Detailed Kinetic Mechanism for the Oxidation of Ammonia Including the Formation and Reduction of Nitrogen Oxides

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Abstract

This work introduces a newly developed reaction mechanism for the oxidation of ammonia in freely propagating and burner stabilized premixed flames as well as in shock tubes, jet stirred reactors and plug flow reactors experiments. The paper mainly focuses on pure ammonia and ammonia-hydrogen fuel blends. The reaction mechanism also considers the formation of nitrogen oxides, as well as the reduction of nitrogen oxides depending on the conditions of the surrounding gas phase. Doping of the fuel blend with NO₂ can result in acceleration of H₂ autoignition via the reaction $NO_2+HO_2 \rightleftharpoons HONO+O_2$ followed by the thermal decomposition of HONO, or in deceleration of H₂ oxidation via $NO_2+OH \rightleftharpoons NO+HO_2$. The concentration of HO₂ is decisive for the active reaction pathway. The formation of NO in burner stabilized premixed flames is shown to demonstrate the capability of the mechanism to be integrated into a mechanism for hydrocarbon oxidation.

1. Introduction

Clean, reliable and renewable fuels are important for future power systems. Alternative fuels for power generation and internal combustion engines of various transportation systems have recently been intensely discussed. Hydrogen has attracted attention as a carbon free transportation fuel with zero-CO₂ emissions. It can be produced in an electrolytic process from overpower of alternative energy sources, i.e. wind or solar power. Its low energy density and its limited storage capabilities are still barriers for a market launch of hydrogen fueled vehicles. Alternatively, Hydrogen is discussed as a reactant for catalytic carbonization processes leading from CO₂ to methane or methanol.

For any hydrogen-atom containing fuel the hydrogen-oxygen chemistry plays a fundamental role. Hydrogen is not only an important fuel, but the chemical kinetics involving H, O, OH, HO₂, H₂O and H₂O₂ also determine the radical pool in hydrocarbon reaction systems ¹. Reactions from this sub-mechanism show the highest sensitivity in almost all hydrocarbon oxidation systems.

Investigations of alternative fuels aim at decreasing the usage of fossil fuels at reasonable transportation costs. Ammonia was considered until the 1960s and recently gained attention again. It is also known as one key species in the de-NOx process ^{2–5}, which can be applied in a narrow temperature window within the combustion chamber or in a catalytic aftertreatment system. Ammonia is recognized as a carbon free fuel and as a hydrogen carrier or storage compound ^{6,7} with a high content of hydrogen atoms per unit volume. Ammonia is catalytically produced from nitrogen and hydrogen. Although the industrial process technology has been steadily improved over the years, it still compares to the process developed by Haber and Bosch in the early 20th century: $N_2(g)+3 H_2(g) \rightarrow 2 NH_3(g) (\Delta H^{0}_{298} = -92.2 kJ/mole)$. As hydrogen still must be produced to obtain ammonia in large quantities, ammonia can be seen as an additional hydrogen energy

vector. The industrial ammonia production process is highly optimized and hard to improve and it is unclear as to whether catalytic carbonization processes can become significantly more efficient in future.

Ammonia is carbon-free and can potentially be burned in an environmentally benign way, exhausting water, nitrogen and nitrogen oxides as only emissions. However, the combustion of ammonia as a fuel in internal combustion engines also has several drawbacks ⁸; i.e. low laminar flame speed, high auto-ignition temperature, high heat of vaporization, narrow flammability limits (16-25 % by volume in air) and high toxicity. It is therefore considered as a dual fuel component. This was demonstrated in modified spark-ignition (SI) and compression-ignition (CI) engines at research ^{9,10} or prototype level ¹¹. Depending on the combustion process ammonia can either lead to reduced or increased NO_x emissions.

To support the engine development process by simulations, detailed mechanisms for the oxidation of ammonia including the formation and the reduction of NO_x are needed. Previous studies ^{12–17} have reported on the development of detailed chemical mechanisms for the NH₃/NO_x system. However the parameter ranges for the validation of these mechanisms was limited due to the lack of experimental data. For example, Mathieu et al. ¹⁷ compared the predictions of nine different mechanisms from the literature against their measured data from shock tube experiments and concluded that further model improvements are needed for spanning the experimentally explored range of conditions. Other authors came to similar conclusions: Hayakawa et al. ¹⁸, for the prediction of laminar flame speeds of ammonia/air blends in a constant volume cylindrical combustion chamber under elevated pressure conditions; Xiao et al. ¹⁹ for the prediction of NH₃/H₂ laminar flame speeds from Li et al. ²⁰ and NH₃ ignition delay times from shock tube experiments ¹⁷. Recently, Zhang et al. ²¹ developed a detailed kinetic mechanism for the H₂/NO_x and

Syngas/NO_x systems. They demonstrated the sensitivity of the thermochemistry on the prediction of shock tube ignition delay times, species concentrations in a jet stirred reactor (JSR) and a flow reactor (FR). However, their study does not include NH₃ containing fuel blends.

This situation calls for the development of a detailed kinetic H₂/CO/C₁/NH₃/NO_x model with a significantly extended validation target range, which is the objective of the present work. To this end the here derived model is critically tested taking into account laminar flame speeds, ignition delay times, speciation in jet stirred reactors (JSR), in plug flow reactors (PFR) and in burner stabilized flames (BSF). The compilation strategy for our new "nested" mechanism is along the lines we have established for C1-C8 hydrocarbon fuels in the past fifteen years (e.g.^{22–27}). The idea is to merge existing, well tested models and adapt the resulting mechanism to simulations comprising a much broader target range than all previous studies. For the identified elementary reaction steps with highly sensitive kinetic data the increased simulation constraints by the augmented target range allow for a more consistent choice of kinetic parameters within the plausible margins. The virtues of our nested mechanism approach were very recently discussed by Westbrook and co-workers in a review type paper on the development and improvement of kinetic models from the beginnings in the 1970ies to present applications in engine simulations ²⁸.

In this paper we concentrate on the $H_2/NH_3/NO_x$ model, and limit the demonstration of the C_1 system to one illustrative experiment by Lamoureux et al.¹⁶. A wider range of experiments is shown in the supporting information, together with the detailed reaction mechanisms used in this work.

2. Development of the kinetic model

The development of the model is discussed here in three subsections. To keep the paper easy to read we focus on the new aspects of the here derived model and refer to the literature where similar compilations for sub-mechanisms exist. In the first subsection the H₂/CO sub-model is only briefly described. Here the kinetic data is mainly taken from existing databases. The subsection on the NH₃/NO_x sub-model introduces a newly merged model and explains the choice of kinetic data from the available references. Here the kinetic data for several highly sensitive reactions were discussed in more detail. The C₁/NO_x mechanisms is literature based and selectively augmented for H₂/NO_x/C₁ cross reactions.

2.1 H₂/CO kinetic model: The detailed chemical kinetic mechanism of H₂ and CO is mostly based on the recommendations of Baulch et al. ²⁹ and the uncertainty boundaries proposed by the authors. We refer Baulch et al. ²⁹ because it is a comprehensive source, which provides a detailed discussion of uncertainties for the compiled reaction rate constants. Elementary reactions which were not available from ²⁹, or which cannot be expressed in the standard Chemkin format are adopted from other publications ^{30–33}. The H₂/CO kinetic model is validated against experimental data from literature which include 87 sets of laminar flame speed, 39 sets of ignition delay times from shock tubes, 16 sets species concentrations in JSRs, 27 sets of species concentrations in PFRs, 8 sets of species concentrations in BSF and 4 sets of species concentrations in shock tube experiments. The choice of kinetic data in the resulting mechanism and its performance is similar to other publications of this decade ^{33–43}. The detailed mechanism for the H₂/CO model is provided in supporting information of this work. The full set of validation calculations is found on ⁴⁴.

<u>2.2 NH₃/NO_x kinetic model</u>: The starting point of the NO formation and reduction mechanism was the development by Lamoureux et al.¹⁶. Though they validated their model against own data

and a large set of experimental data from literature, they did not consider experiments dedicated to ammonia in their model development. We therefore do not demonstrate any extensive comparison of predictions of the model used in this work and the original model from ¹⁶. However to underline why the model update is necessary we will show an ammonia-air laminar flame speed comparison between Lamoureux et al.¹⁶ and the present work. To match the large set of experimental data published in the literature the complete NO_x sub mechanism needed to be revised and updated very carefully.

Moreover we included additional nine nitrogen containing species – N₂H₃, N₂H₄, H₂NN, HNOH, NH₂OH, HNO₂, HONO₂, NO₃ and HNO₃ – and their respective sub-mechanisms. The importance of the inclusion of the N₂H_x chemistry in NO_x modelling is shown by Allen et al.⁴⁵ in their experimental and numerical work on the oxidation of H₂/N₂O mixtures in a flow reactor. The inclusion of these sub-mechanisms has made updates in the original mechanism necessary. Most of the published models in literature are dedicated to particular fuels or mixtures of interest for a limited range of conditions. The novelty of the present model comes from the ability to comprehensively address the pure fuels (H₂, H₂/CO, NH₃, CH₄, CH₃OH, CH₂O) and mixtures at the same time for a large set of experimental conditions. Sub-mechanism of NH₃, NH, NH₂, N₂H₂, NO, NO₂, N₂O etc. which were already present in initial mechanism from ¹⁶ were also updated. The reactions presented in Table 1 have been identified as highly important for the here presented model and its performance.

