Fuel 181 (2016) 358-365

Contents lists available at ScienceDirect

Fuel

journal homepage: www.elsevier.com/locate/fuel



Full Length Article

Ammonia oxidation at high pressure and intermediate temperatures

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ARTICLE INFO

Article history: Received 11 January 2016 Received in revised form 9 March 2016 Accepted 21 April 2016 Available online 10 May 2016

Keywords: NH₃ oxidation High pressure Flow reactor H₂NO + O₂ rate constant Kinetic model

ABSTRACT

Ammonia oxidation experiments were conducted at high pressure (30 bar and 100 bar) under oxidizing and stoichiometric conditions, respectively, and temperatures ranging from 450 to 925 K. The oxidation of ammonia was slow under stoichiometric conditions in the temperature range investigated. Under oxidizing conditions the onset temperature for reaction was 850–875 K at 30 bar, while at 100 bar it was about 800 K, with complete consumption of NH₃ at 875 K. The products of reaction were N₂ and N₂0, while NO and NO₂ concentrations were below the detection limit even under oxidizing conditions. The data were interpreted in terms of a detailed chemical kinetic model. The rate constant for the reaction of the important intermediate H₂NO with O₂ was determined from ab initio calculations to be $2.3 \times 10^2 T^{2.994} \exp(-9510 \text{ K/T}) \text{ cm}^3 \text{ mol}^{-1} \text{ s}^{-1}$. The agreement between experimental results and model work was satisfactory. The main oxidation path for NH₃ at high pressure under oxidizing conditions is NH₃ $\stackrel{+\text{OH}}{\longrightarrow}$ NH₂ $\stackrel{+\text{HO}_2.NO_2}{\longrightarrow}$ H₂NO $\stackrel{+\text{O}_2}{\longrightarrow}$ NO $\stackrel{+\text{O}_2}{\longrightarrow}$ N₂. The modeling predictions are most sensitive to the reactions NH₂ + NO = NNH + OH and NH₂ + HO₂ = H₂NO + OH, which promote the ammonia consumption by forming OH radicals, and to NH₂ + NO = N₂ + H₂O and NH₂ + NO₂ = N₂O + H₂O, which are the main chain-terminating steps.

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1. Introduction

Ammonia is an important combustion intermediate in formation of nitric oxide from nitrogen organically bound in fuels. For most solid fuels, NH₃ is formed directly during devolatilization [1], and it has also been reported as a product of gasification [2]. The selectivity for forming NO or N₂ from N-volatiles in combustion depends largely on the fate of amine radicals such as NH₂, NH, and N [1,3]. Ammonia is also known as an efficient additive for Selective Non-Catalytic Reduction of NO (SNCR) [3,4]. Recently, ammonia has attracted interest as a potential carbon-free energy carrier [5]. Results on oxidation of NH₃ are available from flames [6–16], shock tubes [17–27], and flow reactors [2,28–30]. In addition, a number of modeling studies and reviews of ammonia chemistry have been reported [3,27,28,31–35]. Supplementing the oxidation studies, extensive work has been conducted on the kinetics of the SNCR process [3,4,36–38]. With the interest in amine-based fuels as energy carriers [39], reliable experimental data for oxidation of ammonia at high pressure become important. High-temperature results at increased pressure are available from recent work in shock tubes and premixed flames. Mathieu and Petersen [27] measured the ignition time for ammonia diluted in Ar at 1.4, 11, and 30 bar, respectively, while Hayakawa et al. [15] determined the burning velocity of ammonia/air mixtures at pressures of 1–5 bar. However, studies of ammonia oxidation at low-to-medium temperatures and high pressure are scarce.

The purpose of the present study is to investigate ammonia oxidation at high pressure (30–100 bar) and temperatures up to 925 K. Experiments are conducted with stoichiometric and lean NH₃/O₂ mixtures, highly diluted in N₂, in a laminar flow reactor. The results are interpreted in terms of a detailed chemical kinetic model for ammonia oxidation. The reaction mechanism, which is based on earlier work on nitrogen chemistry [34,38,40,41], is updated in the present work, emphasizing reactions important at high pressure. Under the present conditions, the nitroxide radical H₂NO is an important intermediate, and the rate constant for the



reaction between H_2NO and O_2 has been derived from ab initio calculations.

