RESPONSES TO JOSEPH PITT

We thank the reviewer for his thoughtful comments that were helpful in improving the manuscript. Changes have been made in response to his specific comments listed below (in black). Our responses appear in red, and changes in the revised manuscript are highlighted in yellow.

1) The main issue I have with the study as it stands regards the way the model measurement comparison is performed. It is not clear that comparing the peak mole fraction enhancement is the best way to do this, and at the very least this needs more discussion. For one thing, for narrow plumes the peak measured enhancement is often dependent on the e-fold time of the cell. Looking at Fig. 2 this might not be an issue here (if the actual plume width is much larger than the measurement e-fold time multiplied by the flight speed) but it is hard to be certain. Providing these details in the text might help to clear this up.

Considering that the SPIRIT instrument allows measurements every 1.6 s and the e-fold time of the cell is 5.3 s, and that the average aircraft speed is 118 m s\(^{-1}\) on July 10 (103 m s\(^{-1}\) on July 14) during the period of the peaks of interest, even for the shortest lasting peak on each day (about 16 s on July 10 and about 19 s on July 14), the plume width is larger than the measurement e-fold time multiplied by the flight speed. We add a sentence for the worst case on July 10 (maximum aircraft speed with shortest peak) in the text, p.6 lines 21-24: “Moreover, SPIRIT allows measurements every 1.6 s. Considering the case on July 10 where we have the maximum aircraft speed (118 m s\(^{-1}\)) and the shortest peak (lasting about 16 s), the plume width is larger than the measurement e-fold time multiplied by the flight speed. Thus, for all the narrow peaks, the maximum plume concentration is real, not a plume diluted with its surrounding environment.”

An alternative approach would be to compare the integrated area under each of the plume transects. This would then give a better idea of the total amount of each species within the plume (in the same way as is usually employed for calculating species-species enhancement ratios). Both approaches could be employed alongside each other as long as a suitable discussion of the issues above is included.

This approach seems to be more reliable considering the dispersion modelling error in FLEXPART for example due to horizontal and vertical resolution of the windfield. However, not to complicate the reading of the paper, we decide to keep only this approach and remove the first one used by explaining why it is not possible to do a peak-to-peak comparison. The text (1st paragraph of section 4.2) was modified as follows:

“Concerning the second and the fourth peak (Fig. 2a), the measurements show two close peaks that FLEXPART cannot simulate individually, leading to a single and broader simulated peak. This is probably due to an error in the dispersion modelling induced by the horizontal and vertical wind field resolution that prevents us from comparing peak-to-peak concentrations. Even with a finer wind field grid mesh of 0.125°×0.125° (simulation not shown) such close peaks cannot be distinguished, suggesting a still insufficient spatial resolution. Instead, the integrated area under each of the measured and simulated plume transects will be compared and presented in Figure 3 with the percentages representing the relative differences with respect to SPIRIT measurements…”
According to this new approach, sensitivity tests with new fluxes were performed. They are summarized in Table 1. All the results of the simulations given now correspond to the integrated area under each peak (measured and simulated). We decided to summarize the results of all these new sensitivity tests by a figure instead of a table. This is illustrated in the new figure 3.

<table>
<thead>
<tr>
<th>Run name</th>
<th>Date of flight</th>
<th>NO₂ Flux (kg s⁻¹)</th>
<th>SO₂ Flux (kg s⁻¹)</th>
<th>CO Flux (kg s⁻¹)</th>
<th>Injection height (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTRL</td>
<td>20160710</td>
<td>0.07</td>
<td>4.23x10⁻³ kg s⁻¹</td>
<td>0.11 kg s⁻¹</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ST1</td>
<td>20160710</td>
<td>0.07</td>
<td>Not included</td>
<td>Not included</td>
<td>68</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td>77</td>
</tr>
<tr>
<td>ST2</td>
<td>20160710</td>
<td>0.035 - 0.05</td>
<td>Not included</td>
<td>Not included</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ST3</td>
<td>20160710</td>
<td>0.035 - 0.05</td>
<td>Not included</td>
<td>Not included</td>
<td>68</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td>77</td>
</tr>
</tbody>
</table>

Table 1. Flux and injection height for the reference control run (CTRL) and for the sensitivity tests (ST) for each day of flight.

To interpret the results of figure 3 and show the sensitivity of FLEXPART to the input parameters (flux or injection height), simple statistical tests were made and a paragraph was added:

“To determine whether the observed linear relationship between the percentages and the flux or the injection height occurs by chance, a simple F-test is performed, assuming that the variances are homogeneous and the results follow a Gaussian distribution. F statistic coefficients are calculated and compared to the 95% or 90% confidence interval with (1, N-2) (N: total number of results) as degrees of freedom (see values in brackets [u;+∞] in Figure 3). If the value F is included in the confidence interval then the relationship can be considered as linear. The standard errors on the slope are also added in the plots of Figure 3.

For the flight on July 10, the standard error of the slope coefficients and the F-test (95% of confidence) show linear relationships between the percentage difference and the flux. Only the results on July 14 for the plot with the injection height of VDI 3782 (1985, panel B-2) show a positive F-test but with a 90% of confidence. No conclusions can be drawn for the results on July 14 with the injection height of Briggs (1965, panel B-1). In order to show the response of FLEXPART to the injection height, panels A-3 and B-3 (Fig. 3) show the percentages versus the injection height (Briggs (1965) or VDI 3782 (1985)). FLEXPART shows similar results regardless of the injection height used as input whatever the flux used. All the cases show standard error on the slope coefficients larger than the slope itself and a F- value not included in the confidence interval (at 95% of confidence as shown in the Figure, and even 90%, not shown) . These results suggest that the differences between the two injection heights are not significant enough with respect to the vertical resolution of the model or the
measurements are too far to be influenced by the changes in this parameter. However, to really conclude about the injection height and to evaluate the flux, more measurements are needed at different altitudes and distances from the emission source. Besides the weather conditions and the functioning of the platform, the flight location is also an important parameter to be able to evaluate our measurements. Figures 2 b and e show the NO$_2$ plume simulated with FLEXPART as a function of distance from the source and altitude on July 10 and 14, respectively. The aircraft measurements, represented by the colored circles, are located at the upper part of the plume, away from the strongest concentrations. The work carried out is thus limited by the flight trajectories which were too high and too far from the FPSO platform to catch the part of the plume with the highest concentrations. The operational conditions during the flights were complex for the pilots, and safety concerns forced us to respect a minimum flight level (300 m) and a minimum distance from the source. Finally, we found that the NO$_2$ concentration difference between the measurements and the simulations does not seem to depend on the distance from the source since the measurements are already too far.

![Figure 3: Differences (in %) between SPIRIT integrated measurements and FLEXPART simulations depending on flux or injection height used as input in the model for A: the flight on July 10 and B: the flight on July 14. Panels A-1 and B-1 represent the change in the percentage with the flux by using the injection height from Briggs’ algorithm (1965; blue data; i.e. 27 m) and Panels A-2 and B-2 with injection height from VDI 3782 (1985; orange data; i.e. 68 m (A-2) or 77 m (B-2)). Panels A-3 and B-3 represent the change in the percentage with the injection height for the flux from Deetz and Vogel (2017; blue data; 0.07 kg.s$^{-1}$) and for the flux used in the sensitivity tests (green data, 0.04 kg.s$^{-1}$ for July 10 (A-3) and 0.035 kg.s$^{-1}$ for July 14 (B-3)). For all](image-url)
panels, triangles represent the data for all the peaks measured and squares represent the mean from these data. The slope, standard error values for the slope coefficients and the F statistic are added for all the plots.

