Decarboxylation of Hydrogencarbonatopentaamminecobalt(III) in Aquo-organic Solvent Media[†]

Anadi C. Dash,* Neelamadhab Dash, Prafulla K. Das and Jyotsnamayee Pradhan Department of Chemistry, Utkal University, Bhubaneswar-751004, India

The decarboxylation of hydrogencarbonatopentaamminecobalt(III) has been investigated in aqueous, 99% D₂O and aquo-organic solvent media (0–70 wt.% of cosolvent) at $15 \le t/^{\circ}C \le 40$ ($l = 0.02 \text{ mol dm}^{-3}$), using methanol, propan-2-ol, *tert*-butyl alcohol, ethylene glycol, acetone, acetonitrile, DMSO and ethylene carbonate as cosolvents. The solvent isotope effects on rate ($k_{H_2O}/k_{D_2O} = 1.0$ at 15–35 °C) and activation parameters ($\Delta H^{\neq} = 77.7 \pm 1.0$, $77.8 \pm 0.9 \text{ kJ mol}^{-1}$ and $\Delta S^{\neq} = 16 \pm 3$, $16 \pm 3 \text{ J} \text{ K}^{-1} \text{ mol}^{-1}$ for aqueous and 99% D₂O media, respectively) were negligible.

The decarboxylation rate constant increased with increasing mole fraction (X_{org}) of the cosolvent and the effect was pronounced at relatively high values of X_{org} for the dipolar aprotic cosolvents. This was attributed to a greater degree of destabilisation of the initial state as compared to the transition state with increasing mole fraction of the cosolvent. The $\ln k_s vs. 1/\varepsilon_s$ plots (k_s is the rate constant and ε_s the bulk relative permittivity) showed marked dependence on the nature of the cosolvents; the gradients of such plots generally increased with increasing dipole moment of the cosolvent molecules, indicating thereby that the solvation of the initial state and the transition state of the substrate is governed by the ion-dipole interactions between the water and cosolvent molecules. The relative transfer free energy of activation, $[\Delta\Delta G_t^+]_{(s+w)}$, decreased linearly with X_{org} for all mixed-solvent media, indicating that the preferential solvation effect is not significant. The activation enthalpy and entropy $vs. X_{org}$ plots displayed extrema suggesting that these thermodynamic parameters are sensitive to the structural changes in the bulk solvent phase. The solvent effects on ΔH^* and ΔS^* are mutually compensatory.

Previous studies from our laboratory and elsewhere have dealt with the solvolytic aquation of halogenoaminecobalt(III) complexes in mixed-solvent media.^{2,3} It is believed that such reactions undergo a dissociative interchange mode of ligand substitution at the cobalt(III) centre. The solvation of the ionic leaving groups strongly affects the reactivities of the substrates. Invariably the solvolysis rate decreases with increasing amount of non-aqueous component in the aquoorganic solvent media. The variation of rates and activation parameters was interpreted in terms of electrostatic and non-electrostatic medium effects, as well as solvent structural effects. In continuation of our work, we now report on the effect of solvent on the decarboxylation of the hydrogencarbonatopentaamminecobalt(III) ion.

It has been established by Harris et al.4 that the decarboxylation of $(NH_3)_5CoCO_3^+$ is acid catalysed. However, the rate is independent of the acidity of the medium once the carbonato complex is fully protonated $[pK \text{ of } (NH_3)_5 \text{CoCO}_3 \text{H}^{2+} = 6.7 \text{ at } 25 \,^{\circ}\text{C}, I = 0.5 \text{ mol } \text{dm}^{-3}].^5 \text{ The}$ decarboxylation of the hydrogencarbonato complex occurs with retention of a Co-O bond.⁶ This is a typical reaction in which the leaving group is essentially a non-dipolar molecule (CO_2) . Hence this reaction provides an opportunity to examine the medium effects on the differential solvation of the like charged initial state and the transition state for which there is likely to be little perturbation around the $(NH_3)_5Co^{3+}$ moiety, *i.e.* at a site several atoms away from the reaction site. Besides, there is currently wide interest in understanding the molecular dynamics of processes in mixedsolvent media of low water content which are of relevance to chemistry, biology and physiology. One such reaction of biochemical importance is the reversible hydration of CO_2 .⁷ From these considerations, investigation of the medium effects on the decarboxylation of the model carbonato complex was considered worthwhile.

In the present work we report a thorough study of the decarboxylation of $(NH_3)_5CoCO_3H^{2+}$ in mixed aquoorganic solvent media using dipolar protic and aprotic cosolvents. In contrast to the retarding effects of aquoorganic solvent media on the rates of aquation of halogenoamine-cobalt(III) substrates, we observed appreciable acceleration of the rate of decarboxylation of this hydrogencarbonato complex in all mixed-solvent media, the effect being relatively more significant for dipolar aprotic cosolvents.

Experimental

[Co(NH₃)₅CO₃]NO₃·H₂O, the carbonatopentaamminecobalt(III) nitrate, was prepared by the published method⁴ and recrystallised from water. The purity of the sample was further checked by Co and NH₃ analyses. (Found: Co, 20.6; NH₃, 29.3%. Calc. for [(NH₃)₅CoCO₃](NO₃)·H₂O: Co, 20.7; NH₃, 29.9%.) The UV-VIS spectrum displayed λ_{max} at 505 nm with $\varepsilon = 94.0$ dm³ mol⁻¹ cm⁻¹ [literature data,⁴ $\lambda_{max}/nm (\varepsilon/dm^3 mol^{-1} cm^{-1}): 505 (94)].$

AnalaR-grade (purity >99%) solvents[‡] were stored over molecular sieve (4A) and distilled before use. AnalaR-grade ethylene carbonate (purity >99.5%) was used without further purification. Water from a deioniser was redistilled from alkaline permanganate. D₂O (99.4%) was procured from BARC, Trombay, India. Solvent mixtures were prepared in wt.%. The UV-VIS spectra were recorded on a double-beam UV-VIS Hitachi spectrophotometer.

Kinetics

The rate of decomposition of the carbonato complex was followed at 510 nm using a HITECH SF 51 stopped-flow spectrophotometer equipped with a fully computerised data

⁺ This paper is part 14 of a series entitled Effect of Solvent on the Reactions of Coordination Complexes; Part 13 is ref. 1.

