Homogeneous conversion of NO\textsubscript{x} and NH\textsubscript{3} with CH\textsubscript{4}, CO, and C\textsubscript{2}H\textsubscript{4} at the diluted conditions of exhaust-gases of lean operated natural gas engines

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Abstract
Understanding gas-phase reactions in model gas mixtures approximating pre-turbine heavy-duty natural gas engine exhaust compositions containing NO, NH\textsubscript{3}, NO\textsubscript{2}, CH\textsubscript{4}, CO, and C\textsubscript{2}H\textsubscript{4} is extremely relevant for aftertreatment procedure and catalyst design and is thus addressed in this work. In a plug-flow reactor at atmospheric pressure, five different model gas mixtures were investigated in the temperature range of 700-1200 K, using species analysis with electron ionization molecular-beam mass spectrometry. The mixtures were designed to reveal influences of individual components by adding NO\textsubscript{2}, CH\textsubscript{4}, CO, and C\textsubscript{2}H\textsubscript{4} sequentially to a highly argon-diluted NO/NH\textsubscript{3} base mixture. Effects of all components on the reactivity for NO\textsubscript{x} conversion were investigated both experimentally as well as by comparison with three selected kinetic models. Main results show a significantly increased reactivity upon NO\textsubscript{2} and hydrocarbon addition with lowered NO conversion temperatures by up to 200 K. Methane was seen to be of dominant influence in the carbon-containing mixtures regarding interactions between the carbon and nitrogen chemistry as well as formaldehyde formation. The three tested mechanisms were capable to overall represent the reaction behavior satisfactorily. On this basis, it can be stated that significant gas-phase reactivity was observed among typical constituents of pre-turbine natural gas engine exhaust at moderate temperature.

KEYWORDS
ammonia, exhaust gas aftertreatment, gas-phase kinetics, natural gas engines, nitric oxide

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1 | INTRODUCTION

Today, internal combustion engines operated with liquid fuels are already highly optimized; hence, a significant further reduction of fuel consumption is technologically challenging. Therefore, gaseous fuels such as methane and hydrogen seem to become attractive alternatives. Methane, the major component of natural gas, exhibits the highest hydrogen-to-carbon ratio of all hydrocarbons (HCs) and therefore a superior energy-carbon balance. In addition, natural gas engines emit comparably low levels of toxic CO and only moderate amounts of NOx.1 These rather low emissions of the greenhouse gas CO2 and of local pollutants make natural gas engines a promising technology for a wide range of applications. Up to now, they are widely used in stationary applications, that is, in combined heat and power plants for energy production, and in some countries also for transportation vehicles, ships, and even passenger cars. The long-term trend toward natural gas-operated combustion engines, often being interfered by legislative decisions, is supported by an existing natural gas infrastructure in many countries. Although fossil sources currently represent the main feedstock for natural gas, promising alternative concepts like biomethane and power-to-gas technologies can open up further sustainable accesses to the highly valuable energy source methane. In addition, such carbon-neutral approaches have the ability to substantially improve the cradle-to-grave analysis of methane as climate-friendly fuel.

Small natural gas engines are commonly operated under stoichiometric conditions and use the well-established three-way catalyst as exhaust gas aftertreatment to control the emissions of CO, HCs, and NOx. Operation at lean conditions would however further improve the engine efficiency leading to a greenhouse gas advantage of up to 35% compared with a diesel engine. A very crucial issue with lean-operated natural gas engines is the release of unburnt HCs, mainly the major fuel component CH4 with up to over 3 000 ppm. In addition, other small alkanes occurring in natural gas as well as combustion intermediates such as small olefins and formaldehyde (CH2O), a carcinogenic substance, need to be taken care of. All these HCs can be converted by suitable oxidation catalysts with noble metals as active substances. Even though NOx emissions are at least slightly lower in comparison with diesel engines,2 the tightening environmental legislation will make a dedicated NOx abatement system imperative. The reduction of NOx emissions from gas engines is commonly achieved by the well-established technology of selective catalytic reduction (SCR), using NH3 as reducing agent, which is provided by thermolysis and hydrolysis of a urea-water solution known as AdBlue and DEF (Diesel Exhaust Fluid).

