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Pressure-dependent kinetic analysis of the N₂H₃ potential energy surface†

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The pressure-dependent reactions on the N_2H_3 potential energy surface (PES) have been investigated using CCSD(T)-F12/aug-cc-pVTZ-F12//B2PLYP-D3/aug-cc-pVTZ. This study expands the N_2H_3 PES beyond the previous literature by incorporating a newly identified isomer, NH_3N , along with additional bimolecular reaction channels associated with this isomer, namely $NNH + H_2$ and $H_2NN(S) + H$. Rate coefficients for all relevant pressure-dependent reactions, including well-skipping pathways, are predicted using a combination of *ab initio* transition state theory and master equation simulations. The dominant product of the $NH_2 + NH(T)$ recombination is $N_2H_2 + H$, while at high pressures and low temperatures, N_2H_3 formation becomes significant. Similarly, collisions involving $H_2NN(S) + H$ predominantly produce $N_2H_2 + H$. Secondary reactions such as $H_2NN(S) + H \rightleftharpoons NNH + H_2$ and $H_2NN(S) + H \rightleftharpoons NH_2 + NH(T)$ are found to play a significant role at high temperatures across all examined pressures, while $H_2NN(S) + H \rightleftharpoons NH_3N$ becomes prominent only at high pressures. Notably, none of these four $H_2NN(S)$ reactions have been included with pressure-dependent rate coefficients in previous NH_3 oxidation models. The rate coefficients reported here provide valuable insights for modeling the combustion of ammonia, hydrazine, and their derivatives in diverse environments.

1. Introduction

Nitrogen-based fuels, including ammonia (NH_3) , hydrazine (N_2H_4) , hydrazine derivatives such as monomethylhydrazine (MMH, CH_3NHNH_2) , and aqueous formulations, horm a versatile class of energy carriers. Hydrazine and its derivatives are widely used as satellite and spacecraft propellants, while ammonia receives an increasing interest as a potential energy storage vector. In the envisioned future nitrogen economy, ammonia is expected to play a key role. Despite differences in their specific applications, the chemical kinetic modeling of

Hydrazine, a principal component of diamine-based rocket propellants, stands out as an efficient mono- and bi-propellant. Paired with oxygen as an oxidizer, it is second only to hydrogen in terms of specific impulse—the thrust generated per unit mass of fuel over time. Hydrazine can be used as a monopropellant, typically undergoing decomposition using a catalyst, leading to autoignition and detonation. Huttermore, hydrazine and hydrazine-based fuels, including MMH, function as bipropellants when combined with nitrogen tetroxide (N_2O_4) as the oxidizer. These fuels are ubiquitous as thruster engine propellants for satellite and spacecraft attitude control and in-orbit maneuvers, where they are stored and used at a maximum pressure of ~ 25 bar. $^{12-14}$

Nitrogen-based alternative fuels are particularly attractive due to the abundance of molecular atmospheric nitrogen as a feedstock. Among them, ammonia is the simplest molecule and it shows significant promise as an energy storage vector. ^{1,6,15} The advantages of ammonia as a fuel include a relatively high power-to-fuel-to-power (PFP) efficiency, ⁷ a large-scale distribution infrastructure that is already in place, ^{16,17} a high octane rating, ¹⁸ and a narrow flammability range that makes it relatively safe in terms of explosion risks. Although kinetic modeling of NH₃ oxidation received significant attention lately, ^{19–35} there is still substantial disagreements on thermodynamic data and rate coefficients between different authors in this relatively small chemical system. ^{8,36–39}

these nitrogen-based fuels relies on a shared core chemistry and a set of fundamental reactions.⁸

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[†] Electronic supplementary information (ESI) available: Stationary points geometries, T1 diagnostic factors, and imaginary frequencies (Table S1). Torsional scan of N_2H_3 at a resolution of 10° at the B2PLYP-D3/aug-cc-pVTZ level (Fig. S1). Representation of the N_2H_3 PES along with energy values from previous studies (Fig. S2). NH₃N Thermodynamic properties, Chemkin format (Table S2). NH₃N Thermodynamic properties, Cantera format (Table S3). Computed rate coefficient, Chemkin format (Table S4). Computed rate coefficient, Cantera format (Table S5). 3D representations of species (Fig. S3–S7). Species internal coordinate comparisons (Tables S6–S10). An unsuccessful TS search for the $N_2H_3 \rightleftharpoons H2NN(S) + H$ reaction (Fig. S8). A scan of the N−−H distance in the $N_2H_3 \rightleftharpoons H2NN(S) + H$ reaction (Fig. S9). Estimated reaction rate uncertainty based on computed energies (Fig. S10). Comparison of experimental and computed frequencies (Table S11). See DOI: https://doi.org/10.1039/d4cp03837a

