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Effect of temperature on the gas-phase reaction of CH₃CN with OH radicals: experimental ($T = 11.7$ – 177.5 K) and computational ($T = 10$ – 400 K) kinetic study†

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Acetonitrile (CH₃CN) is present in the interstellar medium (ISM) in a variety of environments. However, at the ultracold temperatures of the ISM, radical-molecule reactions are not widely investigated because of the experimental handicap of getting organic molecules in the gas phase by conventional techniques. The CRESU (French acronym for Reaction Kinetics in a Uniform Supersonic Flow) technique solves this problem. For this reason, we present in this work the kinetic study of the gas-phase reaction of CH₃CN with one of the most ubiquitous radicals, the hydroxyl (OH) radical, as a function of temperature (11.7–177.5 K). The kinetic technique employed to investigate the CH₃CN + OH reaction was the pulsed laser photolysis-laser induced fluorescence. The rate coefficient for this reaction $k(T)$ has been observed to drastically increase from 177.5 K to 107.0 K (about 2 orders of magnitude), while the increase in $k(T)$ from 107.0 K to 11.7 K was milder (around 4 times). The temperature dependent expressions for $k(T)$ are provided in the two distinct T -ranges, excluding the upper limit obtained for $k(177.5$ K):

$$k(11.7\text{--}135.0\text{ K}) = (1.50 \pm 0.12) \times 10^{-11} \left(\frac{T}{300\text{ K}}\right)^{-(0.53 \pm 0.03)} \text{ cm}^3 \text{ s}^{-1}$$

$$k(135.0\text{--}149.9\text{ K}) = (1.50 \pm 1.48) \times 10^{-20} \left(\frac{T}{300\text{ K}}\right)^{-(26.4 \pm 1.5)} \text{ cm}^3 \text{ s}^{-1}$$

In addition, the rate coefficients estimated by the canonical competitive unified statistical (CCUS) theory show a similar behaviour to the experimental results, when evaluated within the high-pressure limit. This is consistent with the experimentally observed independence of $k(T)$ with total gas density at selected temperatures. Astrochemical networks, such as the KIDA database or UMIST, do not include the CH₃CN + OH reaction as a potential depletion process for acetonitrile in the ISM because the current studies predict very low rate coefficients at IS temperatures. According to the model ($T = 10$ K), the impact of the titled reaction on the abundances of CH₃CN appears to be negligible in dark molecular clouds of the ISM (~1% of the total depletion reactions included in UMIST network). With respect to the potential formation of the CH₂CN radical in those environments, even in the most favourable scenario, where this radical could be formed in a 100% yield from the CH₃CN + OH reaction, this route would only contribute around 2% to the current assumed formation routes by the UMIST network.

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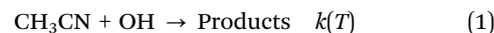


1. Introduction

Acetonitrile, also known as methyl cyanide (CH₃CN), is the simplest organic nitrile molecule. This N-bearing species is present in natural environments such as the Earth's atmosphere, solar system objects, interstellar matter, and even nearby galaxies. This prolate symmetric-top molecule is well-suited for millimetre-wave observations due to its large permanent dipole moment (~4 Debye).¹ Regarding the terrestrial atmosphere, CH₃CN is a trace gas, essentially issued from biomass burning.² Its lifetime in the troposphere is about 6 months with deposition in the ocean being the dominant removal process. Therefore, it can be transported to the stratosphere where it can stay for several decades.³ The reaction with the hydroxyl (OH) radical becomes the main sink.

In the context of the search for the origin of life, a main concern in modern astrochemistry and astrobiology, CH₃CN is of special interest for prebiotic evolution because of the main role played by C–N bonds in the formation of peptide structures, such as amino acids. Then, CH₃CN has been searched and detected routinely in a large variety of extraterrestrial environments, becoming one of the most abundant complex organic molecules (COMs) ever seen in space. In the solar system, CH₃CN has been frequently observed in comets,⁴ for instance in Kohoutek,⁵ Hale-Bopp,^{6,7} P/Halley⁸ or 67P/Churyumov–Gerasimenko.⁹ It has also been identified in one of Saturn's moons, Titan, either by ground-based observations¹⁰ or *in situ* measurements.¹¹ In the interstellar medium (ISM), CH₃CN is one of the few prebiotic molecules seen in all main phases of the lifetime cycle of stars from the early stages to the last remnants. Since its first detection towards SgrA and SgrB molecular clouds in 1971,¹² it has been observed in diffuse and translucent clouds,^{13–15} dark clouds,¹⁶ photodissociation regions,^{17,18} circumstellar envelopes,^{19,20} hot cores,^{21,22} protostellar objects,^{23,24} protoplanetary disks^{25,26} or molecular outflows^{27,28} as well as in extragalactic sources.^{29,30} In addition, CH₃CN is a good

thermometer for interstellar gas^{31,32} and a useful tracer of shocks^{33,34} and kinematic structures.²⁸ Its gas-phase formation routes have been very recently discussed by Giani *et al.*³⁵ As it can be seen, CH₃CN is a quite important molecule in the different surroundings of the ISM, including those where the ultra-low temperatures (10–100 K) prevail. For that reason, it is essential to simulate this cold environment in the laboratory to study the gas-phase reactivity of CH₃CN towards the most abundant radicals in the ISM, like the OH radical which plays a significant role in many chemical processes.^{36–39} Nevertheless, although the CH₃CN + OH reaction (reaction (1)) has been quite extensively studied both experimentally at $T > 250$ K^{40–48} and theoretically at $T > 200$ K,^{49–52} it has never been investigated at the ultralow temperatures of the ISM.



In summary, the experimental kinetic studies of reaction (1) were performed in a wide range of pressures ($P = 1.2$ –760 Torr) and temperatures ($T = 250$ –424 K) using various techniques (see Table 1). Both absolute^{40–45} and relative^{47,48} kinetic techniques were employed to determine the rate coefficient $k(T)$ of reaction (1) at a temperature T . At room temperature, $k(298$ K) has been reported in single temperature experiments^{42,43,46,48} or as part of the temperature dependence study of reaction (1).^{40,41,44,45,47} Most of the studies are in agreement, except the ones by Harris *et al.*⁴⁰ and Andersen *et al.*,⁴⁸ the most recent experimental study, whose $k(298$ K) both differ by a factor of about 2 with previous measurements. As shown in Table 1, there are several studies on the pressure dependence of $k(298$ K).^{43–45} Kurylo and Knable⁴⁴ observed a 10% increase of the rate coefficient in the range 20–50 Torr of either argon or SF₆, while Zetzsch⁴³ found, according to Hynes and Wine,⁴⁵ an increase of $k(298$ K) by a factor of 2.5 in the pressure range 5–100 Torr of argon. However, $k(298$ K) reported by Poulet *et al.*⁴¹ at 1.2 Torr of He is in agreement with high-pressure results. This was confirmed

Table 1 Synthetic view of the available literature for the CH₃CN + OH reaction: experiments. The temperature dependence of the rate coefficient for reaction (1) is mathematically fitted by the modified Arrhenius expression: $k(T) = A \times \left(\frac{T}{298 \text{ K}}\right)^n \times \exp^{-\frac{\gamma}{T}}$

P/Torr	Buffer gas	T/K	Method ^a	$k(298 \text{ K})/10^{-14} \text{ cm}^3 \text{ s}^{-1}$	$A/10^{-13} \text{ cm}^3 \text{ s}^{-1}$	n	γ/K	Products	Ref.
50	Ar	297–424	FP-RF	4.94 ± 0.6	5.86	0	756 ± 126	H-Abst deduced	Harris <i>et al.</i> ⁴⁰
7	Ar	295	FP-RF	2.4 ± 0.3				H-Abst assumed	Fritz <i>et al.</i> ⁴²
5–100	Ar	298	FP-RF	1.9 ± 0.2^b				—	Zetzsch ⁴³
1.2	He	295	DF-EPR	2.1 ± 0.3				H-Abst deduced	Poulet <i>et al.</i> ⁴¹
		393							
20; 50	Ar	250–363	FP-RF	1.94 ± 0.37	6.28	0	1030	H-Abst deduced	Kurylo and Knable ⁴⁴
	SF ₆								
50–700	N ₂	256–388	PLP-LIF	2.48 ± 0.38	11	0	1130 ± 90	~50% H-Abst	Hynes and Wine ⁴⁵
30–600	He							~50% OH Ad.	
(298 K)	N ₂ /O ₂								
700	Air	296	Smog chamber FTIR					~50% H-Abst. ^d	Tyndall <i>et al.</i> ⁴⁶
								~50% OH Ad.	
760	Ar	306–393	FP-FTIR	1.82	18.5	0	1377 ± 62	H-Abst deduced	DeMore ⁴⁷
700	Air	296	Smog chamber FTIR	4.07 ± 1.21^c				H-Abst assumed	Andersen <i>et al.</i> ⁴⁸

^a Flash photolysis resonance fluorescence (FP-RF); discharge flow coupled to electron paramagnetic resonance (DF-EPR); pulsed laser photolysis-laser-induced fluorescence (PLP-LIF); Fourier transform infrared (FTIR). ^b From Hynes and Wine⁴⁵ at about 100 Torr. ^c Average rate coefficients using different reference compounds. ^d Indirectly deduced from HC(O)CN quantification.

