Title: Effectiveness of replacing catalytic converters in LPG-fueled vehicles in Hong Kong
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Responses to the comments from Anonymous Referee #2

General comments:

This paper provides information on the impact of the replacement of catalytic converters in vehicles fuelled by LPG in Hong Kong. The policy was based on the need to maintain the effectiveness of the catalysts; it was extensive and conducted over a period of several months over 2013 – 2014. The paper reports roadside and background measurements of VOCs, NOx, O3 and CO and a set of analyses to investigate the effectiveness of the programme. The measurements and analysis are of considerable interest and of relevance to the development of effective urban air quality policies. The reduction in LPG-related VOCs was substantial over only a 9 month period. The paper, however, has a number of deficiencies related either to the clarity of the presentation or to the methodologies used in the analysis and interpretation.

We thank the reviewer very much for his/her valuable comments and suggestions, which helped improve the manuscript substantially. Apart from the technical/typographical comments, the referee’s concerns mainly focused on three aspects: (i) the background of this study - more information about the program, the physico-chemical processes and previous O3 modeling in Hong Kong. (ii) the methods, including the method to obtain the primary emissions, the photochemical simulation with the backgrounds being subtracted, and the subtraction method to obtain O3 production by LPG and the atmospheric reactivity (i.e. OH, HO2 and their formation/loss pathways). (iii) the results, mainly the source apportionment results and the simulation of OH, HO2 and pathways of HOx. To answer these questions, we have supplemented and revised the manuscript accordingly. The method removing the backgrounds and the subtraction method were abandoned in the source apportionment and in the simulation of O3, OH, HO2 and the pathways of OH and HO2, which resulted in reasonable source apportionment results and the reactivity simulations. Our responses to the reviewer are as follows, along with indications of how the manuscript has been further revised for consideration by ACP. We hope
that these changes will further strengthen the main points and make them clearer in the revised manuscript.

Specific comments:

1. More information should be given to aid the understanding of the general situation in Hong Kong and the basis of this intervention programme. (a) A brief review of the regional physico-chemical processes and atmospheric transport and their extent in influencing air quality in Hong Kong. There have been several such studies and a brief discussion would be helpful. (b) A tabulation of typical VOC concentrations at the roadside and background sites – at present it is difficult to assess the importance of the LPG-related VOCs – a number of values are given for the fractional contribution to the total atmospheric VOC burden, but real data on all significant VOCs measured, perhaps with the pseudo first order rate constant for OH removal (i.e. kOH + VOC [VOC]), would be of value.

Many thanks for the good suggestion. A brief review of the regional physico-chemical processes and atmospheric transport and their extent in influencing air quality in Hong Kong was added in the revised manuscript:

Apart from local emissions, air quality in Hong Kong is also affected by regional transport. In autumn and winter, prevailing northerly and northeasterly winds bring anthropogenic emissions from mainland China to Hong Kong, enhancing local pollution burden. In summer, clean oceanic air masses dilute the levels of air pollutants under the influence of southerly winds (Ling et al., 2013; Guo et al., 2009, Wang et al., 2005; So and Wang, 2003; Chan and Chan, 2000). Ling et al. (2013) quantified the contribution of regional transport as 0-61% for summer O₃ and 0-56% for autumn O₃ on high O₃ days in Hong Kong. Guo et al. (2006) found that over 45% of VOC samples collected from August 2001 to December 2002 at Tai O, Hong Kong, were influenced by regional transport from inland China. Based on numerical simulations, Wang et al. (2006) quantified regional contributions of 40-90% to O₃ formation in Hong Kong, and Huang et al. (2006) indicated that the regional contribution was <50% when a trough was over the South China Sea (SCS), and it increased to >50% in the presence of an anticyclone over mainland China and/or a tropical cyclone over SCS. Furthermore, Wang et al. (2006) revealed that the regional contribution was lower during daytime than that at night, while Lam et al. (2006) found that the contribution decreased from >50% during an O₃ episode to 30-50% after the episode. In
addition, Jiang et al. (2008) reported that the contribution of horizontal transport to surface \( \text{O}_3 \) in Hong Kong was negligible during a typhoon-induced \( \text{O}_3 \) episode. However, all the studies focused on the regional contribution at non-roadside sites, and the \( \text{O}_3 \) chemistry at roadside sites in Hong Kong was seldom reported. Given high \( \text{NO}_x \) levels, tall buildings and low wind speeds along the main roads in Hong Kong, it is expected that the regional impact on air pollutants in roadside environment is relatively minor, especially during daytime hours on non-\( \text{O}_3 \) episode days.

For details, please refer to lines 71-92, pages 3-4 in the revised manuscript.

The average VOC concentrations and their OH-reactivity at the roadside and background sites were also provided:

Table S2 summarizes average mixing ratios of the main VOCs and trace gases, and their OH-reactivity at the roadside (MK) and background site (HT) before (October 2012-September 2013) and during (October 2013-May 2014) the program in Hong Kong. The much higher OH-reactivity at MK (“before”: 86.6±6.0 s\(^{-1}\); “during”: 77.9±6.1 s\(^{-1}\)) than at HT (“before”: 5.0±0.3 s\(^{-1}\); “during”: 4.9±0.1 s\(^{-1}\)) meant that the OH lifetime was much shorter at MK (~0.01 s) than at HT (0.2 s), and even shorter than that at an urban site in New York (0.05 s) (Ren et al., 2003), indicating that reactive species were extremely abundant in the roadside environment of Hong Kong. The LPG-related VOCs (i.e., propane and \text{n/i}-butanes) contributed 56.5±2.7% and 46.0±3.1% to total VOCs, and 18.7±1.2% and 15.1±1.1% to the total OH-reactivity of VOCs at MK, before and during the program, respectively. These fractional contributions suggest the importance of LPG-related VOCs in the budgets of VOCs and \( \text{O}_3 \) in Hong Kong. Consistent with the decreases of fractional contributions once the program started, the mixing ratios (29.4±3.3 and 21.0±2.2 ppbv for before and during the program, respectively) and OH-reactivity (1.12±0.04 and 0.79±0.02 s\(^{-1}\) before and during the program, respectively) of LPG-related VOCs significantly declined at MK (\( p < 0.05 \)), while at HT their mixing ratios increased from 1.5±0.1 ppbv to 1.9±0.04 ppbv, and the OH-reactivity increased from 0.06±0.004 s\(^{-1}\) to 0.07±0.003 s\(^{-1}\).

The minor variation of background LPG-related VOCs relative to the decreases at MK (4.8% in mixing ratios and 3.0% in OH-reactivity of VOCs) indicated that the influences of the background concentrations on the variations of LPG-related VOCs and \( \text{O}_3 \) at MK were minimal and can be neglected.
For details, please refer to lines 169-188, page 7 in the revised manuscript and Table S2 in the supplementary material.

2. Regional processes are excluded from the analysis on the basis of the propane /CO ratio at the main roadside site. The atmospheric lifetimes of the LPG VOCs are considerable (For [OH] $10^6$ cm$^{-3}$ and rate constants of $1-3 \times 10^{-12}$ molecule cm$^{-3}$ s$^{-1}$ the lifetime is 100 h) Even for a windspeed of 2 m s$^{-1}$, ozone will be generated from such gases over 1000 km. So while the roadside concentrations of the VOCs are arguably dominated by local sources, chemistry and ozone production are necessarily regional, although titration and the establishment of the NO$_x$/O$_3$ stationary state are local. Since much of the emphasis of the paper is on ozone formation, some discussion of the area over which the new controls operate, and reference to previous Eulerian / Lagrangian analyses of ozone formation in the Hong Kong region, would be of value. Note that on p35949, line 27, the authors refer to the O$_3$ calculation as being “purely local”.

Thanks very much for the excellent comment and suggestion. We agree that the regional transport of O$_3$ can occur, even at the very low wind speed (actual average=1.3 m/s at MK), because O$_3$ can be formed in the source region and then transport to Hong Kong, and/or during the transport of O$_3$ precursors. As suggested, the regional impacts on O$_3$ levels in Hong Kong were discussed based on previous numerical simulation study. In addition, we also provided more information about the consideration of regional transport influence in the present PBM-MCM model used in this study. Since the reviewer considered that the subtraction method used in previous version was inappropriate and led to strange results, we checked both “with” and “without” input of background concentrations of air pollutants, and found that the impact of the background O$_3$ precursors on the O$_3$ simulation at the roadside site was negligible. Hence, in the revised manuscript, we used the whole-air ambient concentrations of VOCs and trace gases at roadside site to simulate O$_3$ formation, rather than the subtraction method. All the revisions were shown as follows. The inappropriate wording such as “purely local” was deleted in the revised manuscript.

Based on numerical simulations, Wang et al. (2006) quantified regional contributions of 40-90% to O$_3$ formation in Hong Kong, and Huang et al. (2006) indicated that the regional contribution was <50% when a trough was over the South China Sea (SCS), and it increased to >50% in the
presence of an anticyclone over mainland China and/or a tropical cyclone over SCS. Furthermore, Wang et al. (2006) revealed that the regional contribution was lower during daytime than that at night, while Lam et al. (2006) found that the contribution decreased from >50% during an O₃ episode to 30-50% after the episode. In addition, Jiang et al. (2008) reported that the contribution of horizontal transport to surface O₃ in Hong Kong was negligible during a typhoon-induced O₃ episode. However, all the studies focused on the regional contribution at non-roadside sites, and the O₃ chemistry at roadside sites in Hong Kong was seldom reported. Given high NOₓ levels, tall buildings and low wind speeds along the main roads in Hong Kong, it is expected that the regional impact on air pollutants in roadside environment is relatively minor, especially during daytime hours on non-O₃ episode days.

Please refer to lines 79-92, pages 3-4 in the revised manuscript for details.

A base case was established to simulate O₃ and the photochemical reactivity (i.e., the formation and loss pathways of OH and HO₂ listed in Table 2). The observed VOCs and trace gases were input to construct the base case, and the simulated O₃ was compared with the observed levels to validate the model and check the influence of regional transport during daytime hours. At low wind speed (1.3±0.01 m/s) and high NOₓ (223.2±1.8 ppbv) at MK, the majority of regional O₃ would be completely consumed before being detected by measurement instruments. In fact, the PBM-MCM model considered the regionally-transported concentrations of O₃ precursors, as the observed concentrations input into the model included the regional fractions. Moreover, since O₃ levels were higher at night (8.1±0.2 ppbv) than that during daytime (5.9±0.1 ppbv) at MK, it suggests the existence of nocturnal regional transport. Hence, the observed O₃ at 07:00 was input into the model to initiate the simulation for daytime hours, i.e., the nocturnal regional transport of O₃ was also considered.

For details, please refer to lines 303-314, pages 12 and 13 and Table 2 in the revised manuscript.

3. The arguments deriving from the PMF analysis, that NOx concentrations can be ascribed to emissions from gasoline vehicles is unconvincing. Diesel Euro5 NOx emissions are still substantial and the controls on gasoline vehicles would have to be very weak if they were to dominate. This problem emphasises the need for more data—we are not told about the magnitude
of the total VOC concentrations in the four factor areas, nor about the numbers and types of vehicles in the gasoline, LPG and diesel categories.

Thanks for the useful comments and suggestions. The source apportionment is a critical part of this study to evaluate the effectiveness of the program and its impact on O₃ production. We fully agreed with the reviewer’s comments, and made some substantial revisions in this part. Firstly, considering this comment and other comments (points 4-6 below) of the reviewer, the full (or whole-air) ambient concentrations of VOCs and trace gases rather than those with the backgrounds being deducted were used, in order to keep consistency with the simulations of whole-air ambient O₃ and photochemical reactivity. Secondly, because of the change of input data for the PMF apportionment simulation, the sources of O₃ precursors were re-identified and re-quantified. Four sources including “Gasoline/diesel vehicle exhaust”, “LPG-fueled vehicle exhaust”, “Aged air masses”, and “Solvent usage” were identified. The majority of NOₓ was assigned to the “Gasoline/diesel vehicle exhaust”. Thirdly, the actual influence of the background deduction on the reduction of LPG-related VOCs and NOₓ was evaluated to ensure the appropriateness of the use of whole-air ambient concentrations as model input, which would not interfere with the assessment of the replacement program. Lastly, the magnitude of the total VOC concentrations in the four sources, and the number and type of vehicles in the gasoline, LPG and diesel categories were provided.

For details, please refer to lines 483-549, pages 21-25 (section 3.2) in the revised manuscript and Table S4 in the supplementary material.

4. The method of establishing VOC concentration from primary processes, discussed in section 2.3.1-3, which are then used extensively in the rest of the paper, is questionable. The background concentrations are subtracted to determine the base case, and the LPG-related concentrations, presumably based on the PMF analysis, are subtracted to form a set of scenarios. We need to be given information on what these sets of concentrations are.

Thanks for the comments and suggestions. In the revised manuscript, the whole-air ambient concentrations (not the subtracted ones) were used for the source apportionment and the simulation of O₃ and photochemical reactivity in the base case. To evaluate the possible impact
with and without the background subtraction on the assessment of the interventional program, we have conducted sensitivity analysis for both PMF and PBM-MCM results. For the source apportionment section, the following revision was made:

During the study period, the LPG-related VOCs and NO increased by ~1.4 and ~2.9 μg/m³, respectively, at the background site, only respectively accounting for 3.0% and 1.3% of the decreased LPG-related VOCs (~46.1 μg/m³) and NO (~226.8 μg/m³) in LPG-fueled vehicle exhaust. Therefore, it is believed that the reductions of VOCs and NO in LPG-fueled vehicle exhaust benefited from the intervention program.

For details, please refer to lines 537-542, page 23 in the revised manuscript.

For the PBM-MCM results with and without the background subtraction, the following revision was made.

