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Pyrolysis and Combustion Chemistry of Pyrrole, a Reference Component for Bio-oil Surrogates: Jet-Stirred Reactor Experiments and Kinetic Modeling

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Cite This: Energy Fuels 2021, 35, 7265-7284



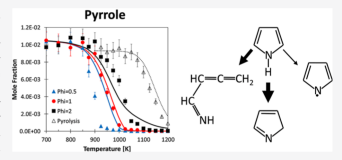
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ABSTRACT: Fast-pyrolysis bio-oils (FPBOs) obtained from lignocellulosic biomass are gaining attention as sustainable fuels for various applications, including the transport sector and power production. A significant fraction of bio-oils is constituted by nitrogen-containing compounds (N fuels) that should be considered when developing surrogate models for FPBOs. Moreover, the content of N fuels in FPBOs is expected to strongly contribute to the production of nitrogen oxides (NO $_x$) directly from fuel-bound nitrogen (fuel NO $_x$), in addition to the thermal NO $_x$ formation pathways typical of high-temperature combustion conditions. This work investigates the pyrolysis and combustion



chemistry of pyrrole (C_4H_5N), a candidate reference fuel component for FPBO surrogate models. Speciation measurements in an atmospheric pressure jet-stirred reactor have been performed for both pyrolysis and oxidation conditions. Pyrolysis experiments have been performed for 1% pyrrole/helium mixtures over the temperature range T = 925-1200 K. Oxidation experiments were carried out for 1% pyrrole/oxygen/helium mixtures at three equivalence ratios ($\varphi = 0.5$, 1.0, and 2.0) over the temperature range T = 700-1200 K. These new data significantly extend the number of experimental targets for kinetic model validation available at present for pyrrole combustion. After a thorough revision of previous theoretical and kinetic modeling studies, a preliminary kinetic model is developed and validated by means of comparison to new experimental data and those previously reported in the literature. The rate of production and sensitivity analyses highlight important pathways deserving further investigations for a better understanding of pyrrole and, more in general, N fuel combustion chemistry. A critical discussion on experimental challenges to be faced when dealing with pyrrole is also reported, encouraging further experimental investigation with advanced diagnostics.

1. INTRODUCTION

Concerns about climate change and energy security are pushing industries and academia to seek alternatives to fossil fuels, pursuing a more sustainable energy scenario. The European Green Deal¹ recently set a roadmap of the key policies and measures needed to meet the United Nations 2030 agenda² in terms of sustainable development goals, aiming at zero net emissions of greenhouse gases by 2050.

Within the different alternatives (e.g., electrification of the transport sector, hydrogen energy, electrofuels, nuclear energy, and hydroelectricity), fast pyrolysis is an effective and promising process to obtain high yields of bio-oils from lignocellulosic biomass. Downstream upgrading of fast-pyrolysis bio-oils (FPBOs) provides valuable fuels for transport and chemicals for industry.³ A great advantage of such a conversion process is that bio-oils have ~5–20 times higher volumetric energy density compared to biomass feedstocks, facilitating transport and distribution to a centralized location for use as feedstock for further downstream processing (e.g., gasification/Fischer—Tropsch synthesis, catalytic hydrotreatment, catalytic cracking,

and hydrodeoxygenation 4,5), therefore driving the development of a sustainable market for lignocellulosic biomass. Bio-oils are very different from fossil fuels in terms of both physical and chemical properties, posing some technical challenge for their effective implementation in existing distribution infrastructures and combustion systems typically used for power or heat generation and in the transport sector [e.g., internal combustion (IC) engines and jet engines]. Indeed, bio-oils are typically non-flammable or non-distillable acidic fuels (pH $\sim 2-3$) with a high water fraction (15–30 wt %), high oxygen content (~ 30 wt % on a dry basis), and significant inorganic fraction (metal, ash, char, and solid particles). Such properties negatively affect both the

Special Issue: In Memory of Mario Costa

Received: November 16, 2020 Revised: February 18, 2021 Published: March 2, 2021





Table 1. Nomenclature of Relevant Species in Pyrrole Pyrolysis and Oxidation^a

[&]quot;Thermodynamic properties are reported in Table S1 of the Supporting Information and compared to literature values where available. Simplified molecular-input line-entry system (SMILES) identifiers are reported in Table S2 of the Supporting Information.

viability of downstream upgrading processes and the direct use of FPBO in combustion processes as a result of generally low heating values, low propensity to ignition, low thermal stability, material incompatibility, corrosion, immiscibility with other hydrocarbon streams, possible fuel pump and nozzle clogging (e.g., in sprays), etc. However, as recently reviewed by Letho et al., accounting for FPBO upgrading and relatively minor technical and operational adjustments (e.g., material selection, air and fuel preheating, co-feeding with support fuel, or feeding to a pilot flame) already allowed for successful testing of FPBOs in large-scale burners, gas turbines, and compression ignition engines for heat and/or power generation.

The key to the solution of the technical challenges related to FBPO use in combustors and IC engines is the knowledge of high- and low-temperature combustion chemistry of FPBOs and their upgraded streams. Indeed, chemical kinetics plays the major role in understanding and optimizing combustion processes, for improved efficiency, improved fuel economy, and reduced pollution. Because it is typical for complex liquid fuels, such as FPBOs, a fuel model requires first the definition of a limited number of reference species accounted for in the surrogate fuel model. For each of these species, a kinetic subset together with thermodynamic and transport properties is then required to describe pyrolysis and high- and low-temperature combustion phenomena, such as ignition, flame propagation, and pollutant formation. FPBOs contain hundreds of organic compounds, such as phenolic components, aldehydes, alcohols, acids, esters, anhydrosugars, furans, and nitrogen-containing compounds, as well as large anhydro-oligosaccharides and lignin-derived oligomers.8 From a pure combustion chemistry perspective, each of these chemical families should be taken into account when formulating a suitable fuel surrogate because each of them carry a specific reactivity strongly related to specific functional groups and the molecular structure⁹ that can be determining in properly predicting the macroscopic target of interest for large-scale applications.

As reported in recent studies, the CRECK kinetic framework was extended to describe a large number of chemical families found in FPBOs. Kinetic subsets for phenol and phenolic compounds (e.g., anisole, catechol, guaiacol, and vanillin) have been recently presented by Pratali Maffei et al. ¹⁰ and Pelucchi et al. ¹¹ Aldehyde chemistry has been recently discussed in refs 12 and 13, with the latter study specifically focusing on benzaldehyde, the simplest aromatic aldehyde. Alcohol chemistry has been systematically investigated in refs 14 and 15. Kinetic subsets for acetic, butanoic, and pentanoic acids have also been developed and validated. ^{16,17} Previous studies by Grana et al., ^{18,19} Saggese et al., ²⁰ and Rodriguez et al. ²¹ addressed the chemistry of methyl esters and fatty acid methyl esters (FAMEs), both of interest as surrogate components for FPBOs and biodiesel fuels (e.g., FAMEs).

This work presents a further extension of the fuel palette of the CRECK kinetic framework 22 to describe the pyrolysis and combustion kinetics of a nitrogen-containing fuel (N fuel): pyrrole (C₄H₅N). Beside the mere necessity of representing the N fuel fraction of FPBOs, understanding pyrrole combustion kinetics is also of relevance for a better assessment of fuel NO_x, namely, the fraction of nitrogen oxides (NO_x) formed during the oxidation of nitrogen contained in a fuel molecule in a combustion environment. 23,24 Fuel NO_x are of interest in not only FPBO combustion but also biomass and coal combustion. Indeed, biomass and coal first undergo devolatilization processes, 25,26 during which a part of fuel-bound nitrogen is

devolatilized and pyrolyzed into NO_x precursors, such as hydrogen cyanide (HCN), ammonia (NH₃), and isocyanic acid (HNCO), ^{23,24} which can be further converted into NO, N₂O, and N₂ as final products. In this regard, this work extends our recently revised model for thermal and prompt NO_x formation as well as high-temperature NO_x reburning phenomena.²⁷

2. PREVIOUS EXPERIMENTAL, THEORETICAL, AND KINETIC MODELING STUDIES ON PYRROLE PYROLYSIS AND OXIDATION

A very limited number of theoretical, experimental, and kinetic modeling studies has been reported for pyrrole in previous studies. Structures and names of the chemical species related to pyrrole chemistry and used in the following discussion are reported in Table 1 to facilitate the reading.

Lifshitz et al. 28 investigated the thermal decomposition of pyrrole in a single-pulse shock tube at T = 1050-1450 K and p = 2.6-3.6 atm (i.e., overall density of $\sim 3 \times 10^{-5}$ mol/cm³). Pyrrole was found to mainly isomerize to cis-crotonitrile (cC₃H₅CN) and allyl cyanide (aC₃H₅CN) or decompose to propyne (C₃H₄-p) and hydrogen cyanide (HCN), in relative ratios of 58, 25, and 17%. Secondary reactivity of these major products yields other species measured in significant quantities, such as acetonitrile (CH₃CN), acetylene (C₂H₂), allene (C₃H₄-a), methane (CH_4) , and ethylene (C_2H_4) . Mackie et al. ²⁹ studied the kinetics of the pyrolysis of highly diluted pyrrole/argon mixtures (0.07 and 0.5 mol % pyrrole) in a single-pulse shock tube at T = 1200-1700 K and p = 7.5-13 atm. A kinetic model composed of 75 elementary steps was presented and found to largely reproduce the experimental observations. The authors suggested that the thermal decomposition of pyrrole is initiated by the reversible isomerization to pyrrolenine (2*H*-pyrrole, PYRLNE), occurring through a 1–2 hydrogen shift. The C-N bond fission in pyrrolenine then leads to ring opening, forming a biradical that is rapidly transformed in cis-crotonitrile or allyl cyanide. H-abstraction reactions by H and CH3 also contribute to pyrrole consumption, forming the resonance-stabilized pyrrolyl radical and secondary products, such as hydrogen (H₂) and methane. Dubnikova and Lifshitz³⁰ theoretically investigated the isomerization pathways of pyrrole using the density functional theory (B3LYP/cc-pvDZ). The theoretical analysis confirmed the previous findings by Mackie et al.,2 according to which the decomposition of pyrrole is initiated by the fast isomerization to pyrrolenine through a 1,2-H migration, reaching a state of equilibrium from which ring-opening and isomerization reactions to cis-crotonitrile (cC3H5CN) and allyl cyanide (aC3H5CN) occur. Rice-Ramsperger-Kassel-Marcus (RRKM) calculations were performed, and the master equation (ME) solved for the highest energy barrier (E_a) steps, i.e., for the first isomerization $(C_4H_5N \leftrightarrow PYRLNE; E_a = 43.1 \text{ kcal/mol})$ and for the ring-opening reaction leading to a biradical intermediate (PYRLNE \leftrightarrow N=CH-CH=CH-CH₂; $E_a = 68.0$ kcal/mol). This intermediate can further isomerize to allyl cyanide ($E_a = 16.3 \text{ kcal/mol}$) or to a less stable conformer ($E_a = 4.1 \text{ kcal/mol}$) accessible by rotating the N=CH moiety. This conformer can then isomerize to *cis*-crotonitrile ($E_a = 7.7$ kcal/mol) and trans-crotonitrile (tC₃H₅CN). Optimized geometries and frequencies for biradical structures were performed using the unrestricted uB3LYP with the same basis set and also optimized with a complete active space multiconfiguration self-consistent field (CASSCF) with CAS(4,4) wave functions for comparison. Each optimized SCF structure was recalculated at a single-point quadratic CI, including single and double substitutions with a triplet contribution to the energy, QCISD(T), using uB3LYP structures as the first guess. An additional theoretical investigation of the pyrolysis mechanisms of pyrrole was presented by Zhai et al.³¹ Equilibrium and transition state structures of the proposed reaction channels were fully optimized by the density functional B3LYP method using the 6-31G(d,p) basis set, and relative energies were evaluated at the QCISD(T)/6-311G(d,p) level of theory or with the unrestricted equivalents for biradical structures [uB3LYP and uQCISD(T)]. Good agreement was obtained in terms of optimized structures, but the channels leading to allyl cyanide turned