Table 1

Selected reactions from the NH₃ subset. Parameters for use in the modified Arrhenius expression $k = AT^{\beta} \exp(-E/[RT])$. Units are mol, cm, s, cal.

Α β Ε	Source							
$1 NH_2 + H(+M) \rightleftharpoons NH_3(+M) \qquad 1.6E14 \qquad 0.000$	0 [43]							
Low pressure limit: 3.6E22 –1.760	0							
Troe parameters: 0.5 1.0E–30 1.0E30								
$2 NH_3 + H \rightleftharpoons NH_2 + H_2 \qquad \qquad 6.4E05 \qquad 2.390 10.17$	71 [44]							
$3 NH_3 + 0 \rightleftharpoons NH_2 + OH \qquad \qquad 9.4E06 \qquad 1.940 \qquad 640$	60 [45]							
$4 \text{NH}_3 + \text{OH} \rightleftharpoons \text{NH}_2 + \text{H}_2\text{O} \qquad \qquad 2.0\text{E06} \qquad 2.040 \qquad 5000 \qquad \qquad 50000 \qquad \qquad 500000 \qquad \qquad 50000 \qquad \qquad 500000 \qquad \qquad 500000 \qquad \qquad 500000 \qquad \qquad 500000 \qquad \qquad 5000000 \qquad \qquad 500000000$	56 [46]							
5 $NH_3 + HO_2 \implies NH_2 + H_2O_2$ 3.0E11 0.000 22,00	00 [3] est							
$ 6 \mathrm{NH}_2 + \mathrm{HO}_2 \ \ \approx \ \mathrm{NH}_3 + \mathrm{O}_2 $	27 [47] ^a							
$7 NH_2 + H \rightleftharpoons NH_2 + H_2 \qquad \qquad 7.2E05 \qquad 2.320 \qquad 75$	99 [48]							
$8 NH_2 + 0 \rightleftharpoons HNO + H \qquad \qquad 6.6E13 \qquad 0.000$	0 See							
	[41]							
$9 NH_2 + 0 \rightleftharpoons NH + OH \qquad \qquad 7.0E12 \qquad 0.000$	0 See							
	[41]							
8.6E-1 4.010 16	73							
$10 NH_2 + OH \implies NH + H_2O \qquad \qquad 3.3E06 \qquad 1.949 \qquad -2^{-3}$	17 [38,49]							
$11 \text{NH2} + \text{HO}_2 \rightleftharpoons \text{H}_2\text{NO} + \text{OH} \qquad 5.0\text{E13} \qquad 0.000$	0 See							
	text							
$12 NH_2 + HO_2 \ \rightleftharpoons \ HNO + H_2O \qquad \qquad 1.6E07 \qquad 0.550 \qquad 52$	25 [47] ^{a,b}							
5.7E15 –1.120 70	07							
$13 \text{NH}_2 + \text{HO}_2 \rightleftharpoons \text{HON} + \text{H}_2\text{O} \qquad 2.1\text{E07} \qquad 0.640 \qquad 8^{-3}$	11 [47] ^a							
$14 NH_2 + O_2 \implies H_2NO + O \qquad \qquad 2.6E11 \qquad 0.487 29,09$	50 [38]							
$15 NH_2 + O_2 \implies HNO + OH \qquad \qquad 2.9E-2 \qquad 3.764 \qquad 18,18$	35 [38]							