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Konnov and De Ruyck⁴⁰ suggested a chemical kinetic model for hydrazine decomposition, and Izato et al. 41 suggested a model for the initial hypergolic reaction of liquid hydrazine with nitrogen tetroxide mixtures. Hydrazine decomposition was also studied by Asatryan et al., 42 and its key pathways in hypergolic ignition with nitrogen dioxide and nitrogen tetroxide were suggested by Raghunath et al. 43 and by Daimon et al., 44 respectively. The thermal decomposition of hydrazine was studied using a reactive force field.45 Hydrazine decomposition on catalytic surfaces was also studied theoretically.46,47 Sun and Law48 developed a model for thermal decomposition of MMH, and Zhang et al.49 performed a theoretical analysis of the kinetics of barrierless recombination reactions in the MMH system as well as secondary channels.⁵⁰ Although chemical kinetic models were previously suggested to predict the reactivity of hydrazine and its derivative, the research on the combustion of these fuels has been based on incomplete kinetic mechanisms, which may not fully capture all the relevant reaction pathways. Despite the development of detailed kinetic models, there is a demand for more advanced modeling approaches.⁵¹

The present work focuses on studying the N2H3 potential energy surface (PES) that is relevant for all the chemical systems mentioned above. This PES was previously studied by Klippenstein et al.52 at the CCSD(T)/CBS//CCSD(T)/aug-cc-pVDZ level of theory when exploring reactions originating from the NH₂ + NH channel. Klippenstein et al. showed that the reaction proceeds through the abstraction of an H atom on the quartet surface (forming NH3 + N(Q)) or by addition followed by H atom elimination on the doublet surface (forming N₂H₂ + H), as shown in Fig. 1(A). They computed a rate coefficient for the barrierless initial addition of NH₂ + NH(T) using variable reaction-coordinate transition state theory (VRC-TST). The authors stated that a transition state (TS) is likely to exist for H₂ elimination from N₂H₃ but was not found in the cited study.⁵² They expected such a TS to lie higher in energy than that for H loss, and thus it should make an insignificant contribution to the kinetics. Raghunath et al. 43 slightly refined the pressure-dependent network by explicitly considering the cis and trans isomers of N2H2, but did

not consider the barrierless addition reaction (Fig. 1(B)). Diévart and Catoire⁵³ used the high-pressure VRC-TST rate coefficient calculated by Klippenstein et al.52 along with an inverse Laplace transform (ILT)⁵⁴ to suggest pressure-dependent reaction rate coefficients for reactions $N_2H_3 + M \rightleftharpoons NH_2 + NH + M$ and $N_2H_3 + M \rightleftharpoons N_2H_2 + H$. However, they also only considered the trans isomer of the N₂H₂ product (Fig. 1(C)) and did not provide a rate coefficient for the wellskipping reaction $NH_2 + NH(T) \rightleftharpoons N_2H_2 + H$.

Pressure-dependent rate coefficients have not previously been reported for the NH₂ + NH(T) \rightleftharpoons N₂H₂ + H and N₂H₂ + $H \rightleftharpoons NNH + H_2$ reactions, nor for any reaction on this PES involving the H₂NN(S) + H well, such as the formation of N₂H₂ + H, $NNH + H_2$, or N_2H_3 . Currently, the literature lacks a comprehensive description of the N2H3 PES, as well as a complete set of pressuredependent reaction rate coefficients for all relevant pathways. The goal of the present work is to address these gaps and provide kinetic modellers of nitrogen-based fuel systems with high-quality kinetic data for the unimolecular N2H3 system.

2. Methods

Ab initio calculations were carried out using the open source Automated Rate Calculator (ARC) software suite, 55 which automates the calculation of species thermochemistry and reaction rate coefficients. Conformer geometry searches were performed based on a random set of force field conformers generated via RDKit⁵⁶ using MMFF94s.⁵⁷ The force field conformers were optimized at the ωB97X-D/Def2-SVP^{58,59} level of theory. The lowest energy DFT (density functional theory) conformer was further used and optimized at the double-hybrid B2PLYP functional⁶⁰ with Grimme's dispersion correction, D3(BI),⁶¹ coupled with Dunning's correlation consistent basis set⁶² augmented with diffuse functions, i.e., B2PLYP-D3/aug-cc-pVTZ. Ro-vibrational analyses were subsequently computed via ARC at the same level of theory. ARC calculated a frequency scaling

