**General comments:** The manuscript “Inter-model comparison of global hydroxyl radical (OH) distribution and their impact on atmospheric methane over the 2000-2016 period” written by Yuanhong Zhao describes the inter-model differences in spatial distribution and temporal evolution of OH concentrations, and elucidates the impacts of simulated OH concentration fields on CH4 using the LMDz chemical transport model. The manuscript contains novel investigation to reveal inter-annual variations in OH and its impact on CH4 over recent decades using multi-model approach. The topic of the manuscript is certainly within the scope of ACP. Overall, the manuscript is well written and easy to follow. I would like to consider the publication of the manuscript from ACP, while I have several comments below which should be addressed before publication.

**Response:**

We thank the reviewer for the helpful comments. All of them have been addressed in the revised manuscript. Please see our itemized responses below.

**Specific comments:**

**Comments:** OH field Is the prescribed biogenic NMVOC emissions (p. 8, l. 187) climatology? Please clarify.

We mean here that some models just prescribed a fix scenario for NMVOC emissions and do not account for time variability.

Text has been clarified: “Biogenic NMVOC emissions in CESM and GEOSCCM are calculated based on the distribution of plant functional types and meteorology conditions with MEGAN, whereas the other models prescribe climatological biogenic NMVOC emissions.”

**Comments:** How did the authors prescribe the ECLIPSE and RCP85 emission inventories in the INCA simulations during the periods before 2004, between 2006-2009, and after 2011?
Response: We clarify by changing this sentence to" Anthropogenic emissions from Short-Lived Pollutants (ECLIPSE) inventory (Stohl et al., 2015) for 2005 and RCP 85 emission inventory (Riahi et al., 2011)) for 2010 are applied to every year of INCA NMHC-AER-S and INCA NMHC simulations, respectively.”

Comments: 2.2.2. Model simulations Please clarify how the OH increasing and decreasing rates are determined in the Run_OH_inc and Run_OH_dec simulations. Why are the rates +1 and −1.

Response: We clarify by add in the text : ” In order to assess the recent change in [OH], we tested two additional scenarios between 2010 and 2016: one with [OH] increase of +0.1% yr⁻¹ (Run_OH_inc) according to the slightly changing of OH calculated by ACCMIP models and one with [OH] decrease of -1% yr⁻¹ (Run_OH_dec) according to obviously decreasing of OH calculated by top-down approaches constrained by observations.”

Comments: 3.1. Spatial distributions of tropospheric OH The authors attributed possible causes of too large interhemispheric differences in OH in the CCMI models to model O3 and CO biases and unaccounted processes in some of the CCMI models, as reported by previous studies. Why is not the model performance on O3 and CO in the CCMI ensembles evaluated or referred? It might be better to cite Strode et al. (2016), Revell et al. (2018), and other papers.

Response: We acknowledge that the depth of analysis of the root causes of what we find here can be increased the lack of evaluation of these models in our paper. We have added in the text:

“Previous studies have attributed the inconsistency between the simulated and the observed OH N/S ratios to a model overestimation of O₃ and underestimation of CO over the Northern Hemisphere (Naik et al., 2013; Young et al., 2013; Strode et al., 2015), which have also been reported for CCMI models (Strode et al., 2016;
We have increased the depth of the analysis of the root causes possibly explaining what we find in the paper, all along with the text (see answers to reviewer 1).

Comments: 3.3. Factors contributing to inter-model differences Why the authors did not assess inter-model differences in tropospheric O3 burden? The tropospheric O3 burden should also affect primary production of OH.

Response: We have calculated global mean O3 mixing ratios averaged over the tropospheric and their pressure altitude levels in table 5 and move the values of O1(D) and reactive humidity, which contribute less to the inter-model difference of [OH] to the supplement (Table S4).

<table>
<thead>
<tr>
<th></th>
<th>CO ppbv</th>
<th>NO pptv</th>
<th>O3 ppbv</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>750</td>
<td>500</td>
<td>250</td>
</tr>
<tr>
<td>CESM1-CAM4Chem</td>
<td>76</td>
<td>71</td>
<td>70</td>
</tr>
<tr>
<td>CESM1-WACCM</td>
<td>75</td>
<td>70</td>
<td>69</td>
</tr>
<tr>
<td>CMAM</td>
<td>77</td>
<td>68</td>
<td>64</td>
</tr>
<tr>
<td>EMAC-L47MA</td>
<td>85</td>
<td>77</td>
<td>70</td>
</tr>
<tr>
<td>EMAC-L90MA</td>
<td>84</td>
<td>76</td>
<td>69</td>
</tr>
<tr>
<td>GEOSCCM</td>
<td>78</td>
<td>74</td>
<td>73</td>
</tr>
<tr>
<td>MOCAFE</td>
<td>67</td>
<td>68</td>
<td>67</td>
</tr>
<tr>
<td>MRI-ESM1r1</td>
<td>93</td>
<td>86</td>
<td>83</td>
</tr>
<tr>
<td>SOCOL3</td>
<td>79</td>
<td>73</td>
<td>74</td>
</tr>
<tr>
<td>Mean ± stand. dev.</td>
<td>79±7</td>
<td>74±6</td>
<td>71±5</td>
</tr>
</tbody>
</table>

And we have added in the text:

"To analyze inter-model differences in OH vertical distributions, we compared CO, NO, and O3 mixing ratios in table 5 as well as O(1D) photolysis rates and specific humidity in Table S4.