out to be much higher in energy compared to the previous calculations as a result of both the different basis sets and some inconsistencies in the calculation methods adopted by Dubnikova and Lifshitz,³⁰ highlighted by Zhai et al.³¹ The latter study proposed an additional closed-shell pathway, not involving the biradical structures, for the formation of cis-crotonitrile. Two low-energy closed-shell intermediates are successively formed from pyrrolenine: 3H-pyrrole (pyrrolenine ↔ 3H-pyrrole; $E_a = 28.2 \text{ kcal/mol}$) through a second 1,2-H migration and cis-isocyanocrotonitrile (3*H*-pyrrole \leftrightarrow cis-isocyanocrotonitrile; E_a = 68.3 kcal/mol) via a concerted transition state, including a C-C bond cleavage and a 1,2-hydrogen migration. The authors concluded that the new low-energy pathway might compete with that investigated by Dubnikova and Lifshitz at low-pressure conditions (i.e., <1 atm), but at a higher pressure the open-shell channel forming the biradical structures dominates the decomposition chemistry as a result of collisional deactivation of the more stable intermediates. Extensive calculations were carried out in the same study seeking decomposition pathways directly generating HCN, as highlighted by the experimental measurements of Dubnikova and Lifshitz.³⁰ However, no such competitive pathway was identified. Martoprawiro et al.³² investigated the pyrolysis kinetics of pyrrole, including the thermochemistry of relevant species, by means of CASSCF, CASPT2, and G2(MP2) calculations. In addition to the two major decomposition channels proceeding through pyrrolenine already reported in the previous studies, a third channel involving the fission of a H atom and formation of the resonance-stabilized cyanoallyl radical ($a\dot{C}_3H_4CN$, $\dot{C}H_2-CH$ = CH−C≡N) was identified. An additional channel forming an allenic imine (HN=CH-CH=C=CH2, HCNPROP) contributes to the formation of HCN and propyne. The lowest energy pathway involves the isomerization of pyrrole to a cyclic carbene through a 4,5-H migration, and successive ring opening produces the allenic imine intermediate.³³ On the basis of the calculated rate constants and the previous studies briefly reviewed above, Martoprawiro et al.³² proposed a kinetic model validated by comparison to the experimental pyrolysis data by Mackie et al.²⁹ Hong et al.³⁴ performed experimental measurements for the pyrolysis of pyrrole (6.46 mol % in argon) with the tunable synchrotron vacuum ultraviolet (VUV) photoionization and molecular-beam mass spectrometry (MBMS) technique in a pyrolysis chamber located in a high-temperature furnace. The measurements were carried out at p = 0.002 atm, over the temperature range T = 1260-1710 K. Formation pathways of the major products (HCN, C₂H₂, CH₃CN, and C₃H₄-p) and radical intermediates (aC₃H₄CN and CH₂CN) were investigated using the composite G3B3 method, highlighting the lowest energy formation pathways. The HCN formation channel proceeding through cyclic carbene discussed above was proposed to be most favored, in accordance with the previous theoretical work by Martoprawiro et al.

Assuming complete oxidation of molecular nitrogen to nitrogen dioxide (NO_2) , the combustion of pyrrole can be defined by the following reaction equation:³⁵

$$C_4H_5N + 6.25O_2 \rightarrow 4CO_2 + 2.5H_2O + NO_2$$

Lumbreras et al.³⁶ presented the first experimental and kinetic modeling study on pyrrole oxidation. Experimental measurements were carried out in an isothermal quartz flow reactor at atmospheric pressure in the temperature range T = 700-1500 K for diluted mixtures of pyrrole (0.01 mol %), oxygen ($\sim 0.05-1.37 \text{ mol }\%$), and water ($\sim 1.0-1.2 \text{ mol }\%$) %). The effects of the temperature, equivalence ratio (φ = 0.04, 0.90, and 1.18, where φ is defined from the above reaction equation), and NO addition (~0.3 mol %) on CO, CO2, HCN, and NO concentrations were evaluated experimentally and modeled by means of a preliminary kinetic model. Rate constants were adopted from previous studies for pyrolysis pathways^{28–33} and based on analogy with phenol/phenoxy chemistry. Ignition delay times of diluted mixtures of pyrrole (0.5 and 1.0 mol %) and oxygen in argon were measured in a low-pressure shock tube by MacNamara and Simmie. 35 Ignition measurements were performed in the temperature range T = 1102-1805 K, pressures p = 220-520 kPa, and equivalence ratios $\varphi = 0.5, 1.0$, and 2.0. Koger and Bockhorn³⁷ investigated the formation of HCN from the oxidation of pyrrole under incinerator conditions (T = 1180

and 1220 K, p = 1 atm, and $\varphi = 0.81$ and 1.04) in a turbulent flow reactor. Tian et al.³⁸ investigated the oxidation of pyrrole, oxygen, and argon mixtures ($\varphi = 0.55$ and 1.84) in premixed laminar flames at p =0.032 atm using tunable synchrotron photoionization and MBMS techniques. Results highlighted that N_2 NO, and NO₂ are the major nitrogen-containing products, while hydrogen cyanide, isocyanic acid (HNCO), and 2-propenenitrile (CH₂CHCN) are the most important nitrogen-containing intermediates. The formation of fuel NO from pyrrole oxidation was studied by Yamamoto et al.³⁹ in a quartz flow reactor in the temperature range T = 800-1400 K at atmospheric pressure. The inlet concentration of pyrrole (0.02 mol %) was kept constant while varying the O₂ content (0.64 and 2.00 mol %) and water content (3 and 8 mol %). The impact of the residence time was also assessed for the mixture containing 2.00 mol % oxygen and 8 mol % water. A kinetic model composed of 89 chemical species and 505 reactions was proposed on the basis of the previous pyrolysis model by Mackie et al.²⁹ and the oxidation pathways proposed by Lumbreras et

The present work reports new experimental measurements for pyrrole pyrolysis and oxidation in an atmospheric pressure jet-stirred reactor (JSR), significantly extending the scarce validation targets presented in the literature thus far. Pyrolysis experiments have been performed in the temperature range T = 925-1200 K for mixtures of ~ 1.0 mol % pyrrole in helium. Oxidation experiments have been performed in the temperature range T = 700-1200 K for 1.0% pyrrole/ O_2 /He mixtures at variable equivalence ratios ($\varphi = 0.5$, 1.0, and 2.0).

In addition, a preliminary model is presented on the basis of previous theoretical, experimental, and kinetic modeling efforts. Thermodynamic data not reported in previous studies or not available in thermodynamic property databases have also been calculated in this work. To the best of our knowledge, this model represents the first comprehensively validated model reported in the literature thus far and, despite evident shortcomings clearly underlined and discussed in the kinetic analysis section, constitutes a useful tool to assess chemical pathways of fuel NO_x formation and to extend the fuel palette of the CRECK kinetic framework to include a representative compound to reproduce the N fuel content in FPBOs.

3. JSR EXPERIMENTS

Different set of experiments have been performed covering both pyrolysis and oxidation conditions. The first section below describes the apparatus used to perform experiments as well as the analytical method. The second section describes the experimental data obtained for pyrrole pyrolysis and oxidation.

3.1. Experimental Method Description. The experimental setup of pyrrole was a laboratory-scale JSR (85 cm³) working close to atmospheric pressure (1.07 bar). This setup is described in detail elsewhere, 12,27,40 and only a brief description is provided here. Experiments were performed in a fused silica JSR, a type of ideal continuously stirred-tank reactor, which is suitable for gas-phase kinetic studies. Reactants, with helium as the carrier gas, entered the spherical JSR through four nozzles located at its center, allowing for the creation of high turbulence resulting in homogeneity in temperature and composition. The reactor was heated using Inconel resistances, and the reaction temperature was measured with a K-type thermocouple positioned in a glass finger close to the center of the reactor (uncertainty of ± 5 K). Pyrrole pyrolysis and oxidation under stoichiometric conditions were carried out at a residence time of about 2 s and at temperatures ranging from 700 to 1200 K with initial fuel mole fraction of $\sim 10\,000$ ppm. The experimental conditions are summarized in Table

The purities of helium and oxygen were 99.99% and provided by Messer. Pyrrole was provided by Sigma-Aldrich with a claimed purity of 98% and used without further purification because a GC analysis did not identify any impurity in the reactant, despite the careful chromatogram analysis. A liquid Coriolis flow controller was used to control the flow of pyrrole, mixed with helium and passed through an evaporator (393 K) before being mixed with oxygen prior to entering the reactor. The relative uncertainty in gas flow rates is about 0.5%.

Table 2. Summary of JSR Experimental Conditions Used in the Present Study

					inlet mole fraction (%)		
set	T (K)	p (bar)	τ (s) ^a	φ^{b}	pyrrole	O ₂	He
1	925-1200	1.067	2	∞	0.93	0.00	99.07
2	700-1200	1.067	2	0.5	1.05	12.95	86.00
3	700-1200	1.067	2	1	1.05	6.20	92.75
4	700-1200	1.067	2	2	1.05	3.08	95.87

^aThe residence time is defined as the ratio between the reactor volume and the gas flow rate (m³/s) under the conditions of the temperature and pressure in the reactor. ^bThe equivalence ratio was defined by considering the following stoichiometric equation: $C_4H_5N + 6.25O_2 \rightarrow 4CO_2 + 2.5H_2O + NO_2$.

Although the boiling point of pyrrole is ~403 K, the temperature of the evaporator was set at a lower temperature to avoid the fouling and clogging of this part of the apparatus (it occurred several times, and the evaporator had to be cleaned; this operation was quite complex given the diameter of the tube inside the evaporator).