The injection of this urea-water solution into the tail pipe requests a minimum exhaust gas temperature of approximately 570 K (200°C) to avoid formation of deposits.3 A much more severe challenge however is the oxidation of methane in the exhaust system because even the best catalysts still require temperatures above 700 K at lean conditions.4,5 Consequently, both the SCR and the oxidation catalysts are positioned closer to the engine, so-called “close-coupled catalysts” in contrast to the conventional “under-floor catalysts.” The catalytic converters are even proposed to be positioned in front of the turbocharger, where also the pressure is significantly higher, up to 5 bar, in addition to the elevated temperatures. Both of these two features would lead to a stronger reduction of pollutant emissions and allow smaller catalytic converters.6 The optimization of the design and operation conditions of those close-coupled catalysts calls for an analysis of potential homogeneous reactions in the exhaust gas due to the rather high temperature. A very crucial point along these lines is the variation of the NH3/NOx ratio due to reactions in the gas phase. In general, the NH3/NOx ratio needs to be unity over the entire inlet cross section of the SCR catalyst to avoid either ammonia or NOx slip; both are not tolerable. Controlling this ratio by adequate dosing of the urea-water solution and additional NH3 and NOx storage-release functions of the catalysts used is a great challenge for the NOx aftertreatment system. Often, ammonia slip catalysts are used in addition, which however lead to the production of significant amounts of N2O, another greenhouse gas. For all these reasons, it is obvious that the kinetics of homogeneous reactions of NH3 and NOx with the other exhaust gas components has to be understood quantitatively, which is the objective of the research study presented here.

There have been only a few studies that deal with the homogeneous reactivity of exhaust gases under those conditions explicitly. Smith et al7 studied the homogenous chemistry in lean-burn exhaust mixtures of internal combustion engines at moderate temperatures, realizing that NO is converted rapidly to NO2, whereas HCs are oxidized without a catalytic converter. Torkashvand et al8 recently found a promotion of the homogeneous oxidation of light alkanes such as CH4 by NOx, which also lead to the additional formation of formaldehyde on a level similar to the amount of HCHO in the engine’s raw emissions. Günter et al9 realized that the performance of SCR catalysts can be significantly influenced at elevated temperature and pressure by homogeneous gas phase reactions due to NO and NH3 oxidation.

The homogeneous reactivity of the respective NH3/NO/O2 mixtures have been investigated in detail both experimentally and numerically.10–15 In some of
these studies, the impact of CO and H₂O on the reactivity was considered. The so-called Thermal DeNOₓ process only works in a narrow temperature range from 900 to 1 400 K, whereas additives such as CO shift the specific NO reduction window toward lower temperatures. The so-called Thermal DeNOₓ process only works in a narrow temperature range from 900 to 1 400 K, whereas additives such as CO shift the specific NO reduction window toward lower temperatures. The kinetic models available were mostly developed for these thermal DeNOₓ conditions. However, these heterogeneous gas-phase reaction mechanisms have not been applied to the best of our knowledge to support the understanding and explain the experimental results of the impact of small HCs on the homogeneous conversion of NOₓ and NH₃ at the specific conditions occurring in the exhaust-gas of lean operated natural gas engines, that is, moderate temperatures (for combustion systems), low concentrations of the reactants, and rather long residence times.

In this work, we experimentally and numerically study the effect of the small HCs CH₄, C₂H₄, and additionally CO on the reactivity of NOₓ and NH₃ in the gas phase under these conditions. Therefore, we used a plug-flow reactor (PFR) coupled to electron ionization molecular-beam mass spectrometry (EI-MBMS) to investigate five gas mixtures (GMs), starting from NH₃/NO/O₂ and subsequently adding NO₂, CH₄, CO, and C₂H₄. The chosen concentrations are close to realistic exhaust GMs of lean burn natural gas engines, and the successive addition of one species at a time allows to individually study the effects of these species on the NH₃/NO reactivity. The experimental results were then analyzed by kinetic simulations using three different kinetic models from literature, which we refer to as CRECK, Glarborg, and Konnov. These mechanisms were developed for high and low temperature HC combustion processes such as the thermal DeNOₓ including the NOₓ chemistry with important HC-nitrogen interaction reactions.

The objectives of this study are twofold: first, the significance of potential gas phase reactions in the presence of the common exhaust components CH₄, CO, and C₂H₄ on the NOₓ and NH₃ concentrations in the exhaust-gas pipe is revealed; second, the prediction capability of three established reaction mechanisms is analyzed for these exhaust-gas relevant conditions.

2 | EXPERIMENTAL

All measurements were performed in a PFR at Bielefeld University. The system, the main features of which have been described in detail previously, was modified slightly regarding preheating and exhaust gas heating to prevent formation of explosive ammonia nitrate. A scheme of the modified PFR is shown in Figure 1.

All gas flows were controlled by calibrated mass flow controllers (MKS Instruments, 5% uncertainty), mixed within a preheated (473 K) gas line and then guided into the PFR. The PFR consists of a heated fused silica tube (7.26 mm diameter, 130 cm length) where a quartz nozzle (100 μm orifice diameter) is placed at the end used for sampling via a two-stage expansion (1st stage ~10⁻⁴ mbar, 2nd stage ~10⁻⁶ mbar). Samples then undergo electron ionization and are analyzed by a time-of-flight mass spectrometer with a mass resolution of m/Δm ~2 000. The experiments were performed at 970 mbar in the temperature regime from 700 to 1 200 K. To ensure plug-flow conditions, a total gas flow rate of 0.5 slm was chosen, which is supported by 2D simulations (see Figure S1 in Supplemental Material 1 (SM1)). This total gas flow leads to residence times between 1.5 and 2.4 s, depending on reactor temperature.