[‡] Abbreviations: methanol, MeOH; propan-2-ol, PrⁱOH; ethylene glycol, EG; *tert*-butyl alcohol, Bu'OH; acetone, AC; acetonitrile, AN; dimethylsulphoxide, DMSO; ethylene carbonate, EC.

acquisition and display system. The cell compartment and the reaction mixtures loaded in Hamilton syringes were thermostatted to ± 0.1 °C by circulating water from a thermostatic liquid circulator C-85 D through a cooler FC-200 (HITECH, UK). Equal volumes of thermostatted solutions of complex and HClO₄, both in the same solvent composition and loaded separately in syringes, were mixed under air pressure using a pneumatic drive mechanism. Final concentrations of the complex and HClO₄ in the reaction mixture were $(1-2) \times 10^{-3}$ mol dm⁻³ and ca. 2×10^{-2} mol dm⁻³, respectively. The exponential decrease of the absorbance with time, displayed as a voltage output was fitted to the relationship $(A_t - A_x) = (A_0 - A_x)\exp(-k_s t)$ where the terms have their usual meaning, by a computer programme available for the Apple II GS computer interface.

The mean value of the first-order rate constant and its standard deviation for a given solvent composition was computed from at least seven runs. The rate measurements were made at five temperatures from 15 to $35 \,^{\circ}$ C ($\Delta T = 20 \,^{\circ}$ C at $5 \,^{\circ}$ C intervals) for all solvent systems, except for ethylene carbonate-water for which the working temperatures were $15 \,^{\circ}$ C (for wt.% EC of 0-40) and 20-40 $\,^{\circ}$ C (for wt.% EC of 0-70) owing to the limited solubility of EC in water.

The activation parameters $(\Delta H^{\neq} \text{ and } \Delta S^{\neq})$ were calculated by fitting the rate data to eqn. (1)

$$\ln k_{\rm s}/T = (\ln k_{\rm B}/h + \Delta S^{\neq}/R) - \Delta H^{\neq}/RT \tag{1}$$

where $k_{\rm B}$ and *h* denote the Boltzmann constant and Planck's constant, respectively. The left-hand side of eqn. (1) was weighted inversely as its variance, which was calculated from the known variances of $k_{\rm s}$ and *T*.

Results and Discussion

Decarboxylation of $(\rm NH_3)_5\rm CoCO_3\rm H^{2+}$ in Aqueous and $\rm D_2\rm O$ Media

The rate and activation parameters for the decarboxylation of $(NH_3)_5CoCO_3H^{2+}$ in water and D_2O (*ca.* 99%) are collected in Table 1. The first-order rate constant is also independent of $[H^+]$ in the range 0.005–1.0 mol dm⁻³ indicating thereby that the hydrogencarbonato complex has little propensity for further protonation. The values of the rate and activation parameters (except for ΔS^{\pm}) reported by Harris *et al.*⁴ for the fully aqueous medium $[k = 1.25 \text{ s}^{-1} \text{ at } 25 ^{\circ}\text{C},$ $\Delta H^{\pm} = 71.4 \pm 2.1 \text{ kJ mol}^{-1}, \Delta S^{\pm} = -2 \pm 4 \text{ J K}^{-1} \text{ mol}^{-1},$ $I = 0.5 \text{ mol dm}^{-3}$ (NaClO₄)] compare satisfactorily with those reported in this work. The comparatively low value of ΔS^{\pm} reported by Harris *et al.*⁴ might be due to the high ionic strength maintained in their study. The observed solvent isotope effect on the rate $(k_{H_2O}/k_{D_2O} = 1.0, 15 \le t/^{\circ}C \le 35)$ and the activation parameters [ΔX^{\pm} (D_2O) $\approx \Delta X^{\pm}$ (H₂O), X = H, S (see Table 1)] further indicates that the differential

Table 1 Rate and activation parameters for the decarboxylation of $[Co(NH_3)_5CO_3H]^{2+}$ in fully aqueous and 99% D_2O media^a

$T/^{\circ}C^{b}$	$k_{\rm H_{2}O}/{\rm s}^{-1}$	k_{D_2O}/s^{-1}	$k_{\mathrm{H_2O}}/k_{\mathrm{D_2O}}$	
15.0 20.0 25.0 30.0 35.0 40.0	$\begin{array}{c} 0.34 \pm 0.03 \\ 0.63 \pm 0.03 \\ 1.10 \pm 0.05 \\ 1.92 \pm 0.06 \\ 3.10 \pm 0.10 \\ 5.10 \pm 0.25 \end{array}$	$\begin{array}{c} 0.342 \pm 0.008 \\ 0.62 \pm 0.02 \\ 1.11 \pm 0.06 \\ 1.94 \pm 0.13 \\ 3.09 \pm 0.07 \end{array}$	$\begin{array}{c} 0.99 \pm 0.09 \\ 1.02 \pm 0.06 \\ 0.99 \pm 0.07 \\ 0.99 \pm 0.07 \\ 1.00 \pm 0.04 \end{array}$	
	$\Delta H^{\neq}/\text{kJ mol}^{-1}$	$\Delta S^{\neq}/J \ K^{-1} \ mol^{-1}$	_	
H ₂ O D ₂ O	77.7 ± 1.0 77.8 ± 0.9	16 ± 3 16 ± 3		

^{*a*} [HClO₄] = 0.02 mol dm⁻³. ^{*b*} $\pm 0.1 \,^{\circ}$ C.

J. CHEM. SOC. FARADAY TRANS., 1991, VOL. 87

solvation effects of D_2O and H_2O are not significant. A twostep mechanism involving the zwitterionic intermediate which decarboxylates *via* rate-determining C-O bond breaking [see reaction (I)] is consistent with our observations.