I also think it would really help interpret what is going on here if Fig. 2 included the results from the other simulations. If you really think it’s getting too cluttered then these could be moved to the supplement, but I think it’s important to include them somewhere.

We think it is better not to overload Fig. 2 as it makes it harder to read. Thus, we added a figure (Fig. S4) in the supplementary material showing the results of the sensitivity tests done with FLEXPART. We show the sensitivity tests ST2 and ST3 with the smallest differences compared to SPIRIT measurements according to Fig. 3, i.e. with a flux of 0.04 kg s$^{-1}$ on July 10 and 0.035 kg s$^{-1}$ on July 14.

S4. (a) July 10: NO$_2$ concentration as a function of time for SPIRIT measurements (black) and FLEXPART sensitivity test (ST) 1 (green), ST2 (purple; with a flux of 0.04 kg s$^{-1}$), ST3 (orange; with a flux of 0.04 kg s$^{-1}$). (b) Similar to (a) for July 14 with a flux of 0.035 kg s$^{-1}$ for ST2 and ST3.

2) The lack of CO in the plumes measured on flight 1, and the subsequent detection of CO in the flight 2 plumes, is a really interesting result. However the discussion in Sect. 4.3 is fairly brief – it would be great to see this expanded. I’m not really clear as to what is meant by "more disturbed weather conditions" – does this mean the boundary layer was more turbulent? If so why does this mean that the combustion is less efficient? I’m not disputing that is the case, it’s just not obvious to me without further explanation. I can see that probing the fluid dynamics within the flare is beyond the scope here, but are there other studies that this finding could be linked to?

As far as we know, this finding could not be linked to any other study. The expression “more disturbed weather conditions” was removed and the explanations were clarified as follows: “The observed difference between July 10 and 14 in terms of CO emissions mostly lies in the different wind conditions between these two days: first, the wind speed was
lower on July 14, which makes less O\textsubscript{2} available to burn with natural gas; second, it appears that the wind direction was not clearly established, as can be seen from the much more dispersed plume in Fig. 1b, resulting in incomplete combustion pockets favoring CO formation. However, a decrease in efficiency should also lead to lower temperatures and NO\textsubscript{x} emissions, which is not observed here. The results of this campaign would require to be analyzed in the light of computational fluid dynamic simulations, accounting for a realistic natural gas composition and its high-temperature chemistry, which are beyond the scope of this study.”

Were there any measurements of CO\textsubscript{2} (and ideally CH\textsubscript{4}) on board? Then something quantitative could be said about the flare efficiency?

A Picarro instrument was brought to measure CO\textsubscript{2} during the campaign but it actually did not work at all because of a technical problem. CH\textsubscript{4} was measured by the SPIRIT instrument. Unfortunately, the flight conditions for these specific days were not ideal with very high temperatures inside the aircraft cabin that did not allow the CH\textsubscript{4} laser to work properly.

Specific points:

P2 L2 – In Nigeria all associated gas may well be flared, but in other places (at least in the UK) this associated gas is exported for use. So I think it would be more accurate to say that gas flaring is used to dispose of this natural gas in cases where the infrastructure to export it does not exist.

Modified following the recommendation: “Gas flaring is used to dispose of this natural gas in cases where the infrastructure to export it does not exist”

P3 L25 – "The concept...deeper water” – sentence reads awkwardly and needs rephrasing

It has been rephrased this way: “The concept of those platforms based on ship structure makes possible the development of small size oil fields and the exploitation of them further from the coast and thus in deeper water”

P3 L28 – I suggest "dispose of” rather than "eliminate"

We followed the suggestion and used “dispose of”.

P3 L30 – "mixture of gas” is ambiguous – presumably this means a mixture of emitted gases?

Yes, it has been modified as “This leads to a mixture of emitted gases”.

P4 L7 – "released along time" needs rewording

We modified it as follows: “It simulates long-range transport and dispersion of atmospheric tracers released over time”.

P4 L11 – If I understand correctly these are just tracer particles, so their assigned mass is just a nominal quantity used in the subsequent calculations (i.e. it does not correspond to a physical mass which impacts on the particle dispersion). If so I think it would be best to clarify this, as particle mass has quite a strong association for people who work with aerosols.

We agree, the model only considered NO\textsubscript{2} as tracer particles. All the particles undergo the same transport. The mass input in the “release” file is considered as a flux of NO\textsubscript{2} (a mass release during time). However, the other parameters such as molecular mass, OH reaction rate constant, Henry law constant… have also to be included and are taken into account for the wet and dry deposition calculations. We removed the sentence: “In this mode, each particle is associated to a given mass of tracer released during the time of the simulation” and add complementary information in the text and in Table 1: “The particles are released with the chemical properties of NO\textsubscript{2}, CO and SO\textsubscript{2} using constant emissions from Deetz and Vogel (2017) inventory during 7 h with a spin-up of 5 h, allowing the model to be balanced independently from the initial conditions. During the simulation, the NO\textsubscript{2} and SO\textsubscript{2} like particles mass is lost by wet and dry deposition and by OH reaction
(concentrations from GEOS-CHEM model; Technical note FLEXPART v8.2, http://flexpart.eu/downloads/26), which allows a lifetime of about 3 h at 298 K in the MBL for NO₂. CO like particles mass is only lost by OH reaction.”

P4 L20 – “for the DACCIWA project”

Modified following the suggestion.

P5 L2 – I think it would be useful to add a sentence in here explaining both the buoyancy and momentum effects. This would make it easier to understand the subsequent assertion that the momentum effect can be ignored.

We added a small definition for each effect in the text: “Indeed, the buoyancy corresponds to a density ratio between the air parcel and its colder surrounding environment and leads to the rise of this parcel under the influence of gravity. This effect is to be distinguished from the momentum effect defined as the product of an element mass by its velocity, which can be neglected for such high temperature plume (Briggs, 1965).”

P5 L31 – P6 L3 – The description of the terms in Eq. 4 is not easy to read. It might make it clearer to define the units of the constituent terms rather than the coefficient 74.4? Also the phrase “f as the fraction of radiated heat equals to 0.27” confuses me.

The units of all constituents, in particular Q and H (in MW), have been added in the manuscript.

The phrase has been rewritten as follows: “f (as) the fraction of radiated heat set to 0.27 by Deetz and Vogel (2017), after having averaged the f values given in Guigard et al. (2000)”.

P6 L30 – How is the background calculated – presumably by averaging the measured data outside the plumes? If so then it’s worth stating this.

It is true that the background has been calculated by averaging the measured data outside the plumes. We added a sentence in the text explaining it: “This value is an average of the measurements taken outside the plume”.

P7 L27 – "only a few quantities of pollutants" needs rewording

It has been modified as follows: “only a small fraction of pollutants”.

P8 L7 – "The turbulence increases the mixing and affects..." might be better

Yes, modified accordingly.

Fig. 1 – I trust your word that there was no SO₂ measured, but you might as well add an SO₂ trace to the CO plots just to demonstrate this point.