The reactants and reaction products leaving the reactors were then transported by a heated transfer line maintained at 393 K to avoid condensation to a Fourier transform infrared (FTIR) spectroscopy device and two gas chromatographs (GCs). The first GC, equipped with a Carbosphere-packed column, a thermal conductivity detector (TCD), and a flame ionization detector (FID), was used to quantify lightweight species. The second GC, fitted with a Q-Bond capillary column and a FID preceded by a methanizer, is used for the quantification of compounds containing two carbon atoms. The methanizer (nickel catalyst for hydrogenation) allows for the detection of species, like CO and CO2, and allows for the detection species, like CH₃CHO, with better sensitivity. Moreover, FTIR spectroscopy is also used to quantify CO, CO₂, and HCN. Identification of species was performed using a GC (with a Q-Bond capillary column) coupled to a mass spectrometer with electron impact ionization at 70 eV. The FTIR spectrometer from Thermo Scientific Antaris is equipped with a mercury cadmium telluride photoelectric detector. Spectra were recorded over the wavelength range of 400-4000 cm⁻¹ with a resolution of 0.5 cm⁻¹. The cell (optic path of 10 m) was heated to 373.15 K, and measurements were performed at a pressure of 150 Torr. An average of 32 scans was considered for a spectrum measurement. The detection limit depends upon the species, the absorption line used for the quantification, and possible interferences with other species. In the present study, it was about 35 ppm for HCN, 25 ppm for CO, and 2 ppm for CO₂. Note that FTIR and GC analyses were performed in separated experiments. Excellent agreement was observed for carbon monoxide (a species with some isolated lines in the absorption spectra) mole fractions obtained with the two diagnostics.

GC calibrations were performed using gaseous standards provided by Messer and Air Liquide for small species, like carbon monoxide, carbon dioxide, oxygen, methane, and HCN. The calibration was performed for the fuel by injecting synthetic gas mixtures of pyrrole and helium. Other species detected with the FID were calibrated using the effective carbon number (ECN) method (their calibration factors were deduced from those of species calibrated taking into account their number of effective carbon atoms). FTIR calibrations were performed for all species, which were detected using this technique (CO, CO₂, and HCN) using gaseous standards provided by Messer and Air Liquide. Relative uncertainties in mole fractions of species detected by GC and calibrated using gaseous standards provided by Messer and Air Liquide were estimated to $\pm 5\%$. The relative uncertainty in the mole fractions of pyrrole was estimated to $\pm 10\%$, although it was calibrated as a result of the difficulties in handling such a species. Relative uncertainties in mole fractions of species calibrated using the ECN methods were estimated

3.2. Experimental Results. Reaction products detected during the pyrolysis of pyrrole are hydrocarbons and N-containing species. Hydrocarbons are methane, acetylene, ethylene, ethane, propene,

allene, propyne, a C_4H_x species, which could not be clearly identified (but highly unsaturated), benzene, and toluene. N-containing species are hydrogen cyanide (HCN), acetonitrile (CH $_3$ CN), 2-propenenitrile (also called acrylonitrile, CH $_2$ CHCN), and three isomers for butenenitrile [but-3-enitrile, (2E)-but-2-enenitrile, and (2Z)-but-2-enenitrile, also named allyl cyanide, *trans*-crotonitrile, and *cis*-crotonitrile, respectively]. Note that the mass spectra for the three isomers are similar and that the peak attribution could not be performed with certainty. The conversion of pyrrole becomes significant from ~1050 K. Mole fractions of most of species increase over the studied temperature range. The few species for which a maximum in mole fraction is observed are allene, propyne, propene, and all N-containing species, except HCN.

Reaction products detected during the oxidation experiments are the same as those detected during pyrolysis (except for toluene, only observed for pyrolysis). In addition, small oxygenated compounds, like carbon monoxide and carbon dioxide, are observed. The reactivity is enhanced in comparison to pyrolysis, with significant pyrrole conversions observed from ~850 K. The equivalence ratio has an effect on the reactivity, with the leaner case being the most reactive and the richer case being the least reactive. At mid conversion, the three pyrrole mole fraction profiles are shifted by about 50 K under the conditions of the present study. In a general way, all detected intermediates see their mole fractions going through a maximum, except for carbon dioxide, which is an end product of combustion reactions. For the rich case, the deficit of $\overline{O_2}$ leads to higher mole fractions of intermediates and significant mole fractions of CO are still observed at 1200 K, which is the highest temperature considered in this study. For the stoichiometric and lean mixtures, some of the intermediates observed under pyrolysis and rich conditions are not detected (e.g., ethane, propene, allene, propyne, etc.).

Atomic balances were performed for all experiments. For pyrolysis, C, N, and H atomic balances are satisfactory up to 1100 K (they lie in between 1 and 0.85). Above 1100 K, the three atomic balances tend to decrease monotonously, reaching ~0.5 for C and N and ~0.7 for H. This was not surprising given the fouling that was observed in the lines between the outlet of the reactors and the analytical devices. The fouling is likely due to the condensation of some heavy aromatic species (possibly containing N atoms). For oxidation experiments, the carbon atom balance is satisfactory over the whole range of temperatures, usually lying in between 0.8 and 1, except in the range of 850-1000 K, where it falls to ~0.8. In a general way, it is slightly better for the lean and stoichiometric mixtures than for the rich mixture. H and O atom balances are not meaningful because water, one of the main reaction products, was not quantified during this study. The N atomic balance is satisfactory up to 875 K, and then it decreases monotonically to 0 for the lean and stoichiometric cases and up to ~0.4 for the rich case, indicating that one or several N-containing species were not detected during these oxidation experiments. The potential presence of some species, such as N2, NH3, NO, NO2, and N2O, was then further investigated. If present, N₂ could be detected if above the detection limit (~500 ppm) using GC with detection using the TCD. This species was not observed during oxidation nor during pyrolysis of pyrrole. If present in a sufficiently high concentration, NH₃, NO, NO₂, and N₂O could be detected using the FTIR spectroscopy tool because they have very characteristic absorption structures in the wavenumber range investigated (detection limits of ~2, ~100, ~10, and ~50 ppm, respectively). None of these species could be identified in recorded spectra during oxidation experiments, even in traces.

4. THEORETICAL METHODS FOR THERMODYNAMIC PROPERTY ESTIMATION

Thermodynamic properties of pyrrole, pyrrolenine, and pyrrolyl radical were calculated by first-principle calculations using the Gaussian 16 revision B suite of programs 41 at the CBS-QB3 42 and G4 43 levels of theory as implemented. Both methods use B3LYP geometries and frequencies, although using different basis sets, and contain several energy calculation steps to extrapolate the electronic energy to a CCSD(T)/CBS level. Electronic energies are converted with the

Table 3. Rate Coefficients for Relevant Reactions in the Following Discussion on Pyrrole Pyrolysis and Oxidation^a

