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An experimental, computational, and uncertainty analysis study of the rates of iodoalkane trapping by DABCO in solution phase organic media†

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NMR spectroscopy was used to measure the rates of the first and second substitution reactions between iodoalkane (R = Me, 1-butyl) and DABCO in methanol, acetonitrile and DMSO. Most of the reactions were recorded at three different temperatures, which permitted calculation of the activation parameters from Eyring and Arrhenius plots. Additionally, the reaction rate and heat of reaction for 1-iodobutane + DABCO in acetonitrile and DMSO were also measured using calorimetry. To help interpret experimental results, *ab initio* calculations were performed on the reactant, product, and transition state entities to understand structures, reaction enthalpies and activation parameters. Markov chain Monte Carlo statistical sampling was used to determine a distribution of kinetic rates with respect to the uncertainties in measured concentrations and correlations between parameters imposed by a kinetics model. The reactions with 1-iodobutane are found to be slower in all cases compared to reactions under similar conditions for iodomethane. This is due to steric crowding around the reaction centre for the larger butyl group compared to methyl which results in a larger activation energy for the reaction.

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Introduction

Nuclear magnetic resonance (NMR) spectroscopy is an important tool in structural biology, for example, where it can be used to study protein folding and dynamics.¹ NMR spectroscopy is also an invaluable quantitative technique that is well-established in analytical and physical chemistry research.^{2,3} Here, we have employed this technique to follow the time dependent behaviour of the reactants, intermediates and products^{3–5} of the single and double S_N2 substitution of 1,4-diazabicyclo[2.2.2]octane (DABCO) reaction with iodoalkanes in solution.^{6–8} DABCO is a tertiary amine and strong nucleophile. This Menshutkin reaction is of interest, not only from the fundamental science perspective,^{6–25} but also for nuclear safety applications, since DABCO-saturated materials are used in reprocessing plant ductwork filters,^{26–30} and to clean up iodine spills from nuclear reactors.^{31–36} During nuclear accidents, volatile iodine radioisotopes can be released as inorganic iodines (I₂, I_xO_y), and as organic iodides that are created *via*

reactions with hydrocarbons and other volatile organic chemicals.^{37–46} These compounds react in the environment in different ways: homogeneous and heterogeneous reactions in the atmosphere,^{47–50} and substitution reactions with aquatic, soil, and plant materials in terrestrial ecosystems.^{51–55} The deposition rate of organic iodides is approximately 200 times smaller than that of molecular iodide⁵⁰ and, consequently, these spread farther in the environment. One form of organic iodide of particular interest is iodomethane, which is the most abundant (as CH₃¹²⁷I) iodo-organic compound in the atmosphere,^{56,57} where it can influence the chemistry of ozone.^{47,49} An especially important radioisotope is ¹³¹I (*t*_{1/2} = 8.05 days), which poses a danger to human health,⁵⁸ and needs to be captured immediately after release, since it accumulates in the thyroid gland and influences metabolic processes and presents a considerable cancer risk.^{58–60} The iodoalkane-DABCO reaction serves as a simple chemical model for the interaction of iodoalkanes with a wide variety of organic materials that may be found in the environment.

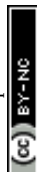
Here we present a multifaceted approach to determining kinetics and thermodynamic parameters for iodoalkane-DABCO S_N2 reactions. Unlike previously reported DABCO reaction investigations,^{6–8} we consider both first and second substitutions with iodoalkanes. We report reaction enthalpies, activation energies and kinetics for the first and second substitution of DABCO with iodomethane and 1-iodobutane in the perdeuterated

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solvents acetonitrile- d_3 (AcN), dimethyl sulfoxide- d_6 (DMSO), and methanol- d_4 (CD_3OD). *Ab initio* calculations have also been performed to determine transition state and product thermodynamic parameters, and transition state and product structures. The polarizable continuum model was added to the calculation to account for solvent shell effects which are known to have a considerable effect on stabilizing molecular species compared to the gas phase.⁶¹

Our approach illustrates how mechanistic kinetic modelling can be used to predict product and reactant concentrations forward in time,⁶² so that concentration levels at longer reaction times may, for example, serve to inform public health decisions. Also, this modelling can be run 'backward' in time to predict the concentration of materials at a potential origin of a reactant in time and location. We have used 1H NMR to illustrate how *in situ* methods available to track the concentrations of multiple species simultaneously increases the certainty of the kinetic modelling and provides an opportunity to increase and quantify uncertainty predictions.

A Markov chain Monte Carlo (MCMC) sampling method^{63–65} was used to determine the distribution of rate constants and initial concentrations for a kinetic model. This uncertainty quantification (UQ) method was used to propagate the uncertainty in the integrated concentration measurements onto the kinetics parameters and allows us to interpolate and extrapolate reactant and product concentrations and their uncertainties forward and backward in time. In environmental studies, this UQ method is an invaluable tool for tracing effluent sources and final deposition in time and location.^{66–69}

Methodology

Experimental

General. Perdeuterated solvents were purchased as anhydrous and sealed in ampoules from either Sigma-Aldrich or Cambridge Isotope Laboratories. Iodomethane was purchased from Acros Organics, deoxygenated during three freeze–pump–thaw cycles, and stored over copper chips under inert atmosphere at $-33\text{ }^\circ\text{C}$. 1-Iodobutane was purchased from Sigma-Aldrich, deoxygenated during three freeze–pump–thaw cycles, and stored in the dark in an inert atmosphere. 1,4-Diazabicyclo[2.2.2]octane (DABCO) was purchased from ACROS Organics and stored under an inert atmosphere at $-33\text{ }^\circ\text{C}$. 1-Methyl-1,4-diazabicyclo[2.2.2]octyl-iodide (Me-DABCO-I) was synthesized according to the literature.⁷⁰ The 1-butyl-1,4-diazabicyclo[2.2.2]octyl-iodide (Bu-DABCO-I) starting material was made in a similar manner. 1H NMR Spectra were recorded on a Varian INOVA 500 MHz spectrometer and referenced to residual proton signal of the appropriate deuterated solvent (i) CD_3HCN , (ii) $S(O)CD_3CD_2H$, and (iii) CHD_2OD .