More precisely, we wrote that no SO₂ (and CO) peaks were detected simultaneously to NO₂ peaks. This is shown in the plots we added for the SO₂ measurements in Figure 1. The legend was also modified. SO₂ FLEXPART simulation was also realized and shows insignificant SO₂ concentration at the aircraft sampled location. A sentence has been added in the text: “Considering the very low SO₂ flux, the FLEXPART simulations in the CTRL run induce insignificant SO₂ concentration at aircraft sampled location (results not shown).”
(a) 

(b) 

(c) 

Particle number concentration (cm⁻³) 

NO₂ (ppbv) 

CO (ppbv) 

SO₂ (ppbv) 

Time (UTC) 

-1.0 1.3 1.6 1.9 2.2 2.5 2.8 3.1 3.4 3.7 -4.0 

NO₂ (ppbv) 

Time (UTC) 

-1.0 1.3 1.6 1.9 2.2 2.5 2.8 3.1 3.4 3.7 -4.0 

(d) 

Particle number concentration (cm⁻³) 

NO₂ (ppbv) 

CO (ppbv) 

SO₂ (ppbv) 

Time (UTC) 

-1.0 1.3 1.6 1.9 2.2 2.5 2.8 3.1 3.4 3.7 -4.0 

(1) 4.1 ppbv 

(2) 2.9 ppbv 

(3) 4.7 ppbv 

(4) 4.5 ppbv 

(1) 7.6 ppbv 

(2) 9.2 ppbv 

(3) 5.9 ppbv
Figure 1: (a) NO₂ concentration as a function of the flight trajectory downwind of the FPSO plume for July 10. The black arrows show the wind direction (from ECMWF). (c) NO₂, aerosol, CO and SO₂ concentrations as a function of time zoomed in a part of the flight trajectory in (a). The peaks studied are labelled by a number (from 1 to 4). (b) and (d) are similar to (a) and (c) for July 14.

Fig. 2  – Could you make the circles around the measured data in panels b) and e) more distinct please? At least on my screen it is really hard to make these out, especially in b). See also my main point 1)

The circles on Figure 2 have been highlighted as shown below. Moreover, a small error was detected on this figure. Fig. 2 (e) was not taken at the right time in the first version of the paper. This is modified in this version but it does not modify the conclusions of the article.
RESPONSES TO ANONYMOUS REFEREE #2

We thank the reviewer for his relevant comments that were helpful in improving the manuscript. Changes have been made in response to his specific comments listed below (in black). Our responses appear in red, and changes in the revised manuscript are highlighted in yellow.

General points:

Like reviewer 1, I am a little concerned about the way the model to measurement comparison has been done. It does seem that the peaks in the model are wider than the measured peaks and therefore comparing the maximum mixing rate enhancement of the two could give misleading results. The authors should try comparing the integrated area under the peaks and see if this gives a different result. The effect of this should at least be discussed in the paper.

→ We agree on this point. An error in the dispersion modelling has been mentioned by Referee #1 and leads to simulated peaks wider than the measured ones. We used the approach that consists in comparing the integrated areas and modified the text (2nd paragraph of section 4.2) as follows:

“Concerning the second and the fourth peak (Fig. 2a), the measurements show two close peaks that FLEXPART cannot simulate individually, leading to a single and broader simulated peak. This is probably due to an error in the dispersion modelling induced by the horizontal and vertical wind field resolution that prevents us from comparing peak-to-peak concentrations. Even with a finer wind field grid mesh of 0.125°×0.125° (simulation not shown) such close peaks cannot be distinguished, suggesting a still insufficient spatial resolution. Instead, the integrated area under each of the measured and simulated plume transects will be compared and presented in Figure 3 with the percentages representing the relative differences with respect to SPIRIT measurements.”

According to this new approach, sensitivity tests with new fluxes were performed. They are summarized in Table 1. All the results of the simulations given now correspond to the integrated area under each peak (measured and simulated). We decided to summarize the results of all these new sensitivity tests by a figure instead of a table. This is illustrated in Figure 3 and results are discussed from the 2nd paragraph of section 4.2 onwards.
The text needs to be clearer on what chemistry is used in most of the measurements. The authors also need to expand on how NO/NO₂ chemistry is treated in the model. It is not clear to me whether they are changing the NO and NO₂ emissions in the model to reproduce the NO₂ measurements or just NO. I would have thought most of the emission from the rig would occur as NO, with subsequent conversion to NO₂ before the measurements is made. The text needs to be clearer on what chemistry is used in the model.
FLEXPART model uses a very simple chemistry: the particles are released with chemical properties like NO\textsubscript{2} using constant emissions (mass release during time). During the simulation, the NO\textsubscript{2} like particles mass is lost by wet and dry deposition and by OH reaction, only.

The emissions used to initialize FLEXPART come from Deetz and Vogel (2017) inventory and are expressed in terms of NO\textsubscript{2}. They considered a rapid conversion for a part of the NO emissions into NO\textsubscript{2} very close to the source. This is confirmed with some quick simulations with a box model (AtChem/MCM) showing conversion in the first 5 minutes of the simulation. Most of the atmospheric models consider NO\textsubscript{2} as primary emissions while it is in fact a secondary emission. A sentence has been included in Section 3.2:

“Fourth, for such a temperature, NO\textsubscript{2} is considered as a primary pollutant coming from the rapid conversion of NO close to the source, and the inventory does not include any later transformation of the species”

Does the emission from the rig include non-flaring combustion (e.g. power generation)?

I would have thought that this would also be a significant source of NOx from a collocated but different source? Could this have been picked up in the measured plume but not included in the emission inventory?

The emission inventory (Deetz and Vogel) from the rig includes only flaring emissions. Emissions from power generation for the facility could be also a source of NO\textsubscript{2}. However, considering that the aircraft was flying at an altitude of about 300 m, only the emissions that have been heated up at very high temperature could reach rapidly and directly this altitude and induce a very localized spike in the NO\textsubscript{2} signal. This is why we think that we have only measured flaring emissions.

It would also be good to have a short discussion as to what actual effect the oil rig emissions have on air pollution in West Africa. For instance, if the emissions are doubled in the inventory, what effect does this have on NO2 and O3 levels at the coast? I realize a full study like this is beyond the scope of this paper but some short statement should be made as to the potential impact of underestimated emissions from oil rigs in the area.

The impact of the FPSO emissions on the air quality on the coast is not mentioned in the paper. In fact, the platform is situated 70 km downwind of the coast, which supposes that there is no impact of direct emissions of NO\textsubscript{2}. However, we agree that O\textsubscript{3} and other pollutants such as CO\textsubscript{2}, CH\textsubscript{4}, BC and CO with relatively long lifetime should impact the air quality on the coast. As you mentioned, “a full study like this is beyond the scope of this paper” but as suggested a short statement is included in the paper with more information directly included in the text (end of section 4.2) and in the conclusion:

“We can use the best simulation on each day to estimate the percentage of pollutants transported inside and above the MBL. In both cases, about 90% of the pollutants stay inside the MBL and are susceptible to impact the population living along the coastline. Measurements made along the coastline have shown that NO\textsubscript{2} concentrations are generally greater than 2 ppb for suburban sites and greater than 20 ppb near industrial sites (Bahino et al. 2018). Given the wind velocity (from 6.6 to 9.4 m \textsuperscript{s\textsuperscript{-1}}), the air masses attain the coast in 2 to 3 hours, which does not allow to bring significant NO\textsubscript{2} concentrations to impact air quality in this area. This is confirmed looking at FLEXPART simulations in Fig. 2b and 2e. They show that NO\textsubscript{2} concentrations are already very low (< 1 ppbv) from 40 km from the source on July 10, and even closer (from 20 km from the source) on July 14. The distance between the coast and the emission source following the wind direction being 70 km, only pollutants with a relatively long lifetime or secondary pollutant as O\textsubscript{3} can impact the air quality of the coast.”