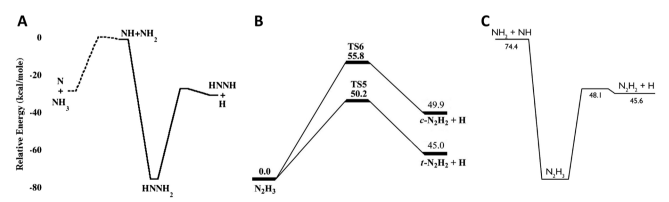


Fig. 1 Representations of the N₂H₃ PES from previous literature works in kcal mol⁻¹. (A) CCSD(T)/CBS//CCSD(T)/aug-cc-pVDZ level of theory calculations by Klippenstein et al. 52 The solid and dashed lines correspond to the doublet and quartet surfaces, respectively. (B) CCSD(T)/CBS//MP2/6-311++G(3df,2p) level of theory calculations by Raghunath et al. 43 on the doublet surface (this inset is not plotted to scale, as in the original manuscript). (C) RCCSD(T)/CBS//M06-2x-D3/aug-ccpVTZ level of theory calculation by Diévart and Catoire⁵³ on the doublet surface. Usage rights: (A) Reprinted (adapted) with permission from *J. Phys. Chem. A* 2009, 113, 38, 10241-10259. Copyright 2009 American Chemical Society. (B) Reprinted from Publication P. Raghunath, N. T. Nghia, M.-C. Lin "Chapter Seven - Ab Initio Chemical Kinetics of Key Processes in the Hypergolic Ignition of Hydrazine and Nitrogen Tetroxide", Adv. Quantum Chem. 69, 2014, 253-301, Copyright (2014), with permission from Elsevier. (C) Reprinted (adapted) with permission from J. Phys. Chem. A 2020, 124, 30, 6214-6236. Copyright 2020 American Chemical Society.

factor of 0.995 for this functional and basis set combination using the method recommended by Truhlar et al.⁶³ Quantum tunneling effects were incorporated using the Eckart correction.⁶⁴ TS verification was performed by intrinsic reaction coordinate (IRC) computations⁶⁵ and automated inspection of the normal mode displacement.

Torsional modes were automatically identified as rotatable single bonds (accounting for relevant resonance structures⁶⁶) and treated with continuous constrained optimizations with all other internal coordinates relaxed using the B2PLYP-D3/aug-ccpVTZ level of theory at a dihedral angle resolution of 10°. The 1D potential energy torsional scans were fitted to truncated Fourier series that were used as input to compute energy levels and the partition function of anharmonic modes using Arkane,⁶⁷ an open source software distributed within the RMG software suite.⁶⁸ All DFT calculations were performed in Gaussian 09.⁶⁹

Single-point energies were computed using three methods: 1. CCSD(T)-F12⁷⁰⁻⁷² with the correlation-consistent aug-cc-pVTZ-F12 base set, 62,73 2. MRCI-F12+Q (internal contracted multiconfiguration reference configuration-interaction with explicit correlation F12 and Davidson correction) $^{74-76}$ with the same basis set, and 3. CCSDT(Q),⁷⁷ which approximates full CCSDTQ calculations⁷⁸ by extending coupled-cluster theory to include full triples excitations (CCSDT) with a perturbative treatment of connected quadruple excitations, using the aug-cc-pVTZ basis set. The MRCI-F12+Q calculations were performed with the full valence active space that included 13 valence electrons and 11 active orbitals for the isomers and TSs on the N₂H₃ PES. The CCSD(T)-F12 and MRCI-F12+O calculations were performed in Molpro 2021.2,79 and the CCSDT(O) calculations were performed in Psi4 v. 1.9.1.80 The frozen core approximation was applied to the CCSDT(Q) computations, wherein the 1s orbitals of nitrogen atoms were kept frozen while the valence and virtual orbitals were fully correlated. This approach effectively reduces computational cost without significantly compromising the accuracy of energy calculations. Empirical systematic errors in atomization energies were corrected using atom energy corrections implemented in Arkane^{67,81} for the respective levels of theory.

Statistical mechanics processing of all electronic structure calculations were done by Arkane⁶⁷ to calculate the thermochemical partition functions and macroscopic parameters, including reaction rate coefficients of interest. Pressure-dependent reaction networks were modeled using a master equation (ME) describing molecular processes on a collisional timescale.⁸² The Modified Strong Collision (MSC) approximation⁸³ was used to solve the ME since the more accurate Chemically Significant Eigenvalues (CSE) method⁸⁴ did not converge for this case. Collision energy transfer rates for deactivating collisions were modeled using the "single exponential down" expression, $\langle \Delta E_{\rm d} \rangle$, with nitrogen as the bath gas. Nitrogen-based systems exhibit a somewhat different trend of $\langle \Delta E_{\rm d} \rangle$ vs. temperature than carbon-based systems. 67 Since both N₄H₆ + N₂ and N₃H₅ + N₂ were shown to have relatively similar values, 67 the parameters computed for the N₃H₅ + N₂ system are assumed to be representative of the present N₂H₃ + N₂ system as well, and were adopted here: $\langle \Delta E_{\rm d} \rangle = 175 \cdot (T/298 \text{ K})^{0.52} \text{ cm}^{-1}$. Lennard-Jones

parameters were taken from RMG's "OneDMinN2" transport library⁸⁵ computed using OneDMin,⁸⁶ and from Glarborg et al.¹⁹

An ILT, 54 a method of deriving microcanonical transition state theory rate coefficients as a function of the total energy from high-pressure limit rate coefficients, was used as implemented in Arkane⁶⁷ for the high-pressure limit recombination rate coefficient of the entrance channel NH2 + NH(T) as determined by Klippenstein et al., 52 and of the recombination channel $H_2NN(S)$ + H as estimated by RMG. 85 Pressure-dependent rate coefficients were fitted to Chebyshev polynomials⁸⁷ over a temperature range of 300-3000 K and a pressure range of 0.01-100 bar. Pressuredependent network sensitivity analysis was performed using Arkane⁶⁷ by perturbing the E0 values of each well and TS in the PES and determining the effect on the calculated pressuredependent rate coefficients.