Variable temperature NMR experiments

In a glove box, 1,4-diazabicyclo[2.2.2]octane (DABCO) was dissolved in the appropriate perdeuterated solvent. Next, 1.0 mL of this solution was drawn and put in an NMR tube and closed

with a septum. The tube contained a reference capillary of known concentration (standardization of the reference capillary is outlined below). The tube was then inserted into the NMR instrument and equilibrated at the appropriate temperature for 30 min. Then, the first spectrum was recorded and the number of moles of DABCO was calculated. Based on this calculation, the volume of iodoalkane required for a given DABCO-to-iodoalkane molar ratio was calculated using the density of the iodoalkane reagent given in Table S1 in ESI.† Addition was accomplished by ejecting the NMR sample tubes from the magnet, injecting the iodoalkane using a microliter syringe, shaking the tube for ~ 10 seconds, and then dropping the sample tube back into the magnet followed by immediate recording of a time series of spectra.

Standardization of the reference capillary ($C_6H_3(CF_3)_3$)

A flame-sealed capillary containing 1,3,5-tris(trifluoromethyl)benzene ($C_6H_3(CF_3)_3$) was put in an NMR tube containing perdeuterated AcN and 5 μL of CH_2Cl_2 . The number of moles of the reference standard in the capillary was calculated using the following equation:^{2,3}

$$\text{moles}_{\text{Standard}} = \frac{\text{Integral}_{\text{Standard}}}{\text{\#protons}_{\text{Standard}}} \times \frac{\text{\#protons}_{CH_2Cl_2}}{\text{Integral}_{CH_2Cl_2}} \times \text{moles}_{CH_2Cl_2}$$

Next, another 5 μL of CH_2Cl_2 was added and the standardization procedure was repeated. The results of the standardization procedure showed that there were 7.5×10^{-5} moles of ($C_6H_3(CF_3)_3$) in the reference capillary.

Temperature calibration of the NMR instrument was performed by setting the desired temperature and measuring the difference in the chemical shift of ethylene glycol signals: $T = (4.637 - \Delta)/0.009967$ (instrument operator's manual), where Δ is the shift difference in ppm between the CH_2 and OH signals.

NMR data workup

All NMR spectra were processed using Mestrenova (Mestrelabs version 14.2.0-26256, Mestrelabs Research S.L.) as follows: 1 Hz line broadening, manual phase correction, and automatic baseline correction (splines) were applied to each spectrum. These sets of data were exported for analytical analysis as .csv and .mnova files for further analysis. Concentration *versus* time curves from the MNova processed NMR spectra were fit to a kinetic model using the Berkeley Madonna (version 10.2.8) general purpose differential equation solver. Kinetic rate constants and starting reactant concentrations were fit as unbounded parameters.

Calorimetry

A Setaram C80 Calvet calorimeter was used to measure the heat of reaction for iodoalkane addition to DABCO in the protonated solvents AcN (CH_3CN) and DMSO ($S(O)CH_3CH_3$). In a typical experiment, the two compartments of a C80 vessel were charged with stock solutions of 1.0 mL stock solution of DABCO in AcN or DMSO and 1.0 mL stock solution of iodoalkanes in the same, corresponding solvent. The reference vessel was



charged with 2.0 mL of AcN or DMSO solvent. All samples were prepared in a glove box, sealed, and placed in the calorimeter at 30 °C. Once this temperature was reached and the heat flow had stabilized, the reaction was initiated by reversal mixing and data points were collected until heat flow returned to the baseline. Two experiments were conducted in AcN: 250 mM DABCO + 50 mM 1-iodobutane, and 125 mM DABCO + 50 mM 1-iodobutane. One experiment was conducted in DMSO: 250 mM DABCO + 50 mM 1-iodobutane. An additional measurement was performed using iodomethane in AcN reacting with DABCO, but the high volatility of the iodomethane called into question the results of this measurement. The DMSO stock solutions were used immediately after preparation to minimize the amount of 1-iodobutane-DMSO adduct formed slowly over time at ambient temperatures. The heat flow data was analysed using the Berkeley Madonna (version 10.2.8) general purpose differential equation solver to provide both kinetic and thermodynamic data. The heat flow rate is equal to the product of the reaction volume, the reaction enthalpy and reaction rate. The equation solver software takes into account the mixing time constant (13.7 s) and the instrument time constant (282 s) when calculating the reaction enthalpy and rate. A detailed description of this fitting procedure is given in ref. 71.

Uncertainty quantification

The uncertainty in kinetics parameters was measured by assuming their values are described by probability distributions. A statistical sampling method called Markov chain Monte Carlo (MCMC)^{63–65} was used to investigate the likelihood of a variety of possible values for the kinetics parameters with respect to the uncertainties in the measured concentrations and the correlations between parameters imposed by the kinetics model. These parameter distributions were then used to generate confidence intervals on the concentration profiles predicted by the model, and then the measured concentration uncertainties were calibrated until the confidence intervals explained as much of the variance in the measured data as possible. For each chemical reaction analysed, the time interval of data used to calculate the parameters was adjusted to avoid potentially inaccurate data near time 0, opting instead to allow later datapoints to constrain the initial concentrations. There are existing chemical dynamics analysis packages, such as Berkeley Madonna, which provide uncertainties on reaction rates given probability distributions on model parameters. These uncertainties are assumed and not derived from the data itself. One reason for the custom statistical approach developed herein is the added ability to quantify the impact of data error estimates on the model parameters. An assumption of perfect data can lead not only to poor fits but also underestimates of rate uncertainties.

To compare measured concentrations to the model, the time delays in the collection of NMR spectra were considered. The concentrations were estimated from peak integrals of the mean spectra, and the mean spectra averaged a set of N_e observed NMR spectra taken over a sequence of times $0 \leq t_0, t_1, t_2, \dots$,

$t_{N_e-1} < T$, where T is the collection interval. Due to rapidly changing concentrations due to the reaction kinetics, the mean of the peak integrals of individual spectra is not equal to the integral of the mean peaks. A model of the NMR-to-concentration measurement process was created to relate the measured concentrations to the kinetics model parameters. The kinetics model (a system of nonlinear ordinary differential equations) is integrated numerically using Python or MATLAB and the second order Heun's method to discretize the model solution, a set of time-dependent functions $c_j(t)$, $0 \leq t$ for the concentration of the j th species in the reaction model.⁷² Then, a measurement of the model analogous to averaging the species concentrations obtained from NMR peak integrals is given by

$$\overline{c_{j,d}} = \frac{1}{N_e} \sum_{k=0}^{N_e-1} c_j(t_k)$$

which is the arithmetic average of the concentrations predicted by the model at the times when NMR spectra were observed.