The simulation results for the scenarios with and without the input of background concentrations indicated minor contribution of background concentrations to O₃ for both periods of “before” (0.24 ppbv, accounting for ~3.5% of measured average) and “during” (0.27 ppbv; ~3.7%) the program, suggesting the use of whole-air ambient concentrations without background subtraction for O₃ simulation was appropriate. The slight increase of O₃ (~0.03 ppbv) caused by the background variations of O₃ precursors from before to during the program constituted only ~7.5% of the O₃ enhancement (0.40 ± 0.03 ppbv) due to the replacement program, further confirming a negligible impact of the background on the assessment of the program.

For details, please refer to lines 575-583, page 26 in the revised manuscript.

In addition, the simulation method for the estimation of the contribution of LPG source to O₃ formation was revised to avoid possible confusion. At last, the configurations of the model input for the base case and the three constrained cases were provided in Table S3 in the supplementary material.

To simulate the contribution of LPG source to O₃ formation, the following approach was adopted to avoid altering the real ambient environment for atmospheric chemistry. Briefly, the concentration of species X in the whole air and in the assigned LPG source was defined as Xwhole-air and XLPG, respectively. Hence, the concentration of “Xwhole-air - XLPG” was the input for the
constrained cases while $X_{\text{whole-air}}$ was the input for the base case. The three constrained cases were i) only VOCs in the LPG source were excluded from the whole-air; ii) only NO$_x$ in the LPG source was excluded; and iii) both VOCs and NO$_x$ were removed simultaneously. As such, the differences of the outputs between the base case and the three constrained cases were the contributions of i) VOCs, ii) NO$_x$ and iii) VOCs + NO$_x$ in the LPG source to the O$_3$ formation, respectively. The changes of these contributions from before to during the program represented the impact of the intervention program on O$_3$ formation. Table S3 in the Supplementary material lists the configurations of the model input for the base case and the three constrained cases.

For details, please refer to lines 317-328, page 13 in the revised manuscript and Table S3 in the supplementary material.

5. These concentration sets are then used in box models of the local processes, using the detailed chemistry from the MCM. Is this approach justifiable? The chemistry occurring is unable to distinguish between the sources of the species present in the atmosphere. LPG VOCs are affected by the background concentrations, and those derived from other sources. Subtracting out species concentrations is highly questionable.

Sorry for the confusion. As replied in questions 3&4 above, in the revised version, the full (or whole-air) ambient concentrations of VOCs and trace gases rather than those with the background concentrations being deducted were used, in order to keep consistency with the simulations of whole-air ambient O$_3$ and photochemical reactivity, and to avoid altering the real ambient environment for atmospheric chemistry. The concentration of “$X_{\text{whole-air}} - X_{\text{LPG}}$” was the input for the constrained cases while $X_{\text{whole-air}}$ was the input for the base case. Through this approach, the interference of other sources on the evaluation of the effectiveness of the interventional program was eliminated. This method also guaranteed that the background concentrations and those derived from other sources still existed in the base case and the constrained cases, which would not alter the chemistry of the atmosphere. Indeed, the influence of the background concentrations to the O$_3$ production was minor (~3.5%--~3.7%), as estimated in question 4.

The corresponding revisions were made in the “Methodology” as follows:
A base case was established to simulate $O_3$ and the photochemical reactivity (\textit{i.e.}, the formation and loss pathways of OH and HO$_2$ listed in Table 2). The observed VOCs and trace gases were input to construct the base case, and the simulated $O_3$ was compared with the observed levels to validate the model and check the influence of regional transport during daytime hours.

To simulate the contribution of LPG source to $O_3$ formation, the following approach was adopted to avoid altering the real ambient environment for atmospheric chemistry. Briefly, the concentration of species X in the whole air and in the assigned LPG source was defined as $X_{\text{whole-air}}$ and $X_{\text{LPG}}$, respectively. Hence, the concentration of “$X_{\text{whole-air}} - X_{\text{LPG}}$” was the input for the constrained cases while $X_{\text{whole-air}}$ was the input for the base case. The three constrained cases were i) only VOCs in the LPG source were excluded from the whole-air; ii) only NO$_x$ in the LPG source was excluded; and iii) both VOCs and NO$_x$ were removed simultaneously. As such, the differences of the outputs between the base case and the three constrained cases were the contributions of i) VOCs, ii) NO$_x$ and iii) VOCs + NO$_x$ in the LPG source to the $O_3$ formation, respectively. The changes of these contributions from before to during the program represented the impact of the intervention program on $O_3$ formation. Table S3 in the Supplementary material lists the configurations of the model input for the base case and the three constrained cases.

For details, please refer to lines 303-306, page 12, lines 317-328, page 13 and Table 2 in the revised manuscript, and Table S3 in the supplementary material.

Corresponding revisions were also made in the “Results and discussion”.

Please refer to lines 550-592, pages 25-27 (section 3.3) in the revised manuscript for details.

6. I presume that this approach is responsible for the rather strange results of the HOx calculations that are given: a. The calculated [OH] and especially [HO2] are very low. A rationale is given, but it is highly qualititative. b. What is meant by “net O3 photolysis”, which is found to consume OH. I presume this arises from the subtraction method used and clearly demonstrates the problems such an approach generates. The production and destruction rates are, I presume, calculated from the ambient concentrations multiplied by the rate constant or J value. O3 photolysis as a sink is nonsensical.
Thank you very much for the excellent comments and suggestions.

(a) Indeed, the strange results of OH, HO\textsubscript{2} and the formation/loss pathways of HO\textsubscript{x} were caused by the subtraction of the background concentrations. In the revised manuscript, the whole-air ambient concentrations were used to simulate the OH and HO\textsubscript{2}, and reasonable results were obtained.

For details, please refer to lines 594-634, pages 27-29 (section 3.4.1) in the revised manuscript and Figure S3 in the supplementary material.

(b) Also, the subtraction method generated wrong reaction pathways of HO\textsubscript{x}. With the use of whole-air ambient concentrations for the PBM-MCM simulations, the problems were solved.

For details, please refer to lines 621-634, pages 28-29 in the revised manuscript.

7. The HO\textsubscript{x} calculations are, in principle, of value in assessing the influence of the replacement strategy on the photochemistry but the approach must be revised. The calculations must be conducted with the full ambient concentrations, rather than using the subtraction approach. One way to examine the effects of the replacement programme would then be to compare total rates and concentrations (of ozone, HO\textsubscript{x} etc) before and during the programme. Of course ambient conditions can change, presenting comparison problems. The best way would be to conduct a proper sensitivity analysis, examining the sensitivity of specific objectives, e.g. the ozone formation rate or [OH] on the concentration of LPG VOCs and NO\textsubscript{x}, under conditions prior to and during the replacement process.

Excellent comments. As stated earlier, the subtraction method was not used anymore, \textit{i.e.}, the background concentrations were not subtracted in the revised manuscript. Using the whole-air ambient concentrations, the sensitivity analysis of [OH] and [HO\textsubscript{2}] on the concentration of LPG VOCs and NO\textsubscript{x}, under conditions prior to and during the replacement process was conducted.

For details, please refer to lines 635-658, pages 29-31 (section 3.4.2) in the revised manuscript.

8. It would also be useful to see a more complete analysis of production and destruction, with separate analyses for OH and HO\textsubscript{2}. I presume that Figure 11 refers to the overall HO\textsubscript{x} radical formation and removal rates and this is why OH + VOC reactions do not figure in Figure 11. It
explains why overall production and destruction rates do not balance, the differences relating to other radical formation / destruction. This partial analysis leaves more questions than it answers. Either this table should relate to total radical formation and destruction – a substantial problem in a region with a complex VOC species range. This would allow assessment of the total initiation and termination rates for the reaction system. Or it should concentrate on OH and HO2, as at present, but reporting their total production and removal rates, including the propagation reactions (e.g. reaction with VOCs). Such an analysis is straightforward with the MCM. It would provide a much more useful analysis of the impact of the LPG reductions than does the present analysis, whose objective is obscure.

The comments and suggestions are appreciated. We fully agreed with the reviewer, and accepted the valuable suggestions. In the revised manuscript, the formation and loss pathways of OH and HO2 were calculated separately, and they were balanced for both OH and HO2.

The corresponding revisions in methodology are as follows:

A base case was established to simulate O3 and the photochemical reactivity (i.e., the formation and loss pathways of OH and HO2 listed in Table 2).

For details, please refer to lines 303-304, page 12, and Table 2 in the revised manuscript.

For the revised results and discussion on the complete analysis of production and destruction, with separate analyses for OH and HO2, please refer to lines 594-658, pages 27-31 in the revised manuscript.

Technical / typographical comments

35942; 16: change since to over.

Accepted with thanks. Please refer to line 104, page 4 in the revised manuscript.

35942; 21: initiate rather than initialize. Also elsewhere in the paper.

Accepted with thanks. Please refer to line 109, page 4 in the revised manuscript.

35942; 23. Reword – the LPG vehicles weren’t replaced with catalytic converters – the converters on them were replaced.
Replaced by “….and the catalytic converters in ~75% of LPG-fueled vehicles were renewed…..”

Please refer to lines 110-111, page 4 in the revised manuscript.

35943, 8. Insert the before South.

Accepted with thanks. Please refer to line 123, page 5 in the revised manuscript.

35943, 12. Insert A before marine

Accepted with thanks. Please refer to line 127, page 5 in the revised manuscript.

35945, 20 et seq. There is confusion of emissions / formation and concentrations. The wording should be changed. Concentrations indicate emissions or formation, but aren’t synonymous—they have different units.

Thanks for the comment. We revised the expression as follows:

The measured concentrations of VOCs and trace gases were composed of the backgrounds, primary emissions and secondary formation (applicable to \(O_3\) and \(NO_2\)). To obtain the concentrations of LPG-related VOCs and NO\(_x\) emitted from primary sources at the roadside sites, the background concentrations and the concentrations elevated by the secondary formation (if applicable) in this study were excluded using the equations (1) - (5) (Takekawa et al., 2013):

\[
[\text{VOC}]_{\text{prim.}} = [\text{VOC}]_{\text{obs.}} - [\text{VOC}]_{\text{bg.}} \quad \text{(Equation (1))}
\]

\[
[NO_2]_{\text{sec.}} = [O_3]_{\text{bg.}} - [O_3]_{\text{obs.}} \quad \text{(Equation (2))}
\]

\[
[NO_2]_{\text{prim.}} = [NO_2]_{\text{obs.}} - [NO_2]_{\text{sec.}} - [NO_2]_{\text{bg.}} \quad \text{(Equation (3))}
\]

\[
[NO_x]_{\text{prim.}} = [NO_x]_{\text{obs.}} - [NO_x]_{\text{bg.}} \quad \text{(Equation (4))}
\]

\[
[NO]_{\text{prim.}} = [NO_x]_{\text{prim.}} - [NO_2]_{\text{prim.}} \quad \text{(Equation (5))}
\]

where \([xx]_{\text{obs.}}, [xx]_{\text{prim.}}, [xx]_{\text{sec.}}\) and \([xx]_{\text{bg.}}\) represent the observed concentrations, the concentrations emitted from primary sources, secondary formation and the backgrounds, respectively.

For details, please refer to lines 212-224, pages 8-9 in the revised manuscript.
Accepted with thanks. Please refer to line 234, page 9 in the revised manuscript.

Accepted with thanks. Please refer to line 300, page 12 in the revised manuscript.

Accepted with thanks. It was revised as “RIR (weighted concentration)”. For details, please refer to line 334, page 14 and Equation (9) in the revised manuscript.

Thanks for the suggestion. Since this part was revised substantially, the word “well” was deleted in the revised manuscript. For details, please refer to lines 551-566, pages 24-25 in the revised manuscript.

Thanks for the question. In the original manuscript, O$_3$ simulation was based on the concentrations with the backgrounds being subtracted. Therefore, the difference between the simulated and observed O$_3$ was partially due to the backgrounds, which was defined as background O$_3$. In the revised manuscript, O$_3$ was simulated with the full ambient concentrations, and the expression of “background O$_3$” was deleted. For details, please refer to lines 551-566, pages 24-25 in the revised manuscript.

Thanks for the comment. Since this part was revised substantially, the original description was changed. For details, please refer to lines 621-632, pages 28-29 in the revised manuscript.

Figure 4. Explain the variability during Jun-Aug 2013 – the variations exceed the subsequent downward trend. Define x.
Thanks for the comment. We noticed that the mixing ratios of VOCs were far from the regression line in June and July, 2013. The explanations were provided in the revised manuscript.

The abnormally low alkane levels in late June 2013 might be caused by the anti-cyclone over southern China and a tropical storm over the SCS, leading to extremely high temperature on June 18-20 (i.e., favorable for photochemical reactions), and stronger winds (i.e., conducive to atmospheric dispersion) on the following days, respectively. In contrast, Hong Kong was strongly influenced by an active ocean flow and a low pressure trough over the SCS in late July 2013. The consequently heavy rain and low temperature suppressed the photochemical reactions and caused the unusually high levels of VOCs (Figure S2 in the Supplementary material illustrates these events).

For details, please refer to lines 401-408, page 17 in the revised manuscript and Figure S2 in the supplementary material.

x in Figure 4 was defined.

Figure 4 Variation trends of the daily average propane and n/i-butanes at MK from June 2013 to May 2014. In the equation label x has units of days.

For details, please refer to lines 434-435, page 18 in the revised manuscript.
Responses to comments from Anonymous Referee #3

General:

The authors currently consider only the importance of this action (replacing catalytic converters) for air quality in Hong Kong. It would be valuable, and of wider interest, to include some discussion on the implications of these results for other regions.

Thanks for the suggestion. The implication of this study for other regions was briefly illustrated:

The findings are of great help to future formulation and implementation of control strategies on vehicle emissions in Hong Kong, and could be extended to other regions in China and around the world.

For details, please refer to lines 33-35, page 2 in the revised manuscript.

Past studies have shown that a small number of cars are responsible for a large part of total mobile emissions. Is there any evidence that the cars in Hong Kong with replaced catalytic converters are typical (or not) of the entire fleet?