Results and discussion

The doublet N₂H₃ PES (Fig. 2) involves pressure-dependent reactions among five bimolecular channels interconnected via two isomers, N2H3 and NH3N (Fig. 3). Fig. S2 (ESI†) compares the present computations with the literature values shown in Fig. 1. The N₂H₃ radical is relatively stable, with an endothermic dissociation energy of $\sim 200 \text{ kJ mol}^{-1}$ (Fig. 2), significantly higher than similar beta-scission reactions.88 The association of NH2 + NH(T) (amino radical and triplet imidogen) without further isomerization can result in the formation of N₂H₃ or diazene (N₂H₂) and a hydrogen radical, as suggested by previous works. 43,52,53 However, the previous studies did not consider additional isomers in this PES other than N2H3 (Fig. 1). The NH3N isomer (Fig. 2 and 3(B), SMILES⁸⁹ representation: [NH3+][N-]) is proposed as a conformation that facilitates further transformations on this PES. Thermodynamic properties for this isomer, calculated at the CCSD(T)-F12/cc-pVTZ-F12//B2PLYP-D3/aug-cc-pVTZ level of theory, are provided in Tables S2 and S3 (ESI†). The NH₃N isomer, a stable well in this PES, allows the pressure-dependent network to access two additional bimolecular wells: NNH + H2 and H2NN(S) + H. The N₂H₃ isomer can also produce barrierlessly H₂NN(S) + H (Fig. 2 and Fig. S8, S9, ESI†).

The NH₃ + N(D) bimolecular well, which is not shown in Fig. 2, was found to lie more than 400 kJ mol⁻¹ above the N₂H₃ isomer (Table 1) and was therefore excluded from the ME analysis. This path originates from the NH₃N isomer. A TS for this channel was located using DFT, but its energy at the CCSDT(Q)/aug-cc-pVTZ level was lower than that of the corresponding bimolecular products. This reaction is likely barrierless, as previously determined for a similar reaction, CH₄ + N(D). This channel is not important for the kinetics of N_2H_3 decomposition due to its high energy, although it might be relevant for plasma-assisted ammonia oxidation systems. 91

Klippenstein et al. 52 commented that a TS which leads to the elimination of H2 from N2H3 likely exists, but was not found in their study. Klippenstein et al. also noted that the missing TS is likely to have a higher energy than TSs for H loss and therefore this pathway should not contribute significantly to the

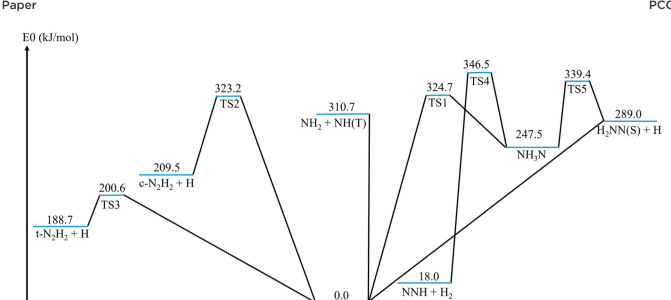


Fig. 2 The pressure-dependent kinetic network of the doublet N₂H₃ PES computed at the CCSD(T)-F12a/aug-cc-pVTZ-F12//B2PLYP-D3/aug-cc-pVTZ level of theory. Zero-point energies (electronic energy plus zero-point energy correction) are in kJ mol⁻¹.

N₂H₃

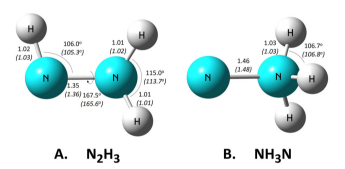


Fig. 3 Geometric representation of 3D structures of the two isomers on the explored PES, (A) N₂H₃ and (B) NH₃N, optimized at the B2PLYP-D3/ aug-cc-pVTZ level of theory. Numbers represent internal coordinate values at the B2PLYP-D3/aug-cc-pVTZ level and in parenthesis at the CCSD/aug-cc-pVDZ level for comparison. The Cartesian coordinates are given in Table S1 (ESI†). Additional internal coordinate comparisons of computed geometries are given in Tables S6-S10 (ESI†).

kinetics. 52 In the present study, we were also unable to identify a TS corresponding to H2 elimination directly from the N2H3 isomer. However, we show here that the elimination of H2 in this PES stems from the NH₃N isomer (TS4, Fig. 2) rather than from N2H3. This TS was indeed found to lie significantly higher in energy than the H loss TSs (TS2 and TS3, Fig. 2). Consequently, this isomerization-mediated H2 elimination channel from N₂H₃ or from NH₂ + NH(T) becomes relevant only at high temperature and low pressure conditions.