Reactions	А	n	Ea	References	
$NH_3(+M) \rightleftharpoons NH_2+H(+M)$	9.000E+16	-0.39	1.103E+05	Baulch et al. ²⁹	
Low pressure limit	2.000E+16	0.00	9.315E+04		
TROE	/ 0.42 4581.0 102.0				
Third body efficiency	H ₂ /1.0/H ₂ O/6.40/CO ₂ /1	Griffiths and Barnard ⁴⁶			
	35/HE/0.35/CO/0.75/C				
$NH_3(+M) \rightleftharpoons NH+H_2(+M)$	7.000E+15	Baulch et al. 29			
Low pressure limit	4.665E+14	0.00	9.315E+04		
TROE	/ 0.42 4581.0 102.0				
Third body efficiency	H ₂ /1.0/H ₂ O/6.40/CO ₂ /1	.50/O ₂ /0.45/	$N_2/0.40/AR/0.$	Griffiths and Barnard ⁴⁶	
	35/HE/0.35/CO/0.75/C	$H_4/3.0/C_2H_6$	/3.0/		
$NH_3+NH_2 \rightleftharpoons N_2H_3+H_2$	1.000E+11	0.50	2.160E+04	Coppens et al. 47	
$NH_3+H \Rightarrow NH_2+H_2$	6.360E+05	2.39	1.017E+04	Klaus ⁴⁸	
NH ₃ +OH≒NH ₂ +H ₂ O	2.040E+06	2.04	5.660E+02	Klaus ⁴⁸	
NNH≒N ₂ +H	6.500E+07	0.00	0.000E+00	Miller and Glarborg ³	
$NNH+NO \Rightarrow N_2+HNO$	5.000E+13	0.00	0.000E+00	Klippenstein et al. 49	
$NH_2+H \rightleftharpoons NH+H_2$	7.300E+13	0.00	5.000E+03	Baulch et al. ²⁹	
NH ₂ +NO≒NNH+OH	3.800E+10	0.425	-8.140E+02	1.65*Miller and	
				Glarborg ³	
$NH_2+NO \Rightarrow N_2+H_2O$	2.800E+20	-2.70	1.258E+03	Miller and Glarborg ³	
$NH_2+NO \Rightarrow N_2O+H_2$	1.000E+13	0.00	3.370E+04	Duynslaegher et	
				al ¹⁴	
$NH_2+NO_2 \leq N_2O+H_2O$	1 600E+16	-1 40	2.680E+02	Park and Lin ⁵⁰	
$NH_2 + NO_2 \Rightarrow H_2O + H_2O$	6 500E+16	-1 40	2.680E+02	Park and Lin ⁵⁰	
NH+H=N+H2	2.011E+13	0.00	0.000E+02	0.67*Baulch et al. ²⁹	
NH+O≒NO+H	5.000E+13	0.00	0.000E+00	Klaus ⁴⁸	
NH+O=N+OH	3.000E+12	0.00	0.000E+00	0.5* Duvnslaagher	
	1 200E+06	1.50	1.000E±02	Vlinnenstein et el. 49	
	1.300E+00	1.30	1.000E+02	Klippenstein et al. 49	
$\frac{1}{1000} = 1000 + 10000 + 1000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 10000 + 1000$	4.000E+03 1 000E+27	2.00	0.300E+03	Straibarg at al. 51	
$N_2H_2+M \Rightarrow N_1H+H$	1.900E+27	-3.03	0.010E+04	Skreiberg et al. ⁵²	
$\frac{NH_2+NH \Rightarrow N_2H_2+H}{NH_2+NH_2+NH_2+H}$	4.300E+14	-0.272	-7.700E+01	Klippenstein et al. 52	
$\frac{NH_2+NH_2=N_2H_2+H_2}{NH_2+NH_2+NH_2+NH_2+NH_2+NH_2+H_2}$	4.000E+15	0.00	1.184E+04	Company at al. 47	
$\frac{1}{1} \frac{1}{2} \frac{1}{1} \frac{1}$	5.000E+10	0.00	$0.000E \pm 04$	Vlingenetain et al. 52	
$\frac{1}{1} \frac{1}{1} \frac{1}$	$3.000E \pm 14$	-0.414	$0.000E \pm 01$	Kippenstein et al.	
	1.000E+34	$1.0E \pm 20/$	1.90/E+05		
	1 520E±15	0.41	0.000E±00	Pagmusson at al 53	
$NO+\Pi(+M) \Rightarrow \Pi NO(+M)$	1.320E+13	-0.41	0.000E+00	Kasiliussell et al.	
	2.400E+14				
HNO+H-NO+H.	70.82 1.0E-30 1.0E+ 4.400E+11	Skraibarg at al ⁵¹			
$HNO+H \rightarrow NO+H_2$	4.400E+11	0.72	0.300E+02	Skreiberg et al. ⁵¹	
$NO+OH(+M) \leftarrow HONO(+M)$	1.100E+13	0.00	0.000E+00	Skreiberg et al. ⁵¹	
Low program limit	1.100E+14	-0.30	0.000E+00	Kasmussen et al. ³³	
TROF	$\frac{5.57211+25}{0.75}$ 1 0F-20 1 0F+				
HONO+OH←NO+H2O	1 700E+12 0 00 5 200E+02			Rasmussen et al 53	
$HONO+H \leftarrow HNO+OH$	5 600E+12	0.00	$= 5.200 \pm 0.02$	Skreiberg et al ⁵¹	
	2 100E+12	0.00	_4 800E±03	Baulch et al 2005 ²⁹	
$NO_2+H = NO_2+OH$	2.100E+12 2.500E+14	0.00	6 760F+02	0.5*Baulch et al 29	

Table 1: List of important reaction identified in H/N/O scheme. Units are cm, mol, s, cal

$NO+O(+M) \rightleftharpoons NO_2(+M)$	2.950E+14	-0.40	0.000E+00	Baulch et al. 2005 ²⁹
Low pressure limit	3.336E+20	-1.60 0.000E+00		
TROE	/0.80 1.0E-30 1.0E+3			
Third body efficiency	H ₂ /1.0/H ₂ O/6.40/CO ₂ /1	Griffiths and Barnard ⁴⁶		
	35/HE/0.35/CO/0.75/C			
$NO_2+HO_2 \Rightarrow HONO+O_2$	1.910E+00	3.32	3.044E+03	Rasmussen et al.53
NO ₂ +H ₂ ≒HONO+H	1.300E+04	2.76	2.977E+04	Rasmussen et al.53
$N_2O(+M) \rightleftharpoons N_2+O(+M)$	1.300E+12	0.00	6.257E+04	Röhrig et al. 54
Low pressure limit	4.00E+14	0.00	5.660E+04	
Third body efficiency	N ₂ / 1.7/ O ₂ / 1.4/ CO ₂ / 3			
N ₂ O+H≒N ₂ +OH	2.530E+10 0.00 4.550E+03		Powell et al. 55	
DUPLICATE				
$N_2O+H \rightleftharpoons N_2+OH$	5.000E+14	0.00	1.810E+04	
DUPLICATE				
$N_2O+H \rightleftharpoons N_2+OH^*$	1.600E+14	0.00	5.030E+04	Hidaka et al. 56
$NO_2+HO_2 \Rightarrow HONO+O_2$	1.910E+00	3.32	3.044E+03	Rasmussen et al.53
$NO_2+HO_2 \Rightarrow HNO_2+O_2$	1.850E+01	3.26	4.983E+03	Rasmussen et al.53
HNO_2 (+M) \Rightarrow HONO(+M)	2.500E+14	0.00	3.230E+04	Rasmussen et al.53
Low pressure limit	3.100E+18	0.00	3.150E+04	
TROE	/1.149 1E-30 3.125E+03 1E+30/			

The thermal NO_x formation is given by the Zeldovich mechanism ⁵⁷. The rate parameters of these reactions are adopted from the recommendation of Baulch et al.²⁹ and optimized within the uncertainty limits provided by the authors to best match experimental data from the literature. A detailed description of other sub-mechanisms which are included in the NH₃/NO_x scheme and the adaptation of the rate parameters are briefly explained below highlighting the important reactions and their kinetics. We follow the kinetic models from Skreiberg et al.⁵¹, Klippenstein et al.^{49,52}, Miller and Glarborg ³, Glarborg et al.^{58,59}, Klaus ⁴⁸, Allen et al.¹², Coppens et al.⁴⁷, Mathieu et al.¹⁷, Mendiara and Glarborg ⁶⁰, Powell et al.⁵⁵, Duynslaegher et al.¹⁴ and Rasmussen et al.⁵³ and sources cited therein.

NO₂ sub-mechanism:

NO₂ kinetics is one of the most important sub-mechanisms in almost all of the cases studied here, which are ignition delay times in shock tube experiments, speciation in flow reactors and jet stirred reactor studies on the H₂/O₂/NO_x and C₁/NO_x systems. The reactions are very important and highly sensitive in the interconversion process of NO \leftrightarrow NO₂ which is discussed in the results and discussion part of this paper (speciation in JSR and PFR). Reactions involving NO₂ are also sensitive on ignition delay time predictions, which is apparent from the sensitivity analysis in Figure 4. Reactions particularly important in the formation/consumption of NO₂ are:

NO+HO₂≓NO₂+OH

NO₂+H≓NO+OH.

In this study, rate parameters for these two reactions are adopted from Baulch et al.²⁹. The importance of these two reactions in promoting the reactivity of the system by forming OH radicals has been discussed in many studies ^{53,61–64}. In this work the rate constant of the NO₂+H \Rightarrow NO+OH reaction has been decreased by 50% to best match the wide range of experimental data taken into consideration which is within the uncertainty proposed by Baulch et al.²⁹.