Many thanks for the comment. Table S4 in the Supplementary material provides the detailed information of registered vehicle number in Hong Kong. Indeed, the LPG-fueled vehicles accounted for only a small part of the registered vehicles (~3.1%). However, the source apportionment results in previous studies and this work all indicated the high contribution of LPG-fueled vehicle exhaust to ambient VOCs. In this study, the LPG related VOCs accounted for 56.5±2.7 % and 46.0±3.1% of the total VOCs, and 18.7±1.2% and 15.1±1.1% of the total OH-reactivity of VOCs at MK before and during the program, respectively. That is, the small number of LPG-fueled vehicles caused significant air pollution in Hong Kong. It is also the motivation of this LPG program.

On the other hand, within the LPG fleet, the catalytic converters in 80% of taxis and 60% of the public light buses needed replacement. The replacement of the catalytic converters in these vehicles significantly reduced emissions of propane, propene, i-butane, n-butane and NO in LPG-fueled vehicle exhausts by 40.8±0.1%, 45.7±0.2%, 35.7±0.1%, 47.8±0.1% and 88.6±0.7%, respectively.
For details, please refer to lines 176-179, page 7 and lines 512-529, page 23 in the revised manuscript and Table S4 in the supplementary material.

Is it possible to estimate the expected reduction from a “bottom-up” approach based on the known number of cars with new catalytic converters and then compare this with what was observed in atmospheric concentrations for this study?

Thanks for the good suggestion. We understood that the bottom-up approach is often used to calculate the total amount of VOCs or NO\textsubscript{x} reductions (in unit of tons). Unfortunately, we do not have all data available for the estimation. We may do this in the future.

Specific:

Page 35940, line 24: replace “matters” with “matter”

Accepted with thanks. Please refer to line 39, page 2 in the revised manuscript.

Page 35942, line 21: replace “initialized” with “initiated”

Accepted with thanks. Please refer to line 109, page 4 in the revised manuscript.

Page 35943, line 24: what is meant by “great”? Do you mean “entire”?

Sorry for the typo. The “great Pearl River Delta (PRD)” was revised as “greater Pearl River Delta (PRD)”.

For details, please refer to line 123, page 5 in the revised manuscript.

Page 35943, line 24: Is HT representative of marine background or continental background?

i.e., does the air tend to come from the ocean or from land?

Thanks for the question. HT is located at the tip of southeastern Hong Kong. It receives air masses from South China Sea in summer and from continental areas including Hong Kong and Pearl River Delta in autumn/winter. Therefore, air masses at HT are the mix of marine and continental air. Whether it is representative of marine background or continental background depends on seasonality. In the revised manuscript, more introductions were provided about this site.
HT is a remote site at the tip of southeastern Hong Kong, ~ 25 km from the urban core. It receives air masses from SCS in summer and from continental areas including Hong Kong and PRD in autumn/winter. Therefore, air masses at HT are a mix of marine and continental background air. Indeed, HT has been serving as a background site and is extensively used for air quality studies in Hong Kong (Wang et al., 2009; Ho et al., 2006; Lee et al., 2002).

For details, please refer to lines 138-142, page 5 in the revised manuscript.

Section 2.2.2: Given the known challenges in making VOC measurements, the stated accuracies seem very high. The approach used to determine the accuracy of these measurements should be described in more detail and include in the assessment not just of the stated accuracy of the calibration standard. Much of the actual uncertainty is due to interferences, peak integrations, etc. VOC intercomparisons published in the scientific literature typically indicate large differences (>30%) in the values of VOC reported by different laboratories. The comparison between the propane data and those from UCI reported here are quite good but are still are beyond the stated accuracy (i.e. they differ by 14% but the accuracy is 10%). What about more reactive VOC?

Thanks for the comment. The detailed description about the accuracy of the measurements and the comparison of more reactive VOCs with UCI was provided.

The accuracy and precision of VOC measurements were 1.0-10% and 2.5-20%, respectively. The accuracy was based on weekly span checks, monthly calibration and annual auto-linearization using the National Physical Laboratory (NPL) span gas. The precision was based on quarterly precision check results (the 95% probability limits for the integrated precision based on weekly precision check results of the latest 3 months). Moreover, the online-measurements of VOCs were regularly compared with whole-air canister samples collected and analyzed by University of California at Irvine (UCI). Good agreements were identified for the alkanes (e.g., \( R^2 = 0.95 \) and 0.85, slope = 1.14 and 0.97 for propane and butanes, respectively), while the agreements for the more reactive alkenes and aromatics were also reasonable (e.g., \( R^2 = 0.64 \) and 0.94, slope = 1.34 and 0.86 for propene and toluene, respectively).

For details, please refer to lines 198-208, page 8 in the revised manuscript.

page 35947, statements about fresh emissions in line 5 and 8 are repetitive
Thanks for the comment. The sub-conclusions drawn from the VOC ratios with and without the backgrounds were integrated.

The ratios of toluene/benzene and xylenes/ethylbenzene at MK were significantly higher than those at the other three sites \((p<0.05)\). Furthermore, these ratios at MK were even higher after the background values were deducted. The results suggested that air masses at MK were dominated by fresh emissions.

For details, please refer to lines 242-245, page 9 in the revised manuscript.

page 35947, line 19: reword to “wind speeds were lower than 2.0 m s\(^{-1}\) more than 95\% of the time”

Accepted with thanks. Please refer to lines 255-256, page 10 in the revised manuscript.

page 35949, line 4: replace “physical” with “transport”

Accepted with thanks. Please refer to line 291, page 11 in the revised manuscript.

section 3.1.1: Some indication of the synoptic scale meteorology in 2013 and 2014 should be given to demonstrate whether any differences in meteorology might explain any of the observed change in VOC concentrations.

Thanks for the good suggestion. The influences of the synoptic scale meteorology on the observed change in the concentrations of LPG-related VOCs and NO\(_x\) were discussed.

Table 3 shows the mixing ratios of the primarily emitted LPG-related VOCs and NO\(_x\) before and during the program. The LPG-related VOCs and NO\(_x\) were reduced significantly from “before” to “during the program” \((p<0.05)\). To investigate the meteorological influences on these reductions, we looked into the average geopotential height (HGT) and wind field on 1000 hPa for East Asia for the periods of “matched” and “during the program”, as shown in Figure S1 in the Supplementary material (the “matched” period (October 2012-May 2013) was defined as the same time span as that “during the program”, but in different years). Noticeably, the meteorological conditions were fairly similar between the two periods, i.e., the pressure decline (indicated by the decrease of HGT) from northern China to SCS led to the prevailing northeasterly winds in Hong Kong, and the differences of HGT and wind speed between the two periods for Hong Kong were only 0-2 gpm and less than 1 m/s, respectively. Ground monitoring
data also indicated insignificant differences of temperature ("matched": 21.9±0.5 °C; “during”: 21.2±0.7 °C) between the two periods (p>0.05). However, the mixing ratios of LPG-related VOCs and NO\textsubscript{x} during the “matched” period were comparable to those “before” the program, and absolutely higher than those “during” the program (p<0.05) (see Table 3). Given the similar meteorological conditions between the “matched” period and “during the program”, the significant decreases of LPG-related VOCs and NO\textsubscript{x} were caused by the interventional program, rather than meteorological variations.

For details, please refer to lines 346-364, pages 14-15, Table 3 and Figure S1 in the revised manuscript.

Page 35955, lines 1-4: The first two sentences indicate that CO should be low for diesel vehicles. This contrasts with the following sentence which suggests that CO should be high for diesel vehicles. I realize that this is discussed in the rest of the paragraph but still the expectation (did you expect CO to be low or high) is unclear here and should be clarified

Thanks for the comment. We have re-done source apportionment simulation using the observed ambient concentrations, rather than the concentrations with the backgrounds being deducted in the original manuscript. Therefore, the source apportionment results were totally changed, which solved this problem.

For details, please refer to lines 484-506, pages 21-22 and Figure 6 in the revised manuscript.

Page 35955, line 14: what is a “goods” vehicle?

Thanks for the comment. “Goods vehicles” means vehicles used to deliver the goods. However, due to the substantial changes of this part, it was only used in the Supplementary material of the revised manuscript.

Page 35956, line 25: was the difference between predicted and observed similar to what the background values are? This information should be provided here.

Thanks for the comment. Since we replaced the concentrations with the backgrounds being deducted by the observed ambient concentrations for model simulations according to the comments of reviewer #2, the differences between the simulated and observed O\textsubscript{3} in the revised
manuscript represented the contribution of regional transport. We added the following in the revised manuscript:

Bearing in mind the uncertainty of the model, the good agreement between the simulated and observed O$_3$ implied a minor regional contribution at this roadside site.

For details, please refer to lines 564-566, page 25 in the revised manuscript.

Page 35958, line25: How can OH be higher than HO2? This is not realistic and indicates a problem in the model simulation.

Thanks for the valuable comment. The question was also raised by the other reviewer. The causes of the problem and revised results of the model simulation were provided in the answer to question 6 of reviewer #2.

For details, please refer to the answer to question 6 of reviewer #2.

For the revised results and discussion, please refer to lines 595-620, pages 27-28 and Figure 9 in the revised manuscript.

Page 35962: How does this compare/contrast with similar studies at roadsides in other cities?

Thanks for the comment. To our best knowledge, this is the first attempt to conduct O$_3$-VOCs-NO$_x$ sensitivity analysis at roadside site.

Page 35963, line 5: replace “that” with “the”

Accepted with thanks.

Figures 3 through 12: Increase the size of the font- it is too small to read

Thanks for the suggestion. The fonts in the corresponding figures were increased.
Effectiveness of replacing catalytic converters in LPG-fueled vehicles in Hong Kong


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Abstract

Many taxis and public buses are powered by liquefied petroleum gas (LPG) in Hong Kong. With more vehicles using LPG, they have become the major contributor to ambient volatile organic compounds (VOCs) in Hong Kong. An intervention program aimed to reduce the emissions of VOCs and nitrogen oxides (NOx) from LPG-fueled vehicles was implemented by the Hong Kong Government in September 2013. Long-term real-time measurements indicated that the program was remarkably effective in reducing LPG-related VOCs, NOx and nitric oxide (NO) in the atmosphere. Receptor modeling results further revealed that propane, propene, i-butane, n-butane and NO in LPG-fueled vehicle exhaust emissions decreased by 40.8±0.1%, 45.7±0.2%, 35.7±0.1%, 47.8±0.1% and 88.6±0.7%, respectively, during the implementation of the program. In contrast, despite the reduction of VOCs and NOx, O3 following the program increased by 0.40 ± 0.03 ppbv (~5.6%). The LPG-fueled vehicle exhaust was generally destructive to OH and HO2. However, the destruction effect weakened for OH and it even turned to positive contribution to
These changes led to the increases of OH, HO$_2$ and HO$_2$/OH ratio, which might explain the positive O$_3$ increment. Analysis of O$_3$-VOCs-NO$_x$ sensitivity in ambient air indicated VOC-limited regimes in the O$_3$ formation before and during the program. Moreover, a maximum reduction percentage of NO$_x$ (i.e., 69%) and the lowest reduction ratio of VOCs/NO$_x$ (i.e., 1.1) in LPG-fueled vehicle exhaust were determined to give a zero O$_3$ increment. The findings are of great help to future formulation and implementation of control strategies on vehicle emissions in Hong Kong, and could be extended to other regions in China and around the world.

**Key words:** LPG; Catalytic converter; VOCs; NO$_x$, Ozone production; PBM-MCM

### 1 Introduction

Vehicular exhaust is a major source of air pollutants such as particulate matter (PM), volatile organic compounds (VOCs) and trace gases including carbon monoxide (CO), carbon dioxide (CO$_2$), sulfur dioxide (SO$_2$) and nitrogen oxides (NO$_x$) (Xie et al., 2003; Barletta et al., 2002; Ruellan and Cachier, 2001). Apart from primary pollutants, secondary pollutants formed from vehicular exhausts, i.e., ozone (O$_3$) and secondary organic aerosols (SOA) are of high concern due to their detrimental effects on atmospheric environment and human health (Ciccone et al., 1998).

Fuel substitution is a universal and effective method to constrain vehicular emissions. As relatively clean energy, liquefied petroleum gas (LPG) has been widely used either alone or combined with petrol and diesel (Lau et al., 2011; Lai et al., 2009; Gamas et al., 1999), for the benefit of emission reduction of CO$_2$, fine PMs and VOCs (Chikhi et al., 2014; Myung et al., 2012). However, one must bear in mind that maintenance status, conversion method, composition and driving speed all influence the emissions of air pollutants (Schifter et al., 2000; Gamas et al., 1999). Therefore, regular on-site measurements are necessary to monitor traffic emission profiles and rates. Although LPG is a relatively clean fuel, the emission of LPG-related VOCs (i.e., propane, propene, and n- and i-butanes) from evaporation and incomplete combustion cannot be neglected. For example, source apportionment analysis indicated that LPG accounted for a considerable percentage of ambient VOCs in Guangzhou (8 - 16%) (Liu et al.,
2008) and Hong Kong (32.6 ± 5.8%) (Ou et al., 2015). Furthermore, the LPG-related VOCs can contribute to O₃ formation and cause photochemical smog. Blake and Rowland (1995) emphasized the importance of LPG leakage in OH reactivity and O₃ formation in urban Mexico. Farmer et al. (2011) reported that the increase of LPG-related VOCs led to O₃ increments even though the total VOCs (TVOCs) were reduced, as the peroxyl radicals (RO₂) generated from VOCs with low vapor pressure had a higher branching ratio leading to NO₂ formation from NO, rather than reserving nitrogen in the form of organic nitrates. On the other hand, NO emitted from LPG combustion may titrate O₃. Therefore, it still remains undecided whether the LPG usage is contributive or destructive to O₃ formation, especially in Hong Kong.