The NNH + H₂ products are the lowest-lying bimolecular well in this PES, whereas the t-N₂H₂ + H products' well is the most kinetically accessible from the NH2 + NH(T) direction due to the relatively low energy barrier (TS3, Fig. 2). Another important entry channel into the N₂H₃ pressure-dependent system is the H2NN(S) + H well that could lead to either of the two isomers or to the $NH_2 + NH(T)$, $N_2H_2 + H$, or $NNH + H_2$ bimolecular wells (Fig. 2).

Table 1 Description of stationary points (wells and TSs) on the N₂H₃ PES. Zero-point energy (E0) comparisons at CCSD(T)-F12a/aug-cc-pVTZ-F12, MRCI-F12+Q/aug-cc-pVTZ-F12, and CCSDT(Q)/aug-cc-pVTZ levels of theory are given in kJ mol^{-1} and are relative to the N_2H_3 isomer. Geometry optimization and frequency calculations for all methods were done at the B2PLYP-D3/aug-cc-pVTZ level of theory. Zero Kelvin enthalpy of formation values in kJ mol $^{-1}$ are also given from the Active Thermochemical Tables (ATcT), 92 and are also relative to the N $_2$ H $_3$ isomer

Stationary point	T1 diagnostic coefficient(s)	CCSD(T)-F12a	MRCI-F12+Q	CCSDT(Q)	АТсТ
N_2H_3	0.025	0.0	0.0	0.0	0.0
NH_3N	0.012	247.5	234.6	243.9	_
$NH_2 + NH(T)$	0.008, 0.001	310.7	287.4	295.1	312.9
c -N ₂ H ₂ + \dot{H}	0.013, 0.000	209.5	194.9	196.3	210.1
t-N ₂ H ₂ + H	0.013, 0.000	188.7	170.7	175.2	188.4
$H_2NN(S) + H$	0.021, 0.000	289.0	270.6	276.5	288.8
$NNH + H_2$	0.028, 0.006	18.0	-2.4	16.5	17.4
$NH_3 + N(D)$	0.008, 0.006	447.4	413.8	415.3	427.2
TS1	0.025	324.7	311.6	324.7	
TS2	0.020	223.2	208.7	220.3	
TS3	0.021	200.6	190.5	197.3	
TS4	0.047	346.5	333.8	293.4	
TS5	0.088	339.4	321.6	321.9	

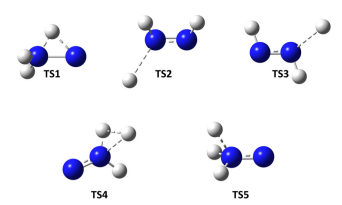


Fig. 4 Geometric representations of TSs on the N₂H₃ PES corresponding to the five non-barrierless path reactions, optimized at the B2PLYP-D3/ aug-cc-pVTZ level of theory. Blue spheres represent nitrogen atoms, smaller light shaded balls represent hydrogen atoms, bond orders are given for illustration purposes only. Cartesian coordinates for all TS structures are available in Table S1 (ESI†).

Table 2 Deviations of E0 values computed at the different levels of theory relative to the N₂H₃ isomer from the corresponding ATcT values, units are kJ mol⁻¹

Stationary point	$\Delta E0$ CCSD(T)-F12a	$\Delta E0$ MRCI-F12+Q	$\Delta E0 \text{ CCSDT(Q)}$
$NH_2 + NH(T)$	-2.20	-25.5	-17.80
c - N_2H_2 + H	-0.59	-15.20	-13.79
t - N_2H_2 + H	0.31	-17.70	-13.19
$H_2NN(S) + H$	0.19	-18.21	-12.31
$NNH + H_2$	0.62	-19.78	-0.88

The geometries of the TSs identified in this PES are shown in Fig. 4. The computed imaginary frequencies of all TSs are provided in Table S1 (ESI†). Table 1 provides the T1 diagnostic coefficients 93-95 for all stationary points on the PES. It also lists the corresponding zero-point energies calculated at three levels of theory: CCSD(T)-F12a/aug-cc-pVTZ-F12, MRCI-F12+Q/aug-ccpVTZ-F12, and CCSDT(Q)/aug-cc-pVTZ. All calculations were performed using B2PLYP-D3/aug-cc-pVTZ geometries and frequencies.

The energy deviations between the CCSD(T)-F12a and the CCSDT(Q) levels were approximately -9 kJ mol⁻¹ on average, with a standard deviation of ~ 15 kJ mol⁻¹. The energy deviations between the MRCI-F12+Q and the CCSDT(Q) levels averaged 6 kJ mol⁻¹, with a standard deviation of ~ 15 kJ mol⁻¹ as well (Table 1). All TSs in this PES, along with N₂H₃, NNH, and H₂NN(S), have T1 diagnostic coefficients exceeding the 0.018-0.020 range, suggesting that a multireference treatment should be used. The MRCI-F12+Q computations account for the system's multireference nature but lack size-consistency, a limitation only partially mitigated by the Davidson (Q) correction. We therefore also computed this PES using the CCSDT(Q) method, which extends coupled-cluster theory to include full triples excitations (CCSDT) with a perturbative treatment of connected quadruple excitations. However, both the MRCI-F12+Q and CCSDT(Q) methods significantly under-predicted the E0 values of nearly all wells relative to the ATcT data (Tables 1 and 2). The CCSD(T)-F12a level, on the other hand, produced highly

accurate zero-point energy predictions when compared to ATcT values (Tables 1 and 2).