The inlet mole fractions of the investigated GMs are summarized in Table 1. All gases were obtained from Linde AG with purities > 99.5 %. To achieve higher accuracy in the experimental compositions despite the low inlet concentrations, NH₃, NO, NO₂, C₂H₄, and CO were added as
TABLE 1 Experimental conditions for the plug-flow reactor experiments

<table>
<thead>
<tr>
<th>Inlet mole fraction</th>
<th>Gas mixture</th>
<th>Ar</th>
<th>O₂</th>
<th>NH₃ (ppm)</th>
<th>NO (ppm)</th>
<th>NO₂ (ppm)</th>
<th>CH₄ (ppm)</th>
<th>C₂H₄ (ppm)</th>
<th>CO (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GM1</td>
<td>0.939</td>
<td>0.06</td>
<td>1000</td>
<td>1000</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>GM2</td>
<td>0.939</td>
<td>0.06</td>
<td>1000</td>
<td>800</td>
<td>200</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>GM3</td>
<td>0.937</td>
<td>0.06</td>
<td>1000</td>
<td>800</td>
<td>200</td>
<td>3000</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>GM4</td>
<td>0.934</td>
<td>0.06</td>
<td>1000</td>
<td>800</td>
<td>200</td>
<td>3000</td>
<td>1000</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>GM5</td>
<td>0.936</td>
<td>0.06</td>
<td>1000</td>
<td>800</td>
<td>200</td>
<td>3000</td>
<td>200</td>
<td>1000</td>
<td>–</td>
</tr>
</tbody>
</table>

diluted mixture (typically 0.5%-5%) of the respective gas in argon (Linde AG, ±2% relative uncertainty).

The data evaluation mainly follows procedures described in Ref. 25. Calibration factors relative to argon were obtained from the elemental balance for main species including reactants and products. Direct calibration was performed for most detected intermediates. Information on the respective calibration method used for a certain species is given in Table S1 in SM1. Since low reactant concentrations were necessary to perform experiments close to pre-turbine conditions, a significant influence of background signals was observed. This was considered by temperature-dependent background measurements and correction methods described in SM1 (see Figure S2) and summarized in Table S2. The resulting experimental uncertainties are about 30%. Absolute uncertainties in temperature control are ±15 K, which are relevant when comparing experimental results with kinetic simulations. When comparing different GMs, a relative temperature error of only about ±5 K based on repeated measurements with identical conditions needs to be considered. A complete set of all quantitative species profiles is given in SM2.

3 | KINETIC MODELING

Numerical simulations were performed using the software packages Cantera, CHEMKIN Pro, and DETCHEM. Since there is no dedicated PFR code available in Cantera, the PFR was simulated using a ReactorNet of ConstPressureReactors (constant pressure 0-dimensional reactors). Integral reaction flow analyses as well as sensitivity analyses were performed using the batch code in CHEMKIN Pro software, whereas simulations of the reactor flow conditions and the NO/NO₂ equilibrium were performed using DETCHEM CHANNEL and DETCHEMEQUIL codes. The temperature profile of the present flow reactor was obtained by an empiric equation based on direct temperature measurements (compare Figure S3 in SM1). This experimental temperature profile was considered in Cantera by using a concatenation of more than 650 reactor segments, which was seen to be a sufficiently high number to obtain an asymptotic invariance to the number of reactor segments. Pre-reactions were considered exemplarily for GM2 and found to be insignificant for the present highly diluted conditions (see Figure S4 in SM1). Three detailed kinetic models from literature were used in their latest original versions including low-temperature chemistry for NO₅ and HCs. The Konnov mechanism is developed for modeling combustion processes of hydrogen, carbon monoxide, formaldehyde, carbon monoxide, formaldehyde, methanol, methane, C₂ to C₃ HCs, and their oxygenated derivatives including H/C/N/O reactions for in-flame NO₂ formation and reburning. The mechanism contains 129 species. The CRECK mechanism includes pyrolysis, partial oxidation, and combustion of HC and oxygenated fuels. The complete mechanism is developed for the high- and low-temperature kinetics for C₁ to C₁₆ HCs including a NO₂ reaction scheme and contains overall 537 species. The Glarborg mechanism is developed for C₁ to C₂ HC, amine, and cyanide chemistry as well as HC-nitrogen interactions. The model contains 151 species and includes detailed nitrogen chemistry. All three mechanisms are validated over a wide range of conditions.

4 | RESULTS AND DISCUSSION

Figure 2 shows the experimental and predicted mole fraction profiles of the main species for the five GMs. The NH₃/NO and NH₃/NO₂ reactivity is studied upon addition of NO₂ (GM2), CH₄ (GM3), CH₄/CO (GM4), and CH₄/C₂H₄ (GM5). The agreement between experimental and simulation results using the three kinetic models is reasonably good for all five GMs. However, some smaller deviations can be observed between experiments and simulations, but also between the different kinetic models.