Integrated spectral peaks often introduce uncertainties on the order of 2% of the peak area. To account for the impact of concentration uncertainties, each concentration measurement $c_{j,d}$ is assumed to have a zero-mean Gaussian error, $\varepsilon_{j,d}$. The error standard deviation $\tau_{j,d}$ is initially taken to be the nominal 0.02 times the concentration magnitude, and then it is calibrated to match the predicted concentration uncertainty to the observed variation in the concentration measurements, which is described below and with key results in a following section. Due to the measurement errors, the relationship between the measured model and the experimental measurement is $c_{j,d} = \overline{c_{j,d}} + \varepsilon_{j,d}$. Then, the log-likelihood which models the probability of observing the measurements $c_{1,d}, \dots, c_{N_s,d}$ from each species involved in the reaction is

$$\log\text{-likelihood}(c_{1,d}, \dots, c_{N_s,d}) = -\frac{1}{2} \sum_{j=1}^{N_s} \frac{(c_{j,d} - \overline{c_{j,d}})^2}{\tau_{j,d}^2}$$

Concentration measurements at more than one time are required to estimate kinetics rates, and the log-likelihood of the full time series is the product of the log-likelihood at each individual time. Markov chain Monte Carlo is used to generate a sequence of predictions of kinetics parameters from correlated samples of the log-likelihood distribution. The statistics of these sequences provide uncertainties for the kinetics parameters, in particular, the mean and covariance. Notably, MCMC analysis allows the correlations between kinetics parameters to be estimated, such as coupling between reaction rates and initial concentrations.⁶³

The uncertainty in concentration measurements is not known *a priori*; the nominal relative error standard deviation of $\tau_{j,d} = 0.02$ is only a starting point. These species-dependent parameters influence the confidence intervals on the model parameters as well as the concentration profiles simulated by the model. If the model is accurate and the error parameters $\tau_{j,d}$ are correct for each species, then the confidence intervals for a given confidence level (e.g., 95%) on the concentration profiles

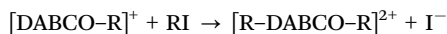
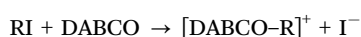


should contain the same percentage of the concentration measurements. An estimate of the true $\tau_{j,d}$ values can be made by iteratively adjusting them until the concentration profile confidence intervals contain the same percentage of concentration measurements as the level of significance of the confidence interval.

Quantum chemistry calculations

Calculations on discrete molecular models of the S_N2 reaction between DABCO and both iodomethane and 1-iodobutane were carried out using the NWChem software version 6.8.1⁷³ using the PBE0 hybrid functional⁷⁴ with Grimme's DFT-D3 corrections to represent dispersive interactions⁷⁵ in order to help interpret experimental observations. The augmented correlation-consistent polarized valence double zeta (aug-ccpvdz) basis set was used for all elements with the associated effective core potential for iodine.^{76,77} The conductor-like screening model (COSMO) of Klamt and Schüürmann was used to describe dielectric screening effects on the reactive species in a solvent environment.^{78,79}

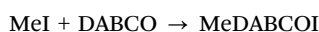
The enthalpies of the following chemical reactions corresponding to the first and second alkylations of the DABCO molecule ($R = \text{Me}, \text{Bu}$):



were determined by performing full geometry optimizations of the reactants and products, followed by a frequency calculation to verify the minimum energy configuration and to calculate the enthalpy. The transition state for each reaction was determined by performing a constrained potential energy scan along the DABCO(N)–R(C) reaction coordinate followed by a saddle point search conducted at the maximum energy on the curve. A frequency calculation was performed at the determined saddle point to obtain the enthalpy and entropy of activation for comparison with experimental data.

Results

The advantage of using time-resolved ^1H NMR spectroscopy to obtain kinetic rate information is the ability to perform the experiment *in situ* at a set temperature and the ability to track starting material, intermediates, and products over a time interval.⁸⁰ This allows for a multi-component analysis to fit a single rate constant; for example, the rate of decrease in DABCO and iodomethane and the rate of increase in the formation of the product, 1-methyl-1,4-diazabicyclo[2.2.2]octyl iodide ($\text{Me-DABCO}^+ + \text{I}^-$) must fit a single rate expression.



$$d(\text{MeI})/dt = -k_1[\text{MeI}][\text{DABCO}]$$

$$d(\text{DABCO})/dt = -k_1[\text{MeI}][\text{DABCO}]$$

$$d(\text{MeDABCOI})/dt = k_1[\text{MeI}][\text{DABCO}]$$

The challenge with using ^1H NMR spectroscopy for kinetic rate data is the limited time resolution on short time scales (<15 seconds). The detectable concentration range is limited by the number of equivalent protons that can be integrated to provide a concentration and the need to signal average a sufficient number of transient pulses to obtain a S/N ratio that provides a minimum of 4 distinct time points to follow kinetics over the first half-life. In our experiments, DABCO solution without added iodoalkane was equilibrated inside of the magnet at the desired temperature, followed by the tuning and shimming of the instrument. Next, the sample was quickly ejected, a measured quantity of iodoalkane was injected into the 5 mm NMR tube. After shaking tube manually to rapidly mix its contents, the NMR tube containing iodoalkane and DABCO was reinserted into the magnet. The whole process required about 15 seconds before accumulation of the first data point. For the NMR measurements, eight transients were coadded, setting the average time resolution for the NMR data sets at 23 seconds per time point. From these limitations, the ability to record 4 data points over the first half-life, and a minimum starting concentration of 0.01 M DABCO, we were limited to measuring rate constants $<1 \text{ M}^{-1} \text{ s}^{-1}$ for an expected half-life of *ca.* 100 seconds for a second order reaction.

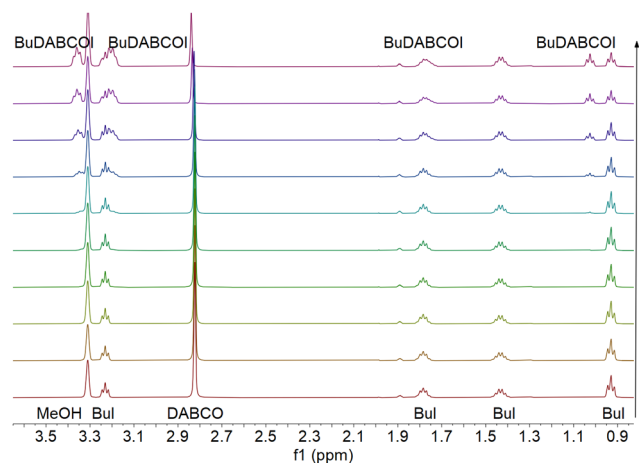
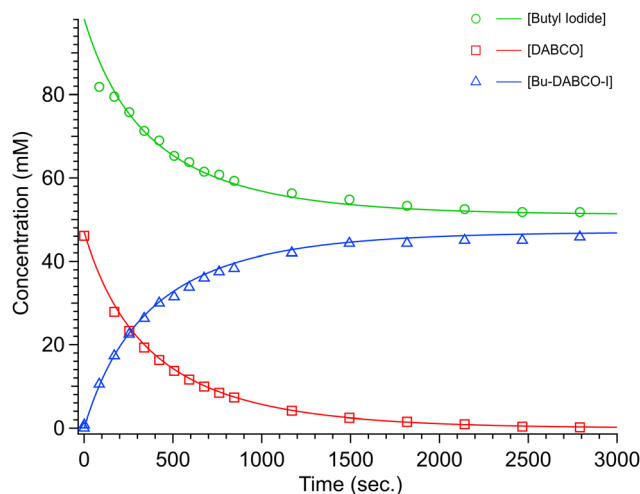
$$t_{1/2} \approx 1/k[\text{DABCO}] = 1/(1 \text{ M}^{-1} \text{ s}^{-1} \times [0.01 \text{ M}])$$