"Tropospheric O3 can also influence primary production of OH, and tropospheric
O₃ burden reflects combined effects of NOₓ, CO, and VOCs. The high O₃ over the lower troposphere simulated by SOCOL3 and the low O₃ over the upper troposphere simulated by MOCAGE can contribute to explain the high and low [OH] simulated the two models over the corresponding altitudes, respectively. ”

Comments: Do inter-model differences in vertical distribution of lighting NO production affect OH vertical distributions?
Response:
Yes indeed. We have added table S3 in the supplement:

Table S3. Lighting NOx emission (Tg N yr⁻¹) over three pressure altitudinal intervals and the total troposphere of CCMI models over 2000-2010.

<table>
<thead>
<tr>
<th>Model</th>
<th>Surface-750hPa</th>
<th>750-500hPa</th>
<th>500-250hPa</th>
<th>250-100hPa</th>
<th>tp</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMAM</td>
<td>0.7</td>
<td>0.4</td>
<td>1.5</td>
<td>1.7</td>
<td>4.2</td>
</tr>
<tr>
<td>EMAC-L90MA</td>
<td>0.2</td>
<td>0.5</td>
<td>1.3</td>
<td>1.8</td>
<td>3.7</td>
</tr>
<tr>
<td>CESM1-WACCMM</td>
<td>0.2</td>
<td>0.6</td>
<td>2.7</td>
<td>0.7</td>
<td>4.2</td>
</tr>
<tr>
<td>GEOSCCM</td>
<td>0.2</td>
<td>1.3</td>
<td>3.3</td>
<td>0.8</td>
<td>5.6</td>
</tr>
<tr>
<td>MOCAGE</td>
<td>0.3</td>
<td>1.2</td>
<td>2.4</td>
<td>1.0</td>
<td>4.8</td>
</tr>
<tr>
<td>MRI-ESM1r1</td>
<td>1.4</td>
<td>0.7</td>
<td>3.2</td>
<td>5.2</td>
<td>10.2</td>
</tr>
<tr>
<td>SOCOL3</td>
<td>0.2</td>
<td>0.8</td>
<td>2.1</td>
<td>1.4</td>
<td>4.4</td>
</tr>
</tbody>
</table>

We have also added in the text:”

L431-L435: “Lighting NOx emissions, which are mainly emitted in the middle and upper troposphere, can contribute to inter-model differences in NO and OH distributions (Murray et al., 2013; 2014). We compare lighting NOx emissions calculated by CCMI models in Table S3. High lighting NOx emissions simulated by MRI-ESM1r1 above 250hPa can explain high NO mixing ratios and increasing OH with altitude over the upper troposphere for this model (Fig. 3). However, High NO in the lower troposphere simulated by MOCAGE and SOCOL3 are not corresponding to high lighting NOx emissions in these models.”

L451: Lighting NOx emissions range from 3.7-10.2 Tg yr⁻¹(table S3)
Comments: ”3.4. Inter-annual variations of OH What is possible cause of significant positive [OH] trends over the tropics (p. 19, l. 454)?"

Response: We add in the text:” By comparing spatial distribution of OH trend with specific humidity (Fig.S6a), NOx and CO emissions (Fig. S6b), and stratospheric O3 (Fig.S6c), we find that positive OH trend over tropical regions are mainly corresponding to increases in water vapor (Fig. S6a)"

And we add figure S6a in the supplement:

**Figure S6a.** Spatial distribution of tropospheric specific humidity trends from 2000 to 2010 (in 10^{-2} \text{g/kg year}^{-1}). Black dots denote model grid-cells with statistically significant trends (p-value < 0.05).

Comments: 4.2.1. Spatial distributions of tropospheric CH4 mixing ratio Could you explain how inter-model differences in spatial and temporal OH variations affect the simulated global CH4 mixing ratio more in depth?

Response: In our paper, we attribute differences in LMDz simulated global mean CH4 mixing ratio to different global OH mean value and trend, and the spatial
distribution of CH4 to multi-model spread in OH spatial and temporal
distributions. To clarify our point, we have re-organized the first paragraph of
section 4.2.1:
" We used the scaled OH fields to perform simulations between 2000 and 2010.
Figure 6 shows the spatial distribution of tropospheric CH4 mixing ratios for the
simulation Run_standard (Table. 2, driven by OH with inter-annual variations)
averaged over 2000-2010. Although all simulations started from the same initial
conditions and OH fields were scaled to give the same global CH4 loss as INCA
NMHC in 2000, LMDz simulations using the different scaled OH fields still
generated a spread of tropospheric mean (8 ppbv) and spatial distribution in CH4
mixing ratios averaged during 2000-2010. Differences between the global
tropospheric mean [OH] cannot explain these differences (see Table 4). Clearly,
the different spatial (horizontal and vertical) and temporal variations of the OH
fields (as described in Sect. 3), which were kept in this experiment by only scaling
[OH] globally, significantly modify the simulated CH4 mixing ratios (Table 7 and
Fig. 6). OH fields with increasing trend will lead to lower LMDz simulated CH4
mixing ratios. The LMDz simulation using the TransCom OH fields (without inter-
annual variability) shows the highest CH4 mixing ratios (1735 ppbv), while the one
using the CMAM OH (with slightly increasing OH trend during the decade) shows
the lowest CH4 mixing ratios (1727 ppbv). "
And we add in the second paragraph:”” The differences in spatial distribution of
OH fields can influence LMDz simulated CH4 spatial distributions.”

Comments: Technical corrections: p. 13, l. 317: publication year is missing.
Response: We add the publication year, thank you very much for pointing out.