In the following sections, we will first analyze the fundamental NH₃/NO₂ system (Section 4.1) and then discuss the influence of HCs and CO (Section 4.2) followed by an investigation of the NH₃/NO₂ ratio which is crucial for the functionality of SCR catalysts (Section 4.3).
FIGURE 2 Experimental mole fraction profiles (symbols) for NH$_3$ (left), NO (center), and N$_2$ (right) for the investigated gas mixtures (GM) compared with simulation results (lines). The experimental uncertainty of 30% is indicated at only one temperature for clarity [Color figure can be viewed at wileyonlinelibrary.com]
4.1 Kinetic analysis for NH₃/NOₓ/O₂ mixtures

In the NH₃/NO system (GM1), the onset temperature of NO and NH₃ is near 1 020 K (Figures 2A–2C). The agreement of experimental and predicted results using the three kinetic models is overall good. However, the Konnov and the CRECK mechanisms show conversion of NO and NH₃ at slightly lower temperatures compared with the Glarborg mechanism by approximately 10 K. N₂ is the main product of this reaction, which is also predicted reasonably well by the models (Figures 2C, 2F, 2I, 2L, and 2O). Nevertheless, a larger scatter is observed in the experimental N₂ mole fraction profile for the HC-containing GM3-GM5 (Figures 2I, 2L, and 2O), since the presence of CO makes it harder to extract a clear N₂ signal from the experimental data due to the small mass difference (0.011 u). At 1 100 K, the Konnov mechanism predicts the highest NH₃ conversion of 99% and the highest N₂ formation, whereas lower conversion can be found using the Glarborg and the CRECK mechanisms. The reduction of NO by NH₃ in the basic NH₃/NO/O₂ mixture has been studied extensively before, both experimentally and numerically, and is well known as SNCR with NH₃ (thermal DeNOₓ). The thermal DeNOₓ process only works in a narrow temperature range from 1 100 to 1 400 K and excess oxygen. Additives such as H₂/CO/CH₄ shift the temperature window to lower temperatures, whereas H₂O has a slightly inhibiting effect. While the binary mixtures of, for example, NH₃+NO or NO+O₂ are inert, the thermal DeNOₓ process works due to chain branching reactions allowing the overall reaction being self-sustaining. Therefore, NH₃ is oxidized by the O/H radical pool, primary OH radicals producing NH₂ which further reacts with NO via two major reaction pathways; one radical producing (R1) and the other chain terminating (R2). The required branching ratio of reactions (R1) and (R2) is defined as α = k_{R1}/(k_{R1} + k_{R2}) with typical values of α = 0.3-0.4 (1 100-1 400 K). In contrast to the thermal DeNOₓ process, highly diluted conditions at moderate temperatures were studied here. A sensitivity analysis was performed at 1 040 K confirming that reaction (R1) is the most sensitive reaction found for both NH₃ and NO conversion. A reaction flow analysis for GM1 is provided in SM1 (Figure S5). The calculated branching ratios α are in the recommended range for the Thermal DeNOₓ process. Figure 3 shows an Arrhenius plot comparing the rate coefficients k for reaction (R1). The origin of the corresponding Arrhenius parameters for reaction (R1) are summarized in Table S3 with calculated k_{R1} and α values at 1 100 K, respectively. The lowest rate coefficient k_{R1} and therefore slower reaction kinetic was found for the Glarborg mechanism which is in accordance with the NH₃ consumption onset shifted slightly toward higher temperatures as shown in Figure 3B. As already discussed by Glarborg et al., small changes to the branching ratio can lead to a shift of onset temperature by 20-50 K.

To underline the importance of the aforementioned reaction and its impact on NH₃ and NO conversion at the highly diluted conditions studied here, the Glarborg mechanism was tentatively modified by increasing the pre-exponential factor A and the temperature coefficient β independently. Both changes led to a higher k_{R1} value and therefore a higher branching ratio indicating a faster reaction of (R1). The impact of this slight modification is visualized in Figure 3A. The k_{R1} values calculated with the Arrhenius parameters from the original Glarborg mechanism are shown as solid dark blue line, whereas the tentatively modified Arrhenius parameters leading to k_{R1} values are presented as a light blue dashed-dotted line. In Figure 3B, the simulation of GM1 was repeated with the adjusted temperature coefficient leading to an earlier conversion compared with the results obtained with the original Arrhenius parameters from the Glarborg mechanism, which agrees better with the experimental data and the simulation results obtained with the CRECK and Konnov mechanisms in Figure 2.

In GM2, NO was partially replaced by NO₂, keeping the overall NOₓ concentration constant (see Table 1). The chosen NO/NO₂ ratio of 4:1 is representative for natural gas engine conditions. Nitrogen dioxide (NO₂) is believed to be a key intermediate in the thermal DeNOₓ process. The modeling predictions for the NO₂ produced at thermal DeNOₓ conditions are according to Glarborg et al in qualitative agreement with measured ones; however, the predicted NO₂ temperature window was extended to higher temperatures, which was not observed experimentally according to Ref. The formation sequence of NO₂ through competitive reactions of HNO and H₂NO with the radical pool are not well characterized but important to predict NO₂ consumption/formation correctly. This might be also important because to the authors'
knowledge, the role of both NO and NO$_2$ in the feed gas as comparable to the exhaust gas of natural gas engines was not investigated before.