$16 \text{NH}_2 + \text{NH}_2 \ \rightleftharpoons \ \text{NH}_3 + \text{NH} \qquad 5.6\text{E00} \qquad 3.530 \qquad 5\text{H}_2 + 1000 \ 5\text{H}_2 + 10000 \ 5\text{H}_2 + 10000 \ 5\text{H}_2 + 10000 \ 5\text{H}_2 + 10000 \ 5$	52 [49]							
$17 \text{NH}_2 + \text{HNO} \ \rightleftharpoons \ \text{NH}_3 + \text{NO} \qquad \qquad 3.6\text{E06} \qquad 1.630 -123$	50 [50]							
$18 NH_2 + NO \rightleftharpoons N_2 + H_2O \qquad \qquad 1.3E16 -1.25$	0 [38]							
-3.1E13 -0.48 118	30							
$19 NH_2 + NO \Rightarrow NNH + OH \qquad \qquad 3.1E13 -0.48 \qquad 118$	30 [38]							
$20 NH_2 + NO_2 \ \rightleftharpoons \ N_2O + H_2O \qquad \qquad 2.6E18 -2.191 \qquad 44$	55 [51]							
$21 NH_2 + NO_2 \rightleftharpoons H_2NO + NO \qquad 9.1E11 \qquad 0.032 -15$	12 [51]							
$22 \text{NH}_2 + \text{HONO} \rightleftharpoons \text{NH}_3 + \text{NO}_2 \qquad 7.1\text{E01} \qquad 3.020 -494$	40 [50]							
$23 H_2 NO + M \rightleftharpoons HNO + H + M \qquad 2.8E24 -2.830 64.9^{\circ}$	15 [33]							
$24 H_2 \text{NO} + \text{H} \rightleftharpoons \text{HNO} + \text{H}_2 \qquad \qquad 3.0\text{E07} \qquad 2.000 \qquad 200$	00 [36] est							
$25 H_2 NO + H \rightleftharpoons NH_2 + OH \qquad \qquad 5.0E13 \qquad 0.000$	0 [36] est							
$26 H_2 NO + O \rightleftharpoons HNO + OH \qquad \qquad 3.0E07 \qquad 2.000 \qquad 200$	00 [36] est							
$27 H_2 NO + OH \equiv HNO + H_2 O \qquad 1.0E14 \qquad 0.000$	0 [52]							
$28 H_2 NO + HO_2 \ \rightleftharpoons \ HNO + H_2O_2 \qquad 2.9E04 \qquad 2.690 -160$	00 [33]							
$29 H_2 NO + O_2 \ \rightleftharpoons \ HNO + HO_2 \qquad 2.3E02 \qquad 2.994 18,992 1$	00 pw							
$30 H_2NO + NH_2 \rightleftharpoons HNO + NH_3 \qquad 3.0E12 \qquad 0.000 \qquad 100$	00 [36] est							
$31 H_2NO + NO_2 \rightleftharpoons HONO + HNO \qquad 6.0E11 \qquad 0.000 \qquad 200$	00 [53] est							
$32 HNO + O_2 \implies NO + HO_2 \qquad 2.0E13 \qquad 0.000 16,000 $	00 [33]							
33 NNH \Rightarrow N ₂ + H 1.0E09 0.000	0 [38]							
$34 NNH + O_2 \ \rightleftharpoons \ N_2 + HO_2 \qquad \qquad 5.6E14 -0.385 -$	13 [38]							