“An estimation of the pollutant distribution above or inside the MBL shows that the pollutants stay mainly inside the MBL, limiting the transport to the coastline located 70 km downwind of the FPSO.
Were there measurements of CH$_4$ made on the aircraft? If so it would have been good to see this included in the study as the rigs could also be an important CH$_4$ source.

→ The SPIRIT instrument measured CH$_4$ during the campaign. Unfortunately, the flight conditions for these specific days were not ideal with very high temperatures inside the aircraft cabin (low altitude make less efficient the air conditioning) that did not allow the CH$_4$ laser to work properly.

Specific points:

P4 L27: Can the authors confirm if this is an NO$_2$ flux or a NOx flux?

→ We confirm that this is an NO$_2$ flux given in the inventory. A NO flux is also given but no direct NO$_x$ flux is provided.

P6 L15: Is this really true. Can it really be said that because no SO$_2$ was measured (on a relatively insensitive instrument) that no H$_2$S was present. The authors should at least put a lower limit on the H$_2$S that could be present.

→ It is well known the oxidation of H$_2$S fully leads to SO$_2$ formation. See any general book about Atmospheric Chemistry; we added the reference (Sonibare and Akeredolu, 2004) already quoted in the present paper. Sentence is added in the paper section 4.1:

“Knowing that SO$_2$ comes from H$_2$S combustion (Sonibare and Akeredolu, 2004), these results suggest that a gas composition of 0.03% of H$_2$S induces an emission of SO$_2$ concentration lower than the detection limit of the instrument from 3 km of our measurements or the natural gas composition given by Deetz and Vogel (2017) for the Niger Delta is different from that in Ghana for those two flights.”

P8 L16: this needs expanding, it is not clear what ‘disturbed weather conditions’ means and how this could effect the CO concentrations in the plume.

We agree this expression was rather confusing so we replaced it by a more specific one which should clarify the reason why CO formation is increased: “the wind direction was not clearly established, as can be seen from the much more dispersed plume in Fig. 1b, resulting in incomplete combustion pockets favoring CO formation.”

P8 L19: How will the results of the campaign improve computational flare fluid dynamics modelling?

→ This sentence was deleted
Local air pollution from oil rig emissions observed during the airborne DACCIWA campaign

Vanessa Brocchi¹, Gisèle Krysztofiak¹, Adrien Deroubaix², Greta Stratmann³, Daniel Sauer³, Hans Schlager³, Konrad Deetz⁴, Guillaume Dayma⁵, Claude Robert¹, Stéphane Chevrier¹, Valéry Catoire¹

¹ Laboratoire de Physique et Chimie de l’Environnement et de l’Espace (LPC2E), CNRS – Université Orléans – CNES, 45071 Orléans cedex 2, France.
² LMD and LATMOS, École Polytechnique, Université Paris Saclay, ENS, IPSL Research University; Sorbonne Universités, UPMC Univ Paris 06, CNRS, Palaiseau, France.
³ Institut für Physik der Atmosphäre, Deutsches Zentrum für Luft und Raumfahrt, Oberpfaffenhofen, Germany.
⁴ Karlsruhe Institute of Technology, Institute of Meteorology and Climate Research, Karlsruhe, Germany.
⁵ Institut de Combustion, Aérothermique, Réactivité et Environnement (ICARE), CNRS, 45071 Orléans cedex 2, France.

Correspondence to: Gisèle Krysztofiak (gisèle.krysztofiak@cnrs-orleans.fr)

Abstract. In the framework of the European DACCIWA (Dynamics-Aerosol-Chemistry-Cloud Interactions in West Africa) project, the airborne study APSOWA (Atmospheric Pollution from Shipping and Oil platforms of West Africa) was conducted in July 2016 to study oil rig emissions off the Gulf of Guinea. Two flights in the marine boundary layer were focused on the floating production storage and offloading (FPSO) vessel operating off the coast of Ghana. Those flights present simultaneous sudden increases of NO₂ and aerosols concentrations. Unlike what can be found in flaring emission inventories, no increase in SO₂ was detected and an increase in CO is observed only during one of the two flights. Using FLEXPART simulations with regional NO₂ satellite flaring inventory in forward trajectory mode, our study reproduces the timing of the aircraft NO₂ enhancements. Several sensitivity tests on the flux and the injection height are also performed, leading to the conclusion that a lower NO₂ flux helps better reproduce the measurements and that the modification of the injection height does not impact significantly the results of the simulations.

1 Introduction

Crude oil extraction from offshore platforms brings raw gas mixed with oil to the surface. Gas flaring is used to dispose of this natural gas in cases where the infrastructure to export it does not exist. This process emits a mixture of trace gases like carbon dioxide (CO₂), carbon monoxide (CO), sulfur dioxide (SO₂), nitrogen oxides (NOₓ) as well as particulate matter. Its impacts concern both the ecosystems (Nwaugo et al., 2006; Nwankwo and Ogagarue, 2011) and the air quality (Osuji and Avwiri, 2005). The pollutants can be transported into the free troposphere (Fawole et al., 2016b) or can reach coastal cities in the marine boundary layer (MBL). Flaring emissions can be derived from remote sensing techniques (Elvidge et al., 2013) by the widely used Visible Infrared Imaging Radiometer Suite (VIIRS) Nightfire satellite product (Deetz and Vogel, 2017).
Oil and gas extraction activities are however uncertain in terms of emitted quantities (Tuccella et al., 2017) and direct measurements are necessary. In Africa, most of the studies focused on the environmental impact of Nigerian oil platform emissions (e.g. Anomohanran, 2012; Hassan and Kouhy, 2013; Asuoha and Osu Charles, 2015; Fawole et al., 2016a) as it is part of the five countries with the highest flaring amounts (Elvidge et al., 2015). Other countries in the Gulf of Guinea are nevertheless affected by environmental problems. The DACCIWA (Dynamics-Aerosol-Chemistry-Cloud Interactions in West Africa; Knippertz et al., 2015) project conducted fieldwork in southern West Africa (SWA) in 2016 to investigate the impact of anthropogenic emissions, notably on air quality. One part of the project included airborne measurements of flaring emissions in order to fill the data gap on oil extraction activities in SWA. Those in situ measurements are, as far as we know, the first reported data on oil extraction activities in this region.

We use here the Lagrangian transport model FLEXPART (FLEXible PARTicle dispersion model; Stohl et al., 2005) in combination with an inventory of emissions dedicated to flaring emissions and created in the framework of the DACCIWA project (Deetz and Vogel, 2017) to reproduce the aircraft measurements. We focus on evaluating the sensitivity of the model to two parameters: the emission flux and the injection height of the flaring emissions. After a brief description of the DACCIWA project and the platform in Sect. 2, we present in Sect. 3 the model and the flaring emission inventory we used, as well as the estimation of the flaring plume injection height. We discuss the modeling results in Sect. 4.