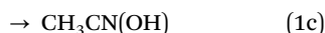


later by Hynes and Wine,⁴⁵ who essentially found no pressure dependence of $k(298\text{ K})$ between 30 and 700 Torr, neither in N_2 nor in He.

Concerning the temperature dependence of $k(T)$, a positive activation energy has been reported between 250 and 424 K, *i.e.*, $k(T)$ decreases when temperature decreases. The Arrhenius parameters reported by Harris *et al.*,⁴⁰ derived from kinetic data obtained at temperatures above room temperature, were slightly lower than those obtained by Kurylo and Knable,⁴⁴ Hynes and Wine⁴⁵ and DeMore *et al.*,⁴⁷ whose temperature dependences were essentially identical. Harris *et al.*⁴⁰ and Kurylo and Knable⁴⁴ interpreted the observed normal Arrhenius behaviour as a prevalence of the H-abstraction from the methyl group of acetonitrile (channel (1a)):



Nevertheless, Hynes and Wine,⁴⁵ from a series of experiments in a mixture of O_2 and N_2 , found indirect evidence that the OH-addition accounted for about 50% of the total reaction process at room temperature. This branching ratio was later confirmed by Tyndall *et al.*⁴⁶ in smog chamber experiments at 700 Torr.



From a theoretical point of view, the potential energy surfaces (PES) of various routes from reactant to products^{49–52} were investigated by diverse methods (see Table 2), and from the obtained PES the temperature dependence of $k(T)$ ^{51–53} was calculated using different variants of the general transition state theory (TST). Li and Wang⁵³ calculated the rate coefficient for the reaction pathway (1a), $k_{1a}(T)$, in the temperature range 250–2500 K using the canonical variational TST (CVT) with a small curvature tunnelling (SCT) correction. The calculated $k(298\text{ K})$ was found to agree with several experiments.^{41,44,45} As mentioned earlier, the contribution of the OH-addition channels to $k(T)$ was not negligible. Thus, the OH-addition channels were theoretically investigated by Galano,⁵⁰ concluding that reaction (1c) was endothermic while reaction paths (1a) and (1b) have similar efficiency, based on close Gibbs free

energy barriers. No rate coefficients were calculated, however. Additional reaction pathways were explored by Tian *et al.*⁵¹ concluding that reaction channel (1a) was the dominant one whereas the two OH-addition channels were not considered as terminal in the reaction pathways. The rate coefficient $k_{1a}(T)$ was calculated using TST with the Wigner correction to take tunnelling into account. At room temperature their calculation agreed well with the measurements from Harris *et al.*⁴⁰ and Andersen *et al.*⁴⁸ More recently, Alihosseini *et al.*⁵² found 12 viable routes along the PES of the OH + CH_3CN reaction. Only the rate coefficient $k_{1a}(T)$ was calculated using TST, at atmospheric pressure over the temperature range of 200–2500 K. In this temperature range, a non-Arrhenius behaviour of $k_{1a}(T)$ was predicted, however the authors fitted the data to a conventional Arrhenius equation which gives a very poor representation of individual $k_{1a}(T)$ (see Fig. 5 in Alihosseini *et al.*⁵²). Furthermore, $k_{1a}(298\text{ K})$ is about two orders of magnitude higher ($2.17 \times 10^{-12}\text{ cm}^3\text{ s}^{-1}$) than previous investigations either experimental or theoretical.

Therefore, there are still matters of controversy concerning the rate coefficient $k(T)$, the branching ratios, and the influence of pressure in the obtained measures. In addition, the extension of experimental and theoretical studies to temperatures lower than 200 K is certainly desirable, especially for the modelling of the chemistry of interstellar objects in which CH_3CN and OH have been observed. Astrochemical networks, such as the KIDA database or UMIST, do not include reaction (1) as a potential depletion process for acetonitrile in the ISM. The present investigation aims at studying the gas-phase kinetics of the $\text{CH}_3\text{CN} + \text{OH}$ reaction between 11.7 K and 177.5 K from an experimental point of view using a pulsed CRESU (French acronym for Reaction Kinetics in a Uniform Supersonic Flow) reactor and from a theoretical point of view using high level quantum chemistry tools such as the density functional theory (DFT) and the competitive canonical unified theory (CCUS) in the temperature range of 10–400 K. The results reported here were used to model the IS abundance of CH_3CN using a pure gas-phase astrochemical model and the implications of our results in the chemistry of IS molecular clouds are discussed.

Table 2 Synthetic view of the available literature for the $\text{CH}_3\text{CN} + \text{OH}$ reaction: theory. The temperature dependence of the rate coefficient is mathematically fitted by the modified Arrhenius expression: $k(T) = A \times \left(\frac{T}{298\text{ K}}\right)^n \times \exp^{-\frac{\gamma}{T}}$

Method	T (K)	$k(298\text{ K})/10^{-14}\text{ cm}^3\text{ s}^{-1}$	$A^a/10^{-13}\text{ cm}^3\text{ s}^{-1}$	n	γ (K)	Products	Ref.
PMP4/6-311_G(2df,2pd)//MP2/6-311G(d,p)	250–2500	2.38 ^a	1.4 ^b	2.89 ^b	620 ^b	H-Abst only considered	Li and Wang ⁵³
QCISD(T)/6-311_G(2df,2pd)//MP2/6-311G(d,p) + CVTST/SCT	250–2500	2.02 ^a	1.3 ^b	2.91 ^b	648 ^b	H-Abst only considered	Li and Wang ⁵³
BHandHLYP/6-311++G(2d,2p) + CCSD(T)/6-311++G(2d,2p)	—	—	—	—	—	~50% H-Abst ~50% OH Ad.	Galano ⁵⁰
B3LYP/CBSB7	250–3000	5.14	7.05	3.045	−780	7 routes obtained	Tian <i>et al.</i> ⁵¹
TST + Wigner correction						H-Abst mainly	
CBS-QB3	250–430	4.79	15.6	2.36	−1038	7 routes obtained	Tian <i>et al.</i> ⁵¹
TST + Wigner correction						H-Abst mainly	
B3LYP + G3B3 or CCSD(T)/6-311++G(3df,3pd) + TST	200–2500	217	489	0	−1277	12 routes obtained but H-Abst only considered	Alihosseini <i>et al.</i> ⁵²

^a Calculated at 298 K (not from fit). ^b Obtained in the present work from the data presented in Table 5 from Li and Wang.⁵³



2. Experimental part

The kinetic study of reaction (1) has been performed at ultralow temperatures (11.7–177.5 K) achieved by a pulsed CRESU apparatus, where the gas is not continuously passing through the Laval nozzle. A detailed description of the technique and the set-up can be found elsewhere.^{54–61} Hence, a schematic diagram with the experimental novelties of the present work is provided in Scheme 1 and only a brief description is given below.

Seven different Laval nozzles and a total of 15 operational conditions were used in this work to fill up the temperature range. The isentropic expansion from a relative high-pressure reservoir ($P_{\text{res}} = 9.97\text{--}366.48$ mbar) to a low-pressure chamber (0.12–6.00 mbar) through the nozzle results in a uniform supersonic cold jet ranging from 11.7 and 177.5 K. A rotary disk with one or two symmetrical apertures was employed to pulse the gas at 5 Hz for $T = 11.7$ K or 10 Hz for the rest of operational conditions, respectively.

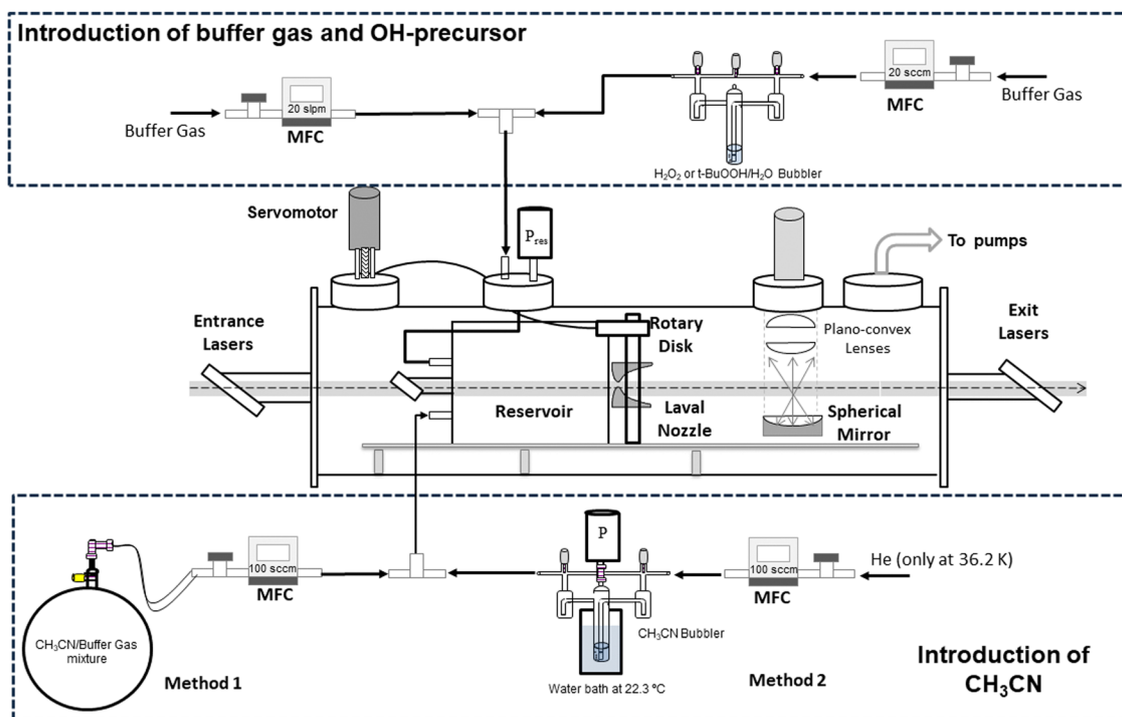
The kinetic technique pulsed laser photolysis-laser-induced fluorescence (PLP-LIF) was also described previously.^{54,55,59,62–68} The production of OH radicals along the supersonic jet was achieved by PLP of gaseous H_2O_2 or *tert*-butyl hydroperoxide (*t*-BuOOH) at 248 nm, radiation generated by a KrF excimer laser (Coherent, model ExciStar XS 200). Both precursors were introduced in the reservoir by flowing a small portion of the buffer gas (He, N_2 , or Ar) through the bubbler containing the concentrated solution of the OH precursor (see Scheme 1). Gaseous CH_3CN was introduced in the reservoir by flowing diluted mixtures in the buffer gas through a calibrated mass flow controller (method 1) or by bubbling liquid samples of CH_3CN with a small portion of the