Based on the rate constants reported in the literature,⁷ the first substitution reaction of $\text{MeI} + \text{DABCO}$ in AcN (k_1 *ca.* $2.05 \text{ M}^{-1} \text{ s}^{-1}$ at 30°C) was expected to be too fast to be reliably followed by NMR spectroscopy; however, kinetics data could be recorded for the reaction run in methanol (k_1 *ca.* $0.014 \text{ M}^{-1} \text{ s}^{-1}$ at 30°C). We were unable to record kinetic data for the second substitution of DABCO with MeI , since the di-substituted ($\text{Me}_2\text{DABCOI}_2$) product precipitates out of AcN solution after reaching saturation. Disubstituted species always precipitate out of solution when acetonitrile is the solvent, no matter the temperature. Therefore, we measure the second substitution rates, k_2 , by starting from the well-defined monosubstituted species so that there is a species whose disappearance can be monitored by NMR with a high degree of confidence. The rate constant for the first substitution of DABCO with BuI was measured in AcN and DMSO at three temperatures (see Table 1). With methanol as the solvent, only the rate at 55°C was measured and was $k_1 = 0.41 \times 10^{-3} \text{ M}^{-1} \text{ s}^{-1}$. Based on this result, we did not attempt to measure the reaction at lower temperatures, hypothesizing that the degree of observable change (*via* the NMR method) would be too small to provide high-confidence results. The second substitution of DABCO with BuI progresses too slowly to measure it confidently in AcN or methanol; it could be measured, however, in DMSO . The rates, for both the mono-substitution and di-substitution reactions, increase as $\text{DMSO} > \text{AcN} > \text{CH}_3\text{OH}$. Fig. 1 shows an example NMR data set collected for the first substitution of DABCO with BuI in methanol at 55°C . Fig. 2 shows an example of a Berkeley Madonna fit of reactants and product concentrations *versus* time for $\text{DABCO} + 1.5 \text{ BuI}$ in AcN at 51°C .



Table 1 Reaction rates ($\text{M}^{-1} \text{s}^{-1}$) (with $\pm 1\sigma$ uncertainties) for DABCO and iodoalkane reactions in various solvents

	AcN-d ₃		DMSO-d ₆		Methanol-d ₄	
	Diff. eq.	MCMC	Diff. eq.	MCMC	Diff. eq.	MCMC
DABCO + MeI	Too fast		Too fast			
k_1					9.33×10^{-3} [30 °C]	$9.65(25) \times 10^{-3}$
k_1					2.081×10^{-2} [45 °C]	$2.156(79) \times 10^{-2}$
k_1					3.309×10^{-2} [55 °C]	$3.400(253) \times 10^{-2}$
DABCO + BuI			Too fast			
k_1	1.468×10^{-2} [30 °C]	$1.389(96) \times 10^{-2}$				
k_1	3.073×10^{-2} [51 °C]	$2.856(38) \times 10^{-2}$			4.1×10^{-4} [55 °C]	$3.0(1.2) \times 10^{-4}$
k_1	7.181×10^{-2} [78 °C]	$6.856(477) \times 10^{-2}$				
MeDABCOI + MeI						
k_2	1.00×10^{-3} [30 °C]	$1.02(7) \times 10^{-3}$	5.098×10^{-2} [30 °C]	$4.981(235) \times 10^{-2}$	5.20×10^{-5} [30 °C]	$5.26(37) \times 10^{-5}$
k_2	1.93×10^{-3} [51 °C]	$1.84(9) \times 10^{-3}$	1.094×10^{-1} [50 °C]	$9.351(525) \times 10^{-2}$	1.31×10^{-4} [45 °C]	$1.251(89) \times 10^{-4}$
k_2	4.28×10^{-3} [78 °C]	$3.55(30) \times 10^{-3}$	1.996×10^{-1} [70 °C]	$1.7804(1127) \times 10^{-1}$	2.37×10^{-4} [55 °C]	$2.152(64) \times 10^{-4}$
BuDABCOI + BuI	Too slow				Too slow	
k_2			2.4×10^{-4} [30 °C]	$2.0(1) \times 10^{-4}$		
k_2			8.4×10^{-4} [50 °C]	$7.4(6) \times 10^{-4}$		
k_2			2.16×10^{-3} [70 °C]	$2.04(5) \times 10^{-3}$		

**Fig. 1** An example of the NMR data collected during first substitution of DABCO with BuI in methanol-d₄ at 55 °C. The progression of spectra moving up is increasing with time. Overall, 102 spectra were recorded over 66 h and only select spectra are shown above.**Fig. 2** Concentration (mM) vs. time (s) plot for DABCO + 1.5 BuI in AcN at 51 °C. The open circles, triangles and squares represent collected experimental data. The solid traces are the Berkeley Madonna fit to the data.

The raw NMR data were processed with MNova as described above to determine concentrations and then analysed using the Berkeley Madonna software and MCMC methods to determine the reaction rate constants and initial concentrations. The temperature-dependent data were used to create Eyring and Arrhenius plots to determine reaction enthalpies and reaction activation parameters, respectively. The rate constants are presented in Table 1, the thermodynamic parameters in Table 2, and the initial reactant concentrations as determined by NMR in Table 3. See the ESI† documents for this article for further results details.

For Berkeley Madonna simulations, reactions were assumed to be second order and both reactants and products were fit to experimental data. Starting concentration of the product was set to 0 and concentrations of both reactants were fit without constraints.