^a Rate constant calculated for 1 atm.

^b Duplicate reaction – the resulting rate constant is the sum of the two expressions.

3.1. Ab initio calculations

The nitroxide radical H₂NO, molecular oxygen and the transition state for their reaction were investigated computationally. First, geometries and frequencies (scaled by a standard factor of 0.954 [54]) were obtained with UQCISD/6-311G(d,p) theory, using spin-unrestricted wavefunctions as implemented within the Gaussian 09 code [55]. At these geometries (see Fig. 2), single-point energies were calculated at the UCCSD(T) level of theory with the aug-cc-pVTZ and aug-cc-pVQZ basis sets, using spin-restricted wavefunctions within the Molpro 2010 program [56], and extrapolated to the complete basis set limit for coupled cluster theory (CCSD(T)/CBS). Corrections were added to this result based on UCCSD(T)/cc-pwVTZ results obtained with core electrons included and excluded from the correlation treatment, and relativistic effects evaluated at the CISD/cc-pwVTZ level of theory. The results are summarized in the supplementary material. The rate constant was then derived via canonical transition state theory as implemented within the Multiwell program suite [57], with tunneling accounted for via the Eckart model.

Some of the prior studies of H_2NO raised the issue of whether this molecule is planar or pyramidal [58,59]. Out-of-plane bending

2. Experimental

The experimental setup was a laboratory-scale high-pressure laminar flow reactor designed to approximate plug flow. The setup is described in detail elsewhere [42] and only a brief description is provided here. The system was used here for investigation of ammonia oxidation at 30 bar and 100 bar pressure, respectively, and temperatures from 450 to 925 K. The reactant gases were premixed before entering the reactor. The reactions took place in a tube under laminar flow conditions. The tube was made of guartz (inner diameter of 7.5 mm) or alumina (Degussit AL23, inner diameter 6 mm). The temperature profile in the flow reactor was measured by a thermocouple positioned in the void between the quartz reactor and the steel shell. Results for 100 bar are shown in Fig. 1 while the 30 bar profiles are available as supplementary material. An isothermal reaction zone of 39-47 cm was achieved in the reactor. All gases used in the present experiments were high purity gases or mixtures with certified concentrations. The total flow rate was $2.8 \,\mathrm{L\,min^{-1}}$ (STP). The product analysis was conducted with an on-line 6890N Agilent Gas Chromatograph (GC-TCD/FID from Agilent Technologies) and an AO2020 NH₃/NO/NO₂ analyser from ABB. The relative measurement uncertainties for the species detected were in the range $\pm 2-6\%$.

3. Chemical kinetic model

The starting mechanism and corresponding thermodynamic properties were drawn from the recent work by Klippenstein et al. [38]. The mechanism was carefully updated, emphasizing reactions of importance under the conditions of the present study. Table 1 lists the key reactions in the NH₃ oxidation scheme with the rate coefficients used in the present work. The full mechanism is available as supplemental material.

At the high-pressure medium-temperature conditions of this work, ammonia oxidation occurs to a significant extent through the H_2NO intermediate. Rate constants for H_2NO reactions are generally quite uncertain. In the present work, we characterize the key reaction of H_2NO with O_2 by ab initio calculations, as described below.



Fig. 1. Measured temperature profiles along the reactor axis for 100 bar conditions.



Fig. 2. UQCISD/6-311G(d,p) geometries of ²A' H₂NO, the planar ²B₁ barrier to inversion, and the transition state for abstraction of H by O₂. Distances are in Å and angles in degrees. Dihedral angles in the TS: HNHO –159.2°, OHNH –49.2°, OOHN –24.2°.

of the C_{2V} structure was explored via scans of the angle between the O–N bond and the NH₂ plane, with the rest of the geometry allowed to optimize. The results are plotted in Fig. 3 that shows a double well potential with a very small barrier of 17 cm⁻¹ to inversion. The harmonic frequency evaluated at either minimum is 481 cm⁻¹, but clearly the potential is significantly anharmonic.



Fig. 3. The double-well potential of H_2NO computed at the UQCISD/6-311G(d,p) level of theory. The barrier height and the first three eigenvalues for out-of-plane bending are relative to the minima. The angle is defined as the deviation from 180° (the planar conformation) for the angle O–N–X where X is a point midway between the two H atoms.

The moment of inertia for bending was estimated to be 0.407 amu Å², by treating the planar geometry like a three-atom linear molecule [60] with a mass of 2 amu at the midpoint of the two H atoms. The eigenvalues obtained via the FGH method [61] for quantized bending motion on this potential are summarized in the supplemental material; the first three are 142, 508 and 1030 cm⁻¹ above the minima. Even the lowest level is well above the inversion barrier so H₂NO explores both sides equally. Because the wavefunction for the ground state is maximized at the planar geometry, we use this structure and a symmetry number of 2 to calculate rotational partition functions. Vibrational partitions were based on the harmonic oscillator model for the remaining 5 modes of H₂NO combined with results derived via a direct count for the out-of-plane bending energy levels. Together with the heat of formation at 298 K from the Active Thermochemical Tables [62], the thermochemistry is summarized within the NASA polynomial included in the supplemental information.

3.2. Reaction mechanism

The reactions of ammonia with the radical pool and O₂ involve hydrogen abstraction to form NH₂. The rate constants for these steps, taken from the mechanism of Klippenstein et al. [38], are mostly well established, but the values for NH₃ + HO₂ (R4) (estimated [3]) and NH₃ + O₂ (R5b) (ab initio calculation [47]) are more uncertain.