2 The DACCIWA/APSOWA project

2.1 Description of the campaign

The EU-funded project DACCIWA focuses on the coupling between dynamics, aerosols, chemistry and clouds (Flamant et al., 2017). The research campaign was set up in June-July 2016 and undertook activities ranging from airborne measurements to running atmospheric numerical models. A map showing the location of the research site is presented in supplementary material (Fig. S1).

Our Atmospheric Pollution from Shipping and Oil Platforms of West Africa (APSOWA) project complementing DACCIWA seeks to characterize gaseous and particulate pollutants emitted by oil platforms off the coast of the Gulf of Guinea by dedicated flights conducted with the DLR (Deutsches Zentrum für Luft- und Raumfahrt) research aircraft Falcon-20. Different instruments for gas and particulate measurements were deployed onboard. We focus on CO, NO$_2$, SO$_2$ and aerosols measurements during two flights on July 10 and 14. Both CO and NO$_2$ were measured by SPIRIT (SPectromètre InfraRouge In situ Toute altitude, Catoire et al., 2017). This infrared absorption spectrometer uses continuous-wave distributed-feedback room-temperature Quantum Cascade Lasers (QCLs) allowing online scanning of mid-infrared rotational-vibrational lines with spectral resolution of $10^{-3}$ cm$^{-1}$. In the present campaign, the ambient air is sampled in a multipass cell with a pathlength of 134.22 m and in micro-windows around the 2179.772 cm$^{-1}$ line for CO and the 1630.326 cm$^{-1}$ line for NO$_2$. A home-made
software using the HITRAN 2012 database (Rothman et al., 2013) is used to deduce the total molecule abundance. The overall uncertainties are 4 ppbv for CO and 0.5 ppbv +5% for NO₂ at 1.6 s time resolution. SO₂ measurements are performed using a pulsed fluorescence SO₂ analyser (Thermo Electron, Model 43C Trace Level). Ultraviolet (UV) light is absorbed by SO₂ molecules in the sample gas which become excited and subsequently decay to a lower energy state. The emitted light is detected by a photomultiplier tube and is proportional to the SO₂ concentration in the sample gas. The time resolution of the measurements is 1 s, with a moving average of 30 s to smooth the data. The lower detection limit is 0.315 ppbv. The instrument was multipoint calibrated before and after the campaign. Total aerosol concentrations were measured with a butanol-based condensation particle counter (CPC TSI 3010, modified for aircraft use; Schröder & Ström, 1997 and Fiebig et al., 2005). The particle counter was mounted inside the fuselage behind the Falcon isokinetic aerosol inlet. The large particle cutoff diameter imposed by the inlet has been found to be between 1.5 and 3 µm depending on flight altitude (Fiebig, 2001). The lower cutoff diameter of the CPC was at ~10 nm. Particle losses due to diffusion effects were minimized by using a 5.75 std l/min bypass flow to the instrument. Using the particle loss calculator described in von der Weiden et al. (2009) we estimate the particle losses in the tubing to be less than 10% for the relevant size range. Flight sequences inside clouds are known to cause sampling artefacts and have therefore been removed from the data set.

2.2 Flight planning and FPSO description

The first Floating Production Storage and Offloading (FPSO) vessels have been installed in Indonesia in 1974. Since this year, their number has steadily increased (Shimamura, 2002). The concept of those platforms based on ship structure makes possible the development of small size oil fields and the exploitation of them further from the coast and thus in deeper water (Shimamura, 2002). Those platforms have other advantages (Shimamura, 2002): they are faster to build than other floating structures, they have inbuilt storage capability and thus do not necessitate pipelines, and they are movable and easily implantable on another oil field. Because of all those reasons, the FPSO systems will continue to develop. Among the techniques used to dispose of the gas associated with the crude oil extraction, one is the flaring, consisting in burning the gas in an open flame through a stack. This leads to a mixture of emitted gases which can reach the free troposphere if the meteorological conditions are favorable (Fawole et al., 2016b). The two targeted flights, on July 10 and July 14, consisted in about 3-4 hours of meandering transects through emission plumes in the MBL about 300 m above sea level (asl) off the coast of West Africa from Ivory Coast to Togo. We focus on the flights in the vicinity of the FPSO Kwame Nkrumah platform, on the Jubilee Field, off the coast of Ghana (4°35′47.04 N, 2°53′21.11 W). It measures 330 m in length by 65 m in width, and its height up to the top of the chimney is estimated to be 112 m asl. During the campaign, no contact with this FPSO could be obtained in order to have more information on its functioning.

3 Method

3.1 Lagrangian particle dispersion modelling

15
The Lagrangian model FLEXPART (FLEXible PARTicle; Stohl et al., 2005) is used to study the transport of the emitted plume from the FPSO in the MBL. It simulates long-range transport and dispersion of atmospheric tracers released over time by computing trajectories of a large number of tracer particles. Model calculations are based on meteorological data from the European Centre for Medium-Range Weather Forecasts (ECMWF), ERA-INTERIM L137 (Dee et al., 2011) extracted every 3 hours and with a horizontal grid mesh size of 0.5°×0.5°. The calculations are performed in forward dispersion mode with the model version 9.0. The particles are released with the chemical properties of NO$_2$, CO and SO$_2$ using constant emissions from Deetz and Vogel (2017) inventory during 7 h with a spin-up of 5 h, allowing the model to be balanced independently from the initial conditions. During the simulation, the NO$_2$ and SO$_2$ like particles mass is lost by wet and dry deposition and by OH reaction (concentrations from GEOS-CHEM model; Technical note FLEXPART v8.2, http://flexpart.eu/downloads/26), which allows a lifetime of about 3 h at 298 K in the MBL for NO$_2$; CO like particles mass is only lost by OH reaction.

3.2 Flaring emission inventory

Our study is based on a new gas-flaring emission inventory developed for the DACCIWA project (Deetz and Vogel, 2017). This inventory provides emissions of CO, CO$_2$, NO, NO$_2$ and SO$_2$ for June-July 2014 and 2015 and we use a 2016 updated version for the period of the APSOWA-DACCIWA campaign. It is based on remote sensing observations using VIIRS nighttime radiant heat in combination with combustion equations from Ismail and Umukoro (2014). The emission estimation method is described in detail in Deetz and Vogel (2017). Only the assumptions of interest for our study are indicated: first, the natural gas composition includes 0.03% H$_2$S (Sonibare and Akeredolu, 2004). Second, this composition measured in the Niger Delta is valid for West Africa in general. Third, the source temperature is deduced from the VIIRS measurements on a monthly mean. For the FPSO platform, it is set to 1600 K, which is a good order of magnitude since the flame temperature can be as high as 2000 K (Fawole et al., 2016b). Fourth, for such a temperature, NO$_2$ is considered as a primary pollutant coming from the rapid conversion of NO close to the source, and the inventory does not include any later transformation of the species. The CO, SO$_2$ and NO$_2$ fluxes estimated with this method for our two days of interest for the FPSO platform are $1.1\times10^{-1}$, $4.23\times10^{-5}$ and $7\times10^{-2}$ kg s$^{-1}$, respectively. Note that the highest uncertainties (+33; -79%) associated to the estimation of gas flaring emissions arise from the parameters required in the combustion equations, e.g. the gas composition, the source temperature and the flare characteristics.