	reaction	A	n	E_{a}	reference	notes
1 C	$C_4H_5N \leftrightarrow PYRLNE$	3.16×10^{13}	0.005	46300	32	
2 C	$L_4H_5N \leftrightarrow HNCPROP$	1.10×10^{14}	0.000	77162	32	
3 P	$PYRLNE \leftrightarrow aC_3H_5CN$	5.24×10^{15}	0.000	75710	32	$A \times 2$
1 P	$PYRLNE \leftrightarrow cC_3H_5CN$	1.65×10^{15}	0.000	70050	32	A/2
5 P	PYRLNE ↔ HNCPROP	2.51×10^{15}	0.000	79474	32	
5 H	$HNCPROP \leftrightarrow HCN + C_3H_{4}p$	5.50×10^{12}	0.000	37740	32	$A/2$, $E_a = +1500$ cal/mol
7 P	$YRLNE \leftrightarrow \dot{H} + a\dot{C}_3H_4CN$	2.04×10^{17}	0.000	86746	32	$A \times 2$
3 a	$C_3H_5CN \leftrightarrow tC_3H_5CN$	7.00×10^{14}	0.000	61969	32	
) a	$C_3H_5CN \leftrightarrow cC_3H_5CN$	7.20×10^{14}	0.000	58863	32	
10 c	$C_3H_5CN \leftrightarrow tC_3H_5CN$	1.40×10^{14}	0.000	57573	32	
l1* a	$C_3H_5CN \leftrightarrow \dot{C}_2H_3 + \dot{C}H_2CN$	3.40×10^{15}	0.000	82640	pw	$*C_4H_8-1 \leftrightarrow C_2H_3 + C_2H_5$
	$C_3H_5CN + \dot{H} \leftrightarrow C_2H_4 + \dot{C}H_2CN$	1.00×10^{13}	0.000	3010	57	$A \times 2$
	$C_3H_5CN + \dot{H} \leftrightarrow HCN + \dot{C}_3H_5-s$	6.00×10^{12}	0.000	4000	pw	$*H + C_2H_2$
	$H + C_4H_5N \leftrightarrow H_2 + PYRLYL$	1.00×10^{6}	2.000	2825	pw	*H-abs. tertiary C-H
15 Ċ	$CH_3 + C_4H_5N \leftrightarrow CH_4 + PYRLYL$	4.50×10^4	2.000	3778	pw	
16 Ċ	$CH_2CN + C_4H_5N \leftrightarrow CH_3CN + PYRLYL$	1.35×10^4	2.000	12460	pw	$*C_3H_3 + C_4H_5N$
17 Ċ	$OH + C_4H_5N \leftrightarrow H_2O + PYRLYL$	9.00×10^{8}	1.000	-695	pw	
	$HO_2 + C_4H_5N \leftrightarrow H_2O_2 + PYRLYL$	3.60×10^6	2.000	14440	pw	
	$O_2 + C_4H_5N \leftrightarrow HO_2 + PYRLYL$	8.00×10^{13}	0.000	37150	pw	
	$O + C_4H_5N \leftrightarrow OH + PYRLYL$	1.10×10^{6}	2.000	1404	pw	
	$H + aC_3H_5CN \leftrightarrow H_2 + aC_3H_4CN$	1.90×10^{2}	3.500	1627	pw	*R + $C_4H_8-1 \leftrightarrow C_4H_71-3 + RH (A/3)$
	$CH_3 + aC_3H_5CN \leftrightarrow CH_4 + aC_3H_4CN$	7.14×10^{0}	3.500	7642	pw	
	$CH_2CN + aC_3H_5CN \leftrightarrow CH_3CN + aC_3H_4CN$	2.00×10^{11}	0.000	12000	57	A/2
	$OH + aC_3H_5CN \leftrightarrow H_2O + aC_3H_4CN$	7.70×10^{5}	2.200	-437	pw	
	$\dot{HO}_2 + aC_3H_5CN \leftrightarrow H_2O_2 + a\dot{C}_3H_4CN$	7.82×10^{-1}	3.970	11702	pw	
	$O_2 + aC_3H_5CN \leftrightarrow H\dot{O}_2 + a\dot{C}_3H_4CN$	5.00×10^{13}	0.000	37190	pw	
	$O + aC_3H_5CN \leftrightarrow OH + aC_3H_4CN$	1.75×10^{11}	0.700	5884	pw	
	$H + tC_3H_5CN \leftrightarrow H_2 + aC_3H_4CN$	3.64×10^{5}	2.455	4361	pw	$*R + C_3H_6 = C_3H_5 - A + RH$
	$CH_3 + tC_3H_5CN \leftrightarrow CH_4 + aC_3H_4CN$	2.21×10^{0}	3.500	5675	pw	
	$CH_2CN + tC_3H_5CN \leftrightarrow CH_3CN + aC_3H_4CN$	5.00×10^{12}	0.000	10989	57	A/2
	$OH + tC_3H_5CN \leftrightarrow H_2O + aC_3H_4CN$	4.46×10^{6}	2.072	1051	pw	$*R + C_3H_6 = C_3H_5 - A + RH$
	$HO_2 + tC_3H_5CN \leftrightarrow H_2O_2 + aC_3H_4CN$	3.07×10^{-2}	4.403	13547	pw	
	$O_2 + tC_3H_5CN \leftrightarrow HO_2 + aC_3H_4CN$	1.20×10^{20}	-1.67	46191	pw	
	$O + tC_3H_5CN \leftrightarrow OH + aC_3H_4CN$	5.24×10^{11}	0.700	5884	pw	
	$H + tC_3H_5CN \leftrightarrow H_2 + C_3H_4CN$	2.25×10^7	1.930	12950	pw	$*R + C_2H_4 = C_2H_3 + RH (A/2)$
	$CH_3 + tC_3H_5CN \leftrightarrow CH_4 + C_3H_4CN$	4.85×10^{2}	2.947	15148	pw	
	$CH_2CN + tC_3H_5CN \leftrightarrow CH_3CN + \dot{C}_3H_4CN$	5.00×10^{12}	0.00	12000	57	A/2
38 Ċ	$OH + tC_3H_5CN \leftrightarrow H_2O + C_3H_4CN$	1.11×10^4	2.745	2216	pw	$*R + C_2H_4 = C_2H_3 + RH (A/2)$
	$\dot{HO}_2 + tC_3H_5CN \leftrightarrow H_2O_2 + \dot{C}_3H_4CN$	2.15×10^{5}	2.000	20243	pw	
	$O_2 + tC_3H_5CN \leftrightarrow HO_2 + C_3H_4CN$	2.11×10^{13}	0.000	57623	pw	
	$O + tC_3H_5CN \leftrightarrow OH + C_3H_4CN$	1.08×10^{7}	2.000	8782	pw	
	$\dot{C}_3H_4CN \leftrightarrow c\dot{C}_3H_4CN$	5.00×10^{13}	0.000	51983	57	
	$PYRLYL \leftrightarrow c\dot{C}_3H_4CN$	1.50×10^{13}	0.000	38987	32	
	$C_3H_4CN \leftrightarrow CH_3 + C_3HN$	6.00×10^{14}	0.000	42000	57	
	$C_3H_4CN \leftrightarrow C_2H_2 + CH_2CN$	1.07×10^{15}	-0.560	36320	pw	$*C_4H_71-1 \leftrightarrow C_2H_2 + C_2H_5$
	$H + C_3HN \leftrightarrow C_2H_2 + \dot{C}N$	1.00×10^{14}	0.000	2000	pw	$*H + C_2H_2$
	$\dot{C}_3H_4CN + H\dot{O}_2 \leftrightarrow C_4H_4N\dot{O} + \dot{O}H$	1.95×10^{18}	-1.060	7852	pw	*C_3H_5 -A + HO ₂ (1 atm)
	$\dot{C}_3H_4CN + O_2 \leftrightarrow C_4H_4NO + \ddot{O}$	2.30×10^{20}	-2.650	6489	pw	$*C_3H_5-A + O_2 (1 \text{ atm})$
	$C_4H_4NO \leftrightarrow C_2H_3CHO + CN$	1.50×10^{13}	0.000	33000	60	C–C β -scission
	$C_4H_4NO \rightarrow C_2H_2 + CH_2O + CN$	1.50×10^{13}	0.000	33000	60	C–C β -scission
	$CH_2CN + CH_2CN \leftrightarrow C_4H_4N_2$	2.30×10^{13}	0.000	0.000	58	
	$CH_3CN (+M) \leftrightarrow CH_2CN + H (+M)$	9.20×10^{12}	0.850	95770	58	high-pressure limit
	$CH_2CN + C_4H_4N_2 \leftrightarrow CH_3CN + C_4H_3N_2$	3.50×10^{12}	0.000	5000	58	
	$C_4H_3N_2 \leftrightarrow CH_2CHCN + \dot{C}N$	4.40×10^{14}	0.000	55000	58	
	$NO + \ddot{O} (+M) \leftrightarrow NO_2 (+M)$	1.30×10^{15}	-0.750	0.000	61	
	Ö + C₃HN ↔ CO + HÖ−CN	7.40×10^{8}	1.280	2472	pw	$*C_2H_2 + O \leftrightarrow CH_2 + CO$
	$O_2 + H\ddot{C} - CN \leftrightarrow CO_2 + HCN$	1.10×10^{12}	0.000	0.000	56	
	$OH + CH_3CN \leftrightarrow H_2O + CH_2CN$	2.00×10^{7}	2.000	5000	pw	$*OH + C_3H_4-p \leftrightarrow C_3H_3 + H_2O$
	$CH_2CN + \ddot{O} \leftrightarrow \dot{H} + OCHCN$	3.00×10^{11}	0.640	0.000	56	
61 C	OCHCN ↔ HCN + CO	3.50×10^{14}	0.000	66300	62	
	OH + HCN ↔ H + HNCO	1.71×10^{11}	0.000	8744	63	

Table 3. continued

	reaction	A	n	E_{a}	reference	notes
R63	$\dot{O}H + HCN \leftrightarrow H_2O + \dot{C}N$	1.45×10^{13}	0.000	10900	64	
R64	$\dot{O}H + HNCO \leftrightarrow H_2O + \dot{N}CO$	3.50×10^{6}	1.500	3600	65	
R65	$O_2 + \dot{N}CO \leftrightarrow CO_2 + NO$	2.00×10^{12}	0.000	20000	66	
R66	$C_2N_2 + M \leftrightarrow \dot{C}N + \dot{C}N + M$	1.60×10^{34}	-4.32	130000	67	
R67	$C_2N_2 + \dot{H} \leftrightarrow HCN + \dot{C}N$	3.10×10^{14}	0.000	7860	58	
R68	$\dot{\text{HCCO}}$ + NO \leftrightarrow HCN + CO ₂	2.23×10^{14}	-0.750	400	68	

"Rate coefficients refer to an Arrhenius expression of the rate constants as $k = AT^n \exp(-E_a/RT)$. Units are cal, mol, K, cm, and s. Reactions noted as "pw" have been estimated in the present work. Analogy assumptions are noted with an asterisk.

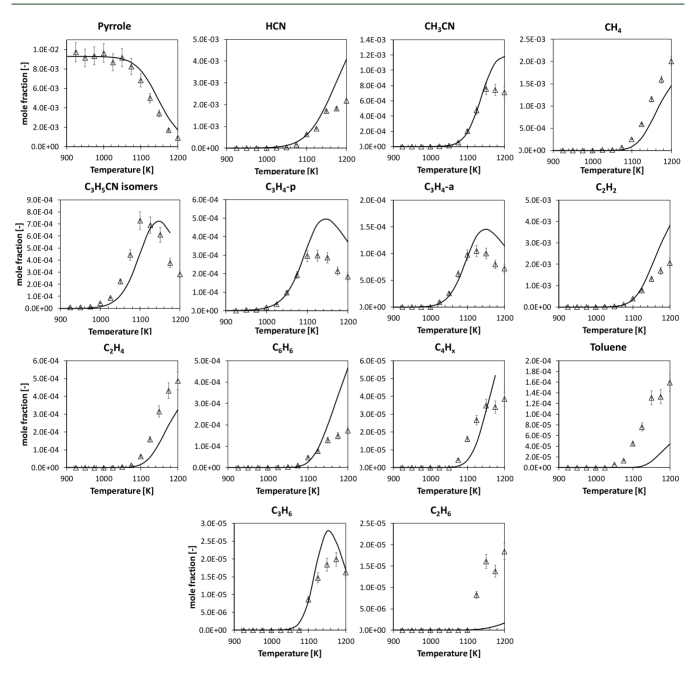


Figure 1. Pyrolysis of pyrrole (\sim 1 mol % in helium) in a JSR at p=107 kPa and $\tau=2.0$ s. Comparison between experimental (symbols) and predicted (lines) mole fraction profiles of intermediate and product species.

atomization method to the corresponding heats of formation. In the case of CBS-QB3, corrections for additive bond errors (BACs)⁴⁴ were applied.

The thermal contributions to the enthalpy, the entropy at 298 K, and the heat capacities as a function of the temperature have been calculated with methods of statistical mechanics. 45 The required input data

Figure 2. Rate of production analysis at T = 1100 K for a pyrrole/helium (1/99 mol %) mixture at p = 107 kPa and $\tau = 2.0$ s. Arrow width qualitatively represents the importance of each reactive flux. Pathways with a flux going from or to an intermediate of <1% have been disregarded for clarity.

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Figure 3. Ring-opening reaction of pyrrolenine to form crotonitrile isomers and allyl cyanide (C₃H₅CN isomers) through a biradical intermediate.

(rotational constants, molecular weight, and scaled frequencies) are readily available from the CBS-QB3 calculations. The harmonic oscillator, rigid rotor assumption is applied because the species of interest do not contain internal rotations. The calculated total entropies contain corrections for the symmetry if needed. The thermodynamic data are converted to NASA polynomials and used in the kinetic model.

The National Institute of Standards and Technology (NIST) WebBook contains two very different entries for the enthalpy of formation of pyrrole: $\Delta_t H^{298} = 143.2 \ \text{kJ/mol}^{46}$ and $108.3 \ \text{kJ/mol}^{47}$ The current CBS-QB3 result of $\Delta_t H^{298} = 106.1 \ \text{kJ/mol}$ agrees well with the older experimental value. With G4, a $\Delta_t H^{298}$ value of 109.7 kJ/mol is obtained, which supports CBS-QB3 as well as the lower experimental enthalpies of formation. The data are also in agreement with the previous works by Simmie, 48 who reports $109.2 \pm 2.3 \ \text{kJ/mol}$ through isodesmic reactions and $110.9 \ \text{kJ/mol}$ via the atomization method, and Lo and Lau, 49 who report $\Delta_t H^{298} = 110.9 \ \text{kJ/mol}$ using a CCSD(T)/CBS approach.

5. KINETIC MODEL

The kinetic model to describe pyrrole pyrolysis and oxidation accounts for 189 chemical species and 2888 reactions and is available as Supporting Information together with thermodynamic properties (model 1). Specifically, the pyrrole pyrolysis and oxidation subset contains 33 species and 456 reactions. Chemical structures and names of relevant chemical species in the pyrrole kinetic subset are reported in Table 1 (section 2).

 for a hierarchical development of the CRECK kinetic modeling framework, modifications to relevant kinetic subsets, such as those describing NO_x or the core $\mathrm{C_0-C_4}$ mechanism that have already been addressed in recent efforts, 27,55 are outside the scope of this study. Thermodynamic properties of relevant species have been determined as described in section 4 or taken from previous studies. $^{27,32,57-59}$ Table 3 lists important reactions in the pyrrole pyrolysis and oxidation subset, with detailed references to the source of selected rate coefficients and notes on minor modifications applied for improved agreement based on insights gained from the kinetic analysis below. Sources of rate constants for remaining reactions of the pyrrole/pyrrolenine subset not reported in Table 3 are provided in detail in the Supporting Information.

