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We report *in situ* high-pressure NMR kinetic studies of catalytic oxidations inside heterogeneous catalysts exploiting distortionless enhancement by polarisation transfer (DEPT) ¹³C NMR. ¹H NMR diffusion and relaxation time measurements are then used to elucidate trends in reaction kinetics in different solvents. The work shows the feasibility of non-invasively monitoring intraparticle kinetics, transport and adsorption in porous catalysts using a comprehensive NMR toolkit.

The application of NMR spectroscopy to study chemical reactions in solid catalysts has grown in recent years. The approach offers several advantages as it allows one to carry out reaction studies non-invasively, exploiting the intrinsic ability of NMR to resolve the structure of chemical species formed during reaction and, unlike conventional approaches based on analysis of bulk fluids surrounding the catalyst particles, allows probing directly chemical kinetics and composition within the catalyst particles. Using such an approach it becomes possible to resolve chemical compositions of species inside the pore space, as for example in heterogeneous catalysis. This has recently been demonstrated when monitoring the evolution of conversion and product distribution during the heterogeneous catalytic ethene oligomerisation reaction at 110 °C and 28 barg over a 1 wt% Ni/SiO₂-Al₂O₃ catalyst.¹

Several other examples of such studies have been reported in the literature. For example, Koptyug *et al.*² have used ¹H NMR to produce spatially resolved spectra within a 2D slice along the axial direction of a fixed bed of 1% Pd/Al₂O₃

In situ high-pressure ¹³C/¹H NMR reaction studies of benzyl alcohol oxidation over a Pd/Al₂O₃ catalyst[†]

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catalyst pellets. The reaction considered was that of the hydrogenation of α -methyl styrene to cumene. Although the spectra showed evidence of changes in chemical composition along the length of the bed, no attempt was made to quantify conversion. This was most likely due to problems in deconvolving the broad ¹H resonances associated with the reactant and product species. In recent work, Leutzsch et al.3 have used ²H NMR combined with neutron scattering to monitor the kinetics of the catalytic hydrogenation of benzene in 3 wt% Pt/MCM-41. Catalytic hydrogenation reactions over solid catalysts have also been ^{1}H studied using NMR spectroscopy, exploiting parahydrogen-induced polarisation (PHIP) to enhance the NMR signal.⁴

There are cases in which the use of ¹H NMR becomes prohibitive when studying reactions in solid catalysts as the relatively narrow chemical shift range of the ¹H nucleus and the line broadening of ¹H resonances, due to contrast in magnetic susceptibility, give rise to a large number of overlapping resonances, making the resulting NMR spectrum almost featureless and any further analysis difficult if not impossible. In such cases, ¹³C NMR spectroscopy is undoubtedly the preferred approach. The principle disadvantage of ¹³C NMR is the low signal-to-noise ratio, due to both low natural abundance (~1%) and low sensitivity of the 13 C nucleus relative to the 1 H nucleus. Polarisation transfer techniques such as the disortionless enhancement by polarisation transfer (DEPT) ¹³C NMR spectroscopy technique⁵ overcomes some of these disadvantages, enhancing the ¹³C signal by up to a factor of four. Moreover, the pulse sequence relies on the ¹H spin-lattice relaxation time (T_1) rather than ¹³C spin–lattice relaxation time, which is usually three to five times longer. Therefore, data acquisition can be much faster and this implies a higher signalto-noise ratio, for a fixed acquisition time. Mantle et al.⁶ used DEPT ¹³C NMR spectroscopy to study hydrogenation and isomerisation of pentenes over commercial Pd/Al₂O₃ catalyst. When trans-2-pentene and hydrogen were both adsorbed onto the support, partial hydrogenation to *n*-pentane was observed in

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addition to the presence of both *cis*-2- and *trans*-2-pentenes. All pentene isomers hydrogenated over the Pd/Al_2O_3 catalyst to give predominantly *n*-pentane and a small amount of the *trans*-2-pentene isomer. ¹³C DEPT-MRI was also used by Sederman *et al.*⁷ to spatially map the isomerisation and hydrogenation of 1-octene over Pd/Al_2O_3 occurring within a trickle-bed reactor.

Whilst hydrogenation reactions over solid catalysts have so far been benefitting from *in situ* NMR spectroscopy, according to our knowledge there has not been any report on catalytic oxidations occurring in porous catalysts. The methodology, if successfully validated, would represent an invaluable tool to study such reactions. Indeed, in recent years there has been growing attention to catalytic oxidations that make use of atmospheric oxygen as the oxidant. This type of reaction is of particular interest as it represents a more sustainable and alternative route to conventional processes using hazardous inorganic oxidizing agents (*e.g.*, $H_2Cr_2O_7$, KMnO₄), which tend to produce large amounts of hazardous waste.

In this work, we studied the aerobic catalytic oxidation of benzyl alcohol over a Pd/Al₂O₃ catalyst (supplied by Johnson Matthey, 0.5% Pd by weight, 2.5 mm trilobes, surface area of approximately 100 m² g⁻¹) at room temperature (20 °C) using air at high pressure (21 bar) as source of oxygen. The reaction was carried out in a home-built batch reactor, made with NMR-compatible materials, and exploiting in situ DEPT-45 ¹³C NMR spectroscopy using ¹³C natural-abundance species. Oxidation of pure benzyl alcohol as well as benzyl alcohol dissolved in primary aliphatic alcohol solvents (methanol, ethanol and 1-propanol) was chosen as the model system to validate the proposed methodology. The NMR batch reactor was built using polyether ether ketone (PEEK), a thermoplastic polymer with excellent mechanical and chemical resistance properties. The reactor was designed to withstand pressures up to 40 bar and was machined in a cylindrical shape with an external diameter of 20 mm (see Fig. S1 in ESI[†]). The Pd/Al₂O₃

catalyst trilobes were first soaked in the reaction mixture, previously purged with nitrogen, for at least 20 hours to make sure the catalyst pores were saturated. The trilobes were then dried on a pre-soaked filter paper to remove any excess liquid on the external surface and transferred into the NMR batch reactor; the pre-soaking of the filter paper prevented liquid from being drawn out of the pores. The cell was then closed, pressurised to 21 bar with air, placed into the magnet and the NMR acquisition started. DEPT-45 ¹³C NMR spectra during reaction were acquired over a period of 15 h. Typical acquisition parameters for the DEPT-45 ¹³C NMR experiments were: ¹³C and ¹H 90° pulses of 20 µs and 25 µs, respectively