For lean and high pressure conditions NO₂ can also be formed via the reaction NO+O(+M) \Rightarrow NO₂(+M); the rate parameters of this reaction are taken from the recommendation of Baulch et al.²⁹ in the temperature range of 200-2000 K. The rate constant of the reaction NO₂+O \Rightarrow NO+O₂ is adopted from experimental work of Bemand et al.⁶⁵ who performed direct measurements in the temperature range of 298-1055 K. The suggested rate is consistent with the experimental studies by Estupiñán et al.⁶⁶, Avallone ⁶⁷ and the theoretical study of Shiekh et al.⁶⁸. Another reaction, which is important under typical combustion conditions, is NO₂+HO₂ \Rightarrow HONO/HNO₂+O₂. Glarborg et al.⁶⁹ in their formaldehyde-NOx interaction study pointed out that the reaction NO₂+HO₂≓HONO+O₂ competes for the HO₂ radical with the reaction HO₂+OH≓H₂O+O₂. Also in the shock tube study of Mathieu et al.⁶³ on the H₂/O₂/NO₂/Ar system NO₂+HO₂≓HONO+O₂ is significant in the sensitivity analysis when 100 ppm and 400 ppm of NO₂ are present in the initial mixture. Rasmussen et al.⁵³ performed a theoretical study on (NO₂+HO₂≓HONO+O₂, NO₂+HO₂≓HNO₂+O₂, NO₂+H₂≓HNO₂+H, HNO₂(+M)≓HONO (+M), HNO₂+OH≓NO₂+H₂O) and confirmed that the HNO₂ isomer HONO, is formed the via same reaction channel. They further discussed that HNO₂ may act as an OH sink through the reaction HNO₂+OH≓NO₂+H₂O inhibiting the system reactivity. Furthermore it may isomerize to HONO and decompose to NO and OH. The latter case is confirmed by the simulation results in our present study. Therefore we have adopted the rate parameters from the work of Rasmussen et al.⁵³. The direct reaction of NO₂ with H₂ has two routes forming HONO+H/HNO₂+H which was proposed in ⁵³. The reaction NO₂+H₂ ≓HONO+H appeared in the sensitivity analysis for the ignition delay time in the work of Mathieu et al.⁶³ and also in this work (see Figure 4). The rate constants of these reactions (NO₂+H₂=HONO+H/HNO₂+H) are also adopted from Rasmussen et al.⁵³.

N₂O sub-mechanism:

N₂O is an important intermediate in the thermal DeNO_x process ⁷⁰ which is mainly formed through the amine radical NH+NO \rightleftharpoons N₂O+H. Another potential route for N₂O formation is NH₂+NO₂ \rightleftharpoons N₂O+H₂O. N₂O can also work as an oxidizing agent ^{56,71–74} which dissociates to produce O atoms *via* N₂O(+M) \rightleftharpoons N₂+O(+M). The backward of this reaction becomes significant under lean and high pressure conditions, and opens an alternative pathway to NO formation. Atomic oxygen further takes part in the chain branching step H₂+O \rightleftharpoons H+OH increasing the radical pool. The reaction (N₂O(+M) \rightleftharpoons N₂+O(+M)) is also one of the channels consuming N₂O in the thermal DeNOx process ⁷⁰. We have adopted the rate constant of this reaction from the work of Röhrig et al.⁵⁴ who studied the pressure dependence of the thermal decomposition of N₂O in shock tube experiments in the pressure range of 0.3-450 atm and at temperatures ranging from 1570-3100 K. The reaction N₂O+H \rightleftharpoons N₂+OH is another important reaction in the N₂O chemistry. This reaction competes with the reaction N₂O+H **⇒**NH+NO for the consumption of H atoms. Powell et al.⁷² studied the laminar flame speeds of H₂/N₂O and C₁-C₃/N₂O mixtures. They adopted the kinetic scheme of Allen et al. 1998 ⁴⁵ to perform an accompanying numerical study. Here they modified the rate parameter of $N_2O+H \rightleftharpoons N_2+OH$ for a better match with their measurements. In their subsequent study Powell et al.55 performed a kinetic model evaluation on a large set of published experimental data (ignition delay times obtained in shock tubes, speciation in a burner stabilized flame, speciation in a flow reactor) and their own previous work (laminar flame speeds) with a large fraction of N₂O in the reactant mixture. They found the reaction N₂O+H \rightleftharpoons N₂+OH as one of the most important and sensitive reactions for accurately predicting the wide set of experimental data which they considered. Powell et al.⁵⁵ modified their previous rate parameters ⁷² to best match the range of experimental data taken into consideration. In our present study we have adopted the rate parameters for the reaction (N₂O+H \rightleftharpoons N₂+OH) from the work of Powell et al.⁵⁵ and this rate constant value is within the uncertainty of a factor 2 of the rate constant recommended by Baulch et al.²⁹. We also find this reaction to be highly sensitive in predicting the H₂/N₂O laminar flame speed (Figure S2), ignition delay times (Figure S5 and Figure S7) and predicting the speciation in a burner stabilized flame (Figure S23).

Other reactions in the N_2O sub-scheme which are of particular interest are the reactions of N_2O with O atoms:

N₂O+O≓NO+NO

 $N_2O+O \rightleftharpoons N_2+O_2$.

In a burner stabilized flame (Figure 10) where N₂O is present in the reactant mixture as an oxidizing agent the NO formation is mainly controlled by the branching ratio between these two reactions. In our present work we have adopted the rate constants of these reactions from the recommendation of Baulch et al.²⁹. The O atom of N₂O can be abstracted when reacting with OH and NO to form N₂+HO₂ and NO₂+N₂. Rate parameters of these reactions are adopted following the suggestion of Mebel et al.⁷⁵, who calculated rate constants using ab initio transition state theory. The rate constant of the reaction N₂O+N \rightleftharpoons N₂+NO is taken from Mathieu et al.¹⁷. Mevel et al.⁷⁶ and Mathieu et al.^{17,77} suggested that inclusion of the excited hydroxyl radical (OH*) in the mechanism, particularly for the reaction N₂O+H ⇒ N₂+OH*, improves predictions for the calculated ignition delay times compared to using the ground state OH radical. Mevel et al.⁷⁶ and Mathieu et al.¹⁷ compared there experimentally determined OH* profile with the numerically predicted OH* formation profile and found a good agreement. They also showed the difference when using the ground state OH profile. Following there suggestion we included this reaction $(N_2O+H \rightleftharpoons N_2+OH^*)$ in our scheme adopting the rate parameter from the experimental work of Hidaka et al. ⁵⁶ who performed a OH* chemiluminescence study in N₂O/H₂/Ar mixtures in a shock tube at 2 atm.

NO₃ sub-mechanism:

The only NO₃ formation routes are *via* reactions NO₂+O(+M) \rightleftharpoons NO₃ and NO₂+NO₂ \rightleftharpoons NO₃+NO. In our present study the inclusion or exclusion of the NO₃ reaction scheme does not have any significant effect on predicted ignition delay times, speciation in flow a reactor and a jet stirred reactor for the H₂/O₂/NO_x system. However, this subset is included for the sake of completeness of the kinetic scheme. The NO₃ reaction sub-mechanism is adopted from the model of Mendiara and Glarborg ⁶⁰ who studied the effect of CO₂ concentration on the ammonia oxidation during oxy-fuel combustion of methane in a flow reactor at atmospheric pressure and temperatures ranging from 973-1773 K. Additionally the reactions $NO_3+NO_3\Rightarrow NO_2+NO_2+O_2$ and $NO_3+HO_2\Rightarrow HNO_3+O_2$, not considered by Mendiara and Glarborg ⁶⁰, were included in our present work. The rate parameters of these two addition reactions were adopted from Coppens et al.⁴⁷ who measured the laminar burning velocity of CH₄/H₂/O₂/N₂ mixtures at varying H₂ fractions in a heat flux burner and NO formation in the studied flames.

HNO₃ sub-mechanism:

In the flow analysis for the H₂/O₂/NO_x system in jet stirred and flow reactors we observe that a small amount of HNO₃ is formed *via* the reaction NO+HO₂+M \Rightarrow HNO₃+M. The formed HNO₃ dissociates to NO₂ and OH *via* the reaction NO₂+OH(+M) \Rightarrow HNO₃(+M) which participates in the NO/NO₂ interconversion process. Other reaction channels that are involved in HNO₃ consumption are the reactions with H, OH and NH₂ radicals. The complete HNO₃ reaction scheme is adopted from the kinetic model of Coppens et al.⁴⁷.

NH₃ sub-mechanism:

The thermal decomposition of NH₃ features two product channels forming NH₂ and H radicals as the major route and NH and H₂ as the minor pathway. The rate parameters have been adopted from the compilation of Baulch et al.²⁹. In our scheme we have introduced the pressure dependent rate parameters for the thermal decomposition reaction of NH₃ in the TROE format which is not included in other published models ^{15,17,47,49,78,79}. Hydrogen abstraction from NH₃ occurs in reactions with H, O, OH and HO₂ radicals mainly forming the NH₂ radical. The NH₃+H/OH \rightleftharpoons NH₂+H₂/O reaction rate parameters are adopted from the modeling work of Klaus 1997 ⁴⁸ who performed a detailed kinetic modeling study of NO_x formation in a burner stabilized flame for NH₃ and C₁-C₄ hydrocarbons. The NH₃+O \Rightarrow NH₂+OH rate parameters are adopted from Baulch et al. ²⁹. Kinetic data for the NH₃+HO₂ \Rightarrow NH₂+H₂O₂ reaction rate are taken from the kinetic model of Skreiberg et al.⁵¹ who a performed kinetic study of ammonia oxidation in a flow reactor based on experiments by Hasegawa and Sato ⁸⁰. The reaction NH₃+NH₂ \Rightarrow N₂H₃+H₂ and its kinetic data are adopted from the model of Coppens et al.⁴⁷.