3.3 Estimation of the flaring plume injection height

The oil flaring emissions are generally emitted at higher temperatures than the temperature of the surrounding environment, which implies an important role for the buoyancy at the stack exit. Indeed, the buoyancy corresponds to a density ratio between the air parcel and its colder surrounding environment and leads to the rise of this parcel under the influence of
This effect is to be distinguished from the momentum effect defined as the product of an element mass by its velocity, which can be neglected for such high temperature plume (Briggs, 1965). The buoyancy raises the plume above its initial injection height and can lead to a source height that can be several times higher than the real height of the stack (Arya, 1999). The calculation required to determine the rise of a plume depends on the wind conditions and the atmospheric stability. Before determining those parameters, we determine the MBL height that defines in which part of the troposphere we flew. The MBL is about 680 m asl on July 10 and 582 m asl on July 14 according to the European (ECMWF) and US (NCEP) operational forecasts. The wind conditions are determined by the Falcon-20 measurement system. An average of the wind speed measurements is calculated during the flight period in the vicinity of the platform (see Sect. 4.1). For July 10, an 8 minute mean wind speed of $9.4 \pm 0.5 \text{ m s}^{-1}$ has been calculated, where the standard deviation represents the natural variability, which is larger than the measurement uncertainty. For July 14, a 7 minute wind speed mean of about $6.6 \pm 0.7 \text{ m s}^{-1}$ has been calculated. Thus, for both days the conditions are windy.

Concerning the potential temperature for July 10 and 14, the atmosphere is considered as stable with a positive potential temperature gradient. With the previous parameters defined, it is possible to calculate the plume injection height $\Delta H$ (in m) by using Eq. (1) reported by Briggs (1965, 1984):

$$\Delta H = 2.6 \left(\frac{F_b}{u s^3}\right)^{1/3}$$

with $F_b$ as the buoyancy effect in $\text{m}^4 \text{s}^{-3}$ as defined in Eq. (2), $u$ as the wind speed (in $\text{m s}^{-1}$) and $s$ as the stability parameter in $\text{s}^2$ defined in Eq. (3) (Briggs, 1965):

$$F_b = g \frac{\Delta T}{T_s} w r^2$$

$$s = \frac{g}{T} \left(\frac{\partial T}{\partial z}\right) + \Gamma$$

$T, T_s, w$ and $r$ are the absolute temperature of the ambient air, the absolute temperature of the stack gases, the vertical velocity of the effluent at the stack exit and the radius of the stack, respectively. $\Delta T$ is the difference between the two temperatures. In Eq. (3), $g$ is the gravitational acceleration, $z$ the altitude and $\Gamma$ the adiabatic lapse rate. Both the temperature at the stack exit and the effluent velocity are taken from Deetz and Vogel (2017). The calculated plume injection height $\Delta H$ is 27 m for July 10 and 14. Briggs’ algorithm underestimates the plume rise in stable atmospheric condition (Akingunola et al., 2018). Therefore, another method has been tested to determine the injection height, based on VDI 3782 (1985). For a stable atmosphere, the injection height $\Delta H$ (in m) of the plume is determined by Eq. (4):

$$\Delta H = 74.4 \times Q^{0.333} x u^{-0.333}$$

with $u$ as the wind speed in $\text{m s}^{-1}$ and the unit of the coefficient (74.4) in $\text{s}^4 \text{ kg}^{-1} \text{ m}^{-2}$. The heat flow $Q$, in units of MW, (Eq. 5) is defined in Deetz and Vogel (2017) as:

$$Q = H^\frac{1}{f}$$
with \( H \) as the radiant heat observed by VIIRS, in units of MW, and \( f \) as the fraction of radiated heat set to 0.27 by Deetz and Vogel (2017), after having averaged the \( f \) values given in Guigard et al. (2000). The calculated plume injection heights are, according to Eq. (4), 68 m and 77 m for July 10 and 14, respectively.

The two methods of injection height calculation do not give similar results. The control run (CTRL; Table 1) will use Briggs’ algorithm with an injection height of the particles of 27 m for both days. A sensitivity test (ST1; Table 1) will be performed to see the impact on the results of the injection height in the model by using the injection height from VDI 3782 (1985).

4 Results and discussion

4.1 Description of the measurements

Figure 1 presents the part of the flights in the vicinity of the FPSO platform. During the flight on July 10, the flaring plume was crossed several times (Fig. 1a). It led to four NO\(_2\) peaks between 12:33 and 12:45 UTC at an altitude of around 300 m (Fig. 1c). Four simultaneous peaks of aerosols were measured (Fig. 1c). No simultaneous CO peaks (within its lower detection limit 0.3 ppbv; Catoire et al., 2017), neither SO\(_2\) peaks (within its lower detection limit 0.3 ppbv (see section 2.1) were detected in the plume of the FPSO platform for this flight. No peak was measured during a second series of transect at a higher altitude (around 600 m, Fig. 1a).

During the flight on July 14, three NO\(_2\) peaks were measured during the first transects between 10:48 and 11:00 UTC at around 300 m of altitude downwind the FPSO vessel (Fig. 1b). Those peaks were simultaneously measured with aerosol and CO peaks (Fig. 1d), but still no SO\(_2\) peak was detected. Knowing that SO\(_2\) comes from H\(_2\)S combustion (Sonibare and Akeredolu, 2004), these results suggest that a gas composition of 0.03% of H\(_2\)S induces an emission of SO\(_2\) concentration lower than the detection limit of the instrument from 3 km of our measurements or the natural gas composition given by Deetz and Vogel (2017) for the Niger Delta is different from that in Ghana for those two flights. The presence (or no) of CO peaks is discussed in Sect. 4.3.

Moreover, SPIRIT allows measurements every 1.6 s. Considering the case on July 10 where we have the maximum aircraft speed (118 m s\(^{-1}\)) and the shortest peak (lasting about 16 s), the plume width is larger than the measurement e-fold time multiplied by the flight speed. Thus, for all the narrow peaks, the maximum plume concentration is real, not a plume diluted with its surrounding environment.