The oxidation rate for ammonia and the products of reaction are largely determined by the fate of NH₂. The NH₂ radical is mainly consumed by reaction with the HO₂ radical (R5b, R11–R13), which builds up in considerable concentrations under the present conditions, and with the stable species NO (R18, R19) and NO₂ (R20,

R21). The $NH_2 + HO_2$ reaction has been studied theoretically by several groups [33,47,63–65], while there are only indirect room temperature measurements available for the rate constant [66– 68]. It has a number of product channels and may occur both on the singlet and triplet surfaces. We choose to rely on the work of Sumathi and Peyerimhoff [47] who calculated rate constants for formation of the adduct H₂NOOH and its isomers HN(OH) OH and $H_2N(OH)$ O, as well as the products $NH_3 + O_2$, $NH_2O + OH$, HNO + H_2O , and HON + H_2O . The predominant contribution to the total rate constant is predicted to be dissociation of energized H₂-NOOH to NH₂O + OH over the temperature range 300–2000 K and pressures from 0.001 to 10 atm. At pressures above 10 atm, stabilisation of H₂NOOH becomes important according to Sumathi and Peverimhoff. However, little is known about the fate of the H₂-NOOH adduct. In the present work, we take $H_2NO + OH$ (R11) to be the primary products of reaction and assume the rate constant to be equal to the high-pressure limit of the overall reaction, which we estimate to be $5 \times 10^{13} \text{ cm}^3 \text{ mol}^{-1} \text{ s}^{-1}$. This value is only slightly higher than the indirect measurement for $NH_2 + HO_2$ by Lozovskii et al. [67] at 570 torr, but a factor of two higher than the calculation of Sumathi and Peyerimhoff.

The rate constants for the reactions of NH₂ with NO and NO₂,

$$\mathrm{NH}_2 + \mathrm{NO} \rightleftharpoons \mathrm{N}_2 + \mathrm{H}_2\mathrm{O} \tag{R18}$$

$$NH_2 + NO \Rightarrow NNH + OH$$
 (R19)

 $NH_2 + NO_2 \rightleftharpoons N_2O + H_2O \tag{R20}$

$$NH_2 + NO_2 \rightleftharpoons H_2NO + NO$$
 (R21)

have been discussed in detail in recent work by Klippenstein and coworkers [38,51]. We have adopted their values without modifications. For NH₂ + NO₂, formation of the adducts H₂NNO₂ and H₂-NONO is pressure dependent, but Klippenstein et al. predict little effect of pressure below 100 atm because the complex lifetimes are too short to permit significant collisional relaxation [51].

Nitroxide may be formed in significant quantities from reaction of NH₂ with HO₂ and NO₂. The H₂NO subset of the reaction mechanism consists largely of reactions with estimated rate constants. While the most important reaction of H₂NO is that with O₂, discussed above, also the reactions with HO₂ (R28) and NH₂ (R30) are important consumption steps. The value of k₂₈ is a rough estimate [36] and k₃₀ was derived from QRRK calculations [33]. Similarly to H₂NO, HNO is mostly consumed by reaction with O₂ (R32); also for this step we use a rate constant based on QRRK theory [33].

4. Results and discussion

Experiments for NH_3 oxidation with high dilution in N_2 as a function of temperature from 450 K to 900 K were conducted under stoichiometric and oxidizing conditions. Table 2 lists the experimental conditions. The fuel-air equivalence ratio, defined

from the overall reaction $NH_3 + 1.25O_2 \rightarrow NO + 1.5H_2O$, ranged from approximately 1.0 to 0.02. Simulations with the full temperature profile were conducted using the CHEMKIN PRO software package [69]. Calculations restricted to the isothermal zone of the reactor showed similar trends, but with a smaller conversion at the highest temperatures (oxidizing conditions). The experimental results are compared with modeling predictions in Figs. 4–7. Symbols denote the experimental data and lines denote numerical results.

Figs. 4 and 5 show results for NH_3 oxidation under stoichiometric conditions and pressures of 30 and 100 bar, respectively. At 30 bar the onset of reaction occurs at 800 K. However, even above this temperature consumption of NH_3 is very slow; at 925 K 730 ppm NH_3 (95% of the inlet concentration) and about 900 ppm O_2 (97% of the inlet concentration) are still unreacted in the reactor outlet. At 100 bar, NH_3 consumption also starts around 800 K, but as the temperature increases oxidation is slightly faster than at 30 bar. This can partly be attributed to a longer residence time. However, at 900 K still 675 ppm NH_3 (i.e., 95% of the inlet) remains unreacted.