Similar to the NH$_2$ + NO reaction channels (R1) and (R2), two further reaction pathways are according to the literature\textsuperscript{16} important at large excess oxygen due to NO$_2$ formation via NO + HO$_2$ = NO$_2$ + OH. Again, one is chain terminating (R3) and the other one is radical producing (R4). The radical producing path (R4) forms NO via the reaction sequence NO$_2$ $\rightarrow$ H$_2$NO $\rightarrow$ HNO $\rightarrow$ NO.

\[
\text{NH}_2 + \text{NO}_2 = \text{N}_2\text{O} + \text{H}_2\text{O} \quad \text{(R3)}
\]

\[
\text{NH}_2 + \text{NO}_2 = \text{H}_2\text{NO} + \text{NO} \quad \text{(R4)}
\]

As can be observed in Figures 2D–2F, NH$_3$ conversion starts about 200 K and NO conversion about 50 K earlier compared with GM1. For both species, the mole fraction profiles of GM2 are flatter compared with the mole fraction profiles of GM1. The flatter shape must be directly correlated to the NO$_2$ addition in the feed gas. Figure 4 shows a reaction path analysis performed for GM2 at 1 040 K. This temperature was chosen to see direct differences in reaction pathways due to NO$_2$ addition compared with GM1 (Figure S4), which was also performed at 1 040 K. The major reaction paths also important for GM1 are shown with solid arrows, whereas the additional reaction paths important due to NO$_2$ addition in the feed gas are denoted with dashed arrows. In the NH$_3$/NO GM, NH$_3$ is mainly oxidized by OH radicals, whereas NH$_3$ can additionally be oxidized by inlet NO$_2$ in GM2. The NO$_2$ addition in the feed gas seems to allow for different reaction pathways producing important radicals which enhance, for example, NH$_3$ conversion at low temperature.

According to the NO/NO$_2$ equilibrium (Figure S6), NO$_2$ is converted to NO with increasing temperature. However, no increase in NO mole fraction can be observed at temperatures below 1 000 K (Figure 2E) and NO consumption starts significantly later compared with NH$_3$ (Figure 2E). At the same time, an increase of N$_2$ was measured experimentally above 800 K and is confirmed by simulation results (Figure 2F). These observations suggest that the formed NO via the NO/NO$_2$ equilibrium and reaction (R4) must be rapidly consumed via reaction (R1) so that no NO conversion can be observed experimentally and numerically below 1 000 K. A sensitivity analysis is shown in Figure 5A for NO and Figure 5B for NH$_3$ at 1 040 K. NO is formed via the NO/NO$_2$ equilibrium and reaction (R4), which was found to be sensitive. According to the mole fraction profile, NO must be directly consumed, possibly via reaction (R1) (NH$_2$ $\rightarrow$ NNH $\rightarrow$ N$_2$), which was again found to be most sensitive for NO and NH$_3$ consumption in all three kinetic models producing N$_2$, whereas quantitative influences are seen to differ for most reactions (Figure 5).

The shapes of NH$_3$ and NO species mole fraction profiles differ from the experimentally measured ones, especially at higher temperatures (Figures 2D–2F). This might indicate that the importance of NO$_2$ as reactant (NO/NO$_2$ equilibrium) and its intermediate species formed are underestimated at temperatures above 1 050 K in all three kinetic models. It can be concluded that the presence of NO$_2$ in the feed gas has a retarding effect on NO conversion (Figure 2E) and hence a negative impact on
**FIGURE 4** Major reaction pathways at 1040 K in the oxidation of NH$_3$ for GM2. The solid arrows indicate the major reaction pathways of GMI and additional pathways due to NO$_2$ addition are denoted with dashed arrows. Simulations with the Konnov (normal), Glarborg (bold), and CRECK (italic) mechanisms [Color figure can be viewed at wileyonlinelibrary.com]

**FIGURE 5** Sensitivity analysis for (A) NO and (B) NH$_3$ at 1040 K with the Konnov, Glarborg, and CRECK mechanisms [Color figure can be viewed at wileyonlinelibrary.com]
4.2 Kinetic analysis of CO, CH₄, and C₂H₄ addition

GM3-GM5 all contain additional HC species representative for natural gas engines, namely CH₄, CO, and additionally C₂H₄ in lesser amounts in GM5. These species...
The decomposition of the HC additives in the fuel mixtures is shown in Figure 7. Again, the three kinetic models can capture the species mole fraction profiles very well. However, the onset with the Glarborg and CRECK mechanisms is shifted about 30 K toward higher temperatures.