Hong Kong has experienced rapid development during the past decades, and increasingly suffers from crowded traffic and population, with nearly 0.7 million vehicles running on the total road length of 2.1 × 10³ km and 7.2 million people living on the territory of 1.1×10³ km². The road usage rate in Hong Kong is among the highest in the world, transporting approximately 1.2 million passengers per day. As such, traffic emissions significantly influence the air quality in Hong Kong. Apart from local emissions, air quality in Hong Kong is also affected by regional transport. In autumn and winter, prevailing northerly and northeasterly winds bring anthropogenic emissions from mainland China to Hong Kong, enhancing local pollution burden. In summer, clean oceanic air masses dilute the levels of air pollutants under the influence of southerly winds (Ling et al., 2013; Guo et al., 2009, Wang et al., 2005; So and Wang, 2003; Chan and Chan, 2000). Ling et al. (2013) quantified the contribution of regional transport as 0-61% for summer O₃ and 0-56% for autumn O₃ on high O₃ days in Hong Kong. Guo et al. (2006) found that over 45% of VOC samples collected from August 2001 to December 2002 at Tai O, Hong Kong, were influenced by regional transport from inland China. Based on numerical simulations, Wang et al. (2006) quantified regional contributions of 40-90% to O₃ formation in Hong Kong, and Huang et al. (2006) indicated that the regional contribution was <50% when a trough was over the South China Sea (SCS), and it increased to >50% in the presence of an anticyclone over mainland China and/or a tropical cyclone over SCS. Furthermore, Wang et al. (2006) revealed that the regional contribution was lower during daytime than that at night, while Lam et al. (2006) found that the contribution decreased from >50% during an O₃ episode to 30-50% after the episode. In addition, Jiang et al. (2008) reported that the contribution of horizontal transport to surface O₃ in Hong Kong was negligible during a typhoon-induced O₃ episode.
However, all the studies focused on the regional contribution at non-roadside sites, and the O\textsubscript{3} chemistry at roadside sites in Hong Kong was seldom reported. Given high NO\textsubscript{x} levels, tall buildings and low wind speeds along the main roads in Hong Kong, it is expected that the regional impact on air pollutants in roadside environment is relatively minor, especially during daytime hours on non-O\textsubscript{3} episode days.

To improve the air quality, the Hong Kong Environmental Protection Department (HKEPD) launched a series of programs including a stepwise LPG replacement program for taxis and light buses from 1997. By the end of 2010, >99% of the registered taxis (i.e., 17,950 vehicles) and 51% of the registered public and private light buses (i.e., 3,280 vehicles) were powered by LPG (HKCSD, 2010). Due to the usage of LPG as vehicle fuel, the ambient VOC profile in Hong Kong changed substantially with the most abundant species switching from toluene to propane and n/i-butanes (Ou et al., 2015; Guo et al., 2007; Ho et al., 2004). In addition, studies consistently confirmed an increased contribution of LPG-fueled vehicle emissions to ambient VOC levels, which was 11% at an urban site in 2001 (Guo et al., 2004), 15% at a suburban site during 2002-2003 (Guo et al., 2007), and 26.9 ± 0.6% and 40.8 ± 0.8% at an urban site during 2002-2003 and 2006-2007, respectively (Lau et al., 2010). Moreover, O\textsubscript{3} has been experiencing an increasing trend in Hong Kong over the last two decades (Wang et al., 2009; Guo et al., 2009). As O\textsubscript{3} formation was generally VOC-limited in Hong Kong (Cheng et al., 2010; Zhang et al., 2007), the contribution of LPG-fueled vehicle emissions to O\textsubscript{3} was not negligible, in view of the dominance of propane and n/i-butanes in ambient air and the high contribution of LPG-fueled vehicle emissions to ambient VOC levels. As such, an intervention program aimed to reduce VOCs and NO\textsubscript{x} emitted by LPG-fueled vehicles was initiated in September 2013. This subsidy program promoted the replacement of aging catalytic converters in LPG-fueled vehicles, and the catalytic converters in ~75% of LPG-fueled vehicles were replaced by the end of May 2015. Although laboratory studies identified a high removal efficiency of the new catalytic converter on VOCs and NO\textsubscript{x} emissions (Table S1 in the Supplementary material), the impact of the program on roadside air quality, which is more complex, required investigation.

In this study, the effectiveness of the intervention program on the reduction of LPG-related VOCs and NO\textsubscript{x}, on O\textsubscript{3} production, and on the photochemical reactivity driving the O\textsubscript{3} formation was evaluated. Furthermore, the sensitivity of O\textsubscript{3} variation (i.e., the difference of net O\textsubscript{3}}
production by LPG before and during the program) relative to the reduction percentage of LPG-related VOCs and NO\textsubscript{x} was studied, and subsequently a maximum NO\textsubscript{x} reduction percentage and a reasonable reduction ratio of VOCs/NO\textsubscript{x} were proposed for achieving the best O\textsubscript{3} reduction.

2 Methodology

2.1 Site description

Hong Kong, located on the coast of the South China Sea (SCS), constitutes the greater Pearl River Delta (PRD) together with Macao and nine mainland cities in Guangdong Province (i.e., Guangzhou, Shenzhen, Zhuhai, Foshan, Dongguan, Zhongshan, Huizhou, Jiangmen and Zhaoqing). Hong Kong has a typical coastal hilly terrain with large water area (~60%) and large vegetation coverage (~70% of land area). A marine subtropical monsoon climate dominates, and the prevailing winds are north and northeast in winter, east in spring and autumn, and south and southwest in summer.

This study involves three roadside sites (Mong Kok (MK, 22.32° N, 114.17° E), Causeway Bay (CWB, 22.28° N, 114.19° E) and Central (22.28° N, 114.16° E)), a general site (Tap Mun (TM, 22.47° N, 114.36° E)), and a background site (Hok Tsui (HT, 22.22° N, 114.25° E)). MK, CWB and Central are in mixed residential/commercial areas with heavy traffic and surrounded by dense tall buildings. TM is a rural site surrounded by country parks, and is upwind of Hong Kong in autumn/winter seasons. These four sites have Air Quality Monitoring Stations deployed by Hong Kong Environmental Protection Department (HKEPD) (more details are provided at http://www.aqhi.gov.hk/en/monitoring-network/air-quality-monitoring-stations9c57.html). HT is a remote site at the tip of southeastern Hong Kong, ~ 25 km from the urban core. It receives air masses from SCS in summer and from continental areas including Hong Kong and PRD in autumn/winter. Therefore, air masses at HT are a mix of marine and continental background air. Indeed, HT has been serving as a background site and is extensively used for air quality studies in Hong Kong (Wang et al., 2009; Ho et al., 2006; Lee et al., 2002).

Figure 1 shows the geographical location of the sampling sites.
Figure 1 Geographical location of the sampling sites. The yellow line represents the border between Hong Kong and mainland China.

2.2 Continuous measurements

2.2.1 Trace gases

Five trace gases (i.e., CO, SO\(_2\), NO, NO\(_2\) and O\(_3\)) were continuously measured at all the sites from 2011 to 2014. The analyzers were deployed in the air quality monitoring stations with a height of 3.0 m (MK and CWB), 4.5 m (Central), 11.0 m (TM) and 20.0 m (HT) above the ground level. Air samples were drawn through a perfluoroalkoxy (PFA) Teflon tube with outside diameter (OD) of 12.7 mm and inside diameter (ID) of 9.6 mm. The inlet of the tube was approximately 0.5 m above the rooftop of the monitoring station, and the outlet was connected to a PFA manifold with a bypass pump drawing air into the analyzers at a rate of 5 L/min. Details about the analyzers are provided elsewhere (HKEPD, 2014; So and Wang, 2003).

Briefly, CO was analyzed using the method of non-dispersive infra-red absorption with gas filter correlation (TECO 48C/API 300); SO\(_2\) was detected with UV fluorescence analyzers (TECO 43A/API 100E/TECO 43I); NO-NO\(_2\)-NO\(_x\) were measured by chemiluminescence technique (API 200A); and O\(_3\) was monitored with the UV absorption method (API 400/API 400A). The detection limits for CO, SO\(_2\), NO, NO\(_2\) and O\(_3\) were 50.0, 1.0, 0.5, 0.5 and 2.0 ppbv, respectively.
2.2.2 VOCs

Thirty C\textsubscript{2}–C\textsubscript{10} non-methane hydrocarbons (NMHCs) including eleven alkanes, ten alkenes and nine aromatics were continuously measured at MK and HT from 2011 to 2014. The VOC samples were collected and analyzed every 30 minutes with an online analytical system (Syntech Spectra GC 955, Series 600/800, the Netherlands). The data were averaged into hourly values. The detection limits varied by VOC species and were in the range of 0.002-0.787 ppbv.

Table S2 summarizes average mixing ratios of the main VOCs and trace gases, and their OH-reactivity at the roadside (MK) and background site (HT) before (October 2012-September 2013) and during the program (October 2013-May 2014) in Hong Kong. The much higher OH-reactivity at MK (“before”: 86.6±6.0 s\textsuperscript{-1}; “during”: 77.9±6.1 s\textsuperscript{-1}) than at HT (“before”: 5.0±0.3 s\textsuperscript{-1}; “during”: 4.9±0.1 s\textsuperscript{-1}) meant that the OH lifetime was much shorter at MK (~0.01 s) than at HT (0.2 s), and even shorter than that at an urban site in New York (0.05 s) (Ren et al., 2003), indicating that reactive species were extremely abundant in the roadside environment of Hong Kong. The LPG-related VOCs (i.e., propane and n/i-butanes) contributed 56.5±2.7 % and 46.0±3.1% to total VOCs, and 18.7±1.2% and 15.1±1.1% to the total OH-reactivity of VOCs at MK, before and during the program, respectively. These fractional contributions suggest the importance of LPG-related VOCs in the budgets of VOCs and O\textsubscript{3} in Hong Kong. Consistent with the decreases of fractional contributions once the program started, the mixing ratios (29.4±3.3 and 21.0±2.2 ppbv for before and during the program, respectively) and OH-reactivity (1.12±0.04 and 0.79±0.02 s\textsuperscript{-1} before and during the program, respectively) of LPG-related VOCs significantly declined at MK (\(p<0.05\)), while at HT their mixing ratios increased from 1.5±0.1 ppbv to 1.9±0.04 ppbv, and the OH-reactivity increased from 0.06±0.004 s\textsuperscript{-1} to 0.07±0.003 s\textsuperscript{-1}.

The minor variation of background LPG-related VOCs relative to the decreases at MK (4.8% in mixing ratios and 3.0% in OH-reactivity of VOCs) indicated that the influences of the background concentrations on the variations of LPG-related VOCs and O\textsubscript{3} at MK were minimal and can be neglected.

2.2.3 Quality assurance and control (QA/QC)

To guarantee the quality of the data acquired from the online monitoring systems, the instruments and QA/QC procedures for trace gases measurements were identical to those in the US air quality monitoring program (http://epic.epd.gov.hk/ca/uid/airdata). The measurements
have been accredited by the Hong Kong Laboratory Accreditation Scheme (HOKLAS). All the
instruments were regularly calibrated, tested and audited by standards with known traceability
for trace gases measurements. According to HOKLAS criteria, a quality system was established
to control the accuracy and precision within the limits of ±15% and ±20%, respectively (HKEPD,
2014). For VOC analysis, a built-in computerized program, including auto-linearization, auto-
calibration and calibration with span gas, was adopted to control the quality. The accuracy and
precision of VOC measurements were 1.0-10% and 2.5-20%, respectively. The accuracy was
based on weekly span checks, monthly calibration and annual auto-linearization using the
National Physical Laboratory (NPL) span gas. The precision was based on quarterly precision
check results (the 95% probability limits for the integrated precision based on weekly precision
check results of the latest 3 months). Moreover, the online-measurements of VOCs were
regularly compared with whole-air canister samples collected and analyzed by University of
California at Irvine (UCI). Good agreements were identified for the alkanes (e.g., \( R^2 = 0.95 \) and
0.85, slope = 1.14 and 0.97 for propane and butanes, respectively), while the agreements for the
more reactive alkenes and aromatics were also reasonable (e.g., \( R^2 = 0.64 \) and 0.94, slope = 1.34
and 0.86 for propene and toluene, respectively).

2.3 Theoretical calculation and model description

2.3.1 Primary emissions of VOCs and NO\(_x\)

The measured concentrations of VOCs and trace gases were composed of the backgrounds,
primary emissions and secondary formation (applicable to O\(_3\) and NO\(_2\)). To obtain the
concentrations of LPG-related VOCs and NO\(_x\) emitted from primary sources at the roadside sites,
the background concentrations and the concentrations elevated by the secondary formation (if
applicable) in this study were excluded using equations (1) - (5) (Takekawa et al., 2013):

\[
[VOC]_{prim.} = [VOC]_{obs.} - [VOC]_{bg}. \quad \text{(Equation (1))}
\]

\[
[NO_2]_{sec.} = [O_3]_{bg} - [O_3]_{obs}. \quad \text{(Equation (2))}
\]

\[
[NO_2]_{prim.} = [NO_2]_{obs} - [NO_2]_{sec.} - [NO_2]_{bg}. \quad \text{(Equation (3))}
\]
\[ [NO_x]_{\text{prim.}} = [NO_x]_{\text{obs.}} - [NO_x]_{\text{bg.}} \]  \hspace{1cm} \text{(Equation (4))}

\[ [NO]_{\text{prim.}} = [NO]_{\text{obs.}} - [NO]_{\text{bg.}} \]  \hspace{1cm} \text{(Equation (5))}

where \([xx]_{\text{obs.}}, [xx]_{\text{prim.}}, [xx]_{\text{sec.}}, \text{ and } [xx]_{\text{bg.}}\) represent the observed concentrations, the concentrations emitted from primary sources, secondary formation and the backgrounds, respectively. In this study, the hourly measured VOCs at HT were considered as the background levels for VOCs. For \(O_3\) and \(NO_x\), the 8-hr averages at TM were treated as background values because (1) \(O_3\) and \(NO_x\) are highly reactive, and the use of 8-hr averages would minimize the impact of abrupt changes; (2) TM is located at the upwind location of Hong Kong, and is a rural coastal site; (3) 8-hr averages of \(O_3\) and \(NO_x\) showed good correlations (\(R^2 = 0.75\) and 0.57 for \(O_3\) and \(NO_x\), respectively) between TM and HT, and not all the data were available at HT during the entire study period.