The modeling results for stoichiometric conditions at 30 bar and 100 bar show very little reaction. The agreement with experiment can be considered satisfactory, since the measurements show that only around 40 ppm or about 5% of NH_3 are consumed at the highest temperatures at the two pressures.

Fig. 6 shows the results for NH_3 oxidation at 30 bar for fuel-lean conditions. To investigate the impact of surface effects, these experiments were conducted in both an alumina and a quartz tube. Ammonia is known to decompose on quartz surfaces [70–72], and while surface effects are unimportant in SNCR for experiments carried out at low surface-to-volume ratios [36,73], it has been observed that induction times for oxidation of NH_3 in quartz reactors were influenced by heterogeneous effects [28].

In the figure, the closed symbols indicate experimental results obtained in the quartz tube, while the open symbols denote results from the alumina tube. In the quartz tube, the onset temperature for reaction occurs at 850 K, compared to a value of 875 K observed in the alumina tube. The difference can partly be attributed to the longer residence time in the quartz reactor, but it cannot be ruled out that some surface initiation takes place in the quartz reactor, despite the high pressure.

Upon initiation, NH_3 is oxidized to N_2 (not quantified) and N_2O ; the concentrations of NO and NO_2 in the product gas were below detection limit. Nitrous oxide is formed in significant amounts, reaching 50 ppm in the alumina reactor and levels above 100 ppm in the quartz reactor at 900 K.

At 100 bar (Fig. 7), reaction is initiated already at 800–825 K, partly due to the longer residence time, and NH_3 is fully oxidized at 875–900 K. The N_2O concentration approaches 200 ppm, corresponding to a selectivity for forming N_2O from NH_3 of about 50%. The N_2O profiles from the quartz and alumina reactors indicate a difference in onset temperature, but the peak concentrations are similar.

Table 2						
Experimental	conditions	for	the	NH ₃	oxidation	study

Experiment	Inlet composition ^a	Pressure (bar)	Temperature (K)	Residence time ^b (s)	Reactor
1	772 ppm NH ₃ , 929 ppm O ₂ (ϕ = 1.04)	30	450-925	3100/T[K]	Quartz
2	714 ppm NH ₃ , 864 ppm O ₂ (ϕ = 1.03)	100	450-925	10,330/T[K]	Quartz
3	729 ppm NH ₃ , 3.95%(v) O ₂ (φ = 0.23)	30	450-925	3100/T[K]	Quartz
4	719 ppm NH ₃ , 4.03%(v) O ₂ (φ = 0.22)	30	450-925	1984/T[K]	Alumina
5	789 ppm NH ₃ , 4.07%(v) O ₂ (φ = 0.24)	100	450-925	10,330/T[K]	Quartz
6	789 ppm NH ₃ , 4.07%(v) O_2 (ϕ = 0.24)	100	450-925	6610/T[K]	Alumina

^a Volume basis; balance N₂.

^b The nominal residence time in the isothermal region of the reactor. It is a function of temperature, since the mass flow rate was held constant.



Fig. 4. Comparison of experimental and predicted concentration profiles as a function of the reactor temperature for the stoichiometric experiment at 30 bar in the quartz reactor. The symbols mark experimental data while solid lines denote model predictions obtained at isothermal conditions. Inlet composition: 772 ppm NH₃, 929 ppm O₂; balance N₂. The fuel–air equivalence ratio is $\phi = 1.04$ and the residence time in the isothermal zone is calculated from τ [s] = 3100/T[K].



Fig. 5. Comparison of experimental and predicted concentration profiles as a function of the reactor temperature for the stoichiometric experiment at 100 bar in the quartz reactor. The symbols mark experimental data while solid lines denote model predictions obtained at isothermal conditions. Inlet composition: 714 ppm NH₃, 864 ppm O₂; balance N₂. The fuel–air equivalence ratio is ϕ = 1.03 and the residence time in the isothermal zone is calculated from τ [s] = 10,330/T[K].