buffer gas (method 2). More details of both methods are given in the ESI.† Once the OH radicals were generated in the electronic ground state, they were excited at *ca.* 282 nm, radiation achieved with a frequency-doubled dye laser (Lambda Physik, model Scanmate) pumped by the second harmonic of a Nd:YAG laser (Continuum, model Surelite III). Finally, the temporal evolution of OH radicals in the presence and absence of acetonitrile was monitored by LIF (I_{LIF}) at *ca.* 310 nm, using a photomultiplier tube (Electron Tube, model 9813B) with a bandpass filter centred at 310 nm with a full width at half maximum of 10 nm (Andover Co., model 310FS10-25).

2.1. Kinetic analysis

The kinetic experiments were performed under pseudo-first order conditions, *i.e.*, in large excess of acetonitrile with respect to the initial concentration of OH radicals ($[\text{CH}_3\text{CN}] \gg [\text{OH}]_0$). In such a situation, after an increase due to rotational relaxation (see ESI† and already explained),^{54,57} the temporal profile of I_{LIF} results in an exponential decay primarily caused by the reaction of OH with CH_3CN , but also by the reaction with the OH-precursor (H_2O_2 or *t*-BuOOH) or diffusion out of the detection zone. Some examples of the temporal profile of I_{LIF} recorded in the absence and presence of CH_3CN at three selected temperatures are shown in Fig. S1 of ESI.† From the analysis of the I_{LIF} decays, the pseudo-first order rate coefficient, k' (or k'_0 in the absence of CH_3CN) was obtained. In these experimental conditions, k' is proportional to $[\text{CH}_3\text{CN}]$ at a single temperature, according to the relationship given by eqn (E1):

$$k' = k(T)[\text{CH}_3\text{CN}] + k'_0 \quad (\text{E1})$$



Scheme 1 Experimental set-up scheme including both methods to introduce CH_3CN into the reservoir. MFC: mass flow controller.



where $k(T)$ is the second-order rate coefficient at each temperature and gas density obtained from the slope of the $k' - k'_0$ vs. $[\text{CH}_3\text{CN}]$ plots. Reliable knowledge of $[\text{CH}_3\text{CN}]$ in the jet is essential to obtain accurate $k(T)$ at each temperature. These concentrations were determined from flow rate or vapor pressure measurements (see ESI†), according to the way CH_3CN is introduced in the reservoir. As shown in Fig. S2 (ESI†), $[\text{CH}_3\text{CN}]$ determined by both methods yield $k(T)$ at 36.2 K in excellent agreement.

This linear relationship is not accomplished at high $[\text{CH}_3\text{CN}]$ and the bimolecular plots display a curvature. The observed downward curvature is associated with the dimerization process of CH_3CN , as previously reported for other OH-reactions with COMs (Fig. S2, ESI†).^{54–60,62,63,68–70} As can be seen in Fig. S3 (ESI†), the $(\text{CH}_3\text{CN})_2$ dimer formation is especially enhanced at low temperatures. Then, the onset of such a dimerization, *i.e.*, $[\text{CH}_3\text{CN}]$ from which the bimolecular plots are not linear, increases with temperature for a total gas density of the same order of magnitude 10^{17} cm^{-3} (*e.g.* $[\text{CH}_3\text{CN}] = 1.30 \times 10^{14} \text{ cm}^{-3}$ at 21.7 K, $1.96 \times 10^{14} \text{ cm}^{-3}$ at 76.0 K and $5.30 \times 10^{14} \text{ cm}^{-3}$ at 135.0 K). At $T > 140$ K, $k' - k'_0$ vs. $[\text{CH}_3\text{CN}]$ plots are linear in the entire explored $[\text{CH}_3\text{CN}]$ range, even reaching 1% of the total gas density. Acetonitrile concentration was not increased beyond 1% of the total gas density since it may affect the uniformity of the flow and the jet temperature. In Table 3, $[\text{CH}_3\text{CN}]$ range within which eqn (E1) is valid are listed for every temperature and gas density. Four examples of the $k' - k'_0$ vs. $[\text{CH}_3\text{CN}]$ plots are shown in Fig. 1 at selected temperatures. As shown, the slope of these plots increases

when the temperature decreases, implying that the OH-reactivity of CH_3CN is enhanced at lower temperatures.

2.2. Chemicals

Buffer gases (He, N_2 , and Ar), all of them with 99.999% purity (Nippon Gases Europe), were used as supplied. Liquid acetonitrile ($\geq 99.9\%$, Sigma-Aldrich) was degassed by repeated freeze-pump-thaw cycles prior to use for the experiments in which the storage bulb was used. The aqueous solution of hydrogen peroxide (50% w/w, Sharlab) was pre-concentrated as previously reported,⁶⁶ whereas the *t*-BuOOH solution (70%, Sigma-Aldrich) was used directly without pre-concentration, because its vapour pressure is higher than that of H_2O . When no use, both precursor bubblers were stored in a refrigerator at 2°C to prevent from thermal decomposition.

3. Computational methodology

All geometry optimizations were performed by the M08-HX Density Functional Theory (DFT) method⁷¹ with the MG3S basis set,⁷² as for the recently studied OH + CH_3NH_2 reaction.⁶⁸ All zero-point energies (ZPEs) were scaled by the recommended factor of 0.973.⁷³ The two computed reactions are the hydrogen abstraction from the methyl group by the OH radical (reaction (1a)) and the OH-addition to the C of cyano group (reaction (1b)). It is worth noting that reaction (1a) is part of a much larger reaction network recently studied by some of us.⁷⁴ The OH-addition to the nitrogen atom, as well as other 4 additional channels previously investigated by Tian *et al.*⁵¹ have very high barriers and are not competitive with reactions (1a) and (1b).

Additional geometry optimizations were also carried out at $\omega\text{B97X-D}^{75}$ with the def2-TZVP basis set.⁷⁶ All ZPEs were scaled by the recommended factor of 0.975.⁷³ The geometries obtained by both electronic structure levels, that is, M08-HX/MG3S and $\omega\text{B97X-D}/\text{def2-TZVP}$, were employed as input for the CCSD(T)-F12/cc-pVTZ-F12 single-point calculations^{77,78} to compare the result of both functionals on an accurate and common ground.

The thermal rate coefficients were evaluated employing the competitive canonical unified theory (CCUS). In this formalism, the rate coefficient for each reaction channel, k_i^{CCUS} (where $i = 1\text{a}$ or 1b), is given by eqn (E2),⁷⁹

$$k_i^{\text{CCUS}} = k^{\text{CCUS}} \alpha_i \quad (\text{E2})$$

$$k^{\text{CCUS}} = \left(\frac{1}{k_a} + \frac{1}{k_1} \right)^{-1} \quad (\text{E3})$$

where k_a is the rate coefficient for the association reaction and $k_1 = k_{1a} + k_{1b}$ is the total rate coefficient for the OH + CH_3CN reaction calculated as the forward flux from reactants through transition states TS1a (H-abstraction) and TS1b (OH-addition). The ratio between the individual rate coefficients for the transition state passage through k_i and the total rate coefficient k_1 is given by α_i . Because the association reaction is the same for both competitive reactions, the products branching ratio in percentage is given by $100 \cdot \alpha_i$.