The uncertainty quantification (UQ) method detailed in the Methodology section was used to propagate the uncertainty in the integrated concentration measurements onto the kinetics parameters (rate coefficients and initial concentrations). The kinetics parameters sampled by the MCMC procedure provided estimates and confidence intervals when analysed as a distribution. When these samples were used to generate confidence intervals on the concentration profiles, these bounds were used to scale the assumed concentration measurement uncertainty for each species until the percentage coverage of the measurements approximately matched the confidence level of the confidence interval. The joint probability distribution of the kinetics and initial concentration parameters were evaluated by estimating the histograms of the sampled parameters in Fig. 3. For each pair of parameters, the 1σ and 2σ confidence regions



Table 2 Activation energies, Gibbs energy parameters, and reaction enthalpies, (with $\pm 1\sigma$ uncertainties) for DABCO and iodoalkane reactions in various solvents. See the ESI for a discussion of the uncertainties. Quantities marked with a * were measured using calorimetry

	AcN-d ₃		DMSO-d ₆		Methanol-d ₄		Vacuum
	Exp.	Calc.	Exp.	Calc.	Exp.	Calc.	Calc.
DABCO + MeI	Too fast		Too fast				
E_a (kJ mol ⁻¹)		24.9		23.8	42(4)	30.3	68.7
$\Delta H^\ddagger$ (kJ mol ⁻¹)		26.7		22.6	39(4)	29.1	70.1
$\Delta S^\ddagger$ (J mol ⁻¹ K ⁻¹)		-17.0		-24.3	-154(11)	-35.0	-64.7
ΔH_{rxn} (kJ mol ⁻¹)		-147		-140		-141	+31
DABCO + BuI					Too slow		
E_a (kJ mol ⁻¹)	29(2)	35.2		34.2		38.4	83.2
$\Delta H^\ddagger$ (kJ mol ⁻¹)	26(4)	35.9		37.1		38.7	86.8
$\Delta S^\ddagger$ (J mol ⁻¹ K ⁻¹)	-193(6)	-53.0		-34.0		-40.4	-28.8
ΔH_{rxn} (kJ mol ⁻¹)	-126*/-122*	-128	-123*	-127		-125	-18
MeDABCOI + MeI							
E_a (kJ mol ⁻¹)	26(2)	35.6	30(2)		50(4)	36.5	121
$\Delta H^\ddagger$ (kJ mol ⁻¹)	24(2)	24.6	27(2)		47(4)	24.6	110
$\Delta S^\ddagger$ (J mol ⁻¹ K ⁻¹)	-224(7)	-127	-181(6)		-171(11)	-147	-92.5
ΔH_{rxn} (kJ mol ⁻¹)		-110				-102	+102
BuDABCOI + BuI	Too slow				Too slow		
E_a (kJ mol ⁻¹)			48(2)				+210
$\Delta H^\ddagger$ (kJ mol ⁻¹)			45(2)				
$\Delta S^\ddagger$ (J mol ⁻¹ K ⁻¹)			-166(7)				
ΔH_{rxn} (kJ mol ⁻¹)							

of the distribution were plotted and contain approximately 68% and 95% of the samples, respectively. Qualitatively, thin confidence regions slanted to the upper-right suggest the two parameters are positively correlated, while vertical or horizontal orientations suggest the two parameters are not correlated. Along each axis are the marginal probability distributions of each parameter, from which confidence intervals on those parameters are defined (see Tables 1 and 3).

For example, for BuI + DABCO (first substitution) in AcN at 50 °C, the kinetics rate and concentration confidence intervals are plotted in Fig. 3. The rate coefficients and initial conditions showed at least some correlation. Correlation was most significant among the initial conditions. This is consistent with the coupling of these parameters by the kinetics model and conservation of mass and can be predicted by a perturbation sensitivity analysis of the model. The initial concentration of the product, BuDABCOI, shows significant positive skewness, which is to be expected for a positive-definite parameter with sufficient variance. The confidence bounds on the concentration show coverage over 90–95% of the data (with the notable exception of the initial concentration for the product). This consistency is lost if the first three data points (including the initial concentration) are included in the dataset.

The 30 °C calorimetry results for the 250 mM DABCO + 50 mM 1-iodobutane measurements in AcN and DMSO are shown in Fig. 4. The results for the 125 mM DABCO + 50 mM 1-iodobutane are given in the ESI† for this article. The solid trace in these figures is from the Berkeley Madonna fit (see the Methodology section) and the markers are the measured data points. A datum was collected every second, but for clarity on the plots, markers are shown for every sixteenth data point. The 250 mM DABCO + 50 mM 1-iodobutane and 125 mM

DABCO + 50 mM 1-iodobutane calorimetry measurements gave enthalpies and reaction rates of $\Delta H = -126$ kJ mol⁻¹, $k_1 = 0.015$ M⁻¹ s⁻¹ and $\Delta H = -122$ kJ mol⁻¹, $k_1 = 0.016$ M⁻¹ s⁻¹, respectively, showing good agreement between the fit constants for the reaction in AcN. For the 250 mM DABCO + 50 mM 1-iodobutane in DMSO reaction, the calorimetry data give $\Delta H = -123$ kJ mol⁻¹ and $k_1 = 0.067$ M⁻¹ s⁻¹. These reaction enthalpies are reported in Table 2. The reactant vessels in the C80 are open to one another in the reaction chamber, and when we tried to measure the reaction rates and enthalpies of iodo-methane with DABCO we got results that are far from those of Kondo and co-workers.⁷ We suspect that the volatile iodo-methane was pre-reacting with DABCO before the measurement started.

The computational chemistry results provide several important insights into the reaction mechanism at the molecular level. The mechanism is thought to proceed *via* the reaction of two neutral molecules (iodoalkane and DABCO) reacting through an S_N2-type mechanism to produce two charged species (a quaternary ammonium cation and the iodide anion). The classical view of the S_N2 mechanism involves the attack of a nucleophile (in this case the amine nitrogen) at the backside of the reaction centre followed by the expulsion of an appropriate leaving group (iodide). Our calculations suggest that this is a good model for the process and show reasonable agreement with our calculated thermodynamic and kinetic parameters with available experimental data. The structure of the calculated transition state shows the expected approximate sp²-hybridized planar geometry at the reaction centre (see Fig. 5(a) for iodomethane). Given the significant change in charge during the reaction it might be expected that the polarity of the solvent would play a role in the mechanism,



Table 3 Initial concentrations (mM) of DABCO and monosubstituted DABCO, as determined by NMR method; concentrations of iodoalkane were calculated as outlined in the experimental section. MCMC initial concentration uncertainties are $\pm 1\sigma$