NH₂ sub-mechanism:

This sub-mechanism is an important part of the DeNO_x process where NH₃ is used as reducing agent. The reaction rate for hydrogen abstraction from NH₂ via H atoms, which forms NH and H₂, is adopted from Baulch et al.²⁹. The rate constant of this reaction ($NH_2+H \Rightarrow NH+H_2$) is decreased by 12 % to better predict the ammonia flame speed, which is within the given uncertainty range. The reaction of NH₂ with O atoms has three product channels (HNO+H/NH+OH/NO+H₂) whose rate parameters are adopted from the kinetic model of Klaus 1997⁴⁸. The rate constant of NH₂+OH \Rightarrow NH+H₂O is also taken from the modeling work of Klaus 1997 ⁴⁸. The reaction of HO₂ with NH₂ has two product channels forming H₂NO+OH and NH₃+O₂. Similarly, O₂ reacts with NH₂ forming H₂NO+O and HNO+OH with the first channel being the major path⁴⁹. The rate parameters of these reactions are adopted from the kinetic scheme of Skreiberg et al.⁵¹. In our present study, these reactions are less sensitive compared to other reactions in the NH₂ submechanism. The reaction of NH₂ with NO which is chain branching in the channel to NNH+OH and chain terminating in the channel to N₂+H₂O is very important in the thermal DeNO_x process ^{3,70}. The rate parameters of these two reactions have been adopted from the study of Miller and Glarborg³ who obtained the total rate constant value by fitting the channel branching in simulations of published experimental data. However, the rate constants used in this work are

increased by 65% in order to better match the wide range of published experimental data considered in the mechanism validation. The reaction NH₂+NO₂ has two product channels forming H₂NO+NO and N₂O+H₂O. Glarborg et al.⁸¹ studied the NH₃/NO₂ system in a flow reactor and concluded that the NH₂+NO₂ reaction forming H₂NO+NO is the more important channel recycling NO₂ back to NO. The importance of this reaction was again emphasized by Miller and Glarborg ³ in their experimental and modeling study of H₂/CO/NO_x combustion in a flow reactor. In our present work we adopt the rate parameters of these reactions from the experimental work of Park and Lin ⁵⁰ who performed a mass spectrometric study of the NH₂+NO₂ reaction in the temperature range of 300-910 K . Additionally the reaction NH₂+NO \Rightarrow N₂O+H₂ is included in our present work whose rate constant is adopted from the kinetic model of Duynslaegher et al.¹⁴ who performed a modeling study of NH₃ oxidation in a burner stabilized low pressure premixed flame.

Additional reaction rates for NH₂+NH₂ \Leftrightarrow NH₃+NH, NH₂+NH \Leftrightarrow NH₃+N and NH+NH \Leftrightarrow NH₂+N are adopted from Klippenstein et al.⁵², who calculated rate parameters using ab initio transition state theory. The reaction NH₂+N \Leftrightarrow N₂+H+H which is not included in kinetic schemes of ^{3,15,51,52,78} is here adopted from the kinetic model of Klaus 1997 ⁴⁸. The reaction rate constants for the reactions NH₂+HNO \Leftrightarrow NH₃+NO and NH₂+HONO \Leftrightarrow NH₃+NO₂ are adopted from Mebel et al.⁸², who calculated rate constants using ab initio molecular orbital theory.

NH sub-mechanism:

The rate parameters of hydrogen abstraction from NH by H atoms forming N and H₂ are adopted from Baulch et al.²⁹ and decreased by 33% which is within the given uncertainty. The reaction NH+O has two product channels (NO+H and N+OH). The reaction forming NO is the dominant path while the channel forming N+OH is usually not included in published mechanisms ^{15,17,49,51,78}. We found this channel to be important and included it in our kinetic scheme. The rate parameters

for the reaction NH+O=NO+H are adopted from the modeling work of Klaus 1997 ⁴⁸ while the rate constant of the NH+O \Rightarrow N+OH reaction is adopted from the kinetic scheme of Duynslaegher et al.¹⁴ and reduced by 50% for improving the model predictions. The reaction of OH with NH has three routes forming HNO+H, NO+H₂ and N+H₂O respectively. However, in the models from 15,17,49,78 the second channel (NH+OH \Rightarrow NO+H₂) is not included. The rate parameters of these reactions are adopted from the kinetic model of Klaus 1997⁴⁸. The rate constant of the NH+O₂ reaction forming HNO+O and NO+OH is taken from the model of ⁴⁹. For the reactions $NH+N \Leftrightarrow N_2+H$, $NH+NO \Rightarrow N_2+OH$, $NH+NO_2 \rightleftharpoons N_2O+OH$, NH+NO₂≒HNO+NO, and NH+HONO≒NH₂+NO₂ kinetic data is taken from Klippenstein et al.⁴⁹ They studied the role of NNH in NO formation and reduction. In particular their theoretical study focused on the reaction systems of NNH+O, NNH+O₂ and NH₂+O₂ using ab initio transition state theory. In addition they provided rate constants for the NH+NO and O+N₂O reaction. They concluded that in the thermal DeNO_x process, the role of the NH₂+NO reaction system is significant. Their mechanism is based on the above mentioned study by Miller and Glarborg³. They⁴⁹ validated their model against the published experimental data with emphasis on flow reactor experiments. In their discussion they stated that NH is formed *via* reactions involving NH₂, particularly NH₂+OH≒NH+H₂O. The formed NH is partially oxidized to NO via the NH+O₂ reaction. In addition, they concluded that the NH+NO reaction forming NNH is a minor channel contributing to NO formation. Finally, we included the reaction NH+N₂O \Rightarrow N₂+HNO and took the kinetic data from the model by Duynslaegher et al.¹⁴.

NNH sub-mechanism:

The NNH sub-mechanism is compiled adopting the rate parameter and reactions from Klippenstein et al.⁴⁹, Miller and Glarborg ³, Glarborg et al.⁵⁸ and Allen et al.¹². Miller and Glarborg ³ proposed

the NNH consumption through NNH \Rightarrow N₂+H and NNH+O₂ \Rightarrow N₂+HO₂. In the study of Kasuya and Glarborg⁸³ a higher amount of NO₂ was detected at higher O₂ concentrations which allowed them (Miller and Glarborg ³) to consider NNH+O₂ \Rightarrow N₂+HO₂ as the major channel for NNH consumption. Based on theoretical and experimental studies from the literature they considered a NNH life time of 1.5×10^{-8} s and thus derived the NNH \Rightarrow N₂+H rate constant of 6.27×10^{7} /s. Details on the lifetime of the NNH radical were discussed in the work of Klippenstein et al.⁴⁹. They assumed the NNH life time to be 10^{-9} s and derived the rate constant of NNH \Rightarrow N₂+H to be 1.0×10^{9} /s which is around a factor 15 higher than proposed by Miller and Glarborg ³. In our present work we have adopted the rate constant value from the kinetic model of Miller and Glarborg ³ which gives better results especially for laminar flame speeds of NH₃. The reaction of NNH+O₂ features two channels: N₂+HO₂ and N₂+H+O₂, the former being the major channel with a branching ratio of 80% (see ref.³). The rate parameters of these reactions have been adopted from the kinetic scheme of Glarborg et al.⁵⁸. The recombination reaction of NNH forming N₂H₂ (NNH+NNH \Rightarrow N₂H₂+H₂), which is not included in work of Miller and Glarborg ³ and Klippenstein et al.⁴⁹, is included in the present work for the completeness of the mechanism adopting the rate parameters from Allen et al.¹². NNH is also consumed by the attack of H, OH, NO, NH, and NH₂ radicals mainly forming N₂ as major product. The reaction NNH+O has three product channels forming N₂+OH, NH+NO and N₂O+H and the rate parameters of these reactions are adopted from Klippenstein et al.⁴⁹.

N₂H₂/H₂NN/N₂H₃/N₂H₄ sub-mechanism:

The sub-mechanism of N₂H₂, N₂H₃, N₂H₄ and H₂NN has been largely taken from the kinetic models of Skreiberg et al.⁵¹, Allen et al.¹², Coppens et al.⁴⁷, Klaus 1997 ⁴⁸ and Klippenstein et al.⁵². Allen et al.¹² studied the combustion of CO/N₂O/H₂O/N₂ mixtures in a flow reactor in a pressure

range of 3-15 atm and a temperature range of 950-1123 K. They used the experimentally measured species profiles as reference for deriving a detailed kinetic model for CO/N₂O interaction. In their later study ⁴⁵ they showed and discussed the importance of N_2H_x chemistry for predicting the speciation in the performed flow reactor experiments. In this work we show the effect of N_2H_x chemistry on NH₃ laminar flame speed in Figure 1.

NH₂OH sub-mechanism:

The complete NH₂OH sub-mechanism is adopted from the work of Klippenstein et al.⁵² who investigated the thermal decomposition of NH₂OH in a combined experimental (shock tube) and theoretical study. In their experiments they measured the OH radical time histories over the temperature range of 1355-1889 K and their predicted OH profile was in good agreement with experimental OH profile. They concluded that NH₂OH decomposes to NH₂ and OH representing the major route while the formation of NH₃+O is a minor channel and can be neglected. Using ab initio transition state theory they derived the rate constants of other elementary reactions of relevance for the NH₂OH sub-mechanism. The reaction HNOH+HNO \Rightarrow NH₂OH+NO, which was not included in their mechanism, is adopted from Coppens et al.⁴⁷.