The rate coefficient for the association reaction is given by the following expression:⁸⁰

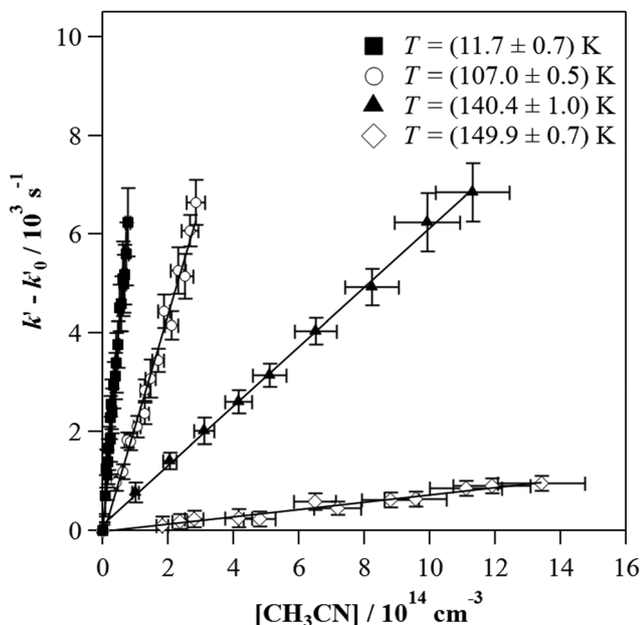


Fig. 1 Examples of $k' - k'_0$ versus $[\text{CH}_3\text{CN}]$ plots at four different temperature conditions. Uncertainties in $[\text{CH}_3\text{CN}]$ are cautiously considered as $\pm 10\%$ and uncertainties in $k' - k'_0$ are the standard deviation obtained from the analysis of the I_{LIF} decays.



$$k_a = C\mu^{-1/2}(d_{\text{OH}}d_{\text{CH}_3\text{CN}})^{2/3}T^{-1/6} \quad (\text{E4})$$

where T is the temperature, $C = 1.83 \times 10^{-9}$ is a constant that provides the rate coefficient in $\text{cm}^3 \text{s}^{-1}$, μ is the reduced mass (in uma), and $d_{\text{CH}_3\text{CN}} = 4.02 \text{ D}$ and $d_{\text{OH}} = 1.66 \text{ D}$ are the dipole moments of acetonitrile and the OH radical, respectively.

The rate coefficients k_i were calculated employing canonical variational transition state theory with small-curvature tunnelling corrections (CVT/SCT),⁸¹ according to the following equation:

$$k_i^{\text{CVT/SCT}} = \Gamma_i^{\text{CVT}} \kappa_i^{\text{CVT/SCT}} k_i^{\text{TST}} \quad (\text{E5})$$

where k_i^{TST} is the conventional transition state theory (TST) rate coefficient, Γ_i^{CVT} is the variational coefficient that accounts for recrossing ($0 \leq \Gamma_i^{\text{CVT}} \leq 1$), and $\kappa_i^{\text{CVT/SCT}}$ is the tunnelling transmission coefficient ($\kappa_i^{\text{CVT/SCT}} \geq 1$) calculated within the SCT approximation. The evaluation of these two parameters requires the calculation of the minimum energy path (MEP), which was evaluated using the Page-McIver algorithm in mass-scaled coordinates.⁸² The MEP for the H-abstraction and OH-addition reactions was followed employing a step size of 0.005 Bohr and Hessian calculations every ten steps. In the case of the reaction through TS1b, we corrected the energies along the MEP to match the coupled-cluster (CC) energy by employing the interpolated single-point energies algorithm.⁸³ The CVT/SCT calculations were performed with Pilgrim.⁸⁴

The CCUS prescription allows calculating the thermal rate coefficients in the low-pressure (LPL) and high-pressure (HPL) limits. The only difference between the two regimes is that in the LPL the lowest energy for tunnelling is the energy of reactants plus its ZPE, whereas in the HPL the lowest energy that allows tunnelling is the energy of the pre-reactive complex plus its ZPE. For the latter, it is assumed that the collisions with other molecules will stabilize the complex, a situation that cannot occur at very low pressures. Therefore, the only factor affected by pressure in eqn (E5) is the tunnelling transmission coefficient (see Section 4.2).

4. Results and discussion

4.1. Experimental kinetic study

The individual rate coefficients $k(T)$ determined in this work are listed in Table 3 as a function of the OH-precursor (H_2O_2 or $t\text{-BuOOH}$), gas density (n), and temperature. As it can be seen in the examples depicted in Fig. 2, no influence of the OH-precursor on $k(T)$ was observed. Thus, in Table 3 the weighted average of $k(T)$ obtained using both precursors is given in bold.

4.1.1. Pressure dependence of $k(T)$. The pressure dependence study of $k(T)$ was carried out for some specific temperatures around 22 K (He as bath gas), 50 K (He, N_2 , or Ar as bath gas), and 106 K (N_2 or Ar as bath gas). The total gas density was explored in the range of $(1.50\text{--}44.4) \times 10^{16} \text{ cm}^{-3}$. n was varied around 5 times at $\sim 22 \text{ K}$, 30 times at $\sim 50 \text{ K}$, and 3 times at $\sim 106 \text{ K}$. As shown in Fig. 3, no pressure dependence of $k(T)$ has been observed at around 22 K, 50 K and 106 K, within the error bars, over the investigated range. This is in agreement with the

observations of Sleiman *et al.*⁸⁶ for the reaction of CH_3CN with CN radicals at 23 K (He as bath gas). These authors reported a strong dependence of the rate coefficient for the CN-reaction at 52 K and 132 K in Ar. The p -dependence was attributed by these authors as evidence of the adduct formation by association of the reactants (CN and CH_3CN). Even though the gas density range investigated by Sleiman *et al.*⁸⁶ is higher ($n = (3.48\text{--}82.1) \times 10^{16} \text{ cm}^{-3}$) than ours, if a pressure dependence of $k(T)$ for reaction (1) prevails it would have been observed at the concentration level used in the present work.

4.1.2. Temperature dependence of $k(T)$. As shown in Table 3, $k(T)$ increases when temperature decreases, resulting in an acceleration of the reaction of CH_3CN towards OH radicals of around 3800 times at 11.7 K and 10 times at 177.5 K with respect to $k(300 \text{ K}) = 2.2 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$.⁸⁷ The value of $k(177.5 \text{ K})$ reported here ($2 \times 10^{-13} \text{ cm}^3 \text{ s}^{-1}$) is an upper limit, since almost no change in k' with respect to k'_0 was observed in the timescale of the experiment and within the $[\text{CH}_3\text{CN}]$ range used. This upper limit of k was estimated as explained by Douglas *et al.*,⁸⁵ *i.e.*, as twice the uncertainty in k' at the maximum CH_3CN concentration, $[\text{CH}_3\text{CN}]_{\text{max}}$, used in the experiment, divided by that $[\text{CH}_3\text{CN}]_{\text{max}}$. In Fig. 4, the log-log plot of average $k(T)$ from Table 3 *versus* T is presented between 11.7 K and 177.5 K. Clearly, two distinct regions in the plot were observed. Between 135.0 K and 177.5 K, an abrupt decrease of $k(T)$ of a factor of around 90 was observed. In this steep region, the expression of $k(T)$ provided by eqn (E6), where the estimated $k(177.5 \text{ K})$ is excluded, is presented as the blue line in Fig. 4.

$$k(135.0\text{--}149.9 \text{ K}) = (1.50 \pm 1.48) \times 10^{-20} \left(\frac{T}{300 \text{ K}} \right)^{-(26.4 \pm 1.5)} \text{ cm}^3 \text{ s}^{-1} \quad (\text{E6})$$

Nevertheless, this expression can be used to extrapolate $k(177.5 \text{ K})$. The extrapolated rate coefficient at 177.5 K is $1.56 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$ (empty black circle in Fig. 4), which is a reasonable value if it is compared with $k(300 \text{ K}) = 2.2 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$.⁸⁷ In contrast, the observed increase of $k(T)$ between 135.0 K and 11.7 K is milder (around 5 times) than that between 135.0 and 149.9 K. The temperature dependence of $k(T)$ in the 11.7–135 K region is well described by eqn (E7) – depicted as the red line in Fig. 4.

$$k(11.7\text{--}135.0 \text{ K}) = (1.50 \pm 0.12) \times 10^{-11} \left(\frac{T}{300 \text{ K}} \right)^{-(0.53 \pm 0.03)} \text{ cm}^3 \text{ s}^{-1} \quad (\text{E7})$$

4.2. Computational study

4.2.1. Energetics. The energy of the M08-HX and ω B97X-D stationary points is compared in Table 4 to that of the CCSD(T)-F12 single point calculations with the cc-pVTZ-F12 basis set performed over the M08-HX and ω B97X-D optimized geometries. The DFT energies are substantially different in the two methods and the stability of the two transition states is



Table 3 Summary of the experimental conditions employed in this work, individual $k(T)$ as a function of the gas density (n) and average $k(T)$ (in bold) as a function of temperature^a