	NMR		MCMC		
DABCO + 1.5BuI	AcN-d ₃		DABCO	BuI	BuDABCOI
30 °C	DABCO	BuI	39.0(2)	68.8(1.7)	1(7)
51 °C	35.3	53.0	42.5(9)	94.4(9)	3.1(7)
78 °C	46.1	69.2	42.7(2.2)	115.5(2.3)	3.7(2.2)
MeDABCOI + MeI	MeDABCOI	MeI	MeDABCOI	MeI	Me ₂ DABCOI ₂
30 °C	57.8	57.8	61.6(3.6)	79.1(2.9)	2.2(1.4)
51 °C	64.5	64.5	63.1(3.0)	92.2(4.4)	3.2(2.5)
78 °C	68.3	68.3	63.8(7)	96(17)	2.0(1.1)
DABCO + 2BuI	DMSO-d ₆		DABCO	BuI	BuDABCOI
30 °C	DABCO	BuI	52.0(2.4)	^a	0.8(1.2)
50 °C	50.1	100.2	47.9(1.9)	^a	0.4(6)
70 °C	47.6	95.2	48.2(2.1)	^a	1.1(9)
MeDABCOI + MeI	MeDABCOI	MeI	MeDABCOI	MeI	Me ₂ DABCOI ₂
30 °C	9.38	15.8	8.7(3)	26.4(4)	0.7(2)
50 °C	8.75	16.5	8.8(3)	19.6(5)	1.8(3)
70 °C	9.75	13.5	10.5(3)	17.0(3)	0.3(2)
DABCO + MeI	Methanol-d ₄		DABCO	MeI	MeDABCOI
30 °C	DABCO	MeI	17.5(2)	27.7(2)	0.05(4)
45 °C	20.4	20.4	19.1(3)	32.6(3)	0.11(9)
55 °C	22.3	22.3	19.3(8)	29.8(5)	0.08(7)
DABCO + BuI	DABCO	BuI	DABCO	BuI	BuDABCOI
55 °C	21.0	21.0	21.0(5)	28.5(8)	0.9(6)
MeDABCOI + 2MeI	MeDABCOI	MeI	MeDABCOI	MeI	Me ₂ DABCOI ₂
30 °C	12.8	25.6	14.1(2.1)	36.0(2.0)	0.2(4)
45 °C	13.9	27.8	13.7(1.9)	44.4(1.7)	0.1(3)
55 °C	13.9	27.8	14.7(2.0)	38.1(1.2)	0.7(4)

^a BuI concentration could not be tracked for this reaction.

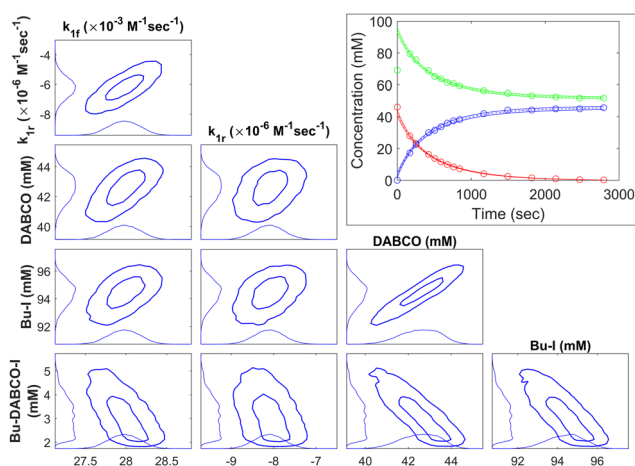


Fig. 3 Distribution of kinetics parameters and concentration profiles for BuI + DABCO (first substitution) in AcN at 50 °C due to uncertainty in concentration measurements: (left) covariances between kinetics parameters in each panel, and marginal distributions for each parameter along each axis, and (right) concentration data (circles) and concentration mean (dashed) and 95%-confidence intervals (solid).

and our calculations predict that is the case. Calculations in the gas phase predict a high barrier and positive or thermoneutral reaction free energies, whereas with the introduction of a continuum solvent model, the computed activation energies are reduced, and the reactions become more exothermic. This is likely due to the stabilization of the transition state and the reaction products, which have significant polar character. The change in activation energies can be related to the dielectric constant of the solvent, with dimethyl sulfoxide showing the largest effect; however, the dependence for the enthalpy of the reaction is less clear. There are some notable differences between the calculated mechanisms for 1-iodobutane compared to iodomethane. The addition of a butyl group to the reaction centre requires considerable conformational reorganization to accommodate nucleophilic attack (Fig. 5(b)) with the attacking amine centre approaching at a different angle. The presence of the additional alkyl chain also introduces increased opportunities for dispersive interactions between the components that can stabilize other geometries. For example, once the reaction is complete, iodide can shift away from the ammonium cation to exploit increased binding interactions with the side of the alkylated species.



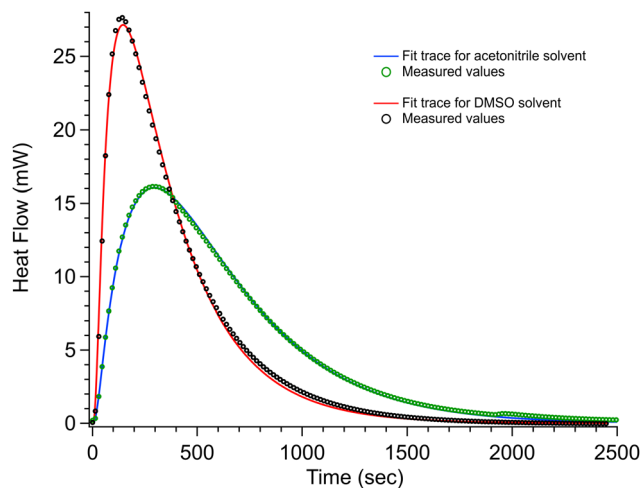


Fig. 4 Calorimetry data for the reaction 250 mM DABCO + 50 mM 1-iodobutane using AcN and then DMSO as solvents. The data were recorded using a Setaram C80 Calvet calorimeter operating at an equilibrium temperature of 30 °C. The markers in the plot are the measured data points and the solid lines are from the Berkeley Madonna fit of the data. For clarity, only 1/16 of the total number of data points are shown on the plot. In AcN, $k_1 = 0.015 \text{ M}^{-1} \text{ s}^{-1}$, $\Delta H = -126 \text{ kJ mol}^{-1}$, and in DMSO $k_1 = 0.067 \text{ M}^{-1} \text{ s}^{-1}$, $\Delta H = -123 \text{ kJ mol}^{-1}$.