HNO sub-mechanism:

The HNO sub-mechanism is compiled mainly from the work of Rasmussen et al.⁵³, Skreiberg et al.⁵¹, Glarborg et al.⁵⁸, Klaus 1997 ⁴⁸ and Coppens et al.⁴⁷. The hydrogen abstraction reactions from HNO by H, O, OH and O₂ forming NO+H₂/OH/H₂O/HO₂ respectively are adopted from the ammonia oxidation modeling work of Skreiberg et al.⁵¹. The recombination reaction HNO+HNO \Rightarrow N₂O+H₂O is adopted from the model of Glarborg et al.⁵⁸, who performed an experimental and kinetic modeling study of C₁/C₂ hydrocarbon interaction with NO_x in a flow

reactor. The reactions HNO+N \Rightarrow NO+NH and HNO+NO \Rightarrow N₂O+OH and its rate constants are adopted from the kinetic model of Klaus 1997 ⁴⁸ while the reactions HNO+N \Rightarrow N₂O+H, HNO+NH \Rightarrow NH₂+NO are adopted from Coppens et al.⁴⁷. Those reactions that we adopted from ^{47,48} were not included in the kinetic schemes of ^{15,49,53,78} and we included these reactions in our scheme for the completeness of the kinetic model. The rate parameters of the reaction of HNO with NO₂ forming HONO+NO and the thermal decomposition reaction of HNO forming NO+H are taken from the kinetic scheme of Rasmussen et al.⁵³.

HON sub-mechanism:

The complete HON sub-mechanism is adopted from the kinetic model of Mathieu et al.¹⁷. They performed an experimental and modeling study of ammonia oxidation in a shock tube spanning wide ranges of pressure, temperature and fuel-oxidizer ratio. In their model HON is mainly formed *via* the combination reaction NO+H+M \Rightarrow HON+M. HON is consumed *via* the reactions: HON+H \Rightarrow HNO+H, HON+H \Rightarrow NH+OH, HON+O \Rightarrow OH+NO, HON+OH \Rightarrow HONO+H and HON+O₂ \Rightarrow NO₂+OH. Among these reactions, HON+H is important for NO formation and consumption via HNO. The reaction forming HNO+H (which is an isomerization) is the dominating channel (branching ratio ~70%) and the formed HNO reacts with NO to yield NH+NO₂ (NH+NO₂ \Rightarrow HNO+NO). Another route (HON+H \Rightarrow NH+OH) which directly forms the amine radical, also has a NO reduction potential by contributing to the amine radical pool.

H₂NO sub-mechanism:

The H₂NO chemistry is an important part of the thermal DeNOx mechanism ³. H₂NO is mainly formed by reactions of the amine radical (NH₂):

 $NH_2+O_2 \rightleftharpoons H_2NO+O$

$NH_2+HO_2 \Rightarrow H_2NO+OH$

$NH_2+NO_2 \rightleftharpoons H_2NO+NO.$

Klippenstein et al. ⁴⁹ performed a numerical analysis of the NH₂+O₂ reaction and confirmed that the H₂NO formation route is the dominant path which competes with the HNO+OH channel. Similarly, NH₂+NO₂ has two product channels forming N₂O+H₂O and H₂NO+NO, the former being the major product ^{3,49,84} with 80% branching ratio. The formed H₂NO either dissociates unimoleculary to HNO+H or is attacked by H, O, OH, HO₂, NO and NH₂ radicals forming HNO and the corresponding products. The reaction of H₂NO with H atoms has another possible channel forming NH₂+OH. If this route is fast enough to produce NH₂ then the amine radical pool will take part in the NO reduction process *via* the reaction NH₂+NO \Rightarrow N₂+H₂O/NNH+OH. H₂NO can also react with O₂ and NO₂ forming HNO+HO₂ and HONO+HNO, respectively. The H₂NO submechanism is adopted from the kinetic model of Glarborg et al.⁵⁹ who performed flow reactor experiments on reduction of NO by CO and H₂ under fuel rich conditions in the temperature range of 1200-1800 K. In their study they found that the H₂NO chemistry was not a significant contributor in reducing NO.

HNOH sub-mechanism:

HNOH is an isomer of H₂NO which is mainly formed *via* the recombination reaction HNO+H+M \Rightarrow HNOH+M. HNOH is consumed in reactions with H, O, OH, HO₂, NH₂, O₂ and NO₂ mainly forming HNO and NH₂. The reaction HNOH+H \Rightarrow NH₂+OH is of particular importance because of its direct contribution to the amine radical pool. Miller and Glarborg 1999 mentioned in their study that inclusion of HNOH chemistry in their model did not have any significant effect on the performance. Glarborg et al.⁵⁹ and Skreiberg et al.⁵¹ suggested in their studies that the

HNOH route can be an alternative path to the amine pool (HNO \rightarrow HNOH \rightarrow NH₂). However, both authors reached the same conclusion that the contribution of HNOH reactions to NO reduction is limited by the rate of formation of HNOH *via* (HNO+H+M \rightleftharpoons HNOH+M). In our present work the complete HNOH sub-mechanism is adopted from the model of Skreiberg et al.⁵¹.

HONO/HNO₂ sub-mechanism:

In the work of Mueller et al.⁶¹ on the $H_2/O_2/NO_x$ system HONO is mainly formed from the HNO+NO2 HONO+NO reaction. At low NO concentrations, the formed HONO rapidly dissociates to NO and OH via NO+OH(+M) \Rightarrow HONO(+M) and takes part in a chain propagating sequence. With increasing NO concentration in the system the reaction HONO+OH \Rightarrow NO₂+H₂O becomes important together with reactions $NO+H(+M) \Rightarrow HNO(+M)$ and and HNO+NO₂≒HONO+NO it forms a chain terminating sequence with net result H+OH≒H₂O. In the experimental and modeling work of Glarborg et al.⁶⁴ on the CO/NO_x combustion in a flow reactor HONO is found to be formed via the recombination reaction NO+OH(+M) \Rightarrow HONO(+M) and the formed HONO reacts mainly with OH to form NO₂ (HONO+OH \Rightarrow NO₂+H₂O) taking part in a chain terminating process. In our present study we find small amounts of NO₂ reacting with the HO₂ radical forming HONO+O₂. The formed HONO thermally decomposes to form NO+OH producing the reactive OH radical. Therefore, it is obvious that depending on the various conditions HONO chemistry can inhibit or promote the reactivity of the system. In our present work the HONO sub-scheme is taken from the model of Rasmussen et al.⁵³. The additional reactions HONO+H≒HNO+OH and HONO+H≒NO+H₂O which were not included in ⁵³ are adopted from Skreiberg et al.⁵¹.

HNO₂ is a structural isomer of HONO and in our present study we find HNO₂ is mainly formed via the reaction NO₂+HO₂ \Leftrightarrow HNO₂+O₂ and NO₂+H₂ \Leftrightarrow HNO₂+H. Since HNO₂ is less stable than HONO ⁵³ it readily isomerizes to more the stable HONO. Other reactions that can take part in HNO₂ consumption are HNO₂+O \Leftrightarrow NO₂+OH and HNO₂+OH \Leftrightarrow NO₂+H₂O. The rate parameters of these reaction are adopted from Rasmussen et al.⁵³ and are based on ab initio calculations on the CBS-QB3 level of theory.

HONO₂ sub-mechanism:

HONO₂ formation can happen *via* two reaction channels: NO₂+OH(+M) \Rightarrow HONO₂(+M) and HONO+NO₂ \Rightarrow HONO₂+NO. Among these, the former route is only important for HONO₂ formation. HONO₂ consumption proceeds mainly via the reactions with H and OH radicals:

HONO₂+H≒H₂+NO₃

 $HONO_2+H \rightleftharpoons H2O+NO_2$

HONO₂+H⇔OH+HONO

 $HONO_2+OH \rightleftharpoons H_2O+NO_3.$

The HONO₂ scheme and the rate parameters are adopted from Rasmussen et al.⁵³.

<u>**2.3** C₁/NO_x mechanism</u>: This part of the mechanism is an extension of the H₂/CO/NH₃/NO_x submechanisms. The H₂/CO submechanism is extended to include C₁ chemistry and the NH₃/NO_x submechanism is extended to include nitrogen chemistry related to carbon species. The target is to include C₁ hydrocarbon species as fuel molecules; i.e. methane (CH₄), methanol (CH₃OH) and formaldehyde (CH₂O). The present model is different from most of the published models in the literature because it is able to predict combustion characteristics of the fuels H_2 , CO, NH₃ to C₁ together with NO_x chemistry. This makes the model more robust for in cylinder combustion modelling of conditions which cannot be studied in standard reactor experiments typically used for model development. The complete reaction mechanism is provided in the supporting information.

2.4 Thermochemistry and Transport properties: The thermochemical properties for the species in the H/C/O mechanism are adopted from the Goos et al. thermochemical database ⁸⁵. The thermochemical properties of nitrogen species are adopted from Lamoureux et al.¹⁶ except for those additional species included in this work. The thermochemistry of additional species are adopted from Goos et al.⁸⁵ while HNO₂ and HONO₂ thermodata (not available from ⁸⁵) is adopted from Rasmussen et al.⁵³. Furthermore the data for NCN are taken from ²⁴.

The transport properties for the species in the H/C/O mechanism are taken from Seidel et al.²⁵. The transport properties of nitrogen species are adopted from Lamoureux et al.¹⁶ while those of the additional species are taken from Coppens et al.⁴⁷.