This mechanism is equivalent to that proposed for the decarboxylation of the protonated hydrogencarbonate.⁸

Variation of Rate Constant with Solvent Composition

The rate data for different aquo-organic solvent media at 15–40 °C are collected in Table 2. The protic cosolvents exert comparable rate acceleration over the composition range studied. The rate acceleration due to the dipolar aprotic cosolvents is much stronger. Using the thermodynamic cycle as depicted in Scheme 1, the activation free energy (ΔG_s^{\pm}) for the mixed solvent (s) is given by eqn. (2)

$$\Delta G_{\rm s}^{\neq} = \Delta G_{g}^{\neq} + \Delta G_{\rm solv, \, t.s.}^{0} - \Delta G_{\rm solv, \, i.s.}^{0}$$
(2)

$$\begin{bmatrix} (NH_{3})_{5} CoOCO_{2}H \end{bmatrix}_{(g)}^{2+} \xrightarrow{\Delta G_{g}^{*}} \begin{bmatrix} (NH_{3})_{5} CoO --- CO_{2} \end{bmatrix}_{(g)}^{2+} \\ (i.s) & \downarrow \\ \downarrow \\ \Delta G_{Solv}^{\circ}(i.s) & \downarrow \\ \Delta G_{Solv}^{\circ}(i.s) & \uparrow \\ (t.s) & \downarrow \\ \begin{bmatrix} (NH_{3})_{5} CoOCO_{2}H \end{bmatrix}_{(s)}^{2+} \xrightarrow{\Delta G_{s}^{*}} \begin{bmatrix} (NH_{3})_{5} CoO --- CO_{2} \end{bmatrix}_{(s)}^{2+} \\ (i.s) & \downarrow \\ (i.s) & \downarrow \\ (i.s) & \downarrow \\ (t.s) & \downarrow \\ (t.s) & \downarrow \\ H & \downarrow \\ (t.s) & \downarrow \\ H & \downarrow \\ (t.s) & \downarrow \\ H & \downarrow \\ (t.s) & \downarrow \\ ($$

Scheme 1

where ΔG_s^{\neq} is the activation free energy in the gas phase and $\Delta G_{\text{solv, t.s.}}^0 - \Delta G_{\text{solv, i.s.}}^0$ denotes the difference in the free energy of solvation of the like charged transition state and the initial state. Thus the rate variation with solvent composition $[k_s = (k_{\beta} T/h)\exp(-\Delta G_s^{\neq}/RT)]$ is attributed to the effect of the differential solvation of the initial $[(NH_3)_5CoOCO_2H^{2+}]$ and transition $\{[(NH_3)_5CoOH\cdots CO_2]^{2+}\}$ states.

The electrostatic contribution to the free energy of solvation of an ion i (i = i.s. or t.s.), $[\Delta G_{solv,.}^0]_{el}$, is inversely dependent on the bulk relative permittivity (ε_s) of the medium.⁹ The plots of ln k_s vs. $1/\varepsilon_s$,¹⁰ and ln k_s vs. X_{org} at a fixed ε_s (=62.5) at 25 °C are shown in Fig. 1. The non-linearity in such plots indicates the importance of the non-electrostatic contribution to the solvation free energy. Note that the order of variation of the rate constant with respect to the cosolvents is

Table 2 Rate data for decarboxylation of $[Co(NH_3)_5CO_3H]^{2+}$ in mixed-solvent media with various wt.% organic solvent at different temperatures