### 2.3.2 Impact of regional and super-regional air masses

Since MK was the only roadside site having VOC and trace gas data, this study mainly focused on the data analysis of this site. As a roadside site, MK was expected to be strongly influenced by fresh vehicular emissions. VOC ratios can indicate the relative ages of air masses and regional transport of air pollutants. For example, the ratios of VOCs with higher reactivity to those with lower reactivity (e.g. toluene/benzene and xylenes/ethylbenzene) imply more fresh air when values are higher, which has been extensively used in previous studies (Guo et al., 2007; Ho et al., 2004). Therefore, ratios of toluene/benzene and xylenes/ethylbenzene were calculated here to roughly estimate the age of air masses at MK. Table 1 summarizes the ratios at MK (roadside site), Tung Chung (TC) and Yuen Long (YL) (both general ambient sites), and HT (background site) in Hong Kong from October 2012 to May 2014. It is noteworthy that ambient VOCs at all the four sites were simultaneously measured. The ratios of toluene/benzene and xylenes/ethylbenzene at MK were significantly higher than those at the other three sites (\(p<0.05\)). Furthermore, these ratios at MK were even higher after the background values were deducted. The results suggested that air masses at MK were dominated by fresh emissions.

However, regional and super-regional transport cannot be completely eliminated only with the VOC ratios, and the influence of regional/super-regional air should be further examined when studying the emissions of local LPG-fueled vehicles. Using the propane/CO ratio method
proposed by Guo et al. (2006), i.e., the ratio range of 5.0-300 pptv/ppbv for air masses in Hong Kong, the influence of regional/super-regional air masses on local air was evaluated. Figure 2 shows the ratios of propane/CO at MK from 2011-2014. The propane/CO ratios of ~96% air masses were between 5.0 and 300 pptv/ppbv. Further inspection of the values of LPG-related VOCs and NO$_x$ with ratios of 5.0-300 showed insignificant differences ($p>0.1$) from those with ratios beyond 5-300, suggesting that the air masses at MK were seldom influenced by regional/super-regional air. Moreover, wind speeds were lower than 2.0 m/s for more than 95% of the time (i.e., calm or light air according to the Beaufort Wind Scale) (http://www.spc.noaa.gov/faq/tornado/beaufort.html), and the values of LPG-related VOCs and NO$_x$ in all air masses were not different from those in the air masses with wind speeds lower than 2.0 m/s ($p>0.1$), confirming a minor impact of regional/super-regional pollution on air at MK. Indeed, the prevailing wind direction was from the east, where local urban areas located.

Table 1 Ratios of toluene/benzene and xylenes/ethylbenzene at MK and other sites in Hong Kong from October 2012 to May 2014 (unit: ppbv/ppbv)

<table>
<thead>
<tr>
<th></th>
<th>toluene/benzene</th>
<th>xylenes/ethylbenzene</th>
</tr>
</thead>
<tbody>
<tr>
<td>MK (roadside site)</td>
<td>4.1 ± 0.04</td>
<td>3.0 ± 0.02</td>
</tr>
<tr>
<td>MK* (roadside site)</td>
<td>11.3 ± 0.9</td>
<td>5.7 ± 0.2</td>
</tr>
<tr>
<td>TC (general site)</td>
<td>0.7 ± 0.1</td>
<td>-</td>
</tr>
<tr>
<td>YL (general site)</td>
<td>3.7 ± 0.04</td>
<td>2.4 ± 0.02</td>
</tr>
<tr>
<td>HT (background site)</td>
<td>2.5 ± 0.1</td>
<td>1.8 ± 0.03</td>
</tr>
</tbody>
</table>

MK* refers to VOCs at MK with the background values being removed.
Figure 2 Hourly ratio of propane/CO at MK during 2011-2014

2.3.3 PMF model

Positive matrix factorization (PMF) is a receptor model for source apportionment and has been extensively used in many fields (i.e., PM, VOCs and sediment) (Brown et al., 2007; Lee et al., 1999). Based on the principle of mass balance, it decomposes the matrix of measurement (X) into the matrices of factor contributions (G) and factor profiles (F) in p sources, as shown in equation (6) (Paatero, 1997; Paatero and Tapper, 1994):

\[ x_{ij} = \sum_{k=1}^{p} g_{ik} f_{kj} + e_{ij} \]  
\hspace{1cm} (Equation (6))

where \( x_{ij} \) is the measured concentration of \( j \) species in \( i \) sample, \( g_{ik} \) represents the contribution of \( k_{th} \) source to \( i \) sample, \( f_{kj} \) indicates the fraction of \( j \) species in \( k_{th} \) source, and \( e_{ij} \) is the residual for \( j \) species in \( i \) sample.

The matrices of G and F are obtained in the case of the minimum of objective function Q, as shown in equation (7):

\[ Q = \sum_{i=1}^{n} \sum_{j=1}^{m} \left[ \frac{x_{ij} - \sum_{k=1}^{p} g_{ik} f_{kj}}{u_{ij}} \right]^2 \]  
\hspace{1cm} (Equation (7))

where \( m \) and \( n \) represent the number of species and samples, respectively, and \( u_{ij} \) is the uncertainty of \( j \) species in \( i \) sample. Q (robust) is automatically calculated by excluding the
points not fit by the model, and the run with lowest Q (robust) value is selected as the optimum solution by the model.

2.3.4 PBM-MCM model

The photochemical box model incorporating the master chemical mechanism (PBM-MCM) is a computing model based on the chemical reactions of observed species. It has been introduced and successfully applied in many previous studies in reproducing the observed values, describing the production and destruction mechanisms, and simulating the photochemical reactivity (Ling et al., 2014; Lam et al., 2013; Cheng et al., 2010). Briefly, the observations of VOCs, trace gases and meteorological parameters are input to construct and constrain the model, which consists of 5,900 reactions and 16,500 species in the latest version (MCM 3.2). However, it is noteworthy that the heterogeneous reactions are not included, and the model does not consider the transport processes (i.e., horizontal and vertical transport).

In this study, CO, SO\textsubscript{2}, NO, NO\textsubscript{2}, O\textsubscript{3}, twenty-seven VOCs and two meteorological parameters (i.e., temperature and relative humidity) were used to construct and constrain the model. Hourly data were used as input (i.e. 24 hours per day) and the outputs were from 07:00 to 19:00 by each hour. To better describe the photochemical reactions in Hong Kong, the model was modified, i.e., the photolysis rates were calibrated using the photon fluxes from the Tropospheric Ultraviolet and Visible Radiation (TUV-v5) Model (Madronich and Flocke, 1997) according to the location of Hong Kong and the modeling period, and the height of mixing layer was set as 300-1400 m. The concentrations of some species in the free troposphere, which may influence the mixing ratios of air pollutants in the lower troposphere with the development of boundary layer, were set according to the real conditions in Hong Kong (Lam et al., 2013).

A base case was established to simulate O\textsubscript{3} and the photochemical reactivity (i.e., the formation and loss pathways of OH and HO\textsubscript{2} listed in Table 2). The observed VOCs and trace gases were input to construct the base case, and the simulated O\textsubscript{3} was compared with the observed levels to validate the model and check the influence of regional transport during daytime hours. At low wind speed (1.3±0.01 m/s) and high NO\textsubscript{x} (223.2±1.8 ppbv) at MK, the majority of regional O\textsubscript{3} would be completely consumed before being detected by measurement instruments. In fact, the PBM-MCM model considered the regionally-transported concentrations of O\textsubscript{3} precursors, as the observed concentrations input into the model included the regional fractions. Moreover, since O\textsubscript{3}
levels were higher at night (8.1±0.2 ppbv) than that during daytime (5.9±0.1 ppbv) at MK, it suggests the existence of nocturnal regional transport. Hence, the observed O₃ at 07:00 was input into the model to initiate the simulation for daytime hours, i.e., the nocturnal regional transport of O₃ was also considered.

Table 2 Formation and loss pathways of OH and HO₂ considered in this study

<table>
<thead>
<tr>
<th>Formation pathways</th>
<th>OH</th>
<th>HO₂</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HO₂ + NO</td>
<td>RO₂ + NO</td>
</tr>
<tr>
<td></td>
<td>HONO + hv</td>
<td>OH + CO</td>
</tr>
<tr>
<td></td>
<td>O (¹D) + H₂O</td>
<td>OH + HCHO</td>
</tr>
<tr>
<td></td>
<td>O₃ + alkenes</td>
<td>HCHO + hv</td>
</tr>
<tr>
<td></td>
<td></td>
<td>O₃ + alkenes</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Loss pathways</th>
<th>OH + VOCs</th>
<th>HO₂ + NO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>OH + NO₂</td>
<td>HO₂ + NO</td>
</tr>
<tr>
<td></td>
<td>OH + CO</td>
<td>HO₂ + NO</td>
</tr>
<tr>
<td></td>
<td>OH + NO</td>
<td>HO₂ + NO</td>
</tr>
</tbody>
</table>

To simulate the contribution of LPG source to O₃ formation, the following approach was adopted to avoid altering the real ambient environment for atmospheric chemistry. Briefly, the concentration of species X in the whole air and in the assigned LPG source was defined as Xwhole-air and XLPG, respectively. Hence, the concentration of “Xwhole-air - XLPG” was the input for the constrained cases while Xwhole-air was the input for the base case. The three constrained cases were i) only VOCs in the LPG source were excluded from the whole-air; ii) only NOₓ in the LPG source was excluded; and iii) both VOCs and NOₓ were removed simultaneously. As such, the differences of the outputs between the base case and the three constrained cases were the contributions of i) VOCs, ii) NOₓ and iii) VOCs + NOₓ in the LPG source to the O₃ formation, respectively. The changes of these contributions from before to during the program represented the impact of the intervention program on O₃ formation. Table S3 in the Supplementary material lists the configurations of the model input for the base case and the three constrained cases.

2.3.5 Relative incremental reactivity (RIR)
RIR, initially proposed by Carter and Atkinson (1989), has been extensively used to describe the relationship between O\(_3\) and its precursors, *i.e.*, VOCs, NO\(_x\) and CO. Although the observation-based models incorporating carbon bond mechanisms were often utilized to simulate the O\(_3\) production rate (Ling et al., 2011; Zhang, et al., 2007; Martien et al., 2003), a more explicit PBM-MCM model was applied in this study. The RIR and RIR (weighted concentration) were calculated using equations (8) and (9) (Ling et al., 2011):

\[
\text{RIR}^S(X) = \frac{[P^S_{O_3-NO}(X) - P^S_{O_3-NO}(X-\Delta X)]/P^S_{O_3-NO}(X)}{\Delta S(X)/S(X)} \quad \text{(Equation (8))}
\]

where, \(P^S_{O_3-NO}(X)\) and \(P^S_{O_3-NO}(X-\Delta X)\) represent the original O\(_3\) production rate, and that in the scenario, with the a hypothetical change \(\Delta X\) (10% in this study) in source/species \(X\), respectively, both of which considered O\(_3\) titration by NO. \(\Delta S(X)\) is the change in the concentration of \(X\) \((S(X))\). The "concentration" refers to the observed or PMF-extracted concentration of source/species \(X\).

\[
\text{RIR (weighted concentration)} = \text{RIR}(X) \times \text{concentration} \quad \text{(Equation (9))}
\]

3 Results and discussion

3.1 Variations of LPG-related VOCs and NO\(_x\) during the intervention program

3.1.1 Concentrations of primary LPG-related VOCs and NO\(_x\)

Table 3 shows the mixing ratios of the primarily emitted LPG-related VOCs and NO\(_x\) before and during the program. The LPG-related VOCs and NO\(_x\) were reduced significantly from “before” to “during the program” \((p<0.05)\). To investigate the meteorological influences on these reductions, we looked into the average geopotential height (HGT) and wind field on 1000 hPa for East Asia for the periods of “matched” and “during the program”, as shown in Figure S1 in the Supplementary material (the “matched” period (October 2012-May 2013) was defined as the same time span as that “during the program”, but in different years.) Noticeably, the meteorological conditions were fairly similar between the two periods, *i.e.*, the pressure decline (indicated by the decrease of HGT) from northern China to SCS led to the prevailing northeasterly winds in Hong Kong, and the differences of HGT and wind speed between the two
periods for Hong Kong were only 0-2 gpm and less than 1 m/s, respectively. Ground monitoring
data also indicated insignificant differences of temperature (“matched”: 21.9±0.5 °C; “during”: 21.2±0.7 °C) between the two periods (p>0.05). However, the mixing ratios of LPG-related VOCs and NO\textsubscript{x} during the “matched” period were comparable to those “before” the program, and absolutely higher than those “during” the program (p<0.05) (see Table 3). Given the similar meteorological conditions between the “matched” period and “during the program”, the significant decreases of LPG-related VOCs and NO\textsubscript{x} were caused by the interventional program, rather than meteorological variations.