$\approx T/K$	T/K	$n/10^{16} \text{ cm}^{-3}$	$[\text{CH}_3\text{CN}]/10^{13} \text{ cm}^{-3}$	$k(T)/10^{-11} \text{ cm}^3 \text{ s}^{-1}$
12	11.7 ± 0.7	6.88 ± 0.62	0.76–7.81	8.18 ± 0.85^b
			1.35–6.15	8.45 ± 0.92^c
				8.25 ± 0.89
22	21.1 ± 0.6	3.37 ± 0.15	0.87–8.93	7.03 ± 0.86^b
			0.87–8.89	5.89 ± 0.62^c
				6.05 ± 1.28
36	21.7 ± 1.4	16.65 ± 1.61	1.63–12.7	6.17 ± 0.71^b
				5.27 ± 0.57^b
				4.91 ± 0.57^c
50	36.2 ± 1.2	17.73 ± 0.86	2.00–14.5	5.14 ± 0.71
			1.37–15.0	
50	50.5 ± 1.6	1.50 ± 0.12	1.94–12.5	4.04 ± 0.50^b
			1.19–8.82	4.29 ± 0.46^c
				4.23 ± 0.52
50	49.9 ± 1.8	3.67 ± 0.32	1.12–9.99	4.33 ± 0.52^b
			1.12–10.1	3.91 ± 0.57^c
				4.18 ± 0.71
50	52.1 ± 0.5	19.52 ± 0.28	0.83–11.7	4.07 ± 0.42^b
			0.92–8.08	4.00 ± 0.61^c
				4.07 ± 0.41
76	76.0 ± 0.8	14.96 ± 0.34	2.19–15.8	2.70 ± 0.37^b
106	106.0 ± 0.6	14.02 ± 0.11	1.26–8.58	2.52 ± 0.38^b
			0.79–6.52	2.70 ± 0.35^c
				2.63 ± 0.36
106	107.0 ± 0.5	4.90 ± 0.06	4.11–30.1	2.20 ± 0.23^b
			4.55–14.8	2.28 ± 0.33^c
				2.21 ± 0.23
135	135.0 ± 0.8	29.45 ± 0.45	6.19–45.4	1.82 ± 0.20^b
			8.47–63.9	1.40 ± 0.17^b
135	136.1 ± 0.8	24.92 ± 0.35		
140	140.4 ± 1.0	21.68 ± 0.40	10.1–113	0.60 ± 0.06^b
			13.4–166	0.32 ± 0.04^b
140	143.3 ± 0.6	17.02 ± 0.17		
150	149.9 ± 0.7	10.69 ± 0.12	18.1–134	0.07 ± 0.01^b
177	177.5 ± 1.2	6.71 ± 0.11	22.8–67.1	$<0.02^{bd}$

^a Uncertainties in $k(T)$ account for statistical ($\pm 2\sigma$) and 10% systematic errors. In n the indicated errors result from the aerodynamic fluctuations in the flow ($\pm 1\sigma$). ^b H_2O_2 as OH-precursor. ^c $t\text{-BuOOH}$ as OH-precursor. ^d Upper limit calculated as Douglas *et al.*⁸⁵

reversed when the ZPE is included. However, CCSD(T)-F12 over the DFT geometries produces very similar results, in particular, the CCSD(T)-F12/M08-HX predict that TS1a is $4.51 \text{ kcal mol}^{-1}$ above TSb, and $4.65 \text{ kcal mol}^{-1}$ when considering CCSD(T)-F12/ $\omega\text{B97X-D}$ calculations. This indicates that both DFT methods supply similar geometries for the stationary points. The potential barrier heights are also in relatively good agreement with the CCSD(T)-F12//B3LYP/6-311++G(3df,3pd) calculations carried out by Alihosseini *et al.*⁵² who reported barriers of 5.30 and $1.36 \text{ kcal mol}^{-1}$ for TSa and TSb, respectively, with a difference between barriers of $3.94 \text{ kcal mol}^{-1}$.

The difference between the two transition states is reduced to 1.22 and $1.23 \text{ kcal mol}^{-1}$ for CCSD(T)-F12/M08-HX and CCSD(T)-F12/ $\omega\text{B97X-D}$, respectively. These values are in good agreement with the results obtained by Galano,⁵⁰ where TS1a is above TS1b by $1.00 \text{ kcal mol}^{-1}$. Of the two DFT methods reported here, M08-HX is the most similar to the coupled-

cluster calculations. Additionally, it does not revert the stability of the two transition states, as is the case of the coupled-cluster calculations. Therefore, hereafter all thermal rate coefficient calculations will involve M08-HX calculations of the MEP and additional single-point energy corrections for reaction (1b), to correct the difference between this DFT method and the coupled-cluster calculations. Fig. 5 plots a schematic reaction network of the CCSD(T)-F12/M08-HX ZPE-corrected level regarding reactants, which is the basic scheme for further dynamics calculations.

The pre-reactive complex (C1) and the transition states that correspond to reactions (1a) and (1b) obtained at M08-HX/MG3S are depicted in Fig. 6. It can be observed that the C1 complex can easily react through both transition states.

4.2.2. Computed rate coefficients. The calculated rate coefficient for the association reaction k_a decreases slightly with temperature and it presents a value of $1.28 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$ at



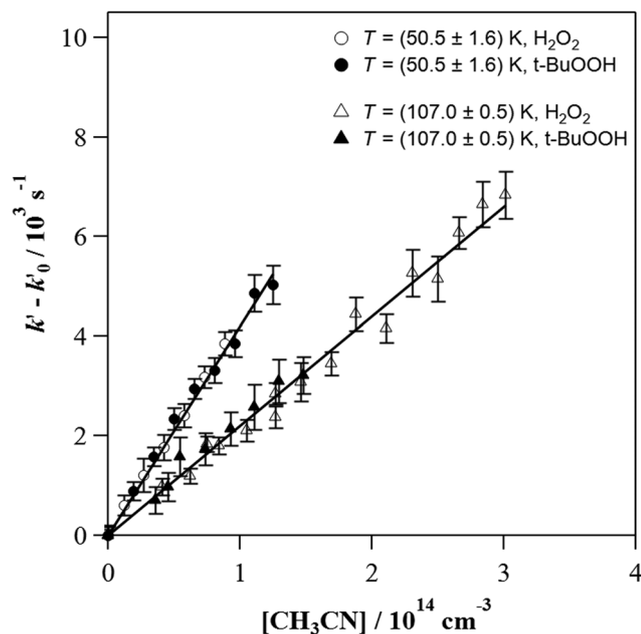


Fig. 2 Examples of the bimolecular plots at 50.5 and 107.0 K using two OH-precursors.

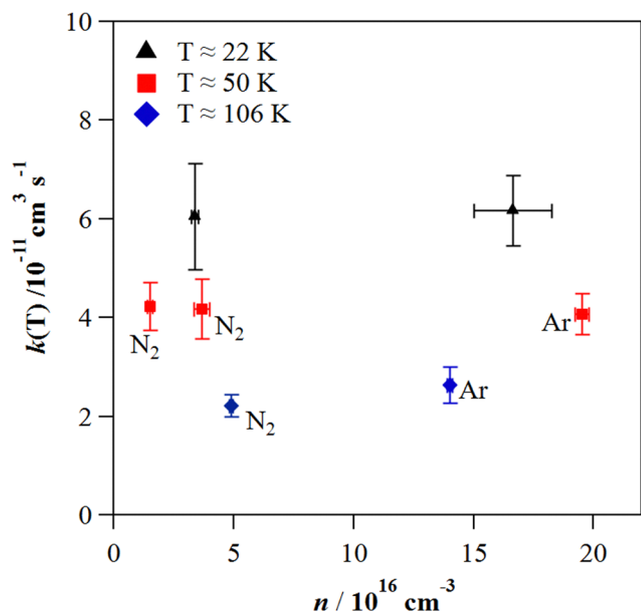


Fig. 3 Dependence of $k(T)$ on the jet gas density (n) at ca. 22, 50, and 106 K. Black triangles correspond to experiments at 22 K in He, while at 50 and 106 K the bath gas is indicated next to each point. Error bars in $k(T)$ are statistical ($\pm 2\sigma$) and 10% systematic errors and in n are only statistical ($\pm\sigma$).

$T = 10 \text{ K}$, and a value of $9.75 \times 10^{-10} \text{ cm}^3 \text{ s}^{-1}$ at $T = 50 \text{ K}$. In Table 5, variational and tunnelling transmission coefficients are presented for selected temperatures between 50 and 300 K. A full account of these parameters and the thermal rate coefficients k_{1a} and k_{1b} is given in the ESI.†

Tunnelling effects are more important for the H-abstraction than for the OH-addition reaction. This is expected as the

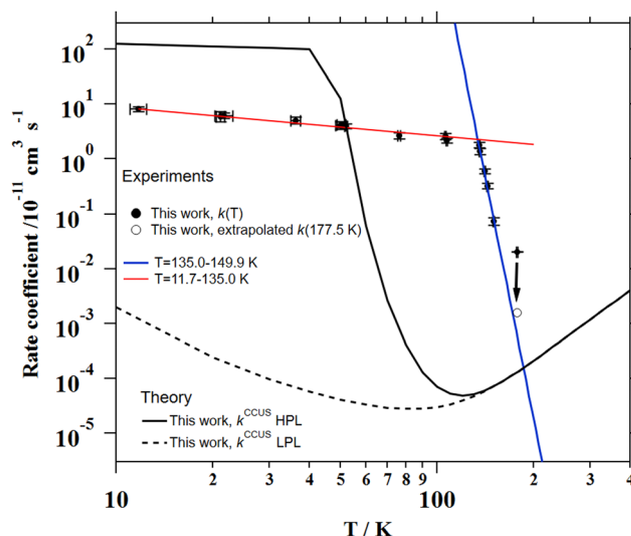


Fig. 4 Experimental and theoretical temperature dependence of the rate coefficient for the $\text{CH}_3\text{CN} + \text{OH}$ reaction. Red and blue lines represent the resulting fit to $k(T) = A (T/300 \text{ K})^n$ in the stated temperature range. Solid and dashed black curves are the computed HPL and LPL rate coefficients.

hydrogen atom is a light particle, whereas the OH-addition involves heavy-atom tunnelling.⁸⁸ However, variational effects are also more relevant in the H-abstraction reaction, so the result of multiplying the variational coefficient by the tunnelling coefficient favours the addition reaction. Moreover, the potential barrier is higher for the H-abstraction reaction and therefore the OH-addition reaction dominates between 10 and 300 K. In fact, it contributes 100% at 10 K and 87% at 300 K.