6. RESULTS AND DISCUSSION

Results from model simulations are compared to the new JSR data discussed in section 3 as well as other targets from the literature. ^{29,35,36,39} Kinetic analyses and discussion highlight relevant reaction pathways for both pyrolysis (section 6.1) and oxidation conditions (section 6.2).

6.1. Pyrolysis. Figure 1 compares model predictions to the newly acquired pyrolysis data in a JSR, operating at p=107 kPa and $\tau=2.0$ s in the temperature range T=900-1200 K. Good agreement is observed for pyrrole conversion, in particular concerning the temperature of onset of reactivity. Pyrrole consumption is slightly underestimated for T>1100 K. Molecular nitrogen is mainly converted into HCN, CH₃CN, and C₃H₅CN isomers, i.e., *cis-* and *trans-*crotonitrile and allyl cyanide.

Figure 2 reports results from a rate of production analysis carried out at $T=1100~\rm K$ for the JSR experiments of Figure 1. Pyrrole is largely consumed by the isomerization to pyrrolenine (reaction R1), followed by the isomerization to allenic imine (reaction R5). This latter step occurs through a 1,4-H migration of the biradical intermediate, resulting from the ring-opening reaction of pyrrolenine (Figure 3), as discussed by Martoprawiro et al. 32

The same intermediate leads to the formation of allyl cyanide and crotonitrile tautomers from pyrrolenine (reactions R3 and R4). For these pathways, we adopted the high-pressure limit rate coefficients from Martoprawiro et al.³² as a result of the lack of a systematic investigation of the pressure dependence in the literature. Pyrrolenine preferentially forms *cis*-crotonitrile (reaction R4) that further isomerizes to allyl cyanide (reaction R9) or tautomerizes to *trans*-crotonitrile (reaction R10), as shown in Figure 3.

H-abstractions by \dot{H} and $\dot{C}H_2CN$ from C_3H_5CN isomers (reactions R21, R23, R28, and R30) lead to the formation of the cyanoallyl radical ($\dot{a}\dot{C}_3H_4CN$) or to its non-allylic isomer ($\dot{c}\dot{C}_3H_4CN$) as well as H_2 and acetonitrile ($\dot{C}H_3CN$). $\dot{c}\dot{C}_3H_4CN$ completely decomposes to methyl radical ($\dot{C}H_3$) and cyanoacetylene (\dot{C}_3HN) through reaction R45. $\dot{a}\dot{C}_3H_4CN$ can further isomerize to another non-allylic isomer (\dot{C}_3H_4CN) through a 1,3-H migration (reaction R42). This latter intermediate undergoes a β -scission reaction (reaction R46), forming acetylene and cyanomethyl radical ($\dot{C}H_2CN$). This pathway constitutes a major source of $\dot{C}H_2CN$, together with the unimolecular decomposition reaction and the H-addition/decomposition reaction of allyl cyanide, forming vinyl radical (\dot{C}_2H_3) and ethylene (\dot{C}_2H_4). The reaction pathways discussed

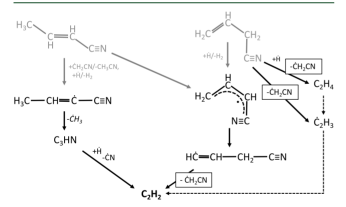


Figure 4. Pathways leading to the formation of acetylene and the cyanomethyl radical in pyrrole pyrolysis.

above ultimately lead to the formation of acetylene, as summarized in Figure 4. The cyanomethyl radical ($\dot{C}H_2CN$) produced by the decomposition channels depicted in Figure 4 is transformed to acetonitrile through H-abstraction reactions (reactions R16, R23, R30, R37, etc.).

Allenic imine is entirely converted into propyne and hydrogen cyanide (HCN) through reaction R6. For this channel, we adopted the rate constant suggested by Martoprawiro et al., with some correction, as reported in Table 3, to better match C_3H_4 -p and HCN profiles in pyrolysis and oxidation experiments. Our modifications are within the uncertainties discussed in ref 32, i.e., 1-2 kcal/mol in single-point energy calculations.

Allene is formed by the isomerization reaction C_3H_4 -p \leftrightarrow C_3H_4 -a and consumed by molecular growth pathways, leading to

cyclopentadiene $(C_2H_2 + C_3H_4 - a \leftrightarrow C_5H_6)$ and benzene (\dot{C}_3H_3) + C_3H_4 -a \leftrightarrow C_6H_6 + \dot{H}). C_4H_x represents the sum of 1,2butadiene (C_4H_6) and but-1-en-3-yne (C_4H_4) , whose peaks were too close to be distinguished in the experimental measurements. Propyne and allene reactions with the methyl radical produce butadiene ($\dot{C}H_3 + C_3H_4$ -p $\leftrightarrow \dot{H} + C_4H_6$ and $CH_3 + C_3H_4$ -a \leftrightarrow H + C_4H_6), while self-recombination reactions produce but-1-en-3-yne $(C_3H_4-p+C_3H_4-p\rightarrow C_2H_4+C_4H_4)$ and C_3H_4 -a + C_3H_4 -a $\rightarrow C_2H_4 + C_4H_4$). H-abstraction reactions by H and CH3 also contribute to fuel consumption, forming the resonance-stabilized pyrrolyl radical. On the basis of the dissociation energy of the N-H bond (BDE_{298 K} = 96.1 kcal/ mol), for H-abstraction reactions leading to the pyrrolyl radical, we adopted the values for the H-abstractions of a tertiary H atom according to the generalized approach by Ranzi et al.⁶⁹ The pyrrolyl radical entirely isomerizes to cC₃H₄CN (reaction R43), further contributing to acetylene and CH₂CN production.

Figure 5 compares model results to the single-pulse shocktube data by Mackie et al.²⁹ Fuel conversion profiles are wellcaptured by the model for both the 700 and 5000 ppm cases.

HCN that was overestimated in JSR experiments (Figure 1) is now underestimated by a similar extent, i.e., factor of \sim 2. The decomposition of allenic imine is still the major source of HCN (Figure 6), thus preventing any optimization of the rate coefficients for this reaction in one direction or the other.

To further highlight the need for reconciling model and experiments for pyrrole pyrolysis and to support the modification of the rate constants for reaction R6, we performed a sensitivity analysis to highlight dominant reaction channels in both shock-tube (ST) experiments and our new ISR measurements. For the ST experiments, we performed a sensitivity analysis at T = 1500 K, corresponding to 80% fuel conversion. A similar conversion is obtained in the JSR experiments at T =1200 K. Figure 7 shows the sensitivity coefficients of the most sensitive reactions controlling pyrrole consumption (top panel) and HCN formation (bottom panel). Concerning ISR simulations, pyrrole consumption is dominated by the isomerization reactions of pyrrolenine to cis-crotonitrile and allyl cyanide (reactions R3 and R4), together with pyrrole isomerization to allenic imine (reaction R2). The same subset of reactions controls fuel conversion in shock-tube pyrolysis. To a minor extent, H-abstractions by H and the decomposition of allenic imine to HCN and propyne (reaction R6) also influence pyrrole consumption. HCN formation is dominated by the reaction series R2 > R6 depicted in Figure 6 in both JSR and ST cases. However, for the JSR case, HCN formation is highly sensitive to successive isomerization and decomposition reactions of pyrrolenine (reactions R4 and R7) and mostly to H ipso-addition reaction of acetonitrile, $H + CH_3CN \leftrightarrow CH_3 +$ HCN. Overall, despite quite different operating conditions, no reactions with opposite effects on model predictions for HCN emerged from this analysis, hampering improved agreement for both JSR and ST pyrolysis data.

The model is once again able to properly predict formation and consumption of C_3H_5CN isomers, which in Figure 5 are resolved in crotonitrile isomers and allyl cyanide. Acetonitrile formation is very well-captured; however, the model underpredicts its consumption at the higher temperature end of both the shock-tube ($T > 1600~\rm K$) and JSR ($T > 1150~\rm K$) experiments. As previously discussed, we adopted rate coefficients for the major consumption pathways of CH₃CN from the recent study by Alzueta et al. ⁵⁶ C_2H_2 is also formed in this case by the decomposition reaction of $c\dot{C}_3H4CN$ (Figure 4),

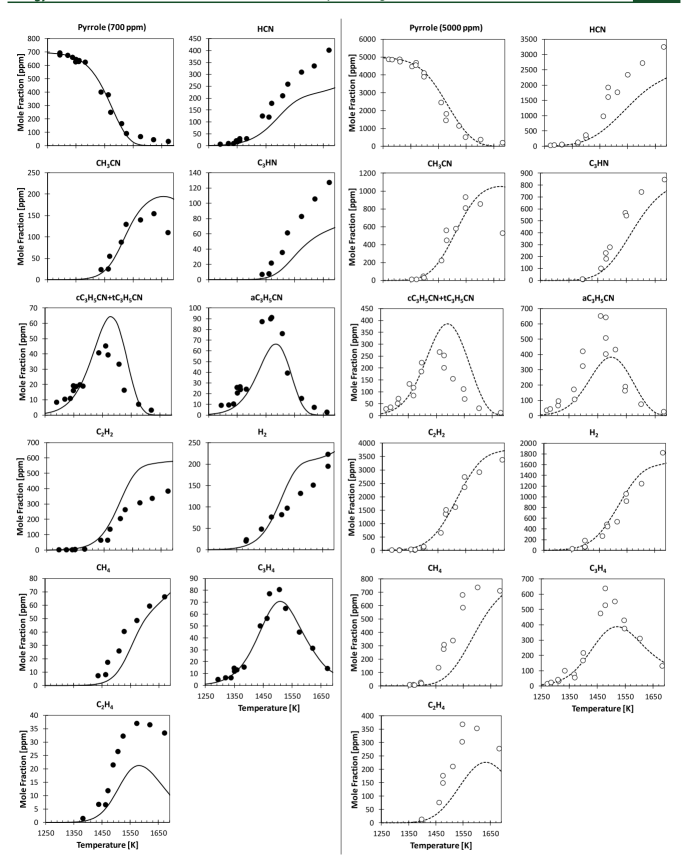


Figure 5. Pyrolysis of pyrrole (700 ppm, left panel, and 5000 ppm, right panel, in argon) in a single-pulse shock tube at p = 13 atm and $\tau = 550 \,\mu s$. Comparison between experimental (symbols)²⁹ and predicted (lines) mole fraction profiles of intermediate and product species.

for which no theoretical estimates exist. We estimated a value by analogy starting from the decomposition of the vinyl radical of 1-

butene ($\dot{C}_4H_71-1\leftrightarrow C_2H_2+\dot{C}_2H_5$) and increasing its activation energy by 6 kcal/mol to better match acetylene and acetonitrile

$$HC = C = CH_2$$

$$HC \longrightarrow HC$$

$$NH$$

$$C_3H_4-p + HCN$$

Figure 6. Pyrrole isomerization to allenic imine (reaction R2) HNCPROP and successive unimolecular decomposition to hydrogen cyanide and propyne (reaction R6).