The mutual promoting effect of HC/NO, HC/NO2, and HC/NOx mixtures has been studied extensively before. Investigations of the CO/NO/NH3 and CH4/CO/NO/NH3 systems remain rather scarce, whereas studies of the whole HC/CO/NH3/NOx system containing both HC and NH3 as reducing agent close to exhaust-gas aftertreatment conditions of natural gas engines are not known. Since simulation results are in quite good agreement with experimental data (Figure 2) at these highly diluted exhaust natural gas conditions, a reaction flow analysis was performed for GM3 as the first mixture containing both N and C species to gain deeper insight into the HC/NH3/NOx reaction behavior. For each mechanism, the reaction flow analysis was performed at two temperatures, namely a lower temperature corresponding to an initial, additional 5% NO formation (ie, −5% NO conversion) and the temperature of maximum NO conversion, which is at the same temperature as the CO and CH2O peak temperature. The maximum NO conversion temperature was chosen because major deviations between the three kinetic models were observed at this temperature (compare Figure 2). In general, major reaction pathways do not differ at these two chosen temperatures except for one aspect. At the maximum NO conversion temperature, all mechanisms have in common that H-abstraction from NH3 forms NH2 as major initial step in NH3 oxidation. At the lower temperature where 5% additional initial NO is formed, however, NH3 can be additionally reduced by NO2 in the Konnov and CRECK mechanisms. Since other major reaction pathways do not differ significantly at low and high temperatures, only the reaction flow analysis at the maximum NO conversion temperature is discussed in detail. In Figure 8, the reaction flow analysis is shown for the NH3/NO/NO2 species consumption in GM3 at both temperatures for each mechanism. The direct interaction of HC/N species is highlighted by reacting HC species written in bold italic characters next to the arrows.

The H-abstraction from NH3-forming NH2 radicals was found as initial step in NH3 oxidation in all three mechanisms. The produced NH2 radicals are mainly consumed by reactions (R1) and (R2) that were also found to be the major NO reduction pathways for GM1 and GM2 (Figure S4). Around 900 K, the direct oxidation pathway of NH3 with NO2 is of minor importance; however, the reaction between NH2 and NO2 might be crucial for NO formation. Comparing the experimental data with the three kinetic models, it is obvious that the Konnov mechanism...
Figure 8: Major reaction pathways for NH₃/NO/NO₂ consumption in GM3. The interacting HC species are written above the arrows highlighted by bold italic letters. Simulations were performed at the temperature where an initial, additional amount of 5% NO is formed (−5% NO conversion, shown in brackets) with the Konnov (normal, red, \(T = 834\) K), Glarborg (bold, blue, \(T = 866\) K) and CRECK (italic, purple, \(T = 894\) K) mechanisms as well as at maximum NO consumption temperature with the Konnov (normal, red, \(T = 900\) K), Glarborg (bold, blue, \(T = 920\) K), and CRECK (italic, purple, \(T = 926\) K) mechanisms [Color figure can be viewed at wileyonlinelibrary.com]

This does not seem surprising since these reaction pathways were already identified as the most important NO oxidizing pathway at low temperature HC oxidation in mixtures doped with NO. 39,40 The promoting effect of NO on HC oxidation was traced back to the formation of chain carriers such as CH₃O and CH₃O₂ radicals, which are no longer chain terminating due to the addition of NO. This finding can be transferred to the present study, since HC radicals promote NOx and NH₃ consumption.

In Figure 9, a reaction flow analysis for CH₄ oxidation is shown with all three kinetic models performed at 5% NO formation (ie, −5% NO conversion) shown in brackets and maximum NO conversion temperature. No differences were observed in major reaction pathways therefore only the reaction flow analysis at maximum NO conversion temperature is discussed in the following. Important nitrogen species are written above the arrows in bold, italic letters. The CH₃O and CH₃O₂ radicals are formed by CH₃
FIGURE 9 Integral reaction flow analysis of CH₄ oxidation pathways for GM3. Simulations were performed at the temperature where an initial, additional amount of 5% NO is formed (−5% NO conversion, shown in brackets) with the Konnov (normal, red, $T = 834$ K), Glarborg (bold, blue, $T = 866$ K), and CRECK (italic, purple, $T = 894$ K) mechanisms as well as at maximum NO consumption temperature with the Konnov (normal, red, $T = 900$ K), Glarborg (bold, blue, $T = 920$ K), and CRECK (italic, purple, $T = 926$ K) mechanisms [Color figure can be viewed at wileyonlinelibrary.com].

oxidation with NO₂ (R7) and O₂, whereas the latter is of minor importance due to high energy barrier,¹⁰ concluding that the methyl radicals are the most important interacting HC-/N-species.

\[
\text{CH}_3 + \text{NO}_2 = \text{CH}_3\text{O} + \text{NO} \quad \text{(R7)}
\]

\[
\text{CH}_3\text{NO}_2 + [M] = \text{CH}_3 + \text{NO}_2 \quad \text{(R8)}
\]

In addition, reaction (R8) is also discussed in the literature,¹³,³⁹ which is particularly important for low temperature HC oxidation in which CH₃NO₂ is produced and competes with reaction (R7). However, this species was not found to have a significant impact on the NOₓ/NH₃/HC reactivity in the reaction flow analysis and was not detected experimentally, possibly due to the highly diluted conditions in the present study.