The variational effect in the H-abstraction shows the magnitude of recrossing in the reaction. In CVT/SCT, this effect is calculated by finding the maximum of the Gibbs free energy along the reaction coordinate s of the MEP (negative in the reactant site, positive in the products site and zero at the transition state), which for the abstraction reaction is located at $s = -0.267 \text{ Bohr}$ at 0 K and at $s = -0.250 \text{ Bohr}$ at 400 K. Variational effects are due to the increasing value of the ZPE as we head toward reactants; it is $31.51 \text{ kcal mol}^{-1}$ at the transition state and $33.17 \text{ kcal mol}^{-1}$ at $s = -0.250 \text{ Bohr}$. This ZPE variation displaces the maximum of the free energy towards the reactants site.

For the evaluation of the normal-mode frequencies along the MEP, we have employed redundant internal coordinates, because it is well-known that an accurate evaluation of the ZPE along the reaction path requires the use of internal coordinates. They provide a more physical description of the normal-mode frequencies than the Cartesian coordinates (notice that the calculated normal mode frequencies in non-stationary points depend on the system of coordinates).⁸⁹ Theoretical methods based on conventional TST that incorporate tunnelling effects (for instance, through the Eckart barrier), completely ignore the recrossing, which in this case has the effect of reversing the contribution of the two channels to the branching ratios.

We have also included torsional anharmonicity in the transition state structures applying the same protocol as for the



Table 4 Relative energies (ΔE) and ZPE-corrected energies (ΔH_0°) with respect to reactants of the complex (C), transition states (TS), and products (P) for reactions (1a) and (1b). Optimizations at the M08-HX and ω B97X-D levels and single-point calculations at the CCSD(T)-F12 level. Energies in kcal mol⁻¹

	$\Delta E_{M08/\omega B97}$	$\Delta E_{CC(M08/\omega B97)}$	$\Delta H_{0,M08/\omega B97}^\circ$	$\Delta H_{0,CC(M08/\omega B97)}^\circ$
Reactants	0.00/0.00	0.00/0.00	0.00/0.00	0.00/0.00
C1	-5.20/-4.04	-3.88/-4.03	-4.24/-3.30	-2.90/-3.09
TS1a	5.67/3.12	5.71/5.77	4.09/1.42	4.13/4.05
P1a	-21.21/20.27	-21.36/-21.40	-21.91/-21.10	-22.06/-22.23
TS1b	2.18/-0.03	1.20/1.12	3.89/1.69	2.91/2.82
P1b	-28.98/-32.31	-28.46/-28.42	-24.90/-27.90	-24.38/-24.01

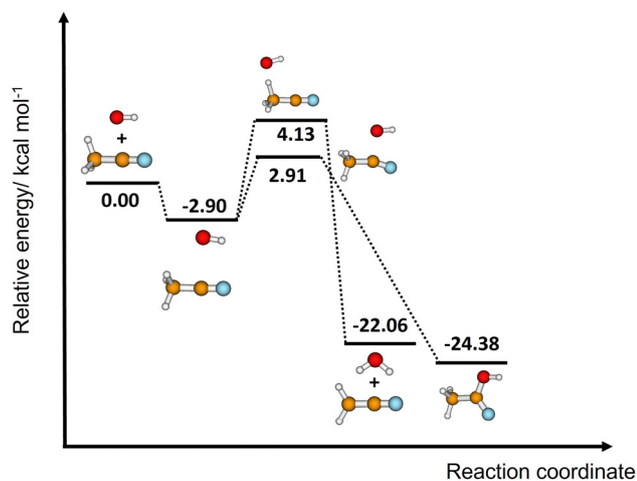


Fig. 5 Schematic ZPE-corrected energy diagram at the CCSD(T)-F12/M08-HX level of theory for reactions (1a) and (1b). See the last column of Table 4.

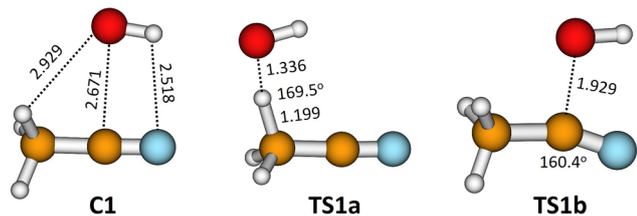


Fig. 6 Pre-reactive complex (C1), TS1a, and TS1b structures with some selected distances (in Å) and bond angles (in degrees). Geometries optimized at the M08/MG3S level.

OH + methylamine system.⁶⁸ In the case of TS1a, the torsional anharmonicity contains the rotation of the OH about the methyl group (with a harmonic frequency $\omega = 144.2$ cm⁻¹). For TS1b, there are two rotors, the internal rotation of the OH (with $\omega = 221.2$ cm⁻¹) and methyl (with $\omega = 140.8$ cm⁻¹) groups. For the two transition states, the treatment of these motions, as one-dimensional hindered rotors, instead of harmonic oscillators has a modest effect in the rate coefficients. The ratio between the anharmonic and harmonic partition functions is always smaller than 2 between 50 and 400 K (see the ESI†).

The total CCUS rate coefficients are plotted in Fig. 4 and summarized in Table 6 for selected temperatures. At the LPL, $k_a \gg k_1$ and the rate-determining reaction is always k_1 , so this rate coefficient and k^{CCUS} coincide. Above $T = 150$ K the thermal

rate coefficients evaluated at the two pressure limits coincide, this is because at this temperature and above the molecules populate the same levels with independence of the depth of the lowest energy level. In other words, the pre-reactive complex is irrelevant for the course of the reaction. However, below $T = 150$ K the HPL rate coefficients start to increase fast and at $T \sim 40$ K k_{HPL} becomes larger than the association rate coefficient, being k_a the rate-determining reaction. Although the agreement between theory and experiment obtained in this work is not perfect, the HPL results capture the wild variation of the rate coefficients with an astonishing increase of about four orders of magnitude in an interval of less than 100 K.

4.3. Comparison with previous studies

4.3.1. Experimental OH-reactivity trend towards CH₃X molecules (X = -CN, -OH, -NH₂, -CHO, CH₂OH, -C(O)CH₃...). The OH-reactivity and its temperature dependence at very low temperatures can be analysed for CH₃X molecules, such as CH₃CN, CH₃OH, CH₃NH₂, CH₃CHO, or CH₃CH₂OH. As it can be seen in Fig. 7, the T -dependence of $k(T)$ for the CH₃CN + OH reaction presents a “pipe-type” shape already observed for the CH₃OH + OH^{55,90,91} and CH₃CH₂OH + OH^{54,92} reactions. In the low temperature regime, these “pipe-type” shape plots are characterized by a drastic change in the OH-reactivity at a certain temperature followed by a slighter increase in $k(T)$ when the temperature approaches 10 K. The change in the trend for the T -dependence of $k(T)$ appears between 177.5 and 250 K for the CH₃CN + OH reaction, while the minimum $k(T)$ was observed at ~ 200 K for the reaction of OH with CH₃OH⁹⁰ or CH₃CH₂OH. For the CH₃C(O)CH₃ + OH reaction, the expected minimum $k(T)$ must lie somewhere between 300 K and 146 K.⁹³ In contrast, for the CH₃NH₂ + OH⁶⁸ and CH₃CHO + OH⁵⁶ reactions, a “hammock-type” shape was observed, *i.e.* once the change in trend in the T -dependence $k(T)$ is produced, it continuously increases at low temperatures. For CH₃CHO, the minimum $k(T)$ was observed at ~ 500 K.⁵⁶

4.3.2. Experimental CH₃CN + R reactions (R = OH, CN, C₂H, C(³P) or O(¹D)). When comparing the OH-reactivity towards CH₃CN, with that of other radicals, such as CN, C₂H, C(³P), or O(¹D), we find scarce kinetic information. Particularly interesting is the comparison between OH and CN reactions,⁸⁶ where a similar behaviour was observed in the low temperature regime. For the CH₃CN + CN reaction,⁸⁶ a “pipe-type” shape was also reported with an expected minimum $k(T)$ between 168 and 258 K. The interpretation of the increase in $k(T)$ when



Table 5 Variational and tunnelling transmission coefficients, and TST thermal rate coefficients (in $\text{cm}^3 \text{s}^{-1}$) at selected temperatures between 50 and 300 K