	$k_{\rm s}/{\rm s}^{-1}$ a								
organic solvent	$T/^{\circ}\mathbf{C}$	5 wt.%	10 wt.%	20 wt.%	30 wt .%	40 wt.%	50 wt.%	60 wt.%	70 wt.%
МеОН	15.0	0.38 ± 0.01	0.37 ± 0.03	0.41 ± 0.01	0.43 ± 0.01	0.46 ± 0.01	0.55 ± 0.04	0.59 ± 0.03	0.68 ± 0.02
	20.0	0.67 ± 0.05	0.64 ± 0.03	0.69 <u>+</u> 0.04	0.73 ± 0.04	0.85 ± 0.02	0.91 <u>+</u> 0.04	1.09 ± 0.05	1.29 ± 0.07
	25.0	1.13 ± 0.08	1.15 ± 0.09	1.21 ± 0.09	1.31 ± 0.09	1.46 ± 0.05	1.69 ± 0.15	1.93 ± 0.06	2.24 ± 0.12
	30.0	2.09 ± 0.07	1.86 ± 0.10	2.19 ± 0.07	2.33 ± 0.09	2.56 ± 0.09	2.96 ± 0.07	3.24 ± 0.11	3.72 ± 0.12
	55.0	(0.029)	(0.059)	(0.123)	(0.194)	(0.273)	(0.360)	(0.457)	(0.569)
Pr ⁱ OH ^b	15.0	0.30 ± 0.02	0.32 ± 0.02	0.34 ± 0.04	0.36 ± 0.02	0.39 ± 0.02	0.44 ± 0.03	0.49 ± 0.03	0.61 ± 0.07
	20.0	0.54 ± 0.04	0.55 ± 0.04	0.64 ± 0.04	0.72 ± 0.05	0.73 ± 0.07	0.77 ± 0.05	0.96 ± 0.04	1.10 ± 0.05
	25.0	0.95 ± 0.04	0.98 ± 0.07	1.15 ± 0.10	1.21 ± 0.09	1.27 ± 0.09	1.34 ± 0.12	1.56 ± 0.06	1.86 ± 0.07
	30.0	1.67 ± 0.08	1.70 ± 0.08	2.00 ± 0.12	2.15 ± 0.14	2.32 ± 0.08	2.23 ± 0.15	3.00 ± 0.17	3.36 ± 0.16
	35.0	3.15 ± 0.24 (0.015)	(0.032)	3.06 ± 0.22 (0.070)	4.08 ± 0.18 (0.114)	4.26 ± 0.18 (0.166)	4.56 ± 0.27 (0.230)	4.68 ± 0.27 (0.309)	5.40 ± 0.13 (0.411)
Bu'OH	15.0	0.36 ± 0.05	0.37 ± 0.05	0.38 ± 0.04	0.36 ± 0.04	0.42 ± 0.06	0.47 ± 0.05	0.40 ± 0.03	
	20.0	0.67 ± 0.05	0.66 ± 0.03	0.67 ± 0.03	0.68 ± 0.04	0.68 ± 0.02	0.75 ± 0.05	0.80 ± 0.07	
	25.0	1.17 ± 0.04	1.19 ± 0.03	1.21 ± 0.04	1.24 ± 0.05	1.23 ± 0.03	1.32 ± 0.02	1.40 ± 0.12	
	30.0	1.87 ± 0.11 3.03 ± 0.10	1.89 ± 0.10 3.13 ± 0.10	1.88 ± 0.08 3.25 ± 0.15	1.91 ± 0.08	2.07 ± 0.11 3.82 ± 0.10	2.33 ± 0.13	2.40 ± 0.08 4.50 ± 0.11	_
	55.0	(0.013)	(0.026)	(0.057)	(0.094)	(0.139)	(0.196)	(0.267)	
EG	15.0	0.30 ± 0.05	0.36 ± 0.06	0.37 ± 0.09	0.40 ± 0.05	0.46 ± 0.05	0.39 ± 0.05	0.44 ± 0.04	0.54 ± 0.05
	20.0	0.63 ± 0.04	0.64 ± 0.05	0.68 ± 0.09	0.73 ± 0.03	0.79 ± 0.06	0.81 ± 0.04	0.88 ± 0.06	1.01 ± 0.03
	25.0	1.10 ± 0.15	1.18 ± 0.04	1.19 ± 0.03	1.20 ± 0.03	1.31 ± 0.06	1.34 ± 0.02	1.51 ± 0.05	1.71 ± 0.05
	30.0	1.99 ± 0.13 3.27 ± 0.16	1.90 ± 0.14 3.38 ± 0.12	1.99 ± 0.03 3.60 ± 0.32	2.11 ± 0.09 3.68 ± 0.21	2.22 ± 0.19 3.84 ± 0.12	2.37 ± 0.04 3.95 ± 0.14	2.07 ± 0.14 4.31 ± 0.07	2.94 ± 0.09 5 10 ± 0.06
	55.0	(0.015)	(0.031)	(0.068)	(0.111)	(0.162)	(0.225)	(0.303)	(0.404)
AC	15.0	0.40 ± 0.01	0.44 ± 0.01	0.47 ± 0.02	0.55 ± 0.03	0.65 ± 0.03	0.94 ± 0.07	1.20 ± 0.03	_
	20.0	0.63 ± 0.05	0.66 ± 0.07	0.80 ± 0.03	0.88 ± 0.07	1.14 ± 0.09	1.71 ± 0.09	2.17 ± 0.12	
	25.0	1.19 ± 0.07	1.19 ± 0.08	1.32 ± 0.06	1.75 ± 0.07	2.23 ± 0.09	3.02 ± 0.11	3.89 ± 0.12	
	30.0	2.02 ± 0.08 3.54 ± 0.27	2.13 ± 0.14 3.49 ± 0.32	2.20 ± 0.10 3.85 ± 0.18	2.96 ± 0.24 5 30 ± 0.44	3.07 ± 0.12 6.70 ± 0.52	5.18 ± 0.29 9.63 ± 0.80	0.30 ± 0.33	
	55.0	(0.016)	(0.033)	(0.072)	(0.117)	(0.171)	(0.237)	(0.317)	
AN	15.0	0.33 ± 0.08	0.39 ± 0.06	0.46 ± 0.04	0.55 ± 0.05	0.65 ± 0.04	0.88 ± 0.06	1.18 ± 0.07	
	20.0	0.64 ± 0.03	0.69 ± 0.03	0.81 ± 0.06	0.94 ± 0.06	1.20 ± 0.04	1.49 ± 0.10	2.06 ± 0.08	_
	25.0	1.09 ± 0.06 1.80 ± 0.13	1.18 ± 0.05 2.02 ± 0.08	1.49 ± 0.05 2.46 ± 0.09	1.73 ± 0.12	2.18 ± 0.10 4.11 ± 0.27	2.82 ± 0.23 5.06 ± 0.30	3.53 ± 0.19 6 37 ± 0.49	
	35.0	1.69 ± 0.13 3.22 + 0.29	2.02 ± 0.03 3 50 ± 0.12	2.40 ± 0.09 4 23 + 0.21	2.82 ± 0.21 4 71 + 0.19	4.11 ± 0.27 676 + 035	3.00 ± 0.39 8 36 ± 0.51	10.37 ± 0.49	
	55.0	(0.023)	(0.046)	(0.099)	(0.158)	(0.226)	(0.305)	(0.396)	
DMSO	15.0	0.36 ± 0.02	0.40 ± 0.03	0.38 ± 0.03	0.46 ± 0.05	0.57 ± 0.04	0.79 ± 0.04	1.17 ± 0.14	1.84 ± 0.11
	20.0	0.64 ± 0.05	0.66 ± 0.06	0.71 ± 0.07	0.83 ± 0.06	0.98 ± 0.09	1.43 ± 0.06	2.08 ± 0.08	3.04 ± 0.15
	25.0	1.17 ± 0.03	1.22 ± 0.05	1.37 ± 0.06	1.51 ± 0.06	1.86 ± 0.07	2.71 ± 0.15	3.54 ± 0.15 5.78 ± 0.25	5.12 ± 0.21
	35.0	2.04 ± 0.08 3 31 + 0.29	2.19 ± 0.03 3.73 ± 0.22	2.40 ± 0.09 3.93 + 0.44	2.08 ± 0.03 4.20 ± 0.39	5.22 ± 0.07 5.56 ± 0.50	4.34 ± 0.20 7.75 ± 0.50	9.78 ± 0.23 9.56 ± 0.50	154 ± 14
	55.0	(0.012)	(0.025)	(0.055)	(0.090)	(0.135)	(0.188)	(0.257)	(0.350)
EC	15.0	0.29 ± 0.05	0.42 ± 0.04	0.45 ± 0.05	0.56 ± 0.10	0.67 ± 0.04	_		
	20.0	0.64 ± 0.05	0.74 ± 0.06	0.83 ± 0.09	1.00 ± 0.05	1.15 ± 0.05	1.42 ± 0.12	1.74 ± 0.11	2.45 ± 0.16
	25.0	1.10 ± 0.04	1.29 ± 0.06	1.37 ± 0.06	1.70 ± 0.09	2.05 ± 0.13	2.63 ± 0.05	3.16 ± 0.21	4.35 ± 0.12 7.14 ± 0.20
	35.0	1.93 ± 0.12 3 30 \pm 0.18	2.03 ± 0.14 3 45 + 0.15	2.30 ± 0.09 4.02 ± 0.12	2.90 ± 0.09 5.01 + 0.17	5.03 ± 0.10 5.82 + 0.17	$\frac{4.34 \pm 0.18}{7.53 \pm 0.44}$	3.01 ± 0.21 9 30 + 0.45	114 ± 0.29
	40.0	5.19 ± 0.18	5.22 ± 0.15	6.46 ± 0.48	7.56 ± 0.35	9.72 ± 0.17	11.5 ± 0.9	14.7 + 1.4	17.4 + 1.1
		(0.010)	(0.022)	(0.049)	(0.080)	(0.120)	(0.169)	(0.234)	(0.323)

^{*a*} Values given in parentheses are mole fractions. ^{*b*} Values of k_s/s^{-1} for 2, 8 and 15 wt.% PrⁱOH at 25 °C are 0.92 ± 0.08, 0.97 ± 0.08, 1.07 ± 0.10, respectively.

