. Overall, crop yield losses of China showed significantly higher values in high-emission scenario (SSP3-7.0 and SSP5-8.5) compared with low-emission scenario (SSP1-2.6). The results suggested that low-carbon policy not only largely weaken O₃ pollution derived from anthropogenic emission in China, but also decrease wildfire-induced O₃ damages to crop yields effectively. The results also confirm that the carbon neutrality policy implemented in China possess sufficient agricultural benefits. In contrast, crop yield losses of SS and SA in low-carbon scenario still showed very high risks. It requires more stringent control measures to further reduce local anthropogenic emission in order to offset the wildfire-induced O₃ contribution.

It should be noted that our study is still subject to some limitations. Firstly, the future wildfire emission inventory still shows some uncertainties because the accuracy of land use types and burned areas in the future scenarios cannot be examined directly. Furthermore, in the historical estimates, we only used a chemical transport model (GEOS-Chem model) to simulate the fire-sourced O₃ concentrations though the ground-level observations were assimilated. However, only one model could increase the uncertainties because the O₃ background might be overestimated. Second, the chemical transport model used in our study did not account for plume rise, which could overestimate the contribution of wildfire emissions to O₃ pollution. Third, the ground-level observations of ambient O₃ are unevenly distributed around the world, which could limit the predictive accuracy of O₃ levels especially in some regions (e.g., SS and SA) lack of monitoring sites. In the future, it is highly necessary to add sufficient ground-level O₃ observations to further improve the accuracy of O₃ estimates. Finally, the zero-out method might suffer from some limitations because O₃ chemistry is highly nonlinear. More other methods such as air pollutant tracing method should be applied to quantify the fire-sourced O₃ concentrations combined with zero-out method. In the GEOS-Chem model, wildfire-emitted precursors (e.g., NO_x, VOCs) could be assigned unique "tags" as separate tracers. These tagged species undergo the same transport, chemistry, and deposition processes as regular emissions but are tracked independently. For ozone (O₃) attribution, the model calculates

345 the fraction of O₃ produced from wildfire-tagged NO_x/VOCs oxidation pathways. The tagged O₃ 346 concentrations are then extracted to quantify the wildfire contribution, while accounting for 347 nonlinear chemical interactions (e.g., NO_x saturation effects). The combination of multiple methods could increase the robustness of fire-sourced O₃ estimates. 348 349 Acknowledgements This work was supported by the National Natural Science Foundation of China (grant no. 350 351 U23A2030). 352 Data availability CMIP6 353 The this publication available https://esgfdataset used in is 354 node.ipsl.upmc.fr/search/cmip6-ipsl. 355 **Author contributions** 356 RL, DT, and HZ designed the study. RL developed the model. DT, YS, YG, and HZ analyzed the observations and model data. RL wrote the paper. RL and YS revised the paper. 357

The contact author has declared that none of the authors has any competing interests.

Competing interests

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Figure 1 The multi-year average concentrations of fire-sourced MDA8 O_3 (Unit: $\mu g/m^3$) during 2015-2019 (2010s) at the global scale (a). The latitudinal variations of fire-sourced MDA8 O_3 levels (Unit: $\mu g/m^3$) (b). The spatial distributions of fire-sourced MDA8 O_3 concentrations (Unit: $\mu g/m^3$) during 2015-2019 (2010s) (c). US, SA, and SS represent the United States, South America, and Sub-Sahara Africa, respectively. The difference of fire-sourced MDA8 O_3 concentrations in different regions (d).

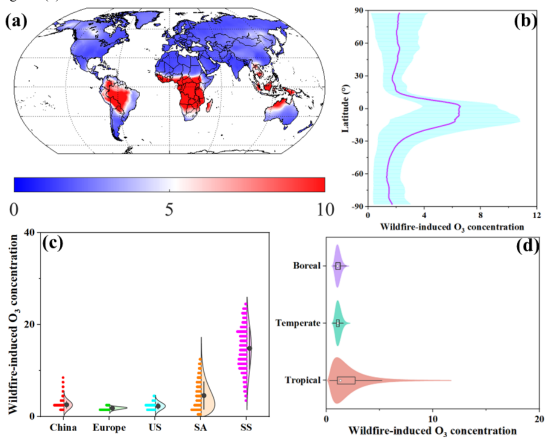


Figure 2 The global variations of fire-sourced MDA8 O_3 levels (Unit: $\mu g/m^3$) in SSP1-2.6 (a), SSP3-7.0 (b), and SSP5-8.5 (c) scenarios during 2040s. The spatial distributions of wildfire-related MDA8 O_3 concentrations (Unit: $\mu g/m^3$) in different regions during 2040s (d). US, SA, and SS represent the United States, South America, and Sub-Sahara Africa, respectively.

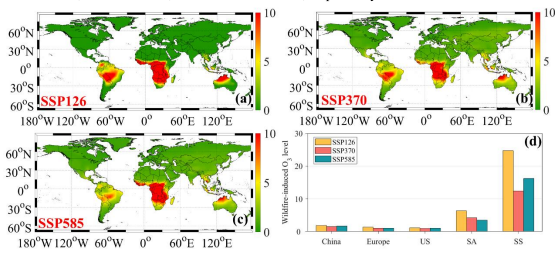


Figure 3 The global variations of fire-sourced O₃-related maize yield losses (Unit: t/km²) during historical (a), SSP1-2.6 (b), SSP3-7.0 (d), and SSP5-8.5 (e) scenarios during 2040s, respectively. The spatial variations of fire-sourced maize yield losses (Unit: t/km²) in major regions during 2040s. US, SA, and SS represent the United States, South America, and Sub-Sahara Africa, respectively.

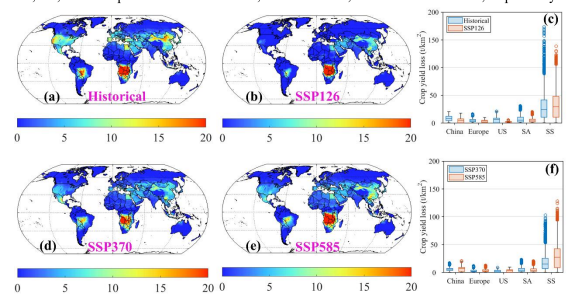
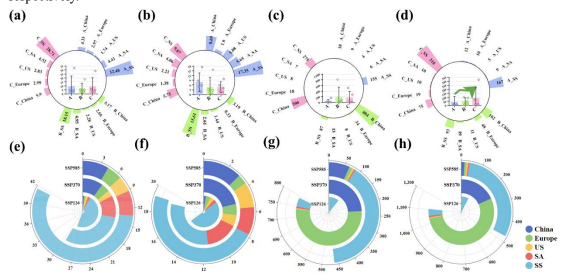


Figure 4 The spatial variations of fire-sourced O₃-related maize (a), rice (b), spring wheat (c), and winter wheat (d) yield losses (Unit: t/km²) during SSP1-2.6, SSP3-7.0, and SSP5-8.5 scenarios during 2040s, respectively. A, B, and C denote SSP1-2.6, SSP3-7.0, and SSP5-8.5 scenarios, respectively. (e)-(h) represent fire-sourced O₃-related maize (e), rice (f), spring wheat (g), and winter wheat (h) yield losses during SSP1-2.6, SSP3-7.0, and SSP5-8.5 scenarios during 2090s, respectively. US, SA, and SS represent the United States, South America, and Sub-Sahara Africa, respectively.



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