T (K)	$I_{1a}^{-\text{CVT}}$	$\kappa_{1a,\text{LPL}}^{\text{CVT/SCT}}$	$\kappa_{1a,\text{HPL}}^{\text{CVT/SCT}}$	k_{1a}^{TST}	$I_{1b}^{-\text{CVT}}$	$\kappa_{1b,\text{LPL}}^{\text{CVT/SCT}}$	$\kappa_{1b,\text{HPL}}^{\text{CVT/SCT}}$	k_{1b}^{TST}
50	1.22×10^{-5}	5.90×10^{12}	1.60×10^{14}	1.10×10^{-29}	0.740	4.73×10^8	1.67×10^{14}	1.18×10^{-24}
100	3.83×10^{-3}	1.13×10^4	1.48×10^4	4.32×10^{-21}	0.838	4.13×10^2	9.64×10^2	8.70×10^{-19}
150	2.63×10^{-2}	56.5	57.8	3.31×10^{-18}	0.854	11.4	11.4	7.33×10^{-17}
200	6.93×10^{-2}	8.87	8.89	9.91×10^{-17}	0.852	3.58	3.58	6.93×10^{-16}
300	0.181	2.56	2.56	3.50×10^{-15}	0.838	1.71	1.71	7.25×10^{-15}

decreasing temperature, given by Sleiman *et al.*⁸⁶ for the CN-reaction, is based on the increase in the probability of stabilization of the pre-reactive complex. The abrupt decrease in the rate coefficient observed at 168 K was related to the lack of stability of the $\text{CH}_3\text{CN} \cdot \text{CN}$ complex at higher temperatures since it decomposes on the same timescale as collision induced vibrational energy relaxation. As shown in Fig. 7, OH and CN radicals surprisingly present a similar reactivity towards CH_3CN and an analogous trend in temperature dependence.

For the $\text{C}_2\text{H} + \text{CH}_3\text{CN}$ reaction ($T = 165\text{--}360$ K),^{94,95} the combination of the kinetic data from Nizamov and Leone⁹⁴ and Hoobler and Leone⁹⁵ yields the following Arrhenius expression $(1.80 \pm 0.35) \times 10^{-11} \exp(-766 \pm 38)/T \text{ cm}^3 \text{ s}^{-1}$. An increase of almost one order of magnitude between 165 K and 296 K was reported by Nizamov and Leone.⁹⁴ However, the complete picture of the temperature dependence of the rate coefficient for the $\text{C}_2\text{H} + \text{CH}_3\text{CN}$ reaction cannot be seen at very low temperatures since no kinetic measurements were performed below 165 K.

For the $\text{C}(^3\text{P}) + \text{CH}_3\text{CN}$ reaction ($T = 50\text{--}296$ K), even though it is very fast, Hickson *et al.*⁹⁶ reported no temperature dependence of $k(T)$ in that temperature range. The same group also investigated the $\text{O}(^1\text{D}) + \text{CH}_3\text{CN}$ reaction at 75, 127 and 296 K,⁹⁷ obtaining rate coefficients close to the collision limit $((2.4\text{--}3.7) \times 10^{-10} \text{ cm}^3 \text{ s}^{-1})$ with a tendency to increase below 127 K. Nonetheless, the authors recommend using a temperature independent value of $(2.85 \pm 0.70) \times 10^{-10} \text{ cm}^3 \text{ s}^{-1}$ in the explored temperature range.

4.3.3. Comparison between experiments and theory. In Fig. 8, the complete picture of the temperature dependence of $k(T)$ between 10 and 3000 K is depicted, which includes both experimental and previous theoretical studies for comparison purposes. As shown in this figure, the LPL rate coefficient reported in this work increases at very low temperatures, but their values are more than three orders of magnitude smaller

than the present experimental results. In other words, these are closer to the HPL than to the LPL. In fact, the experimental results follow the same trend as those of the $\text{CH}_3\text{CN} + \text{CN}$ reaction⁸⁶ (see Fig. 7). At very low temperatures (below 50 K, in the theory, and below 140 K in the experiment) the $\text{CH}_3\text{CN} + \text{OH}$ reaction is controlled by the association reaction, with a very mild-decreasing temperature dependence. After that, there is a very rapid decrease in the rate coefficient with increasing temperature, which is mainly due to tunnelling at energies below the energy of reactants. We may consider that the dynamics of this region is controlled by tunnelling and by the depth of the pre-reactive complex. Finally, above 250 K, the temperature dependence of the rate coefficient k^{CCUS} is positive, and the reaction is dominated by tunnelling above reactants and by overbarrier dynamics. This is consistent with previously reported experimental and theoretical temperature dependencies of $k(T)$, which reveal an Arrhenius-like behaviour, with $k(T)$ decreasing while temperature is lowered,^{40,41,44,45,47,51,53} except for Alihosseini *et al.*⁵² who predicted a non-Arrhenius behaviour. Moreover, these authors overestimated the rate coefficients by several orders of magnitude in the experimental temperature range. Tian *et al.*⁵¹ also overestimated $k(T)$ in the entire T -range. Note that the extrapolated rate coefficients from the work of Li and Wang⁵³ and Tian *et al.*⁵¹ down to 150 K show that the crossing temperature with our extrapolated low-temperature data using eqn (E6) is 183 K (see Fig. 4). Therefore, the minimum $k(T)$ is expected to lie around 180 K.

As mentioned in the Introduction, several studies reported branching ratios for channels (1a) and (1b) at room temperature. More particularly, some evidence of the existence of the OH-addition route was pointed out as briefly discussed in the Introduction section and summarized in Tables 2 and 3.

Experimentally, Hynes and Wyne⁴⁵ claimed that the OH-addition channel (1b) accounted for about 50% of the total reaction process. This is, however, an indirect conclusion based

Table 6 Low- and high-pressure limit CCUS rate coefficients (in $\text{cm}^3 \text{ s}^{-1}$) at selected temperatures between 10 and 400 K

T (K)	k_{1a}^{CCUS} LPL	k_{1b}^{CCUS}	k^{CCUS}	k_{1a}^{CCUS} HPL	k_{1b}^{CCUS}	k^{CCUS}
10	1.083×10^{-37}	2.003×10^{-14}	2.003×10^{-14}	1.123×10^{-16}	1.275×10^{-9}	1.275×10^{-9}
50	7.89×10^{-22}	4.102×10^{-16}	4.102×10^{-16}	2.241×10^{-20}	1.266×10^{-10}	1.266×10^{-10}
100	1.867×10^{-19}	3.013×10^{-16}	3.013×10^{-16}	2.440×10^{-19}	7.032×10^{-16}	7.032×10^{-16}
150	4.192×10^{-18}	7.155×10^{-16}	7.204×10^{-16}	5.024×10^{-18}	7.161×10^{-16}	7.211×10^{-16}
200	6.063×10^{-17}	2.068×10^{-15}	2.129×10^{-15}	6.079×10^{-17}	2.068×10^{-15}	2.129×10^{-15}
300	1.600×10^{-15}	1.030×10^{-14}	1.190×10^{-14}	1.601×10^{-15}	1.030×10^{-14}	1.190×10^{-14}
400	1.152×10^{-14}	2.871×10^{-14}	4.022×10^{-14}	1.152×10^{-14}	2.871×10^{-14}	4.022×10^{-14}



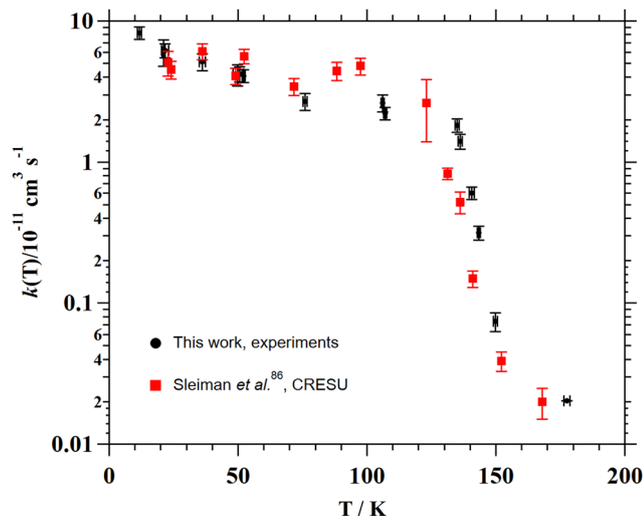


Fig. 7 Comparison between the temperature dependence of $k(T)$ for the OH- and CN-reaction of CH_3CN below 200 K.

on the observation that, in their reactor, the OH temporal decay was smaller in the presence of molecular oxygen than in its absence. The authors analysed this finding by considering that OH regeneration was occurring in their experiment. This was also observed for the OH + CD_3CN reaction in the presence of O_2 and they deduced that the OH-addition channel was necessarily open because the other potential product (CD_2CN radical) could obviously not release OH by reaction with O_2 . By simulating their various OH temporal decays they estimated that the two possible channels for reaction (1) have essentially similar rate coefficients. Later, Tyndall *et al.*⁴⁶ investigated the title reaction in a smog chamber filled with synthetic air. Infrared

observation of the terminal product HC(O)CN with a yield of $(40 \pm 20)\%$ led them to conclude that their measurement was consistent with the conclusions brought by Hynes and Wine⁴⁵ concerning the possible routes of reaction (1).