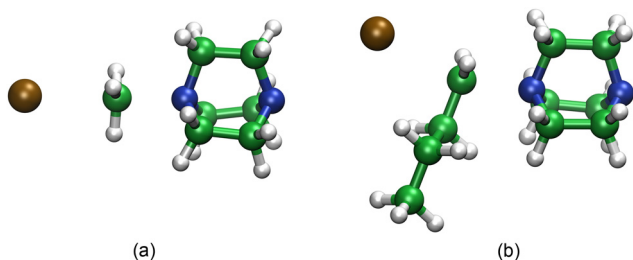


Fig. 5 Computed transition states for (a) MeI and DABCO, and (b) Bul and DABCO.

Discussion

This study details a method for integrating experimental measurements of the kinetics of a relatively simple chemical reaction with computational methods and uncertainty quantification tools to develop predictive models that can be used to extrapolate chemical change both forward and backward in time as a function of temperature when only a limited set of experimental data is available. We also considered the effect of different solvent environments on the reaction rate, which has been known for a considerable time to significantly change kinetics.^{81,82}

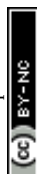
Our incorporation of the techniques of uncertainty quantification in the analysis provide considerable flexibility and value to our method. For example, complications arising from an inability of the NMR measurement to capture the initial data points in the time series at the beginning of the reaction, and also solubility limits causing the possible precipitation of reaction products can be included in the kinetic analysis in a systematic way and used to estimate errors. This approach

allowed us to put an approximate uncertainty on the initial concentration of a reactant to ascertain how precise the measurement was for our experimental technique. Usually, the uncertainty would be obtained by performing multiple measurements under the same conditions, which is not always possible, for example, when performing studies to inform emergency response for a radioactive iodine release.

Computed activation energies, in general, follow the experimentally observed trend, with reactions in DMSO being the fastest and in methanol the slowest. Calculations for reactions performed in vacuum appear to have the most unfavourable enthalpy changes, suggesting that in the absence of solvent the activated complex cannot be sufficiently stabilized. The measured and calculated negative values of the entropy of activation in each solvent support the assumption that the formation of the transition state is *via* an associative mechanism consistent with the S_N2 mechanism.

This study is also consistent with previous studies that have shown the difficulty of using computational methods to obtain accurate entropies of activation (and by extension the Gibbs free energy of activation).⁸³ For example, East and co-workers illustrate the problem using a different S_N2 reaction and show that errors of 8.4 kJ mol^{-1} lead to a change in the rate constants of a factor of 30. The rates depend on the exponential of the thermodynamic parameters (Eyring equation), so small changes in parameters can result in large changes in predicted rate constants.⁸⁴

The monosubstitution of DABCO with iodomethane at 30 °C was studied by Kondo and co-workers,⁷ in the 1980s. The focus of that paper was the influence of the solvent medium on the reaction rates between iodomethane and a base in the Menshutkin reaction. The authors compared single solvents as well as the binary mixtures of protonated AcN and protonated MeOH and concluded that the monosubstitution is about 150 times faster in AcN, even though the dielectric constants of these two solvents are very similar. We observe a similar trend, with good agreement with Kondo and co-workers for MeI reaction with DABCO in methanol- d_4 at room temperature and the rate in AcN is too fast for us to measure by NMR. However, we observe a different temperature dependence with our measured rates, slower at higher temperatures than Kondo reports resulting in different activation parameters, $\Delta H^\ddagger = 39.2(9) \text{ kJ mol}^{-1}$ (Kondo $\Delta H^\ddagger = 59.8 \text{ kJ mol}^{-1}$) and $\Delta S^\ddagger = -154(3) \text{ J K}^{-1} \text{ mol}^{-1}$ (Kondo $\Delta S^\ddagger = -83 \text{ J K}^{-1} \text{ mol}^{-1}$). The resulting difference in rates for MeI addition to DABCO in methanol at 60 °C, as calculated from the activation parameters, has the Kondo reaction rate greater than our rate by a factor of 2.7, which is outside our uncertainty limits obtained from simultaneously fitting the disappearance of DABCO and MeI in addition to the appearance of MeDABCOI. In our case we know that it is difficult to measure the initial concentration of iodomethane due to its high volatility. However, our UQ analysis permits us to obtain the initial concentration of MeI by fitting a series of reactions tied to the observed final concentration of the product, MeDABCOI. Specifically, we assume that all the MeI present in the experiment is trapped by DABCO and



thus the starting concentration of MeI should be equal to the final concentration of MeDABCOI. Furthermore, NMR analysis allows us to rule out the formation of other products since no new unidentified peaks are observed in the spectrum.

Another factor to consider when comparing our measured rates in deuterated solvents and Kondo's rates measured in the corresponding protonated solvents are kinetic isotope effects (KIE) on the reaction rate. We observe k_1 rates of 0.015 and 0.016 $\text{M}^{-1} \text{s}^{-1}$ for two independent BuI + DABCO C80 experiments in CH_3CN and it agrees well with our NMR experiment $k_1 = 0.014$ in CD_3CN . So, there is no KIE in AcN, but there could very well be a KIE in a protic solvent CD_3OD vs. CH_3OH . To follow the evolution of multiple species by ^1H NMR, that is, DABCO, MeI and MeDABCOI, we used deuterated solvents. However, to test the possibility of a solvent KIE as the observed difference in our rates compared to Kondo, we repeated the experiment at 55 °C in CH_3OH , where we observe the greatest difference in rates with Kondo. In this case the protons in the solvent CH_3OH 'swamp' the proton chemical shifts of MeI and MeDABCOI thus we can only follow the reaction with one parameter, the decay of DABCO. The data for the reaction is shown in the ESI† file SUPPLEMENTARY_INFORMATION_KINETICS_PLOTS_v3, and see the DABCO_MeI_CH3OH_55C worksheet. Also, see the MCMC analysis starting on page S3 of the ESI.† The Berkeley Madonna analysis of this data gave a reaction rate value of $k_1 = 6.0 \times 10^{-2} \text{M}^{-1} \text{s}^{-1}$ at 55 °C. The MCMC analysis of the same data set gave a rate value of $5.39(1.01) \times 10^{-2} \text{M}^{-1} \text{s}^{-1}$ at 55 °C. While we do observe a faster rate for disappearance of DABCO in CH_3OH , $k_1 = 5.39(1.01) \times 10^{-2} \text{M}^{-1} \text{s}^{-1}$ at 55 °C compared to the observed rate in CD_3OD , $k_1 = 3.400(253) \times 10^{-2} \text{M}^{-1} \text{s}^{-1}$ at 55 °C, the rate is still about half the rate we calculate from Kondo's activation parameters, *ca.* $1.2 \times 10^{-1} \text{M}^{-1} \text{s}^{-1}$ at 55 °C. Even though the data from a single parameter in CH_3OH has a higher degree of uncertainty, it is faster than the rate we measure in CD_3OD , suggesting a KIE; however, it is still slower than the rate measured by Kondo, so this control was not able to explain the discrepancy in observed rates. However, the focus of this paper is on using multiple parameters to obtain measured rates to enhance the UQ to make predictions both forward and backward in time to understand the initial concentrations and when concentrations are sufficiently low to provide a possible prediction of iodine compound concentrations in the environment.