The model satisfactorily predicts the experimental data. The best agreement is obtained for the alumina reactor data, which are possibly more reliable. At 30 bar, the agreement is very good, while at 100 bar the temperature for onset of reaction is slightly overpredicted. The most important difference is that the concentration of N_2O is underpredicted at the highest pressure. In line with observations, the model predicts that any NO or NO_2 formed are rapidly consumed.

Fig. 8 shows the results of a rate of production analysis under oxidizing conditions. The arrow thickness distinguishes the main and minor routes for ammonia consumption. Results are shown for 100 bar conditions, but the reaction paths are similar for 30 bar. The main oxidation for ammonia is $NH_3 \xrightarrow{+OH} NH_2 - HO_2 MO_2 H_2NO \xrightarrow{+O_2} HNO \xrightarrow{+O_2} NO \xrightarrow{+NH_2} N_2$. Ammonia is mainly consumed through the reaction $NH_3 + OH = NH_2 + H_2O$ (R4). The



Fig. 6. Comparison of experimental and predicted concentration profiles as a function of the reactor temperature for the oxidizing experiment at 30 bar, obtained in a quartz and alumina tube, respectively. The symbols mark experimental data while solid lines denote model predictions obtained at isothermal conditions. Inlet composition: 729 ppm NH₃, 3.95%(v) O₂; balance N₂. The fuel-air equivalence ratio is $\phi = 0.023$ and the residence time in the isothermal zone is calculated from $\tau[s] = 3100/T[K]$ for the quartz tube and $\tau[s] = 1984/T[K]$ for the alumina tube.



Fig. 7. Comparison of experimental and predicted concentration profiles as a function of the reactor temperature for the oxidizing experiment at 100 bar, obtained in a quartz and alumina tube, respectively. The symbols mark experimental data while solid lines denote model predictions obtained at isothermal conditions. Inlet composition: 789 ppm NH₃, 4.07%(v) O₂; balance N₂. The fuel–air equivalence ratio is $\phi = 0.024$ and the residence time in the isothermal zone is calculated from τ [s] = 10,330/T[K] for the quartz tube and τ [s] = 6610/T[K] for the alumina tube.

peroxide radical HO_2 is the main chain carrier under the present conditions with high pressure and low to intermediate temperatures. However, HO_2 is partly converted to OH through the fast steps,

$$NO + HO_2 \rightleftharpoons NO_2 + OH$$

 $NH_2 + HO_2 \rightleftharpoons H_2NO + OH$ (R11)

In addition to (R11), the NH_2 radical reacts mainly with NO_2 ,

$$NH_2 + NO_2 \implies N_2O + H_2O \tag{R20}$$

$$NH_2 + NO_2 \rightleftharpoons H_2NO + NO \tag{R21}$$



Fig. 8. Main reaction pathways for NH_3 oxidation at the investigated conditions. The diagram is based on the 100 bar oxidizing condition at a temperature of 875 K, but reaction paths are similar at 30 bar oxidizing conditions.

Most of the NH_2 is converted to H_2NO (R11, R21). Nitroxide then reacts with O_2 (R29), HO_2 (R28), and NO_2 (R31); these steps all abstract a hydrogen atom from H_2NO to form HNO, which is converted to NO by reaction with O_2 (R32).

$$H_2NO + O_2 \rightleftharpoons HNO + HO_2 \tag{R29}$$

 $H_2NO + HO_2 \rightleftharpoons HNO + H_2O_2 \tag{R28}$

 $H_2NO + NO_2 \rightleftharpoons HNO + HONO \tag{R31}$

 $HNO + O_2 \rightleftharpoons NO + HO_2$ (R32)

Finally, NO reacts with HO₂ or NH₂. The NH₂ + NO reaction leads to formation of N_2 , either directly

 $NH_2 + NO \rightleftharpoons N_2 + H_2O \tag{R18}$

or via

$$NH_2 + NO \rightleftharpoons NNH + OH$$
 (R19)

followed by rapid dissociation of NNH (R33) or reaction with O_2 (R34).