From a theoretical point of view, the OH-addition channel was only investigated by Galano⁵⁰ who also studied the reaction mechanisms in the presence of O_2 . She found that, in atmospheric conditions, reaction (1) will eventually lead to the formation of HC(O)CN in agreement with the smog chamber experiment from Tyndall *et al.*⁴⁶ Further, she demonstrated that molecular oxygen could attack the N of $\text{CH}_3\text{C(OH)N}$ – product from reaction (1b) – generating a hexagonal structure which will eventually release an OH radical agreeing thus with the suggestions of Hynes and Wine.⁴⁵ Although Galano⁵⁰ did not estimate the branching ratios for reactions (1a) and (1b), based on energetic considerations, she stated that reaction (1) should lead to the formation of both species CH_2CN and $\text{CH}_3\text{C(OH)N}$ in similar proportions. In the present work, we computed that the OH-addition reaction contributes 87% at 300 K (see ESI[†]), a temperature for which no pressure dependence is predicted. At lower temperatures, *i.e.* below 150 K, the OH-addition channel becomes the only viable route.

4.4. Astrochemical modelling

Since densities are very small in interstellar molecular clouds (typically 10^4 – 10^6 cm^{-3}), chemical reactions which are pressure dependent can be of potential interest in their LPL regime only. In a recent study concerning the OH + CH_3OH reaction,^{55,98} it has been shown that, at temperatures of typically 10–20 K, the total rate coefficients in the HPL and the LPL regimes were essentially similar. This is not the case for the present reaction according to our theoretical calculations which suggest that experimental measurements below 20 K are closer to the HPL than to the LPL. Hence, to evaluate the impact of the reaction presently studied on the chemical composition of cold dense clouds, where the gas kinetic temperature is as low as 10 K, we carried out chemical modelling calculations assuming the theoretical k_{CCUS} at the LPL ($2 \times 10^{-14} \text{ cm}^3 \text{ s}^{-1}$ at 10 K) and that the main channel is the formation of $\text{CH}_3\text{C(OH)N}$. We adopted typical parameters of cold dense clouds, *i.e.* a H_2 volume density of $2 \times 10^4 \text{ cm}^{-3}$, a visual extinction of 30 mag, and a cosmic-ray ionization rate of $1.3 \times 10^{-17} \text{ s}^{-1}$. We adopted the elemental abundances commonly known as “low-metal” values, in which the abundances of metals, silicon, and sulphur are reduced with respect to the values observed in diffuse clouds.⁹⁹ We used the chemical network rate.2012 from the UMIST database.¹⁰⁰ The results of the model show that reaction (1b) does not have any effect on the abundance of CH_3CN . According to the UMIST network, the main reactions of destruction of CH_3CN are those with cations, such as H_3O^+ , HCO^+ , and C^+ . The reaction with OH has a marginal (1%) contribution to the destruction of CH_3CN . Indeed, the reactions with H_3O^+ and HCO^+ are not fully effective in destroying CH_3CN because they produce the protonated form of CH_3CN , which partially gives back CH_3CN when recombining dissociatively

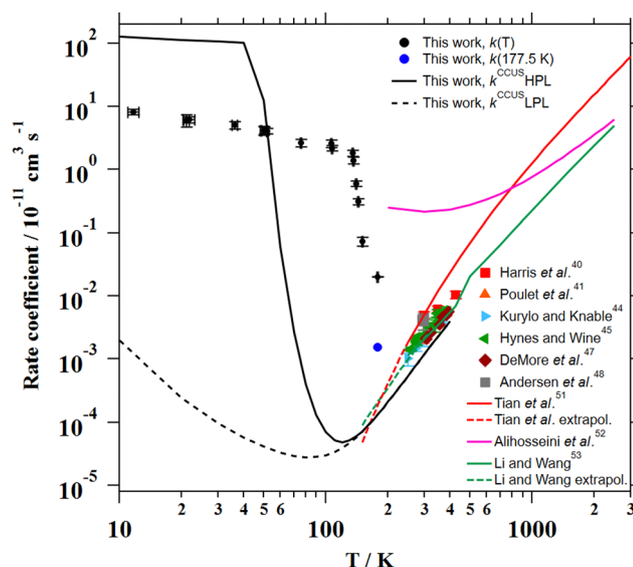


Fig. 8 Experimental and theoretical temperature dependencies of the rate coefficient for the $\text{CH}_3\text{CN} + \text{OH}$ reaction.



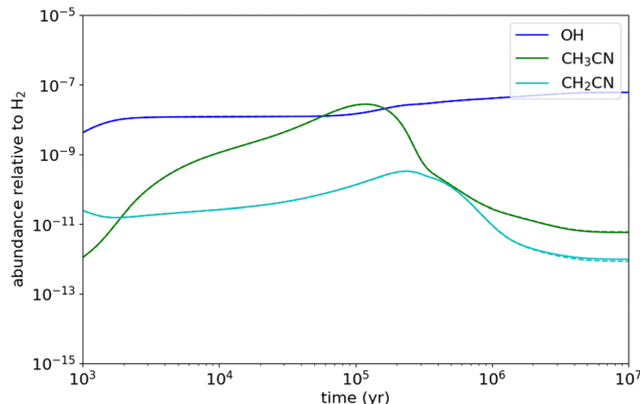


Fig. 9 Abundances relative to H_2 as a function of time calculated with the chemical model when excluding the $CH_3CN + OH$ reaction (dashed lines) and including it (solid lines) with a rate coefficient of $8.25 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$, as measured in this work at 11.7 K.

with electrons. The reaction with C^+ however is an efficient process of removal of CH_3CN . The rate coefficient of this reaction has been measured to be $5.6 \times 10^{-9} \text{ cm}^3 \text{ s}^{-1}$ at 300 K,¹⁰¹ and it is likely even higher at 10 K, following the usual behaviour of ion-polar reactions.¹⁰²

Even though if we assume the total rate coefficient measured at 11.7 K as a higher limit of the reactivity of CH_3CN , the impact of reaction (1) on the abundance of CH_3CN is very little (see Fig. 9). Furthermore, assuming as well that the CH_2CN radical would be the sole product, the $CH_3CN + OH$ reaction would contribute to its gas-phase formation with around 2%. According to the UMIST network, the main reaction forming CH_2CN is the $N + C_2H_3$ reaction. Nevertheless, the $N + C_2H_3$ reaction has been measured to be relatively fast, $7.7 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$, at 298 K, and the main product channel has been inferred to be $CH_2CN + H$.¹⁰³ It is however unknown whether this rate coefficient and product distribution holds also at the very low temperatures of cold dense clouds.

It is desirable to have experimental evidence of the formation of CH_2CN or $CH_3C(OH)CN$ radicals from the $CH_3CN + OH$ reaction at very low temperatures.

5. Conclusions

The present experimental results show that acetonitrile (CH_3CN) reacts very fast with OH radicals in the gas phase at very low temperatures (~ 10 – 100 K). Since the temperature behaviour of the rate coefficient varies in the explored experimental range, we provide two analytical expressions using a modified Arrhenius expression, $(k(T) = \alpha \left(\frac{T}{300 \text{ K}}\right)^\beta \exp\left(-\frac{\gamma}{T}\right))$, with $\alpha = 1.50 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$; $\beta = -0.53$ and $\gamma = 0$ K) between 11.7 K and 135 K and $\alpha = 1.5 \times 10^{-20} \text{ cm}^3 \text{ s}^{-1}$; $\beta = -26.4$ and $\gamma = 0$ K between 135 K and 149.9 K, a temperature within which the dependence of the rate coefficient $k(T)$ is extremely fast, changing more than one order of magnitude in only 15 K.

The computational results indicate that below 150 K, the reaction product is $CH_3C(OH)CN$ and not CH_2CN . Moreover, the

experimental rate coefficients have been found to be pressure independent at 22 K, 50 K and 106 K and appear to be much closer to the calculated high-pressure limit than to the low-pressure regime. Above 150 K, the contribution of the H-abstraction channel begins to increase, reaching 13% at 300 K and 29% at 400 K. It is worthwhile to point out that very efficient heavy-atom tunnelling has been highlighted by the present quantum calculations which is quite unusual.

According to the model ($T = 10$ K), the impact of the titled reaction on the abundances of CH_3CN appears to be negligible in dark molecular clouds of the ISM ($\sim 1\%$ of the total depletion reactions included in UMIST network). Respect to the potential formation of the CH_2CN radical in those environments, even in the most favourable scenario, where this radical could be formed in a 100% yield from the $CH_3CN + OH$ reaction, this route would only contribute around 2% to the current assumed formation routes by UMIST network.

Author contributions

DG and AC performed the experiments and data analysis and wrote the original draft of the paper; EMN and AFR carried out the theoretical calculations; MA and JC run the astrochemical model; BB, AC, and EJ designed and supervised the experiments; EJ got funds for carrying out the present experimental research; all authors contributed to the critical discussion and review of the draft.

Conflicts of interest

There are no conflicts to declare.

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