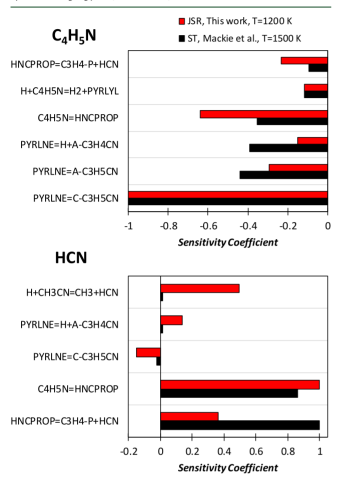


Figure 7. Sensitivity analysis of fuel consumption (top panel) and HCN formation (bottom panel) to model rate constants for the JSR case (Figure 1; T = 1200 K) and the ST case (Figure 5; 5000 ppm of pyrrole and T = 1500 K).

profiles in both pyrolysis and oxidation experiments. Methane and hydrogen are produced by H-abstraction reactions from the fuel, crotonitrile isomers, and propyne. Good agreement is observed for H_2 , while methane is slightly underestimated in both Figures 3 and 5.

6.2. Oxidation. Experimental measurements and model predictions for the oxidation of pyrrole in a JSR are reported in Figure 8 for three equivalence ratios $\varphi=0.5$, 1.0, and 2.0. Fuel consumption is correctly predicted by the model for the $\varphi=1.0$ case. For the leanest case ($\varphi=0.5$), the model strongly underpredicts fuel reactivity, despite correctly capturing the onset of conversion. In the richest case ($\varphi=2.0$), the model captures the start of reactivity, slightly overpredicting pyrrole consumption for T<1050 K and slightly underpredicting the complete conversion at higher temperatures.

Major product formation, such as CO and CO_2 , and O_2 consumption are correctly reproduced, increasing confidence in the model's validity. The effect of the equivalence ratio is

qualitatively captured for all of the intermediates, but some major deviation in quantitative terms is observed. HCN formation is underestimated by ~35% for the lean and stoichiometric cases but is overestimated for the rich case. Good agreement is observed in the case of CH₃CN, with the exception of the rich case, where the model predicts an excessive consumption, underestimating the peak concentration by a factor of ~2. Similar deviations are observed for acetylene, while ethylene peaks are quite nicely reproduced. The acrylonitrile (CH₂CHCN) peak is underestimated at every equivalence ratio, and a delayed formation is observed at $\varphi = 0.5$ because of the underestimation of fuel reactivity in such conditions. Despite methane only being detected for the rich case, the model predicts its formation in significant quantities also for the lean and stoichiometric cases (i.e., 80-200 ppm). Ethane, propene, and propyne are only detected in very low quantities (i.e., 10-20ppm) and only at $\varphi = 2.0$ in the experimental measurements. The model generally underestimates these minor products.

Figure 9 shows the rate of production analysis at T = 950 K for the stoichiometric case of Figure 8 ($\varphi = 1.0$). Pyrrole is consumed through H-abstraction reactions by OH, O, H, and $H\dot{O}_2$ to form the pyrrolyl radical that is largely converted back to pyrrole through the reverse reaction $C_4H_5N + O_2 \leftrightarrow PYRLYL +$ HO_2 (reaction R19). This reaction is highly endothermic (E_a = 46.9 kcal/mol) and most likely proceeds in the backward direction at conditions where the HO2 concentration is high (e.g., T < 1000 K), as observed in the same system for other fuels 70 This channel contributes to 41% of pyrrolyl consumption, while its decomposition to cC_3H_4CN (reaction R43) accounts for 57% of the total flux. At such low temperatures, isomerization of cC₃H₄CN to aC₃H₄CN (cyano allyl radical) through reaction R42 dominates over decomposition pathways, forming acetylene and CH2CN (reaction R46). aC3H4CN reacts with HO₂ in reaction R48, releasing OH and forming a cyano alkoxy radical (C_4H_4NO) that decomposes through β -scission to form a cyano radical (CN) and unsaturated products, such as acrolein (C₂H₃CHO), or acetylene and formaldehyde (reactions R50 and R51). To a lower extent, a recombination/disproportionation reaction with HO2 can also occur, forming allyl cyanide and O₂ (reaction R26), activating the isomerization reactions of C₃H₅CN isomers, until H-abstraction reactions by O form C₃H₄CN (e.g., reaction R41) that further decomposes to the methyl radical and cyanoacetylene through β -scission (reaction R45). The rate coefficients for these pathways were estimated on the basis of analogy with the allyl radical $(C_3H_5-a)/HO_2$ kinetics.

The same cyano alkoxy radical can be formed directly from the interaction of $c\dot{C}_3H_4CN$ with O_2 (reaction R42), releasing \ddot{O} atoms. The rate coefficients for this channel have been adopted in analogy with $\dot{C}_2H_3+O_2$.

HCN is one of the main intermediates in pyrrole pyrolysis and oxidation. Different from the pyrolysis cases discussed above, where the main source was the decomposition of the allenic imine HNCPROP, at $T=950~\rm K$ and in the presence of oxygen, hydrogen cyanide is mostly formed by the reaction of the isocyanate radical (NCO) with acetylene, forming HCCO ($C_2H_2 + \rm NCO \leftrightarrow \rm HCCO + \rm HCN$). Acetonitrile is mainly formed by H-abstraction reactions of $\rm CH_2CN$ on succinonitrile ($\rm C_4H_4N_2$, $\rm CN-CH_2-CH_2-CN$). Succinonitrile (butanedinitrile) is formed by the self-recombination of $\rm CH_2CN$ (reaction R52). H-abstraction reactions on $\rm C_4H_4N_2$, for example by $\rm CH_2CN$ (reaction R54), produce a resonance-stabilized $\rm C_4H_3N_2$ radical, whose decomposition reaction justifies the

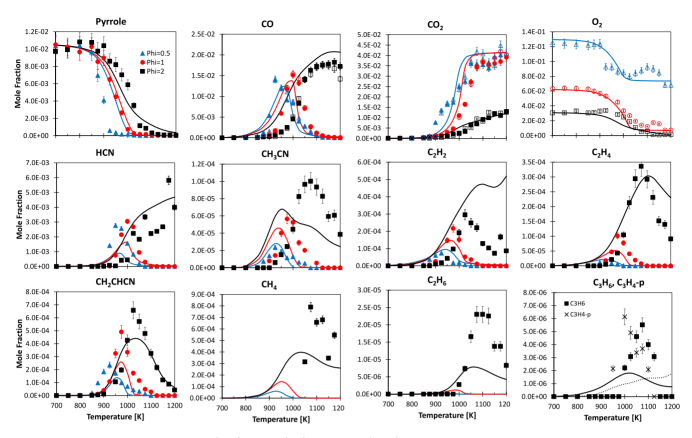


Figure 8. Pyrrole oxidation in JSR at φ = 0.5 (blue), φ = 1.0 (red), and φ = 2.0 (black), p = 107 kPa, and τ = 2.0 s. Comparison between experimental (symbols) and predicted (lines) fuel conversion and mole fraction profiles for intermediate and product species. (Open symbols) GC—TCD—FID with a Carbosphere-packed column and (full symbols) GC—methanizer—FID with a Q-Bond capillary column.

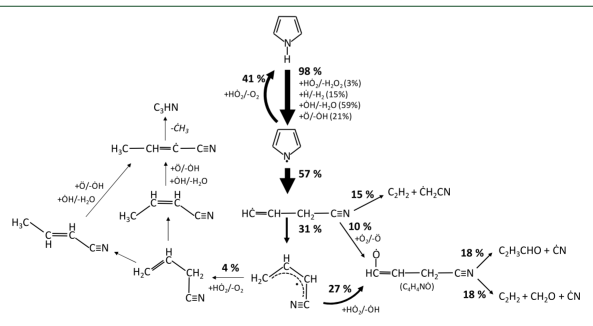


Figure 9. Rate of production analysis at T=950 K for pyrrole oxidation in JSR at $\varphi=1.0$, p=107 kPa, and $\tau=2.0$ s. Arrow width qualitatively represents the importance of each reactive flux. Pathways with a flux going from or to an intermediate of <1% have been disregarded for clarity.

formation of acrylonitrile ($\dot{C}_4H_3N \leftrightarrow CH_2CHCN + \dot{C}N$, reaction R55) as illustrated in Figure 10.

For the succinonitrile subset, we entirely adopted the values proposed by Sendt et al. sand assigned H-abstraction rate coefficients based on analogy with $R + C_4H_8-1 = RH + C_4H_71-3$, accounting for the availability of four H atoms to form a

resonance-stabilized radical rather than two, as in the case of 1-butene.

To investigate possible reasons for model shortcomings in predicting the effect of the equivalence ratio, we performed a sensitivity analysis of rate constants to fuel consumption at T = 950 K for the three mixtures experimentally investigated. As

$$\dot{\text{CH}_2\text{CN}} + \dot{\text{CH}_2\text{CN}} \longrightarrow \text{N=C} \\ + \dot{\text{CH}_2\text{CN}} \downarrow - \text{CH}_3\text{CN}$$

$$\text{CH}_2\text{CHCN} + \dot{\text{CN}} \longrightarrow \text{C=N}$$

$$\text{CH}_2\text{CHCN} + \dot{\text{CN}} \longrightarrow \text{C=N}$$

Figure 10. Succinonitrile $(C_4H_4N_2)$ formation and consumption pathways in pyrrole oxidation.

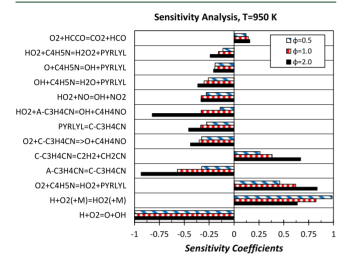


Figure 11. Sensitivity analysis of fuel consumption to model rate constants at T = 950 K for the $\varphi = 0.5$, 1.0, and 2.0 mixtures. Sensitivity coefficients are normalized over that of $\dot{\rm H} + {\rm O_2} \leftrightarrow \ddot{\rm O} + \dot{\rm O} \rm H$. A negative sensitivity coefficient stands for a reaction increasing reactivity (i.e., contributing to fuel consumption) and vice versa.

expected, results provided in Figure 11 do not highlight any possible modification capable of decreasing the reactivity of the

rich mixture ($\varphi=2.0$) while simultaneously increasing that of the lean mixture ($\varphi=0.5$). Sensitivity coefficients have been normalized over that of the most sensitive reaction, $\dot{H}+O_2\leftrightarrow\ddot{O}+\dot{O}H$.