Bu'OH < Pr^iOH < MeOH < EG < AC < AN \leq DMSO < EC for any value of $\varepsilon_s^{-1} \geq 2.0$. The dipole moment of the cosolvent molecules also varies in the same order except for the alcohols ($\mu/D^{\dagger} = 1.70, 1.66, 1.66, 2.28, 2.69, 3.44, 3.90$ and 4.87 for MeOH, PrⁱOH, Bu'OH, EG, AC, AN, DMSO and EC, respectively).^{10,11} This might indicate that solvation (and hence the rate) is at least partly governed by the ion-dipole interactions between the substrate and the cosolvent molecules.

An attempt was made to correlate the rate constant with Dimorth and Richardt's solvatochromic solvent polarity

 $+ 1 D ≈ 3.335 × 10^{-30} C m.$

parameter $E_{\rm T}$ (ref. 10, p. 188). The normalized values of $E_{\rm T}$ ($E_{\rm T}^{\rm N}$) for the mixed-solvent media used here decrease with increasing mole fraction ($X_{\rm org}$) of the cosolvent. The plots of ln $k_{\rm s}$ vs. $E_{\rm T}^{\rm N}$ (see Fig. 2) are non-linear. A similar trend has been reported for the solvolysis reactions of several inorganic and organic substrates including halogenoaminecobalt(III) complexes in mixed-solvent media.¹² The non-linearity in ln $k_{\rm s}$ vs. $E_{\rm T}^{\rm N}$ plots might be due to either (i) preferential solvation of the initial state and transition state of the reactant¹² or (ii) preferential solvation of the probe solute used for the evaluation of $E_{\rm T}$. An attempt is made here to analyse the rate dependence on solvent composition in terms of a simplified preferential solvation model proposed by Marcus (ref. 10, p.



Fig. 1 (a) $\ln(k_s/s^{-1})$ vs. $10^2/\varepsilon_s$ plots at 25 °C for mixed-solvent media: 1, MeOH; 2, PrⁱOH; 3, BuⁱOH; 4, EG; 5, AC; 6, AN; 7, DMSO; 8, EC. (b) $\ln(k_s/s^{-1})$ vs. X_{org} plot at 25 °C for $\varepsilon_s = 62.5$

206) and discussed at length in our earlier papers on the solvolysis of some halogenoaminecobalt(III) complexes in mixed-solvent media.^{2,13} Based on this model, the activation free energy of transfer, $[\Delta\Delta G_t^{\pm}]_{s-w} = (\Delta G_s^{\pm} - \Delta G_w^{\pm})$, where transfer of the reactant takes place from water (w) to the mixed solvent (s), is given by eqn. (3)

$$\begin{split} [\Delta\Delta G_{t}^{\neq}]_{s\leftarrow w} &= X_{\text{org}} [\Delta G_{t, t.s.}^{0} - \Delta G_{t, i.s.}^{0}]_{\text{org}\leftarrow w} \\ &+ \Delta g RT X_{\text{org}} (1 - X_{\text{org}}) \end{split}$$
(3)

where $\Delta g = g_{1.s.} - g_{i.s.}$ (i.e. the difference of the preferential solvation parameter for the transition state and the initial state) and $[\Delta G_{t, t.s.}^0 - \Delta G_{t, i.s.}^0]_{\text{org} \leftarrow w}$ denotes the relative transfer free energy for the transfer of the transition state and initial state from water to the pure organic cosolvent. Note that $[\Delta \Delta G_t^{\neq}]_{s \leftarrow w} \{k_s/k_w = \exp[(-\Delta \Delta G_t^{\neq}]_{s \leftarrow w}/RT\}$ varies linearly with X_{org} in accord with eqn. (4)

$$[\Delta\Delta G_1^{\neq}]_{s\leftarrow w} = a_i X_{org} + \delta_i \tag{4}$$

where the constants a_i and δ_i are sensitive to the nature of the cosolvent. The values of δ_i are only marginally different from zero, being statistically insignificant for most solvent systems at various temperatures (see Table 3). It is thus evident that the preferential solvation of the transition and initial states must be similar (*i.e.* $\Delta g = g_{t.s.} - g_{i.s.} = 0$) in all mixed-solvent media at all temperatures. The negative values of a (Table 3)



Fig. 2 $\ln(k_s/s^{-1}) vs. E_T^N$ plot at 25 °C for MeOH-water (\bigcirc), PrⁱOH-water (\triangle), AC-water (\bigcirc), AN-water (\bigcirc) and DMSO-water (\square) media

indicate that the initial state is destabilised to a greater extent than the transition state $([\Delta G^0_{t, i.s.}]_{org \leftarrow w} > [\Delta G^0_{t, t.s.}]_{org \leftarrow w})$ when transfer takes place from water to pure organic cosolvent.

The transfer chemical potential data¹⁴ for an analogous substrate, $(NH_3)_4CoCO_3^+$, suggest that MeOH-water and PrⁱOH-water media destabilise this substrate relative to water. The substrate used here $[(NH_3)_5CoOCO_2H^{2+}]$ is strongly hydrophilic and its interaction with the solvent is likely to be greater than that of $(NH_3)_4CoCO_3^+$. The solvent cospheres of both the initial and transition states are populated by both H₂O and the molecules of the cosolvent owing to favourable electrostatic interactions and hydrogenbonding effects. The values of the Savage-Wood group interaction parameters indicate that while interaction between alkyl groups will lead to a negative free energy change, favouring the solvation of the initial and transition states by the organic cosolvents, the interactions between the alkyl groups and OH and NH in the solvent cospheres of the substrate will destabilise it in the mixed-solvent media.15-17 This led us to believe that both the initial state and the transition state of the reactant will be destabilised by the mixed-solvent media. The observed rate acceleration is, however, attributed to greater destabilising effects of the mixed-solvent media on the compact initial state as compared to those on the expanded transition state.