4.2 FLEXPART simulations of the flaring emissions

In order to confirm that the peaks detected by the aircraft instruments correspond to the flaring emissions from the platform and to simulate them, forward trajectories are calculated using FLEXPART. First, as a reference, a simulation called “control run” (CTRL) is run, in which the NO\(_2\) flux for the FPSO platform is taken from Deetz and Vogel (2017), i.e. 7×10\(^{-2}\) kg s\(^{-1}\), and the injection height is 27 m, calculated with Briggs’ method (see Sect. 3.3). A comparison of the wind speed and
direction between ECMWF simulations and the Falcon measurements is presented in supplementary material. The wind speed and direction derived from ECMWF agree within 1 m s\(^{-1}\) and \(-1.8^\circ\), respectively, for July 10 (Fig. S2) and within \(-1\) m s\(^{-1}\) and \(-7^\circ\), respectively, for July 14 (Fig. S3). So the transport is well reproduced in FLEXPART. Figure 2a compares observed and simulated time series of NO\(_2\) concentrations for the flight on July 10. The four measured peaks are all remarkably well reproduced in time by FLEXPART. Another study by Tuccella et al. (2017), using WRF-Chem, was able to reproduce the simulated peaks simultaneously to the measured ones for oil facilities emissions in the Norwegian Sea. A NO\(_2\) background concentration has been added to the last two peaks. This value is an average of the measurements taken outside the plume and is representative of the ambient pollution not taken into account in FLEXPART as this model only simulates the pollution coming from the platform. Concerning the second and the fourth peak (Fig. 2a), the measurements show two close peaks that FLEXPART cannot simulate individually, leading to a single and broader simulated peak. This is probably due to an error in the dispersion modelling induced by the horizontal and vertical wind field resolution that prevents us from comparing peak-to-peak concentrations. Even with a finer wind field grid mesh of 0.125°×0.125° (simulation not shown) such close peaks cannot be distinguished, suggesting a still insufficient spatial resolution. Instead, the integrated area under each of the measured and simulated plume transects will be compared and presented in Figure 3 with the percentages representing the relative differences with respect to SPIRIT measurements. Figure 2d also compares the observed and simulated NO\(_2\) concentrations for the flight of July 14. The simulation gives three peaks concomitant with the three measured peaks of interest.

Comparisons between CTRL run and SPIRIT measurements for both flights (Fig. 3, panels A-1 for July 10, B-1 for July 14) show that the simulated concentrations are always overestimated for all the peaks (percentage larger than 0%). Sensitivity tests are performed in order to show the FLEXPART response to the flux. Fluxes lower than 0.07 kg s\(^{-1}\) are tested from 0.035 to 0.05 kg s\(^{-1}\). Figure 3 (panels A-1 and B-1) shows the variations in the differences between the measurements and the FLEXPART simulations with the flux used in FLEXPART for the injection height from Briggs (1965). The plots for both days show a linear response of FLEXPART to the flux increase with a best estimation reached for 0.04 kg s\(^{-1}\) for the flight on July 10 and 0.035 kg s\(^{-1}\) for the flight on July 14. Panels A-2 and B-2 of Figure 3 show similar conclusions but for the injection height from VDI 3782 (1985). To determine whether the observed linear relationship between the percentages and the flux or the injection height occurs by chance, a simple F-test is performed, assuming that the variances are homogeneous and the results follow a Gaussian distribution. F statistic coefficients are calculated and compared to the 95% or 90% confidence interval with (1, N-2) (N: total number of results) as degrees of freedom (see values in brackets \[u;+\infty\] in Figure 3). If the value F is included in the confidence interval then the relationship can be considered as linear. The standard errors on the slope are also added in the plots of Figure 3.

For the flight on July 10, the standard error of the slope coefficients and the F-test (95% of confidence) show linear relationships between the percentage difference and the flux. Only the results on July 14 for the plot with the injection height of VDI 3782 (1985, panel B-2) show a positive F-test but with a 90% of confidence. No conclusions can be drawn for the results on July 14 with the injection height of Briggs (1965, panel B-1). In order to show the response of FLEXPART to the
injection height, panels A-3 and B-3 (Fig. 3) show the percentages versus the injection height (Briggs (1965) or VDI 3782 (1985)). FLEXPART shows similar results regardless of the injection height used as input whatever the flux used. All the cases show standard error on the slope coefficients larger than the slope itself and a F- value not included in the confidence interval (at 95% of confidence as shown in the Figure, and even 90%, not shown) . These results suggest that the differences between the two injection heights are not significant enough with respect to the vertical resolution of the model or the measurements are too far to be influenced by the changes in this parameter. However, to really conclude about the injection height and to evaluate the flux, more measurements are needed at different altitudes and distances from the emission source. Besides the weather conditions and the functioning of the platform, the flight location is also an important parameter to be able to evaluate our measurements. Figures 2 b and e show the NO\textsubscript{2} plume simulated with FLEXPART as a function of distance from the source and altitude on July 10 and 14, respectively. The aircraft measurements, represented by the colored circles, are located at the upper part of the plume, away from the strongest concentrations. The work carried out is thus limited by the flight trajectories which were too high and too far from the FPSO platform to catch the part of the plume with the highest concentrations. The operational conditions during the flights were complex for the pilots, and safety concerns forced us to respect a minimum flight level (300 m) and a minimum distance from the source. Finally, we found that the NO\textsubscript{2} concentration difference between the measurements and the simulations does not seem to depend on the distance from the source since the measurements are already too far.

We can use the best simulation on each day to estimate the percentage of pollutants transported inside and above the MBL. In both cases, about 90% of the pollutants stay inside the MBL and are susceptible to impact the population living along the coastline. Measurements made along the coastline have shown that NO\textsubscript{2} concentrations are generally greater than 2 ppb for suburban sites and greater than 20 ppb near industrial sites (Bahino et al. 2018). Given the wind velocity (from 6.6 to 9.4 m s\textsuperscript{-1}), the air masses attain the coast in 2 to 3 hours, which does not allow to bring significant NO\textsubscript{2} concentrations to impact air quality in this area. This is confirmed looking at FLEXPART simulations in Fig. 2b and 2e. They show that NO\textsubscript{2} concentrations are already very low (< 1 ppbv) from 40 km from the source on July 10, and even closer (from 20 km from the source) on July 14. The distance between the coast and the emission source following the wind direction being 70 km, only pollutants with a relatively long lifetime or secondary pollutant as O\textsubscript{3} can impact the air quality of the coast.

Considering the very low SO\textsubscript{2} flux, the FLEXPART simulations in the CTRL run induce insignificant SO\textsubscript{2} concentration at aircraft sampled location (results not shown). We performed CO simulations as CO peaks were measured in one case out of two (Fig. 2c and 2f). For those simulations, we used the injection height of 27 m of the CTRL run and the flux from Deetz and Vogel (2017) of 1.1x10\textsuperscript{-3} kg s\textsuperscript{-1}. For the flight on July 10, FLEXPART simulates four CO peaks concomitant with the four NO\textsubscript{2} peaks while no increase in CO has been observed. On July 14, three peaks of CO are simulated at the same time as the three measured ones, but underestimated. CO is always included as a gas emitted from flaring in the inventory for this specific vessel while it does not seem to be the case each time. A discussion on the flaring combustion processes is presented below.
4.3 Combustion processes involved in oil flare

From a combustion point of view, flares generate non-premixed highly turbulent flames, characterized by high-frequency fluctuating flow fields. Flares can be air-assisted or steam-assisted in order to achieve a better efficiency. The turbulence increases the mixing and affects the chemical reaction process. Very recent attempts to model such flames (Aboje et al., 2017; Damodara et al. 2017) showed the influence of the composition of the natural gas and the validity of the chemical kinetic mechanism are of primary importance to predict the emitted species. Moreover, the flame characteristics such as temperature, height, and length, have a strong impact on the dispersion of the plume together with wind speed and other meteorological variables (Rahnama et al., 2016). All these parameters need to be taken into account when considering air field campaigns.