The rates of reaction of BuI with DABCO are significantly slower as expected for an $\text{S}_{\text{N}}2$ reaction between a bulkier butyl group of 1-iodobutane compared to iodomethane. For example, the reaction of BuI with DABCO in AcN-d_3 k_1 (30 °C) *ca.* $1.389(96) \times 10^{-2} \text{M}^{-1} \text{s}^{-1}$ (MCMC value) is 2 orders of magnitude slower than the rate of reaction of MeI with DABCO in AcN reported by Kondo ($2.2 \text{M}^{-1} \text{s}^{-1}$).⁷

The time-resolved calorimetry experiments used in this study are slightly different than the approach used by Kondo. Kondo had the iodoalkane (MeI) in a sealed ampule in the calorimeter with a solution of DABCO. After the instrument baseline stabilized, they were able to break the seal of the

ampule to mix the two reactants. For the calorimeter used in this study, the iodoalkane and the DABCO were separated into two different chambers, but these chambers were open to one another in the calorimeter. Due to the volatile nature of MeI, the reaction started before the baseline was able to stabilize and we were unable to measure the ΔH_{rxn} for MeI to compare directly. However, 1-iodobutane is less volatile and a stable baseline was established before any significant reaction could take place. Thus, when the C80 instrument was changed into a mixing mode, the experiment provides a heat of reaction, ΔH_{rxn} *ca.* -123kJ mol^{-1} , for BuI with DABCO that is in good agreement with the heat of reaction for MeI reacting with DABCO in AcN reported by Kondo.⁷ The C80 approach further provides an alternate measurement of the reaction rate that is in good agreement with the NMR measured rate.

Looking in more detail at our data sets, we can see that comparison of the experimental and calculated kinetic parameters for the first substitution reaction of iodomethane and DABCO in methanol shows the best agreement between theory and experiment, and also shows a good comparison with previously reported values in the literature. For example, the reported value of k_1 by Kondo and co-workers is $1.37 \times 10^{-2} \text{M}^{-1} \text{s}^{-1}$,⁷ while the k_1 obtained from our NMR measurement is $9.65(25) \times 10^{-3} \text{M}^{-1} \text{s}^{-1}$, although there appears to be some ambiguity in the treatment of units in the previous study.⁷ Our work also uses *in situ* measurements of concentrations of reactants and products as a function of time rather than the *ex situ* quenching method employed by Kondo and co-workers.⁷

In the case of 1-iodobutane, examination of the calculated MCMC values of k_1 and k_2 rate constants (Table 1) show close correlation to the values obtained by experimental ^1H NMR methods.

In our approach, we included not only protic and aprotic solvents of differing dipole moments (for example, methanol and acetonitrile, Table S1 in ESI†) but also two aprotic solvents of similar dipole moments (acetonitrile and dimethyl sulphoxide). The range of solvent effects on the reactions can make data collection challenging in some cases for our experimental method. For example, in dimethyl sulphoxide, the first substitution of DABCO molecule is too fast to be observed by NMR spectroscopy, while in methanol, the same reaction occurs at a much slower rate. We attribute this result to the ability of DMSO to stabilize both the activated complex and charged products. Other complications can arise from the nature of the solvent. For example, in the case of methanol, the hydroxyl group can hydrogen bond to the basic nitrogen of the DABCO, potentially blocking approach of the iodoalkane required to enable the $\text{S}_{\text{N}}2$ reaction. In addition, dimethyl sulphoxide is known to react with iodomethane forming alkoxysulfonium salts *via* a sulphur ylide.⁸⁵ While the presence of this side product diminishes the concentration of available iodoalkane, it doesn't seem to have a significant influence on the overall kinetics of the reaction. This is probably because iodoalkane is always added in slight excess.



To support the idea that the iodoalkanes react with DMSO, we performed additional, “blank” reactions without the presence of DABCO. In this reaction, a drop of 1-iodobutane was added to DMSO- d_6 and a spectrum was recorded. An additional spectrum was taken after ~ 48 h with the NMR tube standing at room temperature. In addition to the signals from the 1-iodobutane, a new set of ^1H NMR signals appeared (Fig. S1 in ESI†). No attempt was made to assign these or ascertain a structure for the reaction product. For iodomethane reacting with DABCO in DMSO solvent, some signals in our ^1H NMR spectra were consistent with those in the literature where a trimethyl-oxosulfonium iodide ylide was formed when CH_3I was reacted with DMSO at room temperature.⁸⁵

Conclusions

Considering one example of simple substitution reactions, we have illustrated the utility of NMR spectroscopy as a powerful tool to obtain kinetic data.⁸⁰ Using the first and second substitution of DABCO with iodoalkanes in three different solvents and at three different temperatures, we have obtained raw data which was then refined in secondary software and used in kinetic modelling and uncertainty analysis. We have shown that while it is possible to obtain kinetic parameters using programs like Berkeley Madonna, it is not straightforward to ascertain uncertainty of the model using the same set of software. It is possible however, to analyse data using MCMC sampling and create a more accurate reaction profile that accounts for uncertainties of the experiment. Taken together, it is compelling to imagine that such a robust approach could be a useful tool to provide a degree of uncertainty analysis to predict the concentrations of reactants, intermediates, and products in both the forward and reverse directions when a limited amount of time-dependent data are available.

Author contributions

TA conceptualized the study and secured the funding. KG performed the NMR measurements. TA and KK performed the calorimetry measurements and TA analysed this data using Berkeley Madonna. TA and KG performed the initial kinetics analysis of the NMR data using Berkeley Madonna. NJH performed the *ab initio* calculations. WSR performed the uncertainty quantification and kinetics modelling. KG, SF and WSR performed the NMR data reduction. KG and TAB made the Eyring and Arrhenius plots and the uncertainty analysis for these. KG and TAB wrote the initial draft of the paper. KG, WSR, TA, NJH and TAB reviewed and edited the paper. All authors were responsible for data interpretation.

Conflicts of interest

There are no conflicts to declare.

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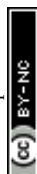
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