The CCSD(T)-F12a/aug-cc-pVTZ-F12 energies computed here are in excellent agreement with the complete basis set extrapolation value based on CCSD(T) energies reported in the literature (Fig. 5). The MRCI calculations, both with and without the explicit F12 correlation⁷⁵ or the Davidson correction,⁷⁶ consistently produced lower energies than the CCSD(T) values (Fig. 5). Similarly, MRCI-F12+Q yielded lower energies than the ATcT values (Tables 1 and 2). The inclusion of the F12 correlation reduced the CCSD(T) energies but increased the MRCI energies. Since the CCSD(T)-F12a/aug-cc-pVTZ-F12 energies showed excellent agreement with both the literature CCSD(T)/CBS values (Fig. 5) and the ATcT values for the wells (Table 2), they were used in this work to compute rate coefficients. The computed pressure-dependent reaction rate coefficients are reported in Tables S4 and S5 (ESI†).

The primary products of NH₂ + NH(T) association at all relevant pressures (1-100 bar) are N₂H₂ + H, primarily the trans N₂H₂ isomer due to the lower energy barrier for its formation (Fig. 2 and 6). At high pressures, N₂H₃ formation becomes relevant but only plays a significant role at temperatures below 1000 K (Fig. 6). The pressure dependence of $NH_2 + NH(T)$ reactions was found to be significant only for the isomer products, N2H3 and NH3N; the rate coefficients of bimolecular products that result from well-skipping reactions showed negligible pressure effect (not shown). As expected, the isomers reach thermal equilibrium more rapidly at higher pressures.

The association of H₂NN(S) + H exhibits more complex pressure-dependent branching (Fig. 7) than the $NH_2 + NH(T)$

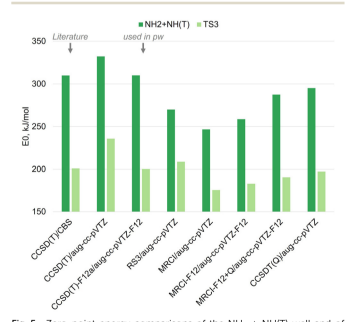


Fig. 5 Zero-point energy comparisons of the NH₂ + NH(T) well and of TS3, both relative to the zero-point energy of the N₂H₃ isomer at the respective level. The CCSD(T)/CBS values were taken from Klippenstein et al.⁵² (annotated as "Literature" in the figure). All other data were computed in the present work, and the CCSD(T)-F12a/aug-cc-pVTZ-F12 level of theory was further used for kinetic computations in the present work (annotated as "used in pw" in the figure).

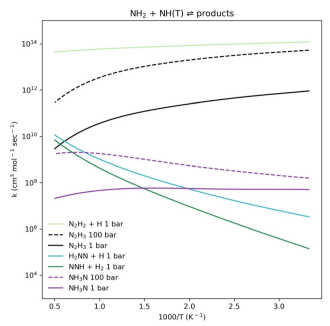


Fig. 6 Comparison of computed rate coefficients for unimolecular NH₂ + NH(T) reactions at 1 and 100 bar. Rate coefficients that do not exhibit a significant pressure-dependence in the examined range are only plotted at 1 bar. The $N_2H_2 + H$ product channel shown here accounts for both trans and cis N2H2 isomers.

channel (Fig. 6), and has not been examined in prior studies of this PES. Similarly to the NH₂ + NH(T) case, the N₂H₂ + H products dominate across almost all relevant temperature and pressure ranges (Fig. 7). The formation of NH₃N becomes the

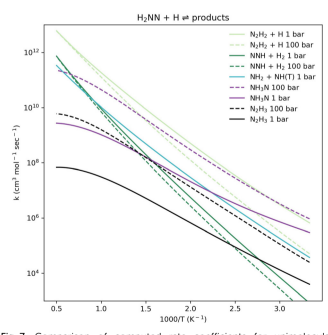


Fig. 7 Comparison of computed rate coefficients for unimolecular H₂NN(S) + H reactions at 1 and 100 bar. Rate coefficients that do not exhibit a significant pressure-dependence in the examined range are only plotted at 1 bar. The N₂H₂ + H product channel shown here accounts for both trans and cis N₂H₂ isomers