McDaniel (1983) clearly demonstrated CO emissions are strongly related to flare efficiency. When the efficiency is ca. 99.8%, CO is below 10 ppmv (Fig. A-5 in McDaniel (1983)) while CO emissions may reach 440 ppmv when the efficiency drops to 64% (Fig. A-8 in McDaniel (1983)). CO emissions can clearly be linked to the quality of the combustion, which is directly impacted by the turbulence. The observed difference between July 10 and 14 in terms of CO emissions mostly lies in the different wind conditions between these two days: first, the wind speed was lower on July 14, which makes less O2 available to burn with natural gas; second, it appears that the wind direction was not clearly established, as can be seen from the much more dispersed plume in Fig. 1b, resulting in incomplete combustion pockets favoring CO formation. However, a decrease in efficiency should also lead to lower temperatures and NOx emissions, which is not observed here. The results of this campaign would require to be analyzed in the light of computational fluid dynamic simulations, accounting for a realistic natural gas composition and its high-temperature chemistry, which are beyond the scope of this study.

5 Conclusions

This study was conducted in the framework of the DACCIWA FP7 European project in July 2016 in southern West Africa. One target of the project was to measure emissions from oil rigs, which were not well estimated until then. With two flights planned in the vicinity of a FPSO platform, we reported the first flaring in-situ measurements in this region. The aim of this work was to evaluate the capacity of FLEXPART model to reproduce the NO2 airborne measurements and to evaluate the inventory of Deetz and Vogel (2017) in the case of point sources of pollution such as oil platforms. The injection height of the plume was estimated by performing different calculations. According to several sensitivity runs, it appears that the emission flux given by Deetz and Vogel (2017) overestimates the concentrations whereas a lower NO2 flux roughly reproduces the measurements. However, we did not know if the FPSO was under standard conditions of functioning. Concerning the injection height, the sensitivity tests are not conclusive, showing the need for more and better-targeted measurements. An estimation of the pollutant distribution above or inside the MBL shows that the pollutants stay mainly inside the MBL, limiting the transport to the coastline located 70 km downwind of the FPSO. Sources of uncertainties are associated with the different calculations and hypotheses but the work is mainly limited by the flights trajectories, too far and too high from the platform. So it remains necessary to better quantify the emissions released
during the flaring processes, locally but also at wider scales. Generally speaking, this study suggests that for flights planned in the heart of a flaring plume, it should be possible to link the flaring observations obtained by satellites with the emissions deduced from the airborne measurements. If this relationship is possible, a general relationship between the emissions and the radiant heat could thus allow estimating the emissions of all flaring processes detected by satellite.

Data availability. The aircraft data used here can be accessed using the DACCIWA database at http://baobab.sedoo.fr/DACCIWA/ (Brissébrat et al., 2017). A two-year embargo period applies after the upload. External users can request the release of datasets before the end of the embargo period.

Author contributions. HS was the mission scientist for the Falcon-20 aircraft and VC the PI of the EUFAR2-APSOWA campaign. VC, SC and VB participated in the SPIRIT measurements onboard the aircraft with remote assistance from CR for the SPIRIT maintenance. VB, GS and DS performed flight data analyses for their respective instruments. KD provided the updated version of his inventory for the period of the campaign and the VDI report. AD performed simulations using WRF-Chem to simulate the FPSO plume and gave advice for the modeling method. GK and VB performed the FLEXPART simulations. GD conducted the combustion part of the paper. VB wrote the manuscript with contribution from all co-authors.

Acknowledgments. We thank P. Jacquet for instrumental support before and during the campaign. The DLR crew is acknowledged for flying operations. This work was funded by the EU FP7 EUFAR2 Transnational Access project and DACCIWA project (Grant Agreement N°603502), the Labex VOLTAIRE (ANR-10-LABX-100-01), the PIVOTS project provided by the Région Centre – Val de Loire (ARD 2020 program and CPER 2015–2020) and the APSOWA project from INSU-LEFE-CHAT program. We thank F. Contino (Vrije Universiteit Brussel, Belgium) for the work undertaken on the simulations of flame. We also thank G. De Coetlogon and A. Weill (Laboratoire Atmosphères Milieux Observations Spatiales, Université Paris-Saclay and CNRS, France) for their help in determining the MBL height.

References


Figure 1: (a) NO$_2$ concentration as a function of the flight trajectory downwind of the FPSO plume for July 10. The black arrows show the wind direction (from ECMWF). (c) NO$_2$, aerosol, CO and SO$_2$ concentrations as a function of time zoomed in a part of the flight trajectory in (a). The peaks studied are labelled by a number (from 1 to 4). (b) and (d) are similar to (a) and (c) for July 14.
Figure 2. Left column: July 10. (a) \(\text{NO}_2\) concentration as a function of time for SPIRIT measurements (black) and FLEXPART CTRL run simulation (red). (b) Vertical section of the simulated \(\text{NO}_2\) plume (CTRL run) as a function of distance from the source and altitude. The colored circles correspond to the measurement peaks. (c) CO concentration as a function of time for measurements (black) and FLEXPART CTRL run simulation (red). Right column: July 14. (d), (e) and (f) similar to (a), (b) and (c).
Figure 3: Differences (in %) between FLEXPART simulations and SPIRIT integrated measurements depending on flux or injection height used as input in the model for A: the flight on July 10 and B: the flight on July 14. Panels A-1 and B-1 represent the change in the percentage with the flux by using the injection height from Briggs’ algorithm (1965; blue data; i.e. 27 m) and Panels A-2 and B-2 with injection height from VDI 3782 (1985; orange data; i.e. 68 m (A-2) or 77 m (B-2)). Panels A-3 and B-3 represent the change in the percentage with the injection height for the flux from Deetz and Vogel (2017; blue data; 0.07 kg.s\(^{-1}\)) and for the flux used in the sensitivity tests (green data, 0.04 kg.s\(^{-1}\) for July 10 (A-3) and 0.035 kg.s\(^{-1}\) for July 14 (B-3)). For all panels, triangles represent the data for all the peaks measured and squares represent the mean from these data. The slope, standard error values for the slope coefficients, the F statistic and the confidence interval (I\(_{95}\%\) or I\(_{99}\%\) only for panel B-1 and B-2) are added for all the plots.
<table>
<thead>
<tr>
<th>Run name</th>
<th>Date of flight</th>
<th>NO\textsubscript{2} Flux (kg s\textsuperscript{-1})</th>
<th>SO\textsubscript{2} Flux (kg s\textsuperscript{-1})</th>
<th>CO Flux (kg s\textsuperscript{-1})</th>
<th>Injection height (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTRL</td>
<td>20160710</td>
<td>0.07</td>
<td>4.23\times10\textsuperscript{-5}</td>
<td>0.11</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ST1</td>
<td>20160710</td>
<td>0.07</td>
<td>Not included</td>
<td>Not included</td>
<td>68</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td>77</td>
</tr>
<tr>
<td>ST2</td>
<td>20160710</td>
<td>0.035 - 0.05</td>
<td>Not included</td>
<td>Not included</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ST3</td>
<td>20160710</td>
<td>0.035 - 0.05</td>
<td>Not included</td>
<td>Not included</td>
<td>68</td>
</tr>
<tr>
<td></td>
<td>20160714</td>
<td></td>
<td></td>
<td></td>
<td>77</td>
</tr>
</tbody>
</table>

Table 1. Flux and injection height for the reference control run (CTRL) and for the sensitivity tests (ST) for each day of flight.