The competition between the branching reaction $\dot{H} + O_2 \leftrightarrow$ \ddot{O} + $\dot{O}H$ and the third-body recombination \dot{H} + O_2 (+M) \leftrightarrow HO₂ (+M) decreasing system reactivity increases for leaner mixtures. The first fuel-specific reaction appearing within the most sensitive reactions is the H-abstraction by O2 on pyrrole $(C_4H_5N + O_2 \leftrightarrow PYRLYL + HO_2$, reaction R19). As already highlighted in the above discussion, under these conditions, the reaction proceeds backward, consuming pyrrolyl and HO₂ and, thus, decreasing fuel consumption. This negative effect is explained by the fact that, in addition to partly restoring the concentration of fuel, produced O2 is then converted through the recombination reaction $\dot{H} + O_2(+M) \leftrightarrow H\dot{O}_2(+M)$ or $O_2 +$ $\dot{HCO} \leftrightarrow \dot{HO_2} + \dot{CO}$, forming once again $\dot{HO_2}$. Reactions belonging to the pyrrole subset, such as $C_4H_5N + HO_2 \leftrightarrow$ PYRLYL + H_2O_2 (reaction R18) and $aC_3H_4CN + HO_2 \leftrightarrow OH +$ \dot{C}_4H_4NO (reaction R48), convert this $H\dot{O}_2$ into more reactive radicals, thus increasing the reactivity. Another key reaction consuming HO_2 is $NO + HO_2 \leftrightarrow OH + NO_2$, for which we adopted the value of Howard et al., 71 as already discussed by Song et al. 27 As expected, H-abstraction reactions by OH and O also have a positive impact on reactivity. The isomerization of the resonance-stabilized pyrrolyl radical to the vinylic radical cC₃H₄CN (reaction R43) clearly favors pyrrole conversion. The important role of C₄H₄N isomer chemistry and, in particular, that of cC₃H₄CN and aC₃H₄CN emerges clearly from the competition between the isomerization (aC₃H₄CN ↔ cC₃H₄CN₁ reaction R42 backward) and oxidation (O₂ + $c\dot{C}_3H_4CN \leftrightarrow \ddot{O} + \dot{C}_4H_4NO$, reaction R49) reactions increasing reactivity, with the decomposition reaction ($c\dot{C}_3H_4CN \leftrightarrow C_2H_2$ + $\dot{C}H_2CN$, reaction R46) consuming $c\dot{C}_3H_4CN$. The oxidation of $a\dot{C}_3H_4CN$ ($a\dot{C}_3H_4CN + H\dot{O}_2 \leftrightarrow \dot{O}H + \dot{C}_4H_4NO$, reaction

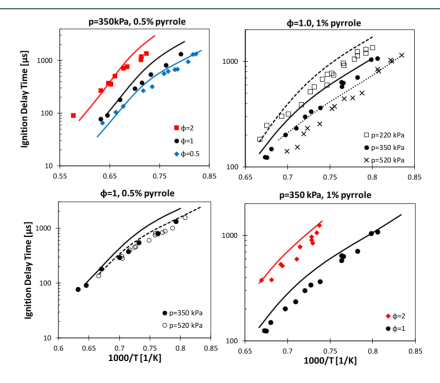


Figure 12. Shock-tube experimental (symbols)³⁵ and simulated (lines) ignition delay times for highly diluted mixtures (>92.7 mol %) of pyrrole (0.5 and 1%) in O_2 and argon at high temperatures.

R48) also contributes to increase the overall reactivity, forming two reactive radicals from relatively stable radicals.

MacNamara and Simmie³⁵ studied pyrrole autoignition in a low-pressure shock tube. Figure 12 compares ignition delay time measurements for different pyrrole/oxygen/argon mixtures with model predictions.

Overall, simulated ignition delay times capture the effect of the pressure, equivalence ratio, and fuel concentration on the ignition propensity of pyrrole. Maximum deviations are as large as a factor of 1.7 in the worst cases. A sensitivity analysis has been

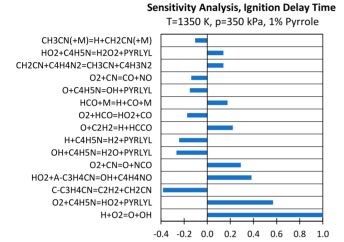


Figure 13. Sensitivity analysis of ignition delay times to rate constants for a stoichiometric 1% pyrrole/oxygen/argon mixture at T=1350 K and p=350 kPa. A positive sensitivity coefficient stands for a reaction promoting ignition and vice versa. Sensitivity coefficients have been normalized over that of the dominating reaction $\dot{\rm H}+{\rm O}_2\leftrightarrow \ddot{\rm O}+\dot{\rm O}{\rm H}$.

Sensitivity Coefficient

carried out for the case of 1% pyrrole, p = 350 kPa, $\varphi = 1$, and T = 1350 K (Figure 13).

The H-abstraction reaction $C_4H_5N + O_2 \leftrightarrow PYRLYL + HO_2$ at this temperature condition proceeds in the forward direction, thus promoting fuel consumption and thereof ignition. Indeed, HO_2 undergoes self-recombination to form H_2O_2 that is rapidly decomposed to form two hydroxyl radicals $[H\dot{O}_2 + H\dot{O}_2 (+M)]$ \leftrightarrow H₂O₂ + O₂ (+M) and H₂O₂ (+M) \leftrightarrow 2 \dot{O} H (+M)] or recombines with H to directly produce two OH radicals (H + $H\dot{O}_2 \leftrightarrow 2\dot{O}H$), promoting ignition. One of the major sources of H atoms is, together with formyl radical decomposition, the addition/elimination reaction involving cyanoacetylene (H + $C_3HN \leftrightarrow C_2H_2 + CN$, reaction R47 backward).⁷² Other Habstraction reactions (\dot{H} + C_4H_5N , $\dot{O}H$ + C_4H_5N , and \ddot{O} + C₄H₅N) producing the pyrrolyl radical show a negative sensitivity coefficient, despite consuming the fuel. This is justified by the fact that pyrrolyl almost entirely isomerizes to cC₃H₄CN that, at such high-temperature conditions, decomposes to acetylene and CH2CN that is resonance-stabilized and acts as a sink of \dot{H} atoms, strongly inhibiting the occurrence of the branching reaction $\dot{H} + O_2 \leftrightarrow \ddot{O} + \dot{O}H$ that dominates hightemperature ignition. As shown in Figure 14, only a minor amount of cC_3H_4CN isomerizes to aC_3H_4CN , whose oxidation $(a\dot{C}_3H_4CN + H\dot{O}_2 \leftrightarrow \dot{O}H + \dot{C}_4H_4NO)$ promotes ignition.

To assess the governing chemistry of NO_x formation from fuel-bound nitrogen, Lumbreras et al. ³⁶ investigated the flow reactor oxidation of pyrrole in experiments with and without NO addition. Figure 15 compares model results to experimental data for the cases without NO (left column) and with NO (right column). The model qualitatively captures both the effect of the equivalence ratio on pyrrole oxidation with and without NO addition. However, except for the CO_2 profiles, for which the reactivity is well-captured, large quantitative deviations (factor of \sim 3 in the worst cases) exist for the peak concentrations of measured species, such as CO and HCN. For example, the model fails to predict the early formation of HCN in the leanest case ($\varphi = 0.05$), both with and without NO. Moreover, predicted CO formation is delayed, while NO consumption is anticipated. Despite a 5% uncertainty in the measurements

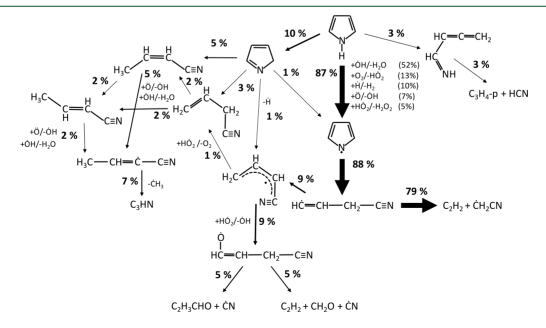


Figure 14. Rate of production analysis for a 1% pyrrole/ O_2 /argon mixture at T = 1350 K, $\varphi = 1.0$, p = 350 kPa, and 20% fuel conversion ($\tau = 3.2 \times 10^{-4}$ s). Arrow width qualitatively represent the importance of each reactive flux. Pathways with a flux going from or to an intermediate of <1% have been disregarded for clarity.

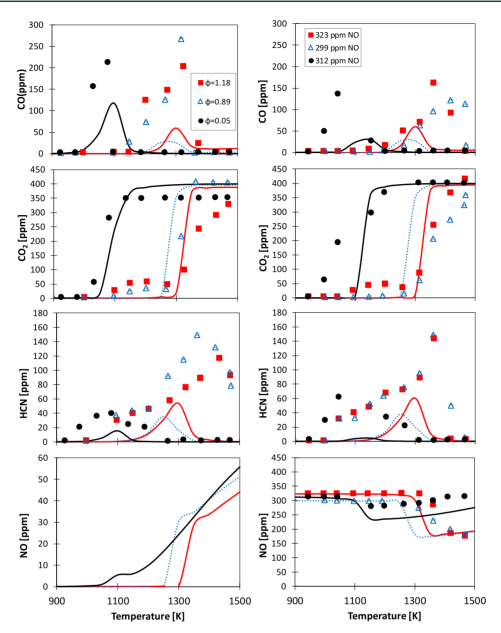


Figure 15. Speciation profiles from pyrrole (100 ppm) oxidation as a function of the temperature for different air excess ratios, without (left column) and with (right column) ~300 ppm of NO addition. Residence time $\tau = 210/T$ s. Comparison between experimental data (symbols)³⁶ and model predictions (lines).

declared by the authors, atomic balances highlight that some species that may be formed in significant quantities were not measured (e.g., C₂H₂, HNCO, C₂H₄, C₃HN, and CH₃CN), thus preventing any quantitative statement on model performances. Moreover, very little impact of model parameters was found when attempting to improve model performances. However, these data, together with those by Yamamoto et al.³⁹ (Figure 17), are valuable to further assess the importance of formation and consumption pathways of HCN and NO, as discussed in the following. Despite not being measured, in Figure 15, we also report the temperature dependences of the NO mole fraction. As expected, the leanest and most reactive mixture shows the earliest and highest production of NO. The change of slope on the higher temperature end is due to the conversion of NO to NO_2 through $NO + O (+M) \leftrightarrow NO_2 (+M)$ (reaction R56), facilitated by the branching reaction $\dot{H} + O_2 \leftrightarrow \ddot{O} + \dot{O}H$ providing high amounts of oxygen atoms.

Figure 16 shows the main formation and consumption pathways forming HCN and NO, also focusing on the conversion of major nitrogen-containing intermediates, such as HCN, CH₃CN, and C₃H₄CN, to NO. In the case without NO addition (left column of Figure 15), HCN is mostly formed by the successive decomposition steps of C_3H_4CN , undergoing β scission to cyanoacetylene (C3HN) that is converted to the cyanomethylene radical (HCCN) through reaction R57 and then to HCN through reaction R58. A secondary pathway, of similar importance in the case of NO addition, is the direct decomposition of allenic imine HNCPROP to HCN and propyne. The channel of lower importance involves acetonitrile (CH₃CN), forming the cyanomethyl radical (CH₂CN) through reaction R59. CH₂CN is subsequently converted to formyl cyanide (OCHCN) via reaction R60. Formyl cyanide then eliminates CO to yield HCN (reaction R61). OH is added to HCN and eliminates H to form isocyanic acid (HNCO, reaction

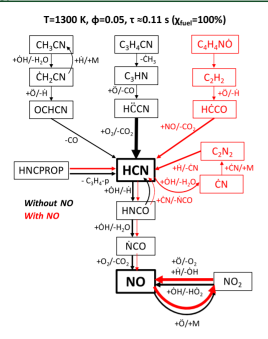


Figure 16. Main HCN and NO formation pathways for $\varphi=0.05$ mixtures of Figure 15 with (red) and without (black) NO addition.