dominant pathway only at relatively low temperatures and high pressures; however, it remains significant across the entire examined temperature range at high pressures. At high temperatures, NNH + H₂ formation becomes important, and NH₂ + NH(T) formation is also a significant minor channel. Vibrationally excited NH₃N species produced by H₂NN(S) + H collisions via TS5 are formed well above the TS1 barrier and near or exceeding the TS4 barrier. From the H₂NN(S) + H association direction, once the reactive complex crosses TS5, it can proceed at low pressures past TSs 1, 2, and 3, all of which are submerged relative to TS5. Lower-energy collisions of $H_2NN(S) + H$ can also lead to reactivity via a barrierless pathway (Fig. 2), forming N₂H₃ at high pressures. Since NNH + H₂ formation requires a concerted motion of two H atoms (TS4, Fig. 4), it becomes competitive only at high temperatures, while the dominant pathway consists of consecutive single H atom motions, e.g., via TS1. At all examined pressures, the significance of NNH + H₂ formation increases with temperature. Consequently, in addition to $H_2NN(S) + H \Rightarrow N_2H_2 + H$, the well-skipping $H_2NN(S) + H \rightleftharpoons NNH + H_2$ reaction should be considered in kinetic models of nitrogen-based fuels that describe hightemperature reactivity.

The N₂H₂ + H products are the dominant pathway for NH₂ + NH(T) association at all relevant temperature and pressure ranges (Fig. 6). Our calculated pressure-dependent rate coefficient for the NH₂ + NH(T) \rightleftharpoons N₂H₂ + H reaction is benchmarked in Fig. 8. All rate coefficients compared in Fig. 8 show good agreement. The calculated rate coefficients for the

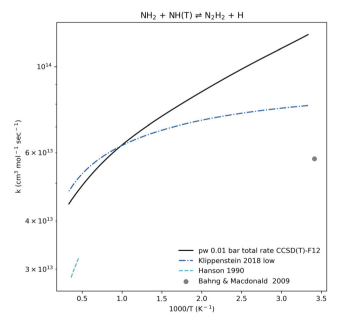
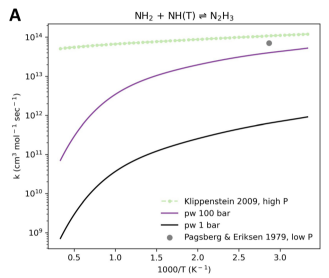


Fig. 8 Rate coefficient comparison for the well-skipping reaction NH₂ + $NH(T) \rightleftharpoons N_2H_2 + H$. The present work (pw) computation shown here accounts for both trans and $cis\ N_2H_2$ isomers at 0.01 bar. The computed rate coefficient is benchmarked against the zero-pressure rate coefficient computed by Kippenstein ${\it et al.}$, 52 a rate coefficient determined by Hanson et al. 96 based on experimental data at 2200-2800 K and 0.8-1.1 bar, and an experimental result by Bahng and Macdonald⁹⁷ at 293 K and pressures up to 0.013 bar.

trans-N₂H₂ product (by Kippenstein et al.) or the combined trans and cis products (reported in the experimental references) are in excellent agreement with values computed in this work, all within a factor of $1.5\times$. This strong agreement between the computed rate coefficient and experimental data is seen across both low- and high-temperature regions when validated against different sources (Fig. 6).

The rate coefficient of the $NH_2 + NH(T) \rightleftharpoons N_2H_3$ reaction is in the fall-off regime in the mid- and high-temperature range, even at 100 bar (Fig. 9(A)). Our computed high-pressure limit



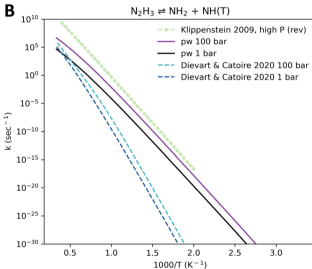


Fig. 9 Rate coefficient comparison for the reaction $NH_2 + NH(T) \rightleftharpoons N_2H_3$ (A) in the forward direction and (B) in the reverse direction. The highpressure limit rate coefficient computed by Klippenstein et al. 52 at CCSD(T)/CBS//CCSD(T)/aug-cc-pVDZ is given in (A) in the original reported direction and in (B) in the reverse direction after being reversed here using thermodynamic data computed by Grinberg Dana et al.⁸ This rate coefficient was also inferred by Pagsberg and Eriksen⁹⁸ at 349 K and a sub-atmospheric pressure on the basis of indirect measurements combined with their model. Diévart and Catoire⁵³ computed a pressuredependent rate coefficient in the reverse direction at CCSD(T)/CBS// M062x-D3/aug-cc-pVTZ.

rate coefficient aligns with the Klippenstein et al.52 highpressure limit rate coefficient (not shown), since it was directly derived from it via an ILT. The pressure-dependent rate coefficient computed here shows good agreement with the previously computed high-pressure limit rate coefficient. The low temperature and low pressure rate coefficient reported by Pagsberg and Eriksen⁹⁸ in 1979 (Fig. 9(A)) is based on an outdated kinetic model manually fitted to low-pressure measurements. It is therefore unreliable, and the experimental result that was interpreted using this model cannot be given a significant weight in this analysis.