Table 3 Mixing ratios of LPG-related VOCs and NO\textsubscript{x} during the periods of “before”, “matched” and “during the program” (Unit: ppbv)

<table>
<thead>
<tr>
<th>Species</th>
<th>“before”</th>
<th>“during”</th>
<th>“matched”</th>
</tr>
</thead>
<tbody>
<tr>
<td>Propane</td>
<td>8.5±0.1</td>
<td>5.8±0.1</td>
<td>9.1±0.1</td>
</tr>
<tr>
<td>i-Butane</td>
<td>6.6±0.1</td>
<td>4.7±0.1</td>
<td>6.9±0.1</td>
</tr>
<tr>
<td>n-Butane</td>
<td>13.2±0.2</td>
<td>8.4±0.1</td>
<td>13.7±0.2</td>
</tr>
<tr>
<td>NO</td>
<td>201.7±2.1</td>
<td>172.0±2.7</td>
<td>201.9±2.7</td>
</tr>
<tr>
<td>NO\textsubscript{2}</td>
<td>25.9±0.3</td>
<td>23.8±0.4</td>
<td>26.8±0.4</td>
</tr>
<tr>
<td>NO\textsubscript{x}</td>
<td>229.4±2.3</td>
<td>197.7±3.0</td>
<td>231.2±2.9</td>
</tr>
</tbody>
</table>

To further understand the effectiveness of the program, the monthly reductions of LPG-related VOCs after the commencement of the converter replacement were calculated and compared with those before the program (Figure 3). Since the program was initiated in October 2013, the averages of VOC species in September in each year were taken as the baselines for the calculation of monthly reduction. Briefly, the monthly reductions of LPG-related VOCs were the differences between their averages in September and those in the other months. Hence, positive and negative values indicate reductions and increments of the corresponding species, respectively. It was found that the monthly averages of LPG-related VOCs consistently decreased from September 2013 to May 2014 except for n-butane in October 2013 when the program was just initiated. Compared to those before the program (i.e. September 2013), the mixing ratios of propane, i-butane and n-butane decreased 3.2 ± 0.2, 2.8 ± 0.2 and 4.9 ± 0.2 ppbv by May 2014, respectively, when 99.2% of catalytic converters participating in the program had been exchanged. Furthermore, the monthly reductions correlated well with the cumulative
converter replacements \( (R^2 = 0.92, 0.93 \text{ and } 0.89 \text{ for propane, } i\text{-butane and } n\text{-butane, respectively}) \). In contrast, no consistent reduction was observed from September 2012 to May 2013 for LPG-related VOCs suggesting the effectiveness of the program on the reduction of LPG-related VOCs. On the other hand, although the average mixing ratios of NO\(_x\) decreased significantly during the program as shown in Table 3, no consistent reductions were found for their monthly averages. This might be due to the fact that NO\(_x\) emitted from LPG-fueled vehicles was minor compared to gasoline- and diesel-fueled vehicles (\(~4.0\%\) from emission inventory, and \(1.1\text{-}7.3\%\) from source apportionment. See section 3.2.2).
Figure 3 Monthly reductions of LPG-related VOCs at MK “before” (blue bar) and “during” (red bar) the program (the average value of VOC species in September is the baseline. The values along the dot line were the cumulative numbers of converters replaced. The bars above and below the X axis refer to the reduction and increase of VOC mixing ratio, respectively)

3.1.2 Temporal variations of primary LPG-related VOCs and NOx

Figure 4 presents the temporal variations of propane and n/i-butanes at MK from June 2013 to May 2014 covering the periods of both before (i.e., June 2013-September 2013) and during the intervention program (i.e., October 2013-May 2014). The abnormally low alkane levels in late June 2013 might be caused by the anti-cyclone over southern China and a tropical storm over the SCS, leading to extremely high temperature on June 18-20 (i.e., favorable for photochemical reactions), and stronger winds (i.e., conducive to atmospheric dispersion) on the following days, respectively. In contrast, Hong Kong was strongly influenced by an active ocean flow and a low pressure trough over the SCS in late July 2013. The consequently heavy rain and low temperature suppressed the photochemical reactions and caused the unusually high levels of VOCs (Figure S2 in the Supplementary material illustrates these events). Generally, the LPG-related VOCs experienced a significant (p<0.05) reduction throughout the year. The daily changing rates of LPG-related VOCs in two periods, i.e., September-December and January-May, in different years at MK are shown in Table 4. It is noteworthy that the two periods were selected based on data availability each year, and the minimized influence of meteorological parameters in the same month of different years. The LPG-related VOCs decreased from September to December in 2011 and 2013, but there was no significant difference in 2012. In general, the levels of VOCs in the atmosphere are associated with source emissions, photochemical reactions
and regional transport. As the regional influence was excluded in this study, source emissions and photochemical reactions became the main factors determining the ambient concentrations of VOCs. The decreasing and unchanged trends in 2011 and 2012 respectively might be related to the integrated influence of reduced photochemical degradation (which increases ambient VOC levels) and temperature decrease (which reduces evaporative emissions of VOCs) from September to December. It is noticeable that the LPG-related VOCs had the highest decreasing rates in September-December 2013, compared to those in previous years. The higher decreasing rates of LPG-related VOCs in September-December 2013 implied the possible effectiveness of the interventional program on VOCs reduction. For the period of January-May, the LPG-related VOCs increased in January-May 2013, except for propane which did not show significant change from January to May. However, significant decreasing trends \((p<0.05)\) were observed for propane and \(n/i\)-butanes during the same period in 2014, which was likely owing to the continuous replacement of catalytic converters on LPG-fueled vehicles. By comparison, the reduction rates of LPG-related VOCs in January-May 2014 were all lower than those in September-December 2013. This was mainly attributable to the fact that a large portion \((i.e. \sim 69\%)\) of the converter replacements were completed by the end of December 2013, and the replaced converters were much fewer \((i.e. \sim 31\%)\) in January-May 2014.

![Graph showing mixing ratio vs. time for propane, isobutane, and normal butane](image-url)
Figure 4 Variation trends of the daily average propane and n/i-butanes at MK from June 2013 to May 2014. In the equation label x has units of days.

Table 4 Changing rates of propane and n/i-butanes in September-December and January-May in different years (ppbv/day)

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Site</td>
<td>2011</td>
<td>2012</td>
</tr>
<tr>
<td>MK</td>
<td>Propane</td>
<td>-0.015</td>
</tr>
<tr>
<td></td>
<td>i-Butane</td>
<td>-0.010</td>
</tr>
<tr>
<td></td>
<td>n-Butane</td>
<td>-0.023</td>
</tr>
</tbody>
</table>

* The changing rate insignificant (p>0.05); the bold numbers are changing rates during the implementation of the program.

Figure 5 shows the temporal variations of primary NOx, NO, NO2 and secondary NO2 at MK from June 2013 to May 2014. The primary NOx and NO mixing ratios decreased significantly (p<0.05), implying the possible effectiveness of the intervention program on NO/NOx reduction. However, the secondary NO2 levels increased, while there was no significant change (p>0.05) for primary NO2. To explore the reasons of NO-NO2-NOx variations, the changing rates of primary NOx, NO, NO2 and secondary NO2 during the same period (i.e., June in the previous year to May in the following year) in different years were compared. Table 5 shows the statistics of changing rates of NOx-NO2-NOx at the roadside sites (i.e., MK, CWB and Central) from June 2011 to May 2014. Taking MK as an example, the primary NOx and NO decreased much faster (i.e., 91 and 94 pptv/day, respectively) from June 2013 to May 2014 than those during the same period in 2011-2012 and 2012-2013, suggesting the effectiveness of the program in reducing NOx and NO. Compared to the decreasing rate in June 2012-May 2013 (i.e., 13 pptv/day), no significant change (p>0.05) was observed for primary NO2 from June 2013 to May 2014. However, it cannot be concluded that the program caused the increase of primary NO2, in view of the same insignificant variation in June 2011-May 2012 and the fact that LPG-fueled vehicles emit negligible NO2 (see section 3.2.2). In contrast, secondary NO2 significantly increased (with a rate of 13 pptv/day) from June 2013 to May 2014, which was also observed in June 2011-May 2012 (i.e., increasing rate of 20 pptv/day). Since secondary NO2 is formed by NO reacting with O3, inspection of the O3 production would provide more comprehensive interpretation on the
secondary NO$_2$ increment during the program. Similar variations were observed at CWB and Central, where primary NO$_x$ and NO showed significant decreasing trends, whereas the variations of primary and secondary NO$_2$ were insignificant ($p > 0.05$).

Figure 5 Variation trends of the daily average primary NO$_x$, NO and NO$_2$ and secondary NO$_2$ at MK from June 2013 to May 2014. In the equation label x has units of days.

Table 5 Changing rates of primary NO$_x$, NO, NO$_2$ and secondary NO$_2$ (ppbv/day)

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>MK</td>
<td>[NO$<em>x$]$</em>{prim}$</td>
<td>-0.055*</td>
<td>0.089*</td>
<td>-0.121</td>
</tr>
<tr>
<td></td>
<td>[NO]$_{prim}$</td>
<td>0.004*</td>
<td>0.053</td>
<td>0.094</td>
</tr>
<tr>
<td></td>
<td>[NO$<em>2$]$</em>{prim}$</td>
<td>0.008*</td>
<td>-0.013</td>
<td>0.003*</td>
</tr>
<tr>
<td></td>
<td>[NO$<em>2$]$</em>{sec}$</td>
<td>0.020*</td>
<td>-0.002*</td>
<td>0.013</td>
</tr>
<tr>
<td>CWB</td>
<td>[NO$<em>x$]$</em>{prim}$</td>
<td>-0.004</td>
<td>0.040*</td>
<td>-0.091</td>
</tr>
<tr>
<td></td>
<td>[NO]$_{prim}$</td>
<td>0.004*</td>
<td>0.053</td>
<td>0.094</td>
</tr>
<tr>
<td></td>
<td>[NO$<em>2$]$</em>{prim}$</td>
<td>0.008*</td>
<td>-0.013</td>
<td>0.003*</td>
</tr>
<tr>
<td></td>
<td>[NO$<em>2$]$</em>{sec}$</td>
<td>0.014</td>
<td>-0.001*</td>
<td>0.006*</td>
</tr>
</tbody>
</table>
The changing rate is insignificant ($p>0.05$); the bold numbers are changing rates during the implementation of the program.

In summary, the LPG-related VOCs were significantly lowered during the program with monthly reductions of $3.2 \pm 0.2$, $2.8 \pm 0.2$ and $4.9 \pm 0.2$ ppbv for propane, $i$-butane and $n$-butane by the end of this study, respectively. Continuous decreasing trends were observed for LPG-related VOCs, and the reduction rates were almost unprecedented, e.g. 39 pptv/day for $n$-butane. Furthermore, the mixing ratios of NO and NO$_x$ decreased as well during the program. The reduction rates of NO and NO$_x$ during June 2013-May 2014 at the three roadside sites were much higher than those in previous years. Overall, the field measurement data indicated that the program was effective in reducing emissions of LPG-related VOCs and NO$_x$.

### 3.2 Variations of LPG contributions to VOCs and NO$_x$

#### 3.2.1 Source identification

To investigate the change of the contributions of LPG-fueled vehicles to VOCs and NO$_x$, the online data of 15 VOCs and 3 trace gases at MK before (i.e., 8,753 samples during October 2012-September 2013) and during the intervention program (i.e., 5,833 samples during October 2013-May 2014) were separately applied to PMF for source apportionments. It is noteworthy that the whole-air ambient concentrations of VOCs and trace gases rather than those with the backgrounds being deducted were used here, in order to keep consistency with the simulations of whole-air ambient O$_3$ and photochemical reactivity. Figure 6 shows the source profiles of the four factors that best reproduced the concentrations of the input species before and during the intervention program, respectively. The first factor had high loadings of C$_3$-C$_5$ hydrocarbons and toluene, and was dominated by CO, NO and NO$_2$, which are all main components of vehicle exhaust (Guo et al., 2011a; Ho et al., 2009). The $n/i$-pentanes and toluene indicated emissions...
from gasoline vehicles, while the high NO$_x$ loading was a signature of diesel exhaust. Since propane and $n/i$-butanes were not prominent, this factor was assigned as gasoline and diesel-fueled vehicle exhaust. Factor 2 was closely associated with LPG-fueled vehicle exhaust, with the dominance of propane, $n/i$-butanes and propene, the major components and combustion product of LPG (Guo et al., 2011b; Blake and Rowland, 1995). Factor 3 was distinguished by NO$_2$ and the long-lived species, i.e., ethane, ethyne, benzene and CO. The long-lived species were the indicators of aged air masses, in which NO$_2$ was accumulated due to photochemical reactions. Hence, this factor represented aged air masses. Indeed, this profile highly coincided with the aged air in Hong Kong identified by Lau et al. (2010). The last factor explained most of the TEX compounds (toluene/ethylbenzene/xylenes), the tracers of solvent usage (Guo et al., 2007; Borbon et al., 2002). Therefore, factor 4 was identified as solvent usage.

Figure 6 Source profiles resolved by PMF before (red bar) and during (green bar) the program. The standard errors are estimated with the bootstrap method in the model.

3.2.2 Source contribution
Based on the VOC loadings in each source, Table 6 summarizes the mass and percentage contributions of the sources to VOCs at MK. LPG-fueled vehicle exhaust was the greatest contributor to VOCs at MK, with the mass contribution of $114.2 \pm 0.1 \mu g/m^3 (51.51 \pm 0.05\%)$ and $64.8 \pm 0.1 \mu g/m^3 (39.07 \pm 0.05\%)$ before and during the program, respectively. These contributions were higher than those quantified at other sites in Hong Kong, i.e., suburban TC (32.6±5.8\%) (Ou et al., 2015), urban TW (21±2\%) (Ling and Guo, 2014) and suburban YL (15\%) (Guo et al., 2007). It is noteworthy that factors such as study period, chemical species, source profiles and models used all influence the source apportionment. Bearing these factors in mind, the higher contributions of LPG-fueled vehicle exhaust in this study were likely due to the fact that MK was a roadside site closer to the emission sources. On the other hand, though gasoline/diesel vehicles emitted considerable VOCs (i.e., 60.5 ± 0.1 and 56.8 ± 0.2 μg/m³ before and during the program, respectively), they were significantly lower than those emitted by LPG-fueled vehicles ($p<0.05$), particularly before the replacement of the catalytic converters. While LPG-fueled vehicles accounted for only a small amount of the registered vehicles (~3.1\%) in Hong Kong (Table S4 in the Supplementary material), it was not unreasonable that LPG made the highest contribution to VOCs, in view of low vapor pressure of the LPG component and high-density flow of the LPG-fueled vehicles at MK. Apart from vehicular exhaust, the aged VOCs and solvent usage also contributed ~24\% to VOCs at MK.