The transition state for the decarboxylation reaction is associative in nature. As such, the principles of additivity,

$$[\Delta G^{0}_{t, t.s.}]_{s \leftarrow w} = [\Delta G_{t, (NH_3)sCoOH^{2+}} + \Delta G^{0}_{t, CO_2}]_{s \leftarrow w}$$

Table 3 Calculated values of $-a_i/kJ \mod^{-1}$ and $\delta_i/kJ \mod^{-1}$ for various solvents at different temperatures

	$-a_i/kJ \text{ mol}^{-1} (\delta_i/kJ \text{ mol}^{-1})^a$							
medium	15 °C	20 °C	25 °C	30 °C	35 °C			
MeOH-H ₂ O	2.39 ± 0.15	3.18 ± 0.20	3.41 ± 0.11	3.20 ± 0.22	2.64 ± 0.08			
-	(0.17 ± 0.05)	(0.14 ± 0.06)	(0.18 ± 0.02)	(0.01 ± 0.07)	(-0.22 ± 0.08)			
Pr ⁱ OH-H ₂ O	4.03 ± 0.13	4.24 ± 0.29	3.91 ± 0.21	4.39 ± 0.38	3.30 ± 0.04			
-	(0.32 ± 0.03)	(0.35 ± 0.07)	(0.32 ± 0.06)	(0.34 ± 0.08)	(-0.12 ± 0.10)			
Bu ^t OH-H ₂ O	1.40 ± 0.80	1.68 ± 0.41	1.48 ± 0.20	2.79 ± 0.32	3.97 ± 0.17			
-	(-0.18 ± 0.13)	(-0.04 ± 0.06)	(-0.14 ± 0.03)	(0.17 ± 0.05)	(0.09 ± 0.03)			
EG-H ₂ O	2.58 ± 0.62	2.78 ± 0.15	2.52 ± 0.21	2.82 ± 0.14	2.73 ± 0.19			
-	(0.01 ± 0.15)	(-0.01 ± 0.04)	(-0.01 ± 0.05)	(0.07 ± 0.03)	(-0.09 ± 0.05)			
AC-H ₂ O	8.69 ± 0.49	10.0 ± 0.7	10.4 ± 0.5	10.1 ± 0.8	10.0 ± 0.6			
-	(-0.22 ± 0.07)	(0.10 ± 0.13)	(0.11 ± 0.05)	(0.17 ± 0.14)	(-0.15 ± 0.12)			
AN-H ₂ O	7.70 ± 0.23	7.59 ± 0.13	7.78 ± 0.25	8.58 ± 0.39	8.08 ± 0.45			
2	(0.09 ± 0.06)	(0.15 ± 0.03)	(0.11 ± 0.05)	(0.23 ± 0.07)	(0.05 ± 0.10)			
DMSO-H ₂ O	11.8 ± 0.5	11.9 ± 0.4	11.1 ± 0.4	10.9 ± 0.5	11.4 + 0.6			
-	(0.15 ± 0.11)	(0.26 ± 0.09)	(0.06 ± 0.06)	(0.08 ± 0.10)	(-0.03 + 0.10)			
EC-H ₂ O ^b	14.1 ± 2.6	10.0 ± 0.5	10.7 ± 0.6	10.5 ± 0.8	10.6 ± 0.7			
-	(0.02 ± 0.21)	(-0.17 ± 0.08)	(-0.10 ± 0.10)	(-0.18 ± 0.13)	(-0.22 ± 0.11)			

^a Values given in parentheses are for δ_i . ^b At 40.0 °C, $a = -10.7 \pm 1.0$ kJ mol⁻¹ and $\delta = -0.10 \pm 0.14$ kJ mol⁻¹ for EC-H₂O.

J. CHEM. SOC. FARADAY TRANS., 1991, VOL. 87

are not applicable in assessing the importance of the transfer free energy of CO₂ on $[\Delta\Delta G_t^*]_{s \leftarrow w}$. $[\Delta G_{t, CO_2}^0]_{s \leftarrow w}^{298.2 \text{ K}}$ for MeOH-water media (molar scale) calculated from the available solubility data¹⁸ is negative at all X_{MeOH} and decreases with increasing X_{MeOH} . This is in keeping with the fact that the observed rate acceleration with increasing X_{MeOH} is at least partly due to the enhanced solvational stabilisation of CO₂ in this mixed-solvent medium relative to water.

From the relation

$$\begin{split} [\Delta G^{0}_{t, t.s.}]_{s \leftarrow w} &= \alpha [\Delta G^{0}_{t, (\mathsf{NH}_{3});\mathsf{CoOH}^{2+}} + \Delta G^{0}_{t, \mathsf{CO}_{2}}]_{s \leftarrow w} \\ &+ (1 - \alpha) [\Delta G^{0}_{t, t.s.}]_{s \leftarrow w} \end{split}$$

where α is a measure of the degree of C—O bond breaking in the transition state, eqn. (5) is obtained

$$\frac{1}{\alpha} \left[(\Delta \Delta G_{t}^{\neq} - \alpha \Delta G_{t, CO_{2}}^{0})_{s \leftarrow w} \right]$$
$$= (\Delta G_{t, (NH_{3})sCoOH^{2+}}^{0} - \Delta G_{t, i.s.}^{0})_{s \leftarrow w} \quad (5)$$

The left-hand side of eqn. (5) decreases steadily with increasing X_{MeOH} up to 0.35 for any value of α in the range 0.5–1.0. Fig. 3 depicts the trend in the variation of the left-hand side of eqn. (5) with X_{MeOH} for different chosen values of α from which it is evident that the initial state [*i.e.* $(NH_3)_5CoCO_3H^{2+}$] is relatively more destabilised than the final product cobalt(III) complex [*i.e.* $(NH_3)_5CoOH^{2+}$] when both are transferred from water to MeOH-water at the same composition. The limited data indicate that the solubility of CO_2 in EG-water at 25 °C (1 atm.) decreases from 0.0316 to



Fig. 3 $(1/\alpha)[\Delta\Delta G_t^{\neq} - \alpha\Delta G_{CO_2}^0]_{s \leftarrow w}/kJ \text{ mol}^{-1} vs. X_{MeOH} \text{ plot at } 25 \,^{\circ}\text{C}: \alpha = 0.5 (a), 0.6 (b), 0.7 (c), 0.8 (d), 0.9 (e), 1.0 (f)$

0.0295 mol dm⁻³ when X_{EG} increases from 0.02 to 0.06. The solubility data for CO₂ in other mixed-solvent media used in this study are not available. However, the solubility of CO₂ in water,¹⁸ acetone,²⁰ and DMSO²⁰ at 25 °C follow the trend: acetone > DMSO \ge water. This might indicate that the observed rate-enhancing effect of AC-water and DMSO-water media is partly due to the solvational stabilisation of CO₂.