3. Results and Discussion

The developed H₂/CO/C₁/NH₃/NO_x kinetic model has been validated against a large set of published experimental data. The validation targets include laminar flame speeds, ignition delay times, speciation in burner stabilized premixed flames, speciation in PFR and JSR. Several numerical and experimental investigations on nitrogen chemistry have been reported in the last decades. However, there are only very few investigations on H₂/NH₃/NO_x and C₁/NO_x chemistry together. This is the focus of this study. The study also addresses the importance of speciation

studies in burner stabilized flames. In all plots shown below the lines represent the simulation results using our present model and symbols represent the experimental data from literature unless stated differently. The modeling results are shown for experiments using nitrogen (NH₃, NO, N₂O, NO₂) containing fuels. All simulations were performed using the LOGEsoft 1.08.00 ⁸⁶ package.

3.1 H₂/CO/NH₃/NO_x model validation

In this section we will validate our proposed model against a wide range of experimental data published in the literature and briefly comment on the model performance. Table 2 summarizes the experimental studies which we used for model development and validation.

Table 2: Ex	perimental	targets u	used for	·H ₂ /CO	/NH ₃ /NC	D _x model	develo	pment and	validation
		<u> </u>							

Experimental devices	Measured	Experimental conditions	References
	Properties		
Cylindrical flow tube ^(Fig.1)	Laminar flame	1.0 atm, 293 K, $\phi = 0.8$ -1.25 for	Zakaznov et al. ⁸⁷
-	speed	NH ₃ /air mixture	
	(1 data set)		
Constant volume	Laminar flame	101 kPa, 295 K, $\phi = 0.72$ -1.12 for	Pfahl et al. ⁸⁸
Combustion vessel ^(Fig.1)	speed	NH ₃ /air mixture	
	(1 data set)		
Constant volume	Laminar flame	1.0 atm, 298 K, $\phi = 0.8-1.78$ for	Ronney ⁸⁹
combustion vessel ^(Fig.1)	speed	NH ₃ /air mixture	-
Constant volume	Laminar flame	1.0 atm, 295 K, $\phi = 0.9-1.3$ for	Jabbour & Clodic 90
combustion vessel ^(Fig.1)	speed	NH ₃ /air mixture	
	(1 data set)		
Constant volume	Laminar flame	$1.05 \text{ atm}, 298 \text{ K}, \phi = 0.9-1.2 \text{ for}$	Takizawa et al. ⁹¹
combustion vessel ^(Fig.1)	speed	NH ₃ /air mixture	
	(1 data set)		
Constant volume	Laminar flame	1.0-5.0 bar, 298 K, $\phi = 0.7$ -1.3 for	Hayakawa et al. ¹⁸
combustion vessel ^{(Fig.1,}	speed	NH ₃ /air mixture	-
Fig.S2)	(3 data set)		
Constant volume	Laminar flame	1.0 atm, 298 K, $\phi = 1.0$ for	Lee et al. ⁹²
combustion vessel ^(Fig.S1)	speed	$NH_3/H_2/air mixture, H_2 = 0.1-0.5$	
	(1 data set)		
Constant volume	Laminar flame	1.0 atm, 298 K, $\phi = 1.0$ for	Li et al. ²⁰
combustion vessel ^(Fig.S1)	speed (1 data set)	$NH_3/H_2/air mixture, H_2 = 0.35$ -	
		0.55	
Constant volume	Laminar flame	$101 \text{ kPa}, 298 \text{ K}, \phi = 0.1-4.0 \text{ for}$	Checkel et al.93
combustion vessel ^(Fig.S1)	speed (1 data set)	NH ₃ /NO mixture	
McKenna flat flame	Laminar flame	$0.8 \text{ atm}, 298 \text{ K}, \phi = 0.8-2.2 \text{ for}$	Powell et al. ⁷²
burner ^(Fig.S2)	speed (1 data set)	$H_2/N_2O/N_2$ mixture	

Shock tube ^(Fig.2,Fig.S3)	Ignition delay	1.4-30.0 atm, 1560-2455 K, $\phi =$	Mathieu and Petersen ¹⁷
	times (12 data sets)	0.5-2.0 for NH ₃ /O ₂ /Ar	
	· · · · · ·	mixtures	
Shock tube (Fig.3,Fig.S4)	Ignition delay	$1.6-32.0$ atm, 940-1675 K, $\phi = 0.5$	Mathieu et al. ⁷⁷
	times (12 data sets)	for $H_2/O_2/N_2O/Ar$	
	· · · · · ·	mixtures	
Shock tube (Fig.3,Fig.S6)	Ignition delay	$1.5-30.0$ atm, 1038-1744 K, $\phi =$	Mathieu et al. ⁶³
	times (12 data sets)	0.3-1.0 for H ₂ /O ₂ /NO ₂ /Ar	
	· · · · · ·	mixtures	
Shock tube ^(Fig.S5)	Ignition delay	2.0 atm, 1400-2000 K, $\phi = 0.5$ -2.0	Hidaka et al. ⁵⁶
	times (3 data sets)	H ₂ /N ₂ O/Ar mixtures	
Shock tube (Fig.S7)	Ignition delay	1.4-10.4 atm, 1654-2221 K for	Kopp et al. ⁷¹
	times (2 data sets)	H ₂ /CO /N ₂ O/Ar mixture	
Jet stirred reactor (Fig.4, Fig.5,	Species profiles	1.0-10.0 atm, 700–1150 K, $\phi =$	Dayma & Dagaut ⁶²
Fig.S8-Fig.S12)	(78 data sets)	0.1-2.5 for H ₂ /O ₂ /NO _x /N ₂	
		mixtures	
Jet stirred reactor	Species profiles	1.0 atm, 800-1400 K, $\phi = 0.1$ -2.0	Dagaut et al. ⁹⁴
(Fig.S13,Fig.S14)	(9 data sets)	for H ₂ /CO/O ₂ /NO _x /N ₂	_
		mixtures	
Flow reactor (Fig.6)	Species profiles	10.0 atm, 802 K, H ₂ /O ₂ /NO	Mueller et al. ⁶¹
	(3 data sets)	mixtures	
Flow reactor (Fig.S15)	Species profiles	3.0 atm, 995 K, H ₂ /N ₂ O/NH ₃ /N ₂	Allen et al. ⁴⁵
	(4 data sets)	mixtures	
Flow reactor (Fig.7,Fig.S16-	Species profiles	1.05 atm, 800-1400 K,	Glarborg et al. ⁶⁴
Fig.S18)	(22 data sets)	CO/O ₂ /NO/NO ₂ /H ₂ O/N ₂	-
		mixtures	
Buner stabilized flame ^(Fig.8)	Species profiles	4.7 kPa, 298 K, $\phi = 1.91$,	Vandooren 95
	(7 data sets)	H ₂ /O ₂ /NH ₃ /Ar mixture	
Buner stabilized	Species profiles	4.6 kPa, 298 K, $\phi = 0.12$ -1.0,	Bian & Vandooren 96
flame ^(Fig.S19,Fig.S20)	(4 data sets)	NH ₃ /H ₂ /O ₂ /Ar mixture	
Buner stabilized	Species profiles	4.6 kPa, 298 K, $\phi = 0.12$ -1.0,	Bian & Vandooren 96
flame ^(Fig.S19,Fig.S20)	(4 data sets)	NH ₃ /H ₂ /NO/O ₂ /Ar mixture	
Buner stabilized	Species profiles	7.2 kPa, 298 K, φ = 1.46,	Vandooren et al.97
flame ^(Fig.S21)	(8 data sets)	NH ₃ /NO/Ar mixture	
Buner stabilized	Species profiles	$5.0 \text{ kPa}, 298 \text{ K}, \phi = 1.0,$	Duynslaegher et al.98
flame ^(Fig.S22)	(9 data sets)	NH ₃ /H ₂ /O ₂ /Ar mixture	
Buner stabilized	Species profiles	4.0 kPa, 300 K, $\phi = 1.08$,	Sausa et al. ⁷³
flame ^(Fig.S23)	(5 data sets)	$H_2/N_2O/Ar$ mixture	
Buner stabilized	Species profiles	$6.66 \text{ kPa}, 298 \text{ K}, \phi = 1.0-1.5,$	Dindi et al. ⁷⁴
flame ^(Fig.9,Fig.S24,Fig.S25)	(18 data sets)	CO/N ₂ O/Ar mixture	

S = in Supporting Information

Laminar flame speed: Predicted NH₃/air laminar burning velocities are shown in Figure 1 in comparison to the available experimental data. Most experimental data resulted from closed vessel experiments. The 1978 measurements by Zakaznov ⁸⁷ were performed by a cylindrical flow tube. Among the measured laminar burning velocities there is a good agreement for fuel lean and

stoichiometric conditions. For fuel rich conditions a significant discrepancy is noticed. Our model captures the experimental trends from lean to stoichiometric condition very well. For lean conditions the best agreement is found with data by Pfahl 2000⁸⁸. For fuel rich conditions the model agrees with the older measurements from Ronney 1988⁸⁹. Newer Experiments by Takizawa 2008 ⁹¹ and Hayakawa 2015 ¹⁸ follow the trend of Zakaznov ⁸⁷. Data by Jabbour 2004 ⁹⁰ are closer to the data by Rooney 1988. Recently, Nakamura et al.⁹⁹ performed an experimental and kinetic modeling study of weak ammonia-air flames in a micro flow reactor. In addition they compared laminar flame speeds of ammonia-air flames available from the literature with predictions of five different mechanisms including their own model. They ⁹⁹ showed that none of the mechanisms was able to predict the experimental laminar flame speeds on the lean and the fuel rich side at the same time. Testing the impact of the thermodynamic data for NH3, NH2 and NH from Bugler et al.¹⁰⁰ we can rule out that the problem is related to thermodynamic properties. However, we are unable to explain where these discrepancies come from. Therefore this problem remains and should be addressed by the combustion community in the future from both the experimental and the modelling side. Flow analyses shows that NH3 undergoes hydrogen abstraction by O, H and OH radicals and decomposes to the Amidogen radical (NH₂), which further decomposes to the Imidogen radical (NH). Both are very reactive species and their reactions with H, O, OH, and NO control the overall reactivity of the system. The thermal decomposition of NH₃ which forms NH₂ and H is also sensitive on the laminar flame speed. We also demonstrate the importance of the N_2H_x chemistry in Figure 1. Without this chemical mechanism we predict significant lower laminar flame speeds for all fuel-air equivalences radius.