R62) that through H-abstraction by $\dot{O}H$ in reaction R64 forms the isocyanato radical ($\dot{N}CO$). $\dot{N}CO$ reacts with O_2 , which is present in large excess, to form NO and CO_2 (reaction R65). NO is further converted to NO_2 through the third-body reaction $NO + \ddot{O}(+M) \leftrightarrow NO_2(+M)$ and then regenerated from NO_2 by means of addition/elimination reactions.

In the case of NO addition (right column of Figure 15), different pathways lead to the formation of HCN. C₄H₄NO decomposition to acetylene according to the lumped reaction R51 ($C_4H_4NO \rightarrow C_2H_2 + CH_2O + CN$) yields the ethynyloxy radical (HCCO) through $C_2H_2 + \ddot{O} \leftrightarrow \dot{H} + HCCO$. HCCO is transformed to HCN by means of reaction R68: HCCO + NO \leftrightarrow HCN + CO₂. HCN then undergoes H-abstraction by OH or OH addition (reactions R62 and R63). The first channel leads to the formation of the cyano radical (CN) that after selfrecombination (reaction R66) produces cyanogen (C_2N_2) . Cyanogen is converted back to HCN by H addition in reaction R67, producing once again the cyano radical. The H-abstraction pathway leads to the same formation route of NO previously discussed for the case without NO. The presence of NO in the feed triggers NO conversion to NO_2 that is once again converted back to produce NO. From this reaction cycle, it is possible to explain the successive consumption and formation of NO observed in the right column of Figure 15.

Figure 17 compares model predictions with the flow reactor measurements by Yamamoto et al. ³⁹ The largest deviations are observed for the lean cases (20 000 ppm of O_2 , i.e., $\varphi = 0.06$ without considering the H_2O content) when 8% H_2O is added to the system. Indeed, the model overestimates the reactivity by ~50 K. The conditions of these experiments are comparable to those of Lumbreras et al., ³⁶ where the model showed a similar deviation in terms of the temperature but in the opposite direction. Therefore, we consider model performances to be in reasonable agreement with the experimental data because no modification to the kinetics would lead to improvement for both sets of data. It should also be noted that H_2O does not play any role in terms of modifications to the radical pool, as evident from the inhibiting effect on reactivity rather than an increase that

would be expected from, for example, higher yields of OH. However, the collisional efficiency of water (i.e., $\sim 6-12$ times higher than that of N₂⁵⁵) strongly promotes the chain propagation reaction $[\dot{H} + O_2 (+M) \leftrightarrow H\dot{O}_2 (+M)]$ and termination reaction $[H\dot{O}_2 + H\dot{O}_2 (+M) \leftrightarrow H_2O_2 + O_2 (+M)]$ over chain branching (i.e., $\dot{H} + O_2 \leftrightarrow \ddot{O} + \dot{O}H$) in the H_2/O_2 subset in the temperature window where the onset of fuel conversion is observed (T = 1000-1100 K), thus reducing the overall reactivity. Model performances would benefit from a better assessment of pressure dependence of primary fuel reactions largely discussed in section 6.1 and from a more rigorous implementation of collisional efficiencies in the current formalism for pressure-dependent rate expressions. Model results agree better for the remaining cases, i.e., 20 000 ppm of O_2 with 3% H_2O addition and 6400 ppm of O_2 ($\varphi = 0.2$) with 8% H₂O addition. Notably, the model correctly predicts fuel conversion for short residence times ($\tau = 136/T$ s), even for the leanest case with 8% H₂O addition. In general, the model captures very well the dependence of HCN and NO formation trends and the relative magnitude of their peaks on operating conditions.

7. CONCLUSION

In this work, the pyrolysis and oxidation of pyrrole were experimentally investigated in an atmospheric pressure JSR, significantly extending the validation targets available for pyrrole kinetic model validation purposes. A preliminary model based on previous research efforts and analogy with kinetic subsets already implemented in the CRECK kinetic framework is presented, showing generally good agreement after some adjustment on available kinetic parameters, within their uncertainties. To the knowledge of the authors, this pyrrole model is the first model comprehensively validated in the literature and allows for the inclusion of pyrrole as a representative nitrogen-containing component in more complex surrogate models of pyrolysis bio-oils. 11 Moving from a detailed kinetic analysis aimed at highlighting reasons for model deviations and existing shortcomings, we believe that theoretical, experimental, and kinetic modeling efforts should be devoted to the following: (1) There should be better assessment of temperature- and pressure-dependent kinetics of isomerization channels to pyrrolenine, crotonitrile isomers, and allyl cyanide that initiate and dominate pyrrole decomposition kinetics. Available theoretical information on the potential energy surfaces are indeed quite accurate and detailed to be reproduced with current state-of-the-art electronic structure methods and multi-well master equation solvers. (2) Habstraction reactions consuming pyrrole, pyrrolenine, and their C₃H₅CN isomers are currently largely based on analogy rules with systems, leading to resonance-stabilized radicals (e.g., allylic type radical) not containing nitrogen, whose influence is expected to be significant. Such rate coefficients should be theoretically re-evaluated with more accuracy. (3) Secondary reactivity of derived radicals of such an unsaturated system can be complicated by resonance stabilization. For example, recombination reactions of the pyrrolyl radical and HO₂ have been included in our kinetic model, largely on the basis of analogy with cyclopentadienyl chemistry and/or allyl radical chemistry. However, no impact of such pathways was observed at the conditions where experimental data are available at present. Clearly, this observation poses some question on the general validity of the analogy rules adopted in this work and in previous kinetic modeling studies. Similar observations apply to

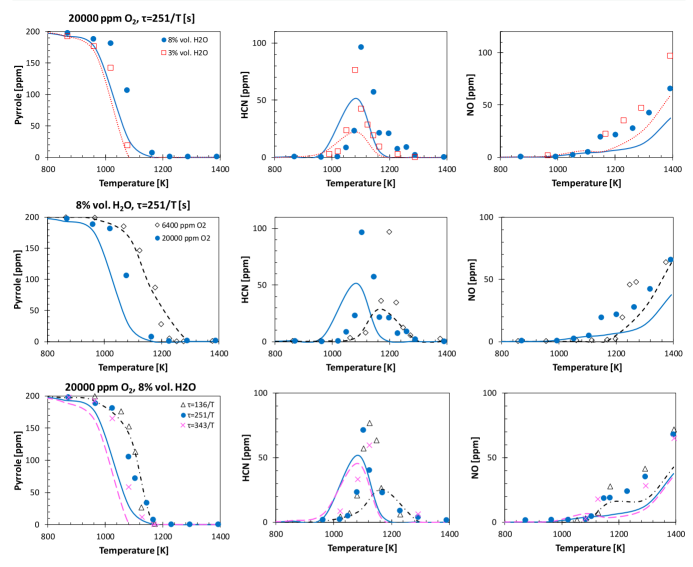


Figure 17. Comparison of experimental data to model predictions for pyrrole, HCN, and NO under different conditions of H₂O concentration (top row), O₂ concentration (central row), and residence time (bottom row). Symbols are experimental data, ³⁹ and lines are model predictions.

the interactions between the cyano allyl radical ($a\dot{C}_3H_4CN$) and its vinyl isomers ($c\dot{C}_3H_4CN$ and \dot{C}_3H_4CN) with O_2 and HO_{24} where further theoretical investigations would be beneficial to increase model predictive capabilities. (4) The deficit in N atoms highlighted by the atomic balances should be investigated thoroughly. A valuable perspective would be to perform such a study with an advanced diagnostic tool, like time-of-flight mass spectrometry with more direct sampling using, for example, a molecular beam to minimize the loss of species during the sampling. This technique would also allow for the detection of species having relatively low stabilities, like pyrrolyl and other resonance-stabilized radicals, which play a central role in the gasphase chemistry of pyrrole. (5) Experimental data on pyrrole combustion at a higher pressure would be useful to extend the confidence of the proposed model at conditions closer to that of real combustion devices (e.g., turbines). In particular, ignition delay time data of fuel/air mixtures in a high-pressure shock tube would be useful, although the low volatility of pyrrole might inhibit tests in non-diluted mixtures. In addition, flame data are only available at a very low pressure (i.e., 0.032 atm³⁸), where pressure-dependent kinetics are extremely important. Indeed, such targets have not been reported in the validation of the

present kinetic model, where we only adopt high-pressure limit rate constants. However, the understanding of N fuel chemistry would benefit from laminar flame speed measurements at atmospheric pressure. (6) On the basis of past tests of bio-oil use at the industrial scale, 6 where pilot flames fed with conventional hydrocarbons are often used, it would be interesting to assess the kinetic effects of hydrocarbon fuels doped with pyrrole. It is important to note that, for this type of test, the possible importance of cross chemical interactions of resonancestabilized radicals from pyrrole decomposition and oxidation chemistry with other components in the mixtures should be assessed. In addition to the impact on macroscopic reactivity targets (e.g., ignition delay times and laminar flame speed), speciation measurements on NO_x formation and polycyclic aromatic hydrocarbon (PAH) growth should be performed. Indeed, as highlighted in this study, acetylene and other unsaturated hydrocarbons are produced in large quantities and may significantly contribute to molecular growth kinetics, posing some question on the pollution potential of bio-oils not only in terms of nitrogen oxides but also for PAH and particulate matter formation.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.energyfuels.0c03874.

Experimental data from JSR measurements (XLSX)

Thermodynamic properties of relevant species, SMILES identifiers, and structural data from theoretical calculations (PDF)

Model 1 kinetics: pyrrole pyrolysis and oxidation (189 species, 2888 reactions, core chemistry + NO_x subset + pyrrole subset) (TXT)

Model 1 thermo: pyrrole pyrolysis and oxidation (189 species, 2888 reactions, core chemistry + NO_x subset + pyrrole subset) (TXT)

Model 2 kinetics: high-temperature CRECK model with NO_x and pyrrole chemistry (500 species and 17 472 reactions) containing the kinetic subset of reference species of possible interest for bio-oil surrogates (e.g., aldehydes, alcohols, organic acids, phenolic species and oxygenated aromatics with multiple substitutions, and methyl esters) (TXT)

Model 2 thermo: high-temperature CRECK model with NO_x and pyrrole chemistry (500 species and 17 472 reactions) containing the kinetic subset of reference species of possible interest for bio-oil surrogates (e.g., aldehydes, alcohols, organic acids, phenolic species and oxygenated aromatics with multiple substitutions, and methyl esters) (TXT)

Pyrrole pyrolysis and oxidation subset with detailed references on selected rate coefficients (TXT)

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Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors at Politecnico di Milano and CNRS Nancy acknowledge the financial support of the IMPROOF Project (H2020-IND-CE-2016-17/H2020-SPIRE-S016) within the European Union's Horizon 2020 Research and Innovation Program (Grant Agreement 723706).

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