Table 4 Mole fractions (X_{org}) at which extrema in the plots of ΔH^* (or ΔS^*) vs. X_{org} are observed for the solvolysis of halogenoaminecobalt(III) complexes and the decarboxylation of $(NH_3)_5 CoCO_3 H^{2+}$ in mixed-solvent media

				$X_{ m org}^{ m max}$	$(X_{ m org}^{ m min})^a$			
complex	MeOH	Pr ⁱ OH	Bu'OH	EG	AC	DMSO	EC	AN
1		0.03-0.07 0.04, 0.07 (0.11)	0.06					
2		(0.11)	0.025 (0.1 plt)				ca. 0.05	
3			0.02-0.03					
4	0.2–0.3	0.05-0.15	0.045-0.075			0.15		
5		(0.03) (0.1)	(0.05-0.06)					
6		(0.05), (0.1)	0.04					
8 7		(0.05) (0.1 plt)	(0.03-0.04) (0.07-0.1)					
8	0.1	(0.1 pit)	(0.07 0.1)					
ğ	0.1			0.07				
10	0.06 (0.28–0.45)			(0.11)	(0.07), (0.17)			
11	(0.05)	0.1 (0.3 plt)	0.1-0.15					
12	0.2 (0.03), (0.12)	0.17 (0.12)	(0.02–0.06)	(0.01)	(0.07-0.23 plt)			
13		. ,	(0.04-0.06)					
14			0.04–0.11 (0.16)					
15		0.05, 0.15 (0.1)	× ,					
16		0.07–0.10	0.02					0.02-0.04 (ca. 0.2 plt)
17	0.050.35	0.12	(0.015)		(0.035)	0.05-0.15 (0.3 plt)	(0.025)	0.22

^a Values in parentheses are for X_{org}^{min} ; plt; plateau. 1, (NH₃)₅CoCl²⁺;^{16,17} 2, trans-Co(en)₂Cl²;^{18,19} 3, cis-Co(en)₂Cl²;²⁰ 4, trans-Co(py)₄Cl²;²¹ 5, trans-Co(en)₂ClN₃⁺;²³ 7, cis-Co(en)₂ClCNS⁺;²⁴ 8, cis-Co(en)₂(bzmH)Br²⁺;²⁵ 9, cis-Co(en)₂(imH)Cl²⁺;²⁶ 10, cis-Co(en)₂ (tzNH₂)Br²⁺;²⁷ 11, cis-Co(en)₂(ethanolamine)Cl²⁺;² 12, cis-Co(en)₂(cyclohexylamine)Cl²⁺;^{11,11} 13, cis-Co(en)₂(2-aminopropan-2-ol)Cl²⁺;²⁸ 14, trans-Co(4EtPy)₄Cl²;²⁹ 15, trans-Co(4MePy)₄Cl²;³⁰ 16, cis-Co(en)₂NH₃Br²⁺;³¹ 17, (NH₃)₅COOCO₂H²⁺ (this work).



Fig. 4 (a) $\Delta H^{\pm}/kJ \mod^{-1} vs. X_{org}$ plot: 1, MeOH; 2, PrⁱOH; 3, BuⁱOH; 4, EG. (b) $\Delta H^{\pm}/kJ \mod^{-1} vs. X_{org}$ plot: 5, AC; 6, DMSO; 7, EC; 8, AN



Fig. 5 (a) $\Delta S^{\neq}/J \text{ K}^{-1} \text{ mol}^{-1} vs. X_{\text{org}} \text{ plot: 1, MeOH; 2, Pr^iOH; 3, Bu'OH; 4, EG. (b) <math>\Delta S^{\neq}/J \text{ K}^{-1} \text{ mol}^{-1} vs. X_{\text{org}} \text{ plot: 5, AC; 6, DMSO; 7, EC; 8, AN}$

Variation of Activation Parameters (ΔH^{\neq} and ΔS^{\neq}) with Solvent Composition

The activation enthalpies (Fig. 4) and entropies (Fig. 5) show distinct non-linear variation with solvent composition for all mixed-solvent systems except for MeOH-H₂O and EG-H₂O for which the trend is subdued by the experimental errors in these thermodynamic parameters. The mole fractions (X_{org}) at which extrema are exhibited in the plots of ΔH^{\neq} (or ΔS^{\neq}) vs. X_{org} (Table 4) also closely agree with those reported for the solvolytic reactions of halogenoaminecobalt(III) complexes^{1,2,13,21-36} in the corresponding mixed-solvent

Downloaded by Korea Advanced Institute of Science & Technology / KAIST on 06 October 2011

Published on 01 January 1991 on http://pubs.rsc.org | doi:10.1039/FT9918703753

media. Note that the extrema are observed for the decarboxylation reaction for which the effects of differential solvation of the initial and transition states in highly aqueous binaries are likely to be minimal. We submit that this is due to variation of the solvational components of the activation parameters with solvent structure which is modified with the addition of the cosolvent to water.^{37–39}

In contrast with all other media, ΔH^{\neq} (or ΔS^{\pm}) for AN-H₂O remained virtually independent of composition up to $X_{AN} = 0.15$ and then increased to attain a sharp maximum at $X_{AN} = 0.225$, indicating a pronounced change in solvent structure around $X_{AN} \approx 0.2$. This is consistent with the suggestion of Esteal *et al.*⁴⁰ that a pronounced structural change due to breakdown of the water structure sets in around $X_{AN} \approx 0.15$ -0.2 after the hydrogen-bonded water network becomes saturated with weakly interacting acetonitrile molecules. Studies of the thermodynamic and transport properties of the DMSO-water system have led to the generally accepted conclusion that the DMSO-H₂O interaction due to hydrogen bonding is at its maximum in the range of X_{DMSO} of 0.3-0.4.³⁹⁻⁴¹ Strikingly, ΔH^{\neq} and ΔS^{\neq} tend to attain plateau minima in the range of X_{DMSO} of 0.25-0.35.