Figure 1: Laminar flame speed for NH₃/air blends at 1 atm and 298K. Symbols: experiments from $^{18,87-91}$. Left figure: dashed line; this model prediction, solid line: Lamoureux et al. 16 model prediction. Right figure: dashed line, this model prediction; solid line, model prediction without N₂H_x scheme.

Ignition delay times: Ignition delay time data from shock tube experiments published in the literature cover pressure ranges which are relevant for practical combustion systems. The ignition delay times of NH₃/O₂ blends with high dilution of Ar were reported by Mathieu et al.¹⁷ in the pressure range of 1.4-30.0 atm for the fuel-air equivalence ratio range, ϕ =0.5-2.0. As can be seen from Figure 2 the present model captures the experimental trends in temperature for all pressures and equivalence ratios. It can also be observed from Figure 2 that with increasing pressure the ignition delay time of ammonia monotonously decreases.



Figure 2: Ignition delay times from shock tube experiments in comparison to model predictions for NH₃/O₂/Ar blends. Symbols: experiments from ¹⁷, dashed lines: model predictions.

To further validate the H/N/O chemistry, experiments investigating the sensitivity of the addition of NO₂ and N₂O to H₂/O₂/Ar blends on ignition delay times are shown in Figure 3. The comparison shows that 1600 ppm of N₂O have almost no sensitivity on the H₂ chemistry (3a with 3b), while 100 ppm of NO₂ (3a with 3b) are enough to change the pressure dependence of H₂ autoignition. We note that in simulations without N₂O the ignition is slightly delayed at high temperatures resulting in deviations from the experiment, especially for the lowest pressure. The experiments with NO₂ still show a significant influence of the chain breaking reaction H+O₂+M \Rightarrow HO₂+M on the H₂ ignition delay times, resulting in the crossing lines for the ignition delays, which are moved at higher pressure to higher temperature. In the supporting information the low sensitivity of N₂O is further demonstrated in figure S4. NO₂ can suppress this effect through direct reactions with HO₂ through reaction NO₂+HO₂ \Rightarrow HONO+O₂ resulting in a chain propagation instead of a chain breaking effect through the thermal decomposition of HONO via HONO \Rightarrow NO+OH. This reaction path only becomes relevant if high enough concentrations of HO₂ are available in the oxidizing gas mixture. The kinetic model predicts this sensitivity in agreement to the experiments. A comparable reaction path does not exist for N₂O, which can explain the week sensitivity of N₂O.



Figure 3: Ignition delay times from shock tube experiments of $H_2/O_2/N_2O/Ar$ and $H_2/O_2/NO_2/Ar$ blends. Left: (a), $H_2(0.01)/O_2(0.01)/N_2O(0.0016)$; (b), $H_2(0.01)/O_2(0.01)/NO_2(0.0001)$, symbols experiments from ^{63,77}, dashed lines: model prediction. Right: (c) and (d) symbols same as in figure (a) and (b) respectively, solid lines model prediction without N₂O and NO₂ doping.

In addition a sensitivity analysis towards ignition delay time was performed at 1160 K for two different pressure: 1.7 atm and 33 atm. The mixture composition for both pressures is the same (as mentioned in Figure 3(b)) including a small amount of NO₂. Figure 4 shows the most sensitive reactions for each pressure. It can be observed that the sensitivity varies strongly with pressure. The most important chain propagating reaction for both pressures is the reaction O_2 +H \Rightarrow OH+O

while at 33.0 atm its sensitivity is 3 times higher compared to that at 1.7 atm. The most sensitive reaction to prolong ignition delay time (negative sensitivity) is $H+O_2(+M) \Rightarrow HO_2(+M)$ at 33.0 atm. This reaction is not found to be sensitive at 1.7 atm. The most sensitive NO_x chemistry reaction is NO+HO₂ \Rightarrow NO₂+OH at 33.0 atm. In general all reactions involving NO₂ are found to have increasing sensitivities with increasing pressure, which is in agreement with the observation made in Figure 3 (b) and 3 (d). The most notable observation from this analysis is that the sensitivity direction of the reaction NO₂+H \Rightarrow NO+OH changes from negative (prolonging ignition delay time) at 1.7 atm to positive (shortening ignition delay time) at 33.0 atm. Reactions involving HO₂ are generally found to be more sensitive at high pressure which can be explained by the increasing importance of the reaction H +O₂(+M) \Rightarrow HO₂(+M) forming HO₂.



Figure 4: Ignition delay sensitivity coefficient (σ) for H₂ (0.01)/O₂ (0.01)/NO₂ (0.0001)/Ar at 1160 K, 33.0 atm and 1.7 atm.

Speciation in JSR and PFR: To further investigate the sensitivity of NO_x on the hydrogen chemistry we simulated experiments published in the literature dedicated to H₂/O₂/N₂/NO_x, H₂/CO/O₂/N₂/NO_x and CO/O₂/H₂O/N₂/NO_x blends in a JSR and PFR. In Figure 5 H₂/O₂/N₂ blends are doped with 220 ppm of NO at 10 atm for a residence time of 1.0 s. NO is consumed and NO₂ is formed in the temperature range of 750-1100 K. The conversion of NO to NO₂ is explained by the chain propagation reaction NO + HO₂ \Leftrightarrow NO₂ + OH, which is accelerating the H₂ chemistry. As can be seen the highest NO₂ concentration occurs at 800 K. At higher temperature H2 oxidation



Figure 5: Species profile comparison between measurements and model prediction for $H_2(0.01)/O_2(0.01)/NO(220 \text{ ppm})/N_2$ oxidation in a JSR at 10 atm, residence time (τ)=1.0 s. Symbols: experiments from ⁶², dashed lines: model prediction.

continues, and NO₂ is reduced to NO via reaction NO₂ + H \rightleftharpoons NO + OH. In the H₂/O₂ system doped with NO₂ (see Figure 6) in the same temperature range as in Figure 5 we do not observe any NO conversion to NO₂. In this case (Figure 6) almost all of the NO₂ is converted to NO *via* the reaction NO₂ + H \rightleftharpoons NO + OH and to a lesser extent *via* NO₂ + O \rightleftharpoons NO + O₂. It can be observed that around 1000 K where almost all the NO₂ is consumed, the peak concentration of NO is also reached. The doping of the H₂/O₂/N₂ blend with NO₂ results in a delayed oxidation of the blend. This is explained by the backward reaction NO + HO₂ \leftrightarrows NO₂ + OH, which now consumes reactive OH to form less reactive HO₂.



Figure 6: $H_2(0.01)/O_2(0.00333)/NO_2(60 \text{ ppm})/N_2$ oxidation in JSR at 10 atm, residence time, τ =1.0 s and in temperature range 700-1100 K. Symbols: experimental data from ⁶²; lines: prediction with present model.

Figure 7 shows the comparison between model predictions and experimental data of Mueller et al. ⁶¹ for H₂/O₂ oxidation in the presence of 532 ppm NO in a flow reactor at 10 atm and 802 K. We observe NO to NO₂ conversion which mainly occurs *via* the same reaction $(NO+HO_2 \rightleftharpoons NO_2+OH)$ as in the JSR shown in Figure 5.



Figure 7: Speciation of $H_2(1\%)/O_2(2\%)/NO(532 \text{ ppm})/N_2$ oxidation in flow reactor at 10 atm and 802 K. Symbols: experimental data from Mueller et al. ⁶¹; lines: prediction with present model. Simulation lines are shifted by 200 ms to match fuel consumption.

Figure 8 shows the results for the mutual oxidation of CO and NO in a flow reactor studied by Glarborg et al. ⁶⁴. The upper figure shows the oxidation of CO to CO₂ which proceeds mainly *via* CO+OH=CO₂+H while the lower figure shows the conversion of NO to NO₂. The oxidation of NO to NO₂ proceeds primarily through the reaction NO+HO₂=NO₂+OH with minor contributions from the reactions NO+O(+M)=NO₂(+M), NO+OH(+M)=HONO(+M) and HONO+OH=NO₂+H₂O. In this moist CO/NO oxidation system HO₂ is formed entirely by recombination of H atoms with O₂, which is the rate limiting step in the NO to NO₂ conversion. The complex effect of NO on the CO oxidation rate can be understood in terms of the competition between NO+HO₂=NO₂+OH which promotes the oxidation process and the reactions NO+O

 $(+M) \rightleftharpoons NO_2$ and NO+OH $(+M) \rightleftharpoons HONO (+M)$ which leads to recombination of the main chain carriers. The model agrees very well with the experimental results. Both CO/CO₂ and NO/NO₂ conversion are predicted well over the range of conditions investigated (see Figure S16 – Figure S18).



Figure 8: Speciation of CO(530 ppm)/NO(522 ppm)/O₂(4.2%)/H₂O(5.2 %)/N₂ oxidation in flow reactor at 1.05 atm, residence time, $\tau = 202$ [K]/T. Symbols: experimental data from Glarborg et al. ⁶⁴; lines: prediction with present model.