The rate coefficient computed by Diévart and Catoire⁵³ and reported by them in the reverse direction relative to Klippenstein et al.'s high-pressure limit rate coefficient exhibits a different slope compared to the values reported here and compared to the reversed high-pressure limit rate coefficient. At both 1 and 100 bar the Diévart and Catoire rate coefficient is several orders of magnitude lower than the reversed highpressure limit rate coefficient in most of the explored temperature range (Fig. 9(B)).

The pressure-dependent rate coefficient computed for the $N_2H_3 \rightleftharpoons NH_2 + NH(T)$ reaction is highly sensitive to the energy of TS2 and TS3 relative to the N2H3 well (Fig. 10). These transition state barriers dictate the primary competing pathways for N₂H₂ + H formation (Fig. 2), making their accurate determination critical for reliable rate predictions. Additionally, the rate coefficient is sensitive to the energy difference between the N_2H_3 isomer and the $NH_2 + NH(T)$ well (Fig. 10), as this difference influences the overall reaction kinetics. Uncertainties in these energy values were propagated to estimate their impact on the high-pressure limit rate coefficients, as shown in Fig. S10 (ESI†). As shown above, the system's energy strongly depends on the level of theory used (Fig. 5 and Table 1).

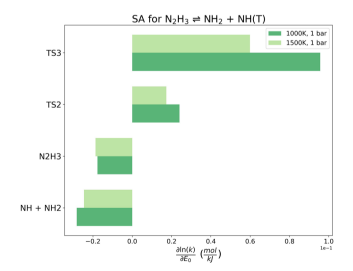


Fig. 10 Normalized sensitivity coefficients of the computed $N_2H_3 \Rightarrow$ NH₂ + NH(T) rate coefficient evaluated with respect to the E0 values of all wells (the two isomers and bimolecular channels) as well as all TSs on the N₂H₃ PES at 1000 and 1500 K, both at 1 bar. Only the highest normalized sensitivity coefficients are reported

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 $N_2H_3 \rightleftharpoons N_2H_2 + H$ Dean & Bozzelli 2000 1 bar 10⁸ pw 100 bar Dievart & Catoire 2020 100 bar 10⁵ pw 1 bar Dievart & Catoire 2020 1 bar 10² 10- 10^{-4} 10^{-7} 10^{-10} 0.5 1.0 2.0 2.5

Fig. 11 Rate coefficient comparison for the $N_2H_3 \rightleftharpoons N_2H_2 + H$ reaction. Calculations done in the present work (pw) are shown in this figure as an overall rate coefficient for the formation of both the trans-N2H2 and the cis-N₂H₂ isomers at 1 and 100 bar. The Dean and Bozzelli⁸⁸ rate coefficient is given here only at 1 bar (Dean and Bozzelli also reported 0.1 bar and 10 bar values), and the Diévart and Catoire⁵³ rate coefficient is plotted at both 1 and 100 bar.

The rate coefficient for the $N_2H_3 \Rightarrow N_2H_2 + H$ reaction reported by Dean and Bozzelli88 is lower than that computed in this work at the same pressure (Fig. 11), since Dean and Bozzelli used a significantly higher barrier for this beta-scission reaction. Dean and Bozzelli reported an endothermicity of 226 kJ mol⁻¹ for this reaction. The reaction barrier they used is estimated by us to be at least $\sim 238 \text{ kJ mol}^{-1}$ based on the reverse energy barrier found here (Fig. 2). This barrier is significantly higher than the E0 barrier of 200.6 kJ mol⁻¹ computed here for the trans-N₂H₂ isomer (Fig. 2). The rate coefficients calculated in the present work and by Diévart and Catoire⁵³ agree reasonably well (Fig. 11).

4. Conclusions

This study provides a deeper understanding of the N₂H₃ potential energy surface (PES), building upon prior work at the CCSD(T)/CBS level. This work identified the NH₃N isomer as an important well in the N₂H₃ PES, which introduces three new bimolecular wells: NNH + H_2 , $H_2NN + H_3$, and $NH_3 + N(D)$. The $N_2H_3 \Rightarrow NH_2 + NH(T)$ rate coefficient was improved compared to previous calculations, and the computed NH₂ + $NH(T) \rightleftharpoons N_2H_2 + H$ rate coefficient shows strong agreement with experimental measurements across both high and low temperatures. The $NH_2 + NH(T)$ reaction predominantly yields N₂H₂ + H on the doublet surface, with N₂H₃ formation becoming significant at low temperatures and high pressures. For $H_2NN(S) + H$ reactions, $N_2H_2 + H$ remains the primary product, with minor contributions from NNH + H₂ and NH₂ + NH(T) at high temperatures; NH₃N formation becomes significant at elevated pressures.

The comprehensive dataset of pressure-dependent rate coefficients presented in this study is valuable for refining models of nitrogen-based fuel combustion and pyrolysis, particularly for ammonia and hydrazine.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare that there are no conflicts of interest to disclose.

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