Table 7 lists the reduction of individual species from LPG-fueled vehicles due to the intervention program. Propane (“before”: 21.15 ± 0.02 μg/m³; “during”: 12.53 ± 0.01 μg/m³), n-butane (“before”: 57.96 ± 0.01 μg/m³; “during”: 30.27 ± 0.02 μg/m³) and i-butane (“before”: 27.20 ± 0.02 μg/m³; “during”: 17.50 ± 0.01 μg/m³) all decreased remarkably throughout the study period ($p<0.05$). Meanwhile, the concentrations of CO and NO were also reduced from 248.4 ± 1.3 and 18.2 ± 0.1 μg/m³ to 228.9 ± 0.6 and 2.08 ± 0.04 μg/m³, respectively. However, NO₂ apportioned in LPG exhaust was extremely minor (i.e., 0.6 ± 0.2 μg/m³ and nil before and during the program, respectively), which might explain the insignificant decrease or even increase of NO₂. During the study period, the LPG-related VOCs and NO increased by ~1.4 and ~2.9 μg/m³, respectively, at the background site, only respectively accounting for 3.0\% and 1.3\% of the decreased LPG-related VOCs (~46.1 μg/m³) and NO (~226.8 μg/m³) in LPG-fueled vehicle exhaust. Therefore, it is believed that the reductions of VOCs and NO in LPG-fueled vehicle exhaust benefited from the intervention program.
Table 6 Mass concentration and percentage contribution of the four sources to VOCs at MK before and during the program

<table>
<thead>
<tr>
<th>Source</th>
<th>“before”</th>
<th>“during”</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mass (μg/m³)</td>
<td>Percentage (%)</td>
</tr>
<tr>
<td>Gasoline/ diesel vehicle exhaust</td>
<td>60.5 ± 0.1</td>
<td>27.30 ± 0.05</td>
</tr>
<tr>
<td>LPG-fueled vehicle exhaust</td>
<td>114.2 ± 0.1</td>
<td>51.51 ± 0.05</td>
</tr>
<tr>
<td>Aged air masses</td>
<td>19.8 ± 0.2</td>
<td>8.9 ± 0.1</td>
</tr>
<tr>
<td>Solvent usage</td>
<td>27.30 ± 0.04</td>
<td>12.31 ± 0.01</td>
</tr>
</tbody>
</table>

Table 7 VOCs and trace gases emitted from LPG-fueled vehicles before and during the intervention program (unit: μg/m³ unless otherwise specified). The standard errors are estimated with the bootstrap method in the PMF model.

<table>
<thead>
<tr>
<th></th>
<th>“before”</th>
<th>“during”</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethane</td>
<td>1.192 ± 0.001</td>
<td>1.19 ± 0.01</td>
</tr>
<tr>
<td>Ethene</td>
<td>1.327 ± 0.005</td>
<td>0.431 ± 0.004</td>
</tr>
<tr>
<td>Ethyne</td>
<td>0.0000±0.0004</td>
<td>0.000 ± 0.001</td>
</tr>
<tr>
<td>Propane</td>
<td>21.15 ± 0.02</td>
<td>12.53 ± 0.01</td>
</tr>
<tr>
<td>Propene</td>
<td>2.860 ± 0.004</td>
<td>1.553 ± 0.001</td>
</tr>
<tr>
<td>n-Butane</td>
<td>57.96 ± 0.01</td>
<td>30.27 ± 0.02</td>
</tr>
<tr>
<td>i-Butane</td>
<td>27.20 ± 0.02</td>
<td>17.50 ± 0.01</td>
</tr>
<tr>
<td>n-Pentane</td>
<td>0.251 ± 0.003</td>
<td>0.039 ± 0.002</td>
</tr>
<tr>
<td>i-Pentane</td>
<td>1.04 ± 0.01</td>
<td>0.786 ± 0.002</td>
</tr>
<tr>
<td>Benzene</td>
<td>0.00 ± 0.01</td>
<td>0.000 ± 0.001</td>
</tr>
<tr>
<td>Toluene</td>
<td>0.66 ± 0.02</td>
<td>0.50 ± 0.01</td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>0.000 ± 0.001</td>
<td>0.000 ± 0.004</td>
</tr>
<tr>
<td>m,p-Xylene</td>
<td>0.416 ± 0.003</td>
<td>0.00 ± 0.01</td>
</tr>
<tr>
<td>o-Xylene</td>
<td>0.144±0.001</td>
<td>0.006 ± 0.001</td>
</tr>
<tr>
<td>CO</td>
<td>248.4 ± 1.3</td>
<td>228.9 ± 0.6</td>
</tr>
<tr>
<td>NO</td>
<td>18.2 ± 0.1</td>
<td>2.08 ± 0.04</td>
</tr>
<tr>
<td>NO₂</td>
<td>0.6 ± 0.2</td>
<td>0.00 ± 0.08</td>
</tr>
<tr>
<td>Sum of VOCs</td>
<td>114.2 ± 0.1</td>
<td>64.8 ± 0.1</td>
</tr>
</tbody>
</table>
### Contribution to VOCs (%)

| 51.5 ± 0.1 | 39.1 ± 0.1 |

#### 3.3 Impact of the intervention program on O$_3$ production

##### 3.3.1 O$_3$ Simulation

As VOCs and NO$_x$ are key O$_3$ precursors, it is essential to examine the impact of the reduction of VOCs and NO$_x$ during the program on the O$_3$ production. Figure 7 compares the observed and simulated O$_3$ during daytime (07:00-19:00 local time (LT)) in base case.

![Figure 7: Hourly simulated and observed O$_3$ during daytime hours (07:00-19:00 LT) at MK.](image)

To quantitatively evaluate the performance of the model, the index of agreement (IOA) was introduced to test the agreement between the simulated and observed O$_3$. Ranging from 0 to 1, higher values of IOA indicate better agreement between the simulation and observation. The IOA was calculated using equation (10) (Huang et al., 2005):

\[
IOA = 1 - \frac{\sum_{i=1}^{n} (o_i - s_i)^2}{\sum_{i=1}^{n} (|o_i - \bar{o}| + |s_i - \bar{s}|)^2}
\]  

(Equation (10))

where $o_i$ and $s_i$ are the hourly observed and simulated values, and $\bar{o}$ represents the average of observations. In this study, the IOA reached 0.75, and the accuracy of the simulation was 16.7 ± 2.1%, suggesting good performance of the model in O$_3$ simulation. Bearing in mind the uncertainty of the model, the good agreement between the simulated and observed O$_3$ implied a minor regional contribution at this roadside site.
3.3.2 Net O₃ production

Figure 8 shows the net O₃ variations induced by VOCs, NOₓ and VOCs + NOₓ in LPG-fueled vehicle exhaust before and during the program. From “before” to “during” the intervention program, the daily average O₃ variation induced by VOCs decreased significantly \( (p<0.05) \), with a rate of \(-9.3 \times 10^{-5}\) ppbv/day, whereas O₃ increased significantly at a rate of \(1.3 \times 10^{-3}\) ppbv/day due to the reduction of NOₓ titration \( (p<0.01) \). As a result, the net contribution to O₃ made by the LPG-fueled vehicle exhaust increased \( (p<0.01) \) from \(-0.47 \pm 0.03\) ppbv before the program to \(-0.06 \pm 0.02\) ppbv during the program, with a rate of \(1.2 \times 10^{-3}\) ppbv/day. Namely, O₃ increased by \(0.40 \pm 0.03\) ppbv, \(\sim5.6\%\) of the observed O₃ at MK \((7.14 \pm 0.21\) ppbv). The simulation results for the scenarios with and without the input of background concentrations indicated minor contribution of background concentrations to O₃ for both periods of “before” \((0.24\) ppbv, accounting for \(\sim3.5\%\) of measured average) and “during” \((0.27\) ppbv; \(\sim3.7\%\) the program, suggesting the use of whole-air ambient concentrations without background subtraction for O₃ simulation was appropriate. The slight increase of O₃ \((\sim0.03\) ppbv) caused by the background variations of O₃ precursors from before to during the program constituted only \(\sim7.5\%\) of the O₃ enhancement \((0.40 \pm 0.03\) ppbv) due to the replacement program, further confirming a negligible impact of the background on the assessment of the program. It is noteworthy that the increase was not a sudden change in reality, but was caused by the segmentation of source apportionment \( (i.e.,\text{ October 2012-September 2013 and October 2013-May 2014}) \). Indeed, the measurement data also revealed an increasing O₃ trend from October 2012 to May 2014, with a rate of \(3.3\) pptv/day \( (p<0.05) \), higher than the O₃ increase caused by LPG. This discrepancy might be related to the O₃ production by other sources and/or other mechanisms not considered in the above simulations, e.g., alkyl nitrate \( (RONO_2) \) chemistry.
3.4 Photochemical reactivity

3.4.1 OH, HO_2 and their formation/loss rates in whole air

As the “detergent” of atmosphere, OH initiates the oxidation of air pollutants including VOCs, CO and NO_x, leading to O_3 formation, and the cycling between OH and HO_2 accelerates the propagation of the chain reactions. Thus, the budget of OH and HO_2 is an important parameter of a photochemical system. Figure 9 presents the average daytime patterns of OH and HO_2 before and during the program. Typical bell-shaped patterns were observed for OH and HO_2, which increased from morning to noon, peaked at 12:00-13:00, and then decreased to low levels in the afternoon.

The average simulated concentrations of OH and HO_2 were $1.64 \pm 0.78 \times 10^5$ and $2.49 \pm 1.28 \times 10^5$ molecules/cm$^3$ before the program, and $1.80 \pm 0.85 \times 10^5$ and $4.18 \pm 2.03 \times 10^5$ molecules/cm$^3$ during the program, respectively. Compared to those modeled at an urban site in Hong Kong ($2.3-3.6 \times 10^6$ molecules/cm$^3$ for OH and $3.4-4.4 \times 10^8$ molecules/cm$^3$ for HO_2) (Ling et al., 2014), and those measured at a VOC-rich site in PRD ($15 \times 10^6$ and $18 \times 10^8$ molecules/cm$^3$ for OH and HO_2, respectively) (Lu et al., 2012; Lou et al., 2010), the OH and HO_2 levels were much lower at MK. This is not surprising because much OH and HO_2 were consumed by high levels of VOCs and NO_x at the roadside MK (Figure S3 in the Supplementary material shows the...
simulated OH and HO₂ at different sites with the PBM-MCM model). In addition, this study covered different seasons of the study years, rather than the O₃ episodes in Ling et al. (2014) and summer only in Lu et al. (2012) and Lou et al. (2010) when photochemical reactivity was usually stronger. Apart from the increase of OH and HO₂ from before to during the program, the ratio of HO₂/OH was significantly higher during the program (i.e., 3.7 ± 0.5) than that before (i.e., 1.9 ± 0.3) (p<0.05), which might partly explain the increase of O₃ throughout the study period. Nevertheless, since the OH, HO₂ and HO₂/OH were simulated using the whole-air concentrations of O₃ precursors, whether their increases were mainly caused by the intervention program requires further investigation, as shown below.

Figure 9 Daytime patterns of OH and HO₂ (a) before and (b) during the program.

Since the changes in the formation/loss rates from before to during the program were minor for each pathway of OH and HO₂, Figure 10 presents their average pathway-specific formation and loss rates over the whole study period. The total formation/loss rates of OH and HO₂ were 5.8 ± 2.4 ×10⁶ and 2.6 ± 1.0 ×10⁶ molecules/cm³/s, respectively. The reaction of HO₂ with NO, HONO photolysis, O₃ photolysis and ozonolysis of alkenes were the main sources of OH, with contributions of 56.7 ± 11.6%, 31.7 ± 10.7%, 6.6 ± 3.3% and 5.0 ± 1.7%, respectively. On the other hand, OH was consumed by reaction with NO (36.5 ± 5.5%), NO₂ (35.1 ± 4.6%), VOCs (14.8 ± 1.2%) and CO (13.6 ± 1.0%). For HO₂ formation, the reaction between RO₂ and NO was the most predominant pathway (54.8 ± 8.8%), followed by the reaction of OH with CO (23.3 ± 3.9%), HCHO photolysis (13.1 ± 4.6%), ozonolysis of alkenes (7.9 ± 2.1%) and the reaction of
OH with HCHO (0.8 ± 0.2%). Meanwhile, HO₂ was almost exclusively consumed by reacting with NO.

![Figure 10 Average formation and loss pathways of OH and HO₂ at MK during the study period](image)

### 3.4.2 Influence of the program on photochemical reactivity

The differences between the simulation outputs of the scenarios with and without LPG VOCs and NOₓ inputs determined the contributions of LPG-fueled vehicle exhaust to the formation/loss rates of OH and HO₂, as summarized in Table 8. Generally, the formation/loss rates of OH and HO₂ contributed by the LPG source decreased from before to during the program, which was caused by the reduction of VOCs and NOₓ in the LPG source. Furthermore, since the sum of the formation rates were lower than the loss rates for OH for both before and during the program, but for only HO₂ before the program, the net effects of the LPG source to OH and HO₂ were destructive. However, the HO₂ formation rate exceeded its loss rate for the period of during the program, indicating a net production of HO₂ by the LPG source.