The reaction $NH_2 + NO_2 = N_2O + H_2O$ (R20) is the main source of N_2O at both 30 bar and 100 bar oxidizing conditions. A minor route for consumption of NH_2 is the recombination reaction to form hydrazine,

$$NH_2 + NH_2(+M) \rightleftharpoons N_2H_4(+M)$$

Fig. 9 shows the results of a sensitivity analysis for oxidizing conditions at 30 and 100 bar. The coefficients were obtained by varying A-factors for all reactions by a factor of two. Reactions of NH_2 with NO and NO_2 are important for the oxidation rate. For both reactions a chain propagating/branching product channel,

$$NH_2 + NO \rightleftharpoons NNH + OH$$
 (R19)



Fig. 9. First order sensitivity coefficients for $\rm NH_3$ for conditions of 30 bar (925 K) and 100 bar (900 K).

$$NH_2 + NO_2 \rightleftharpoons H_2NO + NO$$
 (R21)

competes with a chain terminating channel,

$$NH_2 + NO \rightleftharpoons N_2 + H_2O \tag{R18}$$

$$NH_2 + NO_2 \rightleftharpoons N_2O + H_2O \tag{R20}$$

All these steps show up in the sensitivity analysis with large positive (branching) or negative (terminating) coefficients.

Other important reactions which accelerate ammonia consumption are,

$$NH_2 + HO_2 \rightleftharpoons H_2NO + OH$$
 (R11)

$$H_2NO + O_2 \rightleftharpoons HNO + HO_2 \tag{R29}$$

The reaction $H_2NO + O_2$ (R29) has the largest positive sensitivity coefficients. H_2NO is a significant intermediate species as shown in the reaction path analysis and (R29) is favored by the excess of O_2 in oxidizing conditions. It yields HO_2 radicals, which is converted to the more reactive OH radical by $NH_2 + HO_2$ (R11).

The following chain terminating steps inhibit the ammonia consumption,

$$H_2NO + HO_2 \rightleftharpoons HNO + H_2O_2 \tag{R28}$$

$$\begin{array}{l} H_2 NO + NO_2 \rightleftharpoons HONO + HNO \\ NH_2 + NH_2(+M) \rightleftharpoons N_2 H_4(+M) \end{array}$$
(R31)

Reactions (R28) and (R31) are minor consumption channels for H_2NO , compared to $H_2NO + O_2$ (R29), but sufficiently important to show up in the sensitivity analysis. Similarly, $NH_2 + NH_2(+M)$ is a minor sink for NH_2 , but it is strongly chain terminating.

The results show that to develop a reliable reaction mechanism for NH_3 oxidation at the present conditions, it is desirable to obtain more accurate rate constants for several reactions in the H_2NO subset.

5. Conclusions

Ammonia oxidation experiments were conducted at high pressure (30 bar and 100 bar) and temperatures of 450–900 K in oxidizing and stoichiometric conditions. The data were interpreted in terms of a detailed chemical kinetic model, developed for high-pressure conditions. As part of the work, the rate constant for the reaction of $H_2NO + O_2$ was calculated from ab initio theory. The oxidation of ammonia was slow under stoichiometric conditions in the temperature range investigated. Under oxidizing conditions the onset temperature for reaction was 850–875 K at 30 bar, while at 100 bar it was about 800 K, with complete consumption of NH_3 at 875 K. The products of reaction were N_2 and N_2O , while NO and NO_2 concentrations were below the detection limit even under oxidizing conditions.

The agreement between experimental results and modeling work was satisfactory. The main oxidation path for NH₃ under the present conditions was NH₃ $\xrightarrow{+OH}$ NH₂ $\xrightarrow{+HO_2,NO_2}$ H₂-NO $\xrightarrow{+O_2}$ HNO $\xrightarrow{+O_2}$ NO $\xrightarrow{+NH_2}$ N₂. The modeling predictions were most sensitive to the reactions NH₂ + NO = NNH + OH and NH₂ + HO₂ = H₂NO + OH, which promoted the ammonia consumption by forming OH radicals, and to NH₂ + NO = N₂ + H₂O and NH₂ + NO₂ = N₂O + H₂O, which were the main chain-terminating steps.

Acknowledgments

The work is part of the CHEC (Combustion and Harmful Emission Control) research program at DTU Chemical Engineering. YS wishes to acknowledge funding from CSC (China Scholarship Council). PM thanks the R.A. Welch Foundation (Grant B-1174) for support.

Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fuel.2016.04.100.

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