The importance of CH₃ and therefore CH₃O radicals can be underlined by a reaction flow analysis for CH₃O radicals shown in Figure 10 performed at 920 K using the Glarborg mechanism. The reactions (R7) (#932) and (R5) (#968) with NO/NO₂ as oxidizing species are most responsible for CH₃O radical formation, whereas CH₃O radicals are consumed by oxidation reactions, producing CH₂O as intermediate species of partial HC oxidation.

The dehydrogenation of CH₂O toward CO₂ via the reaction sequence CH₂O → CH₂O → HCO → CO → CO₂ then also contributes to the OH radical pool. Also, for OH radicals, the major formation/consumption pathways are shown in Figure 10. The OH radical formation is mainly driven by NO/NO₂ interchanging reactions (#728, #730) such as reaction (R6). Major consumption pathways of OH radicals are HC oxidation reactions such as CH₄ + OH (#68), CH₂O + OH (#50), and CO + OH (#29). The formation of the intermediate species CH₂O is mainly by oxidation of CH₃ via the reaction path CH₃ → CH₃O₂/CH₃O → CH₂O, whereas its consumption is mainly via the reaction sequence CH₂O → HCO → CO → CO₂. Formaldehyde is not only an important intermediate in CH₄ combustion but also crucial regarding possible emissions of natural gas engines because of its toxicity and carcinogenicity. Simulated mole fraction profiles of CH₂O are shown in Figure 11, which are compared with normalized signals of CHO. Due to the very small mass difference between CH₂O and NO of only 0.013 u, a possible CH₂O signal is overlapped with the stronger signal of NO. However, a signal of CHO at $m/z = 29$ u was detected and corrected for isotopes (¹⁵N¹⁴N and ¹³C¹⁶O). CHO is a very prominent fragment of CH₂O in EI-MBMS, and therefore the normalized CHO signal can be considered directly proportional to the CH₂O mole fraction. Similar profile shapes are observed in Figure 11 in the experiment and for the Konnov, Glarborg, and CRECK mechanisms. Consistent with the HC consumption results in Figure 7, the maximum CHO signal is observed at nearly identical temperatures of about 890 K for GM3 and GM4, whereas the maximum for GM5 is detected at a lower temperature of about 870 K. Maximum formaldehyde mole fractions in the $10^{-4}$ range can be expected according to the simulations that are non-negligible regarding the share of HCs in the initial GMs. The present results lead to the suggestion that CH₄ dominates the CH₂O formation for all three mixtures, since neither CO nor C₂H₄ addition substantially modify the CH₂O concentration.
Formaldehyde is then further oxidized by the reaction sequence CH$_2$O $\rightarrow$ HCO $\rightarrow$ CO $\rightarrow$ CO$_2$. Alzueta et al.\textsuperscript{37} studied a mixture of CO/NH$_3$/NO/O$_2$/H$_2$O experimentally, indicating that the presence of CO affects the radical pool and leads therefore to chain branching reactions but does not contribute directly to NO reduction. As shown in Figure 6, CO peaks at 900 K as well as CO$_2$ and H$_2$, whereas the NO consumption starts rapidly. Since in the present study lean mixtures were investigated, CO is oxidized very fast to CO$_2$ producing important radicals around 900 K and reducing the overall oxygen availability in the system. As already observed for GM2, the NO/NO$_2$ equilibrium strongly impacts the reaction kinetic of these GMs. At temperatures below 900 K, NO is produced which was observed experimentally and is confirmed by simulation results. Equilibrium calculations in Figure 12 for GM3 show that NO$_2$ is still present in the GM below 900 K, which can then be reduced by H or CH$_3$ radicals to NO (see Figure 8). With increasing temperature, the NO/NO$_2$ equilibrium is then shifted toward NO. Therefore, the equilibrium is not only dependent on temperature but also on oxygen concentration. This leads to the conclusion that if HCs are present in the GM, they will be oxidized very fast to CO and H$_2$ and further to CO$_2$, consuming oxygen, which reduces the oxygen availability drastically. This influences the NO/NO$_2$ equilibrium in shifting it further to lower temperatures allowing the sharp NO consumption profile above 900 K (see Figure 6).

The addition of further HC and C species, namely CO in GM4 and CO/C$_2$H$_4$ in GM5 has only minor impact on the NH$_3$/NO$_x$ reactivity. The impact of CO is in comparison with CH$_4$ less pronounced; however, the addition of C$_2$H$_4$ should have a stronger promoting effect than CH$_4$.\textsuperscript{37,39} This can be observed by a slight temperature shift about 10 K toward lower temperature compared with the addition of CH$_4$ in GM3 (Figure 7). This only slight temperature shift in GM5 must be due to the lower concentrations and highly diluted conditions. In the present study, it can be concluded that the influence of additional CO (GM4) and minor amounts of C$_2$H$_4$ (GM5) on NO$_x$ conversion is significantly lower than the effect of CH$_4$ addition at highly diluted exhaust-gas aftertreatment conditions.