Figure 11 (a) shows the net effects of the LPG source on the production of OH and HO₂. From before to during the program, the destruction rate of OH decreased, while the destruction of HO₂ switched to production. These variations led to the increases of OH and HO₂ from before to during the program, as shown in Figure 11 (b). Different from the increases of OH and HO₂ in the whole air as shown in Figure 9, the increases here were caused by the intervention program. The OH and HO₂ levels increased by 6.9±1.1 ×10³ molecules/cm³ and 3.4±1.2 ×10⁴
molecules/cm$^3$, respectively. The higher increase of HO$_2$ than OH led to a higher ratio of HO$_2$/OH during the program, resulting in a consequent O$_3$ increment.

Table 8 Contributions of LPG-fueled vehicle exhaust to the formation and loss of OH and HO$_2$

<table>
<thead>
<tr>
<th>Reaction</th>
<th>“Before” ($\text{molecules/cm}^3/\text{s}$)</th>
<th>“During” ($\text{molecules/cm}^3/\text{s}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>OH formation</td>
<td>$(1.8 \pm 0.8) \times 10^3$</td>
<td>$(2.8 \pm 1.2) \times 10^4$</td>
</tr>
<tr>
<td>O$_3$ + alkenes</td>
<td>$(3.3 \pm 1.2) \times 10^4$</td>
<td>$(3.1 \pm 1.2) \times 10^4$</td>
</tr>
<tr>
<td><strong>Sum of OH formation</strong></td>
<td><strong>$(2.2 \pm 0.9) \times 10^5$</strong></td>
<td><strong>$(5.2 \pm 1.9) \times 10^4$</strong></td>
</tr>
<tr>
<td>OH loss</td>
<td>$(1.4 \pm 0.6) \times 10^4$</td>
<td>$(1.5 \pm 0.6) \times 10^5$</td>
</tr>
<tr>
<td>OH + CO</td>
<td>$(5.0 \pm 2.0) \times 10^3$</td>
<td>-</td>
</tr>
<tr>
<td>OH + NO$_2$</td>
<td>$(1.9 \pm 0.02) \times 10^5$</td>
<td>$(2.4 \pm 1.0) \times 10^4$</td>
</tr>
<tr>
<td>OH + NO</td>
<td>$(2.3 \pm 0.02) \times 10^5$</td>
<td>$(1.5 \pm 0.6) \times 10^5$</td>
</tr>
<tr>
<td><strong>Sum of OH loss</strong></td>
<td><strong>$(5.7 \pm 1.6) \times 10^5$</strong></td>
<td><strong>$(3.2 \pm 1.4) \times 10^5$</strong></td>
</tr>
<tr>
<td>Net OH formation</td>
<td>- $(3.6 \pm 0.8) \times 10^5$</td>
<td>- $(2.7 \pm 1.2) \times 10^5$</td>
</tr>
<tr>
<td>HO$_2$ formation</td>
<td>$(8.1 \pm 3.4) \times 10^4$</td>
<td>$(1.3 \pm 0.5) \times 10^4$</td>
</tr>
<tr>
<td>O$_3$ + alkenes</td>
<td>$(3.3 \pm 1.2) \times 10^4$</td>
<td>$(3.1 \pm 1.2) \times 10^4$</td>
</tr>
<tr>
<td><strong>Sum of HO$_2$ formation</strong></td>
<td><strong>$(1.1 \pm 0.4) \times 10^5$</strong></td>
<td><strong>$(3.6 \pm 1.3) \times 10^4$</strong></td>
</tr>
<tr>
<td>HO$_2$ loss</td>
<td>$(1.8 \pm 0.8) \times 10^5$</td>
<td>$(2.8 \pm 1.2) \times 10^4$</td>
</tr>
<tr>
<td><strong>Sum of HO$_2$ loss</strong></td>
<td><strong>$(1.8 \pm 0.8) \times 10^5$</strong></td>
<td><strong>$(2.8 \pm 1.2) \times 10^4$</strong></td>
</tr>
<tr>
<td>Net HO$_2$ formation</td>
<td>- $(7.0 \pm 3.1) \times 10^4$</td>
<td>$(7.8 \pm 3.1) \times 10^3$</td>
</tr>
</tbody>
</table>
Figure 11 (a) Net production of OH and HO₂ by LPG-fueled vehicle exhaust (negative values indicate net destruction); and (b) increases of OH and HO₂ from before to during the program.

3.5 Improvement of the intervention program

3.5.1 O₃-VOCs-NOₓ sensitivity in the whole air

O₃-VOCs-NOₓ sensitivity can reflect the O₃ variation relative to the change of VOCs and NOₓ, from which VOC-limited regimes and NOₓ-limited regimes in O₃ formation can be indicated. Figure 12 (a) and (b) show the O₃-VOCs-NOₓ sensitivity in the air at MK between 40-100% and 0.5-40% of the observed average NOₓ, respectively. The ranges were selected according to the O₃ levels and behaviors responding to the variations of VOCs and NOₓ (details were given below). This sensitivity diagram was obtained with the aid of the PBM-MCM model, and based on the average diurnal profiles of air pollutants in the atmosphere before the intervention program. The abscissa and longitudinal coordinates represent the percentages of NOₓ and VOCs relative to the real average values measured at MK (i.e., 100% NOₓ = 235.6 ppbv; 100% VOCs = 51.2 ppbv). In other words, they reflect the reduction percentages. For example, 80% NOₓ or VOCs means NOₓ or VOCs was cut by 20%. O₃ was simulated in 220 cases (i.e., 10 VOCs × 22 NOₓ), and the maximum O₃ in each case was extracted.

It was found that within the cutting range of 0-60% of NOₓ (Figure 12 (a)), O₃ increased with the increase of VOCs and the decrease of NOₓ, indicating a VOC-limited regime in O₃ formation. The black curve, perpendicular to the isopleths of O₃, demonstrated the most sensitive response of O₃ to VOCs and NOₓ. Namely, O₃ reduction could be achieved most efficiently towards the abscissa. Using the absolute value of RIR (i.e., |RIR|) as a measure of the O₃-VOCs-NOₓ sensitivity, it increased with the reduction of VOCs and NOₓ. For example, |RIR| for VOCs and NOₓ increased from 0.09 and 0.22 in the scenario of 90% VOCs and 90% NOₓ, to 0.25 and 0.90 in the scenario of 50% VOCs and 50% NOₓ, respectively. From before to during the program, VOCs and NOₓ decreased ~12% and ~15%, respectively (i.e., from point A to B), causing a slight O₃ increment as described in section 3.3.2. The red and green curves in the lower right corner of Figure 12 (a) show the O₃ production as a function of VOCs cut before (NOₓ=100%) and during the program (NOₓ=~85%), respectively. With the reduction of VOCs, O₃ decreased. Since O₃-VOCs-NOₓ sensitivity increased during the program, O₃ decreased by 45.5 and 67.6 pptv with 10% cut of VOCs before and during the program, respectively, which.

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means that O₃ reduction could be achieved more efficiently by further cutting VOCs during the program.

With the continuous reduction of NOₓ, it was expected that the O₃-VOCs-NOₓ sensitivity might change substantially due to the dual role of NOₓ in O₃ formation and titration. Figure 12 (b) shows the O₃-VOCs-NOₓ sensitivity in the cutting range of 60-99.5% of NOₓ. It is noticeable that in the cutting range of ~60-89.5% of NOₓ, O₃ increased with the increase of VOCs and decrease of NOₓ, similar to that in the cutting range of 0-60% of NOₓ. However, a transition area appeared when NOₓ was further cut, where O₃ stayed relatively stable with NOₓ variations, and decreased with VOC reductions. This transition area changed from ~5.5-10.5% (i.e., VOCs =100%) to ~2.5-6% of NOₓ (i.e., VOCs =10%). The appearance of the transition area implied that the titration of O₃ by NOₓ reached the minimum level, and further cutting of NOₓ might actually cause O₃ reduction. As expected, O₃ decreased with the reduction of NOₓ when NOₓ was reduced to lower than ~2.5-5.5% (i.e., ~2.5% and ~5.5% for 10% and 100% VOCs, respectively), and responded weakly to VOC variations. This means that O₃ formation switched to a NOₓ-limited regime. It should be emphasized that this is the first attempt on the study of O₃-VOCs-NOₓ sensitivity at roadsides in Hong Kong, which could be a reference for the formulation and implementation of future air pollution control strategies in Hong Kong.
Figure 12 (a) O$_3$-VOCs-NO$_x$ sensitivity within the cutting range of 0-60% of NO$_x$. The black curve demonstrates the most sensitive response of O$_3$ to VOCs and NO$_x$. Points A and B represent the O$_3$-VOCs-NO$_x$ relationship “before” and “during the program”, respectively. The red and green curves in the small legend show the O$_3$ production as a function of VOC cuts before and during the program, respectively.

Figure 12 (b) O$_3$-VOCs-NO$_x$ sensitivity within the cutting range of 60-99.5% of NO$_x$

3.5.2 O$_3$-VOCs-NO$_x$ sensitivity in LPG

Figure 13 shows the net O$_3$ increment (i.e., positive and negative values indicate the increase and decrease of O$_3$, respectively) as a response of the reduction of VOCs and NO$_x$ in LPG-fueled vehicle exhaust. It was found that the net O$_3$ increment uniformly increased with the increase of NO$_x$ cut and the decrease of VOCs cut. To ensure that O$_3$ would not increase during the program, the reduction of VOCs and NO$_x$ should be within the highlighted area. That is, the highest cutting percentage of NO$_x$ should be less than 68.9% (i.e., point A). Furthermore, when the cutting percentage of VOCs increased from 20% to 100%, the maximum cutting percentage of NO$_x$ for zero O$_3$ increment was between 18.2% and 68.9%, and the cutting ratio of VOCs/NO$_x$ increased from 1.1 to 1.45, suggesting that the cutting ratio of VOCs/NO$_x$ should be > 1.1 in
order to maintain zero O₃ increment. During the program, VOCs and NOₓ in LPG-fueled vehicle exhaust were cut by ~43% and ~89% (i.e., point B), respectively. According to the cutting ratios of VOCs/NOₓ (~1.45) in the high cutting range of NOₓ, the minimum reduction percentage of VOCs should be ~129% when NOₓ was cut by ~89%. In other words, O₃ would inevitably increase in this case.

Indeed, NO₂ experienced an overall increase from 1999 to 2013 at the roadsides in Hong Kong (HKEPD, 2014). According to this study, secondary NO₂ might be more responsible for the increase. Considering nil-emission of NO₂ (section 3.2.2) and low emission of NO from LPG usage (i.e., 4.0% and 1.1-7.3% based on the emission inventory and source apportionment, respectively), an alternative scheme was proposed. Namely, reasonable cutting ratio of VOCs/NOₓ was used to fulfill zero O₃ increase when VOC and NOₓ were cut. Although the O₃ production increase was minor (i.e., 0.4 ppbv or 5.6%) in this study, this scheme could be applicable in future programs. Moreover, since the reactivity and concentration of VOCs influenced their O₃ formation potential, the relative incremental reactivity (RIR) and RIR (weighted concentration) of LPG-related VOCs were calculated. Table 9 shows the RIR and RIR-weighted concentration of propane, propene and n/i-butanes. Propene had the highest RIR (i.e., 5.21×10⁻²), suggesting the highest sensitivity of O₃ production to propene. However, n-butane in LPG was found to have the highest RIR (weighted concentration) (2.61 μg/m³), indicating that cutting n-butane in LPG source was optimal for O₃ pollution control.

![Figure 13 Net O₃ increment as a function the reduction percentages of VOCs and NOₓ in LPG-fueled vehicle exhaust](image)
Table 9 RIR and RIR (weighted concentration) of LPG-related VOCs

<table>
<thead>
<tr>
<th></th>
<th>RIR (%/%×10⁻²)</th>
<th>RIR (weighted concentration) (μg/m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Propane</td>
<td>3.47</td>
<td>1.04</td>
</tr>
<tr>
<td>Propene</td>
<td>5.21</td>
<td>0.30</td>
</tr>
<tr>
<td>i-butane</td>
<td>3.47</td>
<td>1.33</td>
</tr>
<tr>
<td>n-butane</td>
<td>3.47</td>
<td>2.61</td>
</tr>
</tbody>
</table>

4 Conclusions

In this study, an intervention program, initiated in September 2013 and aimed to reduce emissions of VOCs and NOₓ from LPG-fueled vehicles in Hong Kong, was evaluated. The results indicated that LPG-related VOCs and NOₓ decreased significantly during the program, when higher reduction rates were observed for LPG-related VOCs and NOₓ compared to those in previous years. Source apportionment revealed that propane, n/i-butanes and NO in LPG-fueled vehicle emissions were significantly lower during than before the program. It was therefore concluded that the program was remarkably effective in reducing VOCs and NOₓ. To evaluate the program more comprehensively, the variation of O₃ production was simulated with the PBM-MCM model. It was found that LPG-fueled vehicular emission was generally O₃ destructive at the roadside MK site, and the O₃ destruction decreased from 0.47 to 0.06 ppbv due to the intervention program, causing an hourly average O₃ increase of 0.40 ppbv (~5.6%). The LPG-fueled vehicle exhaust generally made negative contributions to the production of OH and HO₂. During the program, the destructive effect weakened and even turned to a positive contribution to HO₂ production, resulting in the increases of OH, HO₂ and HO₂/OH. This was in line with the fact that O₃ increased slightly during the implementation of the program. To improve the program for future application, an O₃-VOCs-NOₓ sensitivity analysis was conducted for ambient air that is not partitioned to sources, and the individual source of LPG-fueled vehicle exhaust. The NOₓ-limited regime in O₃ formation was only found when NOₓ was reduced to less than 5.5%. Furthermore, for the emission reductions in LPG-fueled vehicle exhaust, the maximum NOₓ cutting percentage of 68.9% and the lowest cutting ratio of VOCs/NOₓ (i.e., 1.1) were determined to maintain zero O₃ increment.
Acknowledgements

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