The variations of ΔH^{\neq} and ΔS^{\neq} with solvent composition are mutually compensatory, as is evident from the linearity in the plot of $\Delta H^{\neq}/kJ \text{ mol}^{-1} vs$. $\Delta S^{\neq}/J \text{ K}^{-1} \text{ mol}^{-1}$ (not shown) which satisfactorily accommodates all data points.

We are grateful to Dr. J. Burgess, University of Leicester, for many helpful suggestions on this work. J.P. thanks the U.G.C., New Delhi for a Senior Research Fellowship. N.M.D. thanks the CSIR, New Delhi for a Senior Research Fellowship and P.K.D. thanks the Govt. of Orissa, Education Department, for grant of study leave. Our thanks are due to the Department of Science and Technology, Government of India for generous funding to procure the stopped-flow spectrophotometer.

References

- 1 A. C. Dash and J. Pradhan, Indian J. Chem., Sect. A, 1990, 29, 379.
- 2 A. C. Dash and P. K. Das, J. Chem. Soc., Faraday Trans. 1, 1989, 85, 2405.
- 3 K. H. M. Halawani and C. F. Wells, J. Chem. Soc., Faraday Trans. 1, 1990, 86, 1741; O. Grancicova and V. Holba, Transition Met. Chem. (Weinheim, Ger.), 1984, 9, 322.
- 4 T. P. Dasgupta and G. M. Harris, J. Am. Chem. Soc., 1968, 90, 6360.
- 5 E. Chaffee, T. P. Dasgupta and G. M. Harris, J. Am. Chem. Soc., 1973, 95, 4169.
- 6 J. B. Hunt, A. C. Rutenberg and H. Taube, J. Am. Chem. Soc., 1952, 74, 268.
- 7 D. N. Silverman and S. Lindskog, Acc. Chem. Res., 1988, 21, 36.

- 8 P. Paneth and M. H. O'Leary, J. Am. Chem. Soc., 1983, 107, 7381.
- 9 J. O'M. Bockris and A. K. N. Reddy, Modern Electrochemistry, Planum New York 1970, vol. 1, 77, 80
- Plenum, New York, 1970, vol. 1, pp. 57, 80. 10 Y. Marcus, *Ion Solvation*, John Wiley, New York, 1985, p. 186.
- 11 CRC Handbook of Chemistry and Physics, CRC, Boca Raton, Florida, 60th edn., 1979, p. E-64.
- 12 J. G. Dawber, J. Chem. Soc., Faraday Trans., 1990, 86, 287.
- 13 A.C. Dash and J. Pradhan, J. Chem. Soc., Faraday Trans. 1, 1989, 85, 2797.
- 14 M. J. Blandamer and J. Burgess, Transition Met. Chem. (Weinheim, Ger.), 1988, 13, 1.
- 15 J. J. Savage and R. H. Wood, J. Solution Chem., 1976, 5, 753.
- 16 S. K. Suri, J. J. Spitzer, R. H. Wood, E. G. Abel and P. T. Thompson, J. Solution Chem., 1985, 14, 781.
- 17 J. J. Spitzer, S. K. Suri and R. H. Wood, J. Solution Chem., 1985, 14, 561.
- 18 M. Takahashi, Y. Kobayshi and H. Takeuchi, J. Chem. Engng. Data, 1982, 27, 328.
- 19 A. Yasunishi and F. Yoshida, J. Chem. Eng. Data, 1979, 24, 11.
- 20 E. Wilhelm and R. Battino, Chem. Rev., 1973, 73, 1.
- 21 A. C. Dash, N. C. Naik and R. K. Nanda, Indian J. Chem., Sect. A, 1988, 27, 294.
- 22 K. H. Halawani and C. F. Wells, J. Chem. Res. (S), 1988, 58; J. Chem. Res. (M), 1988, 501.
- 23 G. S. Groves, K. H. Halawani and C. F. Wells, J. Solution Chem., 1987, 16, 399.
- 24 G. S. Groves and C. F. Wells, J. Chem. Soc., Faraday Trans. 1, 1985, 81, 2475.
- 25 G. S. Groves, A. F. M. Nazer and C. F. Wells, J. Chem. Soc., Faraday Trans. 1, 1987, 83, 1119.
- 26 C. N. Elgy and C. F. Wells, J. Chem. Soc., Dalton Trans., 1980, 2405; J. Chem. Soc., Faraday Trans. 1, 1983, 79, 2367; 1985, 81, 2145.
- 27 A. E. Eid and C. F. Wells, J. Chem. Soc., Faraday Trans. 1, 1983, 79, 253.
- 28 A. E. Eid and C. F. Wells, Transition Met. Chem. (Weinheim, Ger.), 1985, 10, 223.
- 29 A. E. Eid and C. F. Wells, J. Chem. Soc., Faraday Trans. 1, 1985, 81, 1401; 1986, 82, 1643.
- 30 A. C. Dash and N. Dash, J. Chem. Soc., Faraday Trans. 1, 1987, 83, 2505.
- 31 A. C. Dash and N. Dash, J. Chem. Soc., Faraday Trans. 1, 1988, 84, 75.
- 32 A. C. Dash and J. Pradhan, J. Chem. Soc., Faraday Trans. 1, 1988, 84, 2387; A. C. Dash and J. Pradhan, Indian J. Chem., Sect. A, 1988, 27, 772.
- 33 A. C. Dash and P. K. Dash, Int. J. Chem. Kinet., 1990, 22, 307.
- 34 I. M. Sidahmed and A. M. Ismail, Transition Met. Chem. (Weinheim, Ger.), 1987, 72, 332.
- 35 I. M. Sidahmed and C. F. Wells, J. Chem. Soc., Dalton Trans., 1983, 1035.
- 36 V. Holba and O. Grancicova, J. Inorg. Nucl. Chem., 1981, 43, 2071.
- 37 F. Franks and D. J. G. Ives, Quart. Rev. Chem. Soc., 1966, 20, 1.
- 38 M. J. Blandamer and J. Burgess, Chem. Soc. Rev., 1975, 4, 55.
- 39 M. J. Blandamer, Adv. Phys. Org. Chem., 1977, 14, 247.
- 40 A. J. Easteal, Aust. J. Chem., 1979, 32, 1379; 1980, 33, 1667; S. Balakrishna and A. J. Easteal, Aust. J. Chem., 1981, 34, 943.
- 41 D. D. MacDonald and J. B. Hyne, Can. J. Chem., 1971, 49, 611.

Paper 1/02213J; Received 10th May, 1991