4.3 | NH$_3$/NO$_x$ ratio

Since the NH$_3$/NO$_x$ ratio of one is crucial for SCR catalytic converters to function properly, experimental and simulation results are shown in Figure 13 for GM2 and GM3, respectively. Please note, that the absolute uncertainty of the experimental NH$_3$/NO$_x$ ratio is shown in
models. Even though these three mechanisms predict the trends quite well, none of them leads to a quantitative agreement in the temperature ranges of 900-1 150 K and 850 and 900 K for GM2 and GM3, respectively. Again, the strong impact of the presence of CH4 in the exhaust is clearly observed. Since the NH3/NOx ratio begins to deviate from unity at rather low temperatures as occurring in the tail pipe close to the engine, it is obvious that homogenous gas phase reactions have to be considered for the development of exhaust gas aftertreatment systems for lean-operated natural gas engines.

5 | CONCLUSION

In this work, the reactivity of the NO/NH3/O2 system has been studied regarding the influence of NO2, CH4, CO, and C2H4, which are important components of engine exhaust in lean-operated natural gas engines, that is, in excess of oxygen. Our study combines PFR experiments coupled to EI-MBMS in the temperature range from 700 to 1 200 K with simulations using three well-established kinetic models from the literature, namely CRECK,19–21 Glarborg,16 and Konnov.22 Methane, which is always present in the exhaust of these engines, was found to have a very strong impact on the reactivity of NOx and NH3; the onset of conversion of these two components was decreased by nearly 200 K in comparison with a mixture free of methane and NO2. This mutual promoting effect was traced back to NO and NO2 first enhancing CH4 oxidation due to CH3O
radical formation (CH₃ oxidation with NO/NO₂). CH₃O is further oxidized to CH₂O and in the end CO₂ producing important radicals, which further enhance NO and NH₃ reactivity. Therefore, a small amount of CH₄ in the exhaust gas might be considered advantageous regarding the required temperature of DeNOₓ processes. The impact of NO₂ is significant as well but not so pronounced. The influences of CO and C₂H₄ added to the mixture on the reactivity remained rather small. Since NH₃ and NOₓ are converted to a very different extent, and NOₓ is even produced by NH₃ oxidation, the NH₃/NOₓ ratio decreases significantly with increasing temperatures. Even for rather moderate temperatures, the function of NH₃ as reducing agent for NOₓ in a SCR system is limited by the onset of gas-phase reactions, which has severe implications on the ammonia (urea-water solution) dosing strategy. Obviously, quantitative models are needed to describe the homogeneous conversion of NH₃ and NOₓ in the presence of CH₄ in the exhaust of lean-operated natural gas engines.

The three mechanisms used to simulate these effects were able to predict all major trends and the experimentally observed temperature dependence of the NH₃/NOₓ ratio qualitatively well for all five mixtures. Since all three mechanisms were not developed using conditions we applied here, we very much appreciate their impressive ability to model the NH₃ and NOₓ reactivity under these scarcely investigated conditions. Both model-based designs of the aftertreatment system and optimization of the ammonia dosing strategy request quantitative model predictions. Therefore, we would like to recommend further developments of these mechanisms specifically for these conditions with highly diluted concentrations (on the order of hundreds of ppm) of NH₃, NOₓ, and CH₄ at moderate temperatures. Here, we began to analyze the kinetics of these mechanisms for the specific conditions of this application in more detail, which led to different behaviors of the individual mechanisms. The reaction flow analyses reveal differences regarding main C-N interaction. A small underprediction of NO mole fractions was found for the Konnov mechanism, which could be influenced by an underestimation of the importance of the reaction sequence NH₂ → H₂NO → HNO → NO. According to the reaction flow analyses, this sequence is apparently more important in both Glarborg and CRECK mechanisms. Regarding the exhaust gas compositions, intense reactions in the gas phase can be expected involving both nitrogen and carbon chemistry. With this analysis, we can give some hints in which direction further mechanism development should be conducted and we would like to encourage further research here.

As we have learnt, homogeneous gas-phase reactions affect the concentrations of NH₃ and NO in the exhaust pipe of lean-operated natural gas engines. They are likely to also occur inside the channels of the catalytic converters, that is, the NH₃-SCR and the oxidation catalysts positioned close to the engine. Hence, we can expect some interference between homogeneous gas-phase reactions and heterogeneous catalytic reactions inside the converter as observed for many other systems such as by the Mantzaras group for catalytically supported combustion. Consequently, we will next study the interaction of gas-phase and surface reactions inside the catalytic converters.

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REFERENCES


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