Speciation in a burner stabilized flame: Speciation predictions for premixed burner stabilized flames receive less attention in the literature than speciation predictions in reactors. However, burner stabilized flames provide important information about species formation and consumption in the reaction zone and subsequently about emission formation pathways. They are the major experimental setup used to study the formation of polycyclic aromatic hydrocarbons (PAHs) and

NO_x from carbon based fuels. For all premixed flames the diffusion of small radicals is the determining process for the flame structure. The concentration of the radical pool species O, H and OH during the oxidation of the fuels are the most important species controlling the reactivity of the system. Therefore we give in this study special attention to the speciation predictions for burner stabilized flames.

Figure 9 shows the measured species profiles in a low pressure, fuel rich (ϕ =1.91) H₂/O₂/NH₃/Ar premixed flame studied by Vandooren ⁹⁵ in comparison to model predictions. Calculations were performed, using the temperature profile from ⁹⁵ (dash lines), and by solving the energy conservation equation (lines). It can be seen that the overall model prediction is in good agreement with the measurements. The proposed kinetic scheme well predicts the intermediates species and the radical pool in the flame. The calculated temperature profile results in a better agreement of the predicted species concentrations. This is also seen for close to equilibrium conditions at larger heights above the burner. As the equilibrium conditions are independent on the kinetic mechanism this indicates that the calculated temperature is accurate. Further simulations of burner stabilized flames are available in the supporting information (Figure S19 – Figure S25).



Figure 9: Speciation comparison between experimental data and model predictions for a fuel rich $H_2/O_2/NH_3/Ar$ (ϕ =1.91) premixed burner stabilized flame at 4.7 kPa. Symbols: experimental data from ⁹⁵. Dashed lines: model predictions imposing the experimental temperature profile, solid line: model predicted temperature profile.

Figure 10 compares the experimental species profiles of a low pressure (6.66 kPa), stoichiometric $(\phi = 1.0)$ CO/N₂O flame studied by Dindi et al. ⁷⁴ against the model predictions from this work. The simulations were performed imposing the experimental temperature profile as provided by ⁷⁴ in their study. As we can see in Figure 10 the species consumption and formation profile is well predicted by the model. It can be noted that the model is also capable to address the kinetics

between NO_x and CO species. Further comparison between experimental data and model predictions for fuel rich conditions of the CO/N₂O flame can be seen in Figure S24 and Figure S25.



Figure 10: Speciation comparison between experimental data and model predictions for a ϕ =1.0 premixed burner stabilized CO (0.5)/N₂O (0.5) flame at 6.66 kPa. Symbols: experimental data from ⁷⁴. Dashed lines: model predictions imposing the experimental temperature profile.

Reaction path analysis in a burner stabilized flame

Figure 11 shows the reaction pathway analysis based on the nitrogen atom for the laminar premixed burner stabilized flames in Figures 9 and 10. In analyzing the H₂(35.4%)/O₂/NH₃(2.9%)/Ar rich flame studied by the Vandooren ⁹⁵ (see Figure 9) with imposing the experimental temperature profile we observe that almost all of the NH₃ is decomposed to form the amidogen (NH₂) radical reacting with H, OH and O radicals. Among these three radicals, the

H atom reacting with NH₃ is the main reaction forming NH₂ while least is contributed by O atoms. Furthermore, the NH₂ radical reacts with H to form the imidogen (NH) radical. The NH radical consumes almost 67 % of NH₂ and almost 27 % of NH₂ is consumed via the reaction NH₂+O \rightleftharpoons HNO+H forming the nitroxyl (HNO) radical. Imidogen radicals are further decomposed forming N atoms via the reaction NH+H \rightleftharpoons N+H₂ and 25 % of these N radicals react with OH and O₂ to form NO while the remaining 75% of N atoms contribute for the formation of N₂ by reacting with NO. The HNO radical, which was initially produced, directly contributes to NO formation by reacting with H atoms *via* the reaction HNO+H \rightleftharpoons NO+H₂.



Figure 11: Flow analysis (a) NH₃ decomposition and the NO formation pathway in the fuel rich flame (ϕ =1.91) shown in Figure 9; (b) N₂O consumption and NO formation in the CO/N₂O flame shown in Figure 10. The numbers in the flow diagram indicate the percentage flow based on the N atom.

For the stoichiometric CO/N₂O flame as shown in Figure 10 the most important reactions are $CO+N_2O \rightleftharpoons CO_2+N_2$, N_2O (+M) $\rightleftharpoons N_2+O$, $N_2O+O \rightleftharpoons N_2+O_2$ and $N_2O+O \oiint NO+NO$. Almost all the CO is consumed *via* the reaction CO+N₂O \rightleftharpoons CO₂+N₂. Most of the N₂O (94%) is consumed to form

N₂ via the first three reactions among which the first reaction $(CO+N_2O \Rightarrow CO_2+N_2)$ contributes most while N₂O+O \Rightarrow NO+NO contributes least. The last two reactions $(N_2O+O \Rightarrow N_2+O_2 \text{ and } N_2O+O \Rightarrow NO+NO)$ control the formation of O₂ and NO.

3.2 Brief discussion of C₁/NO_x model

Figure 12 follows up the discussion in our recent publication Goos et al. ²⁴. The figure shows that the current model agrees with the accurate predictions of our former publication ²⁴. Changes in the base chemistry, and the addition of species in the H/N/O system do not change former conclusions. The thermodata for NCN are highly sensitive on the predicted NO emissions, and we recommend the use of $\Delta_f H_{298}^0 = 457.7 \text{ kJ/mol}^{24}$ for NCN.



Figure 12: Comparison between experimental NO profiles and model predictions in a CH₄/O₂/N₂ premixed flame at 5.3 kPa and different equivalence ratios. Symbols: experiments from Lamoureux et al.¹⁶; dashed lines: this model with NCN heat of formation, $\Delta_f H_{298}^0 = 457.7$ kJ/mol from Goos et al.²⁴; solid lines: this model with NCN heat of formation $\Delta_f H_{298}^0 = 444.1$ kJ/mol suggested by Lamoureux et al.¹⁶.

4. Effect of Nitrogenated Species Thermochemistry on model predictions

Very recently, Glarborg et al.¹⁰¹ investigated the nitrogen chemistry and reevaluated the thermodynamics properties of nitrogenated species and some hydrocarbon radicals using the Active Thermochemical Tables (ATcT) approach ^{102,103}. Klippenstein et al. ¹⁰⁴ studied the prompt NO formation in burner stabilized methane premixed flame and used thermodata of Glarborg et al. ¹⁰¹. To study the effect of nitrogenated species thermochemistry on our model we replaced the complete nitrogenated species thermochemistry with the data from Glarborg et al.¹⁰¹ and performed simulations for some of the key experiments used for the model development.



Figure 13: Laminar flame speed of NH₃/air blends at 1 atm and 298 K, (a) (experiments same as in figure 1); Ignition delay time of NH₃/O₂/Ar at $\phi = 1.0$ in shock tube, (b) (experiments same as in figure 2); Speciation in jet stirred reactor for H₂/O₂/NO/N₂, (c) (experiments same as in figure

5); Speciation in CH₄/O₂/N₂ premixed burner stabilized flame, (d) (experiments same as in figure 12). Dashed lines: predictions of this model, solid lines: model prediction using the nitrogenated species thermochemistry from Glarborg et al.¹⁰¹.

Only a small difference in predicted laminar flame speeds at 1 atm and 298 K (Figure 13(a)) on the lean side was observed. The impact on predicted ignition delay times is negligible (Figure 13 (b)). In Figure 13 (c) using the thermochemistry suggested by Glaborg et al. decreases the reactivity at low temperatures and increases the onset temperature of H₂ consumption by 10 K. We found that this shift in reactivity is particularly due to the 1.9 kJ/mol lower heat of formation at 298 K of the HONO molecule.

The only significant impact we notice using the complete nitrogenated species thermochemistry from Glarborg et al. ¹⁰¹ is for prompt NO prediction in a rich, low-pressure CH₄/O₂/N₂ premixed burner stabilized flame (Figure 13 (d)). This impact on NO prediction was expected due the different NCN thermochemistry with $\Delta_f H_{298}^0 = 450.80 \text{ kJ/mol}$ used by Glarborg et al.¹⁰¹ compared to one adopted in our model ($\Delta_f H_{298}^0 = 457.7 \text{ kJ/mol}$). The sensitivity of NCN thermochemistry on prompt NO prediction has been discussed by many authors ^{16,24,100,105–107} and

is also disscused by Glarborg et al.¹⁰¹.

5. Conclusions

Ammonia has gained growing attention as an alternative fuel or fuel compound, which can be produced from alternative energy sources. As the fuel is carbon free it has zero-CO₂ emissions and

is therefore an alternative to hydrogen as fuel. The advantage of NH₃ is its long term storage capability.

In this paper we present a reaction mechanism for the oxidation of ammonia, considering reaction pathways to NO formation and NO reduction. We selected a number of experiments from the literature that demonstrate important features of the NH₃/NO/H₂/CO chemistry. It is possible to cover all these features with the here derived, broadly validated reaction mechanism. Special emphasize is made on gas conditions, which allow the NO_x kinetics to either accelerate or decelerate the oxidation of H₂. It is finally demonstrated that the mechanism can be combined with hydrocarbon kinetics to predict NO formation in hydrocarbon flames.

Supporting Information

Kinetic mechanism, thermochemistry and transport properties of the species used in the mechanism, brief description and convergence criteria of the numerical model used for simulation, additional mechanism validation plots for mixtures listed in Table 2 and rate constant comparison plots for H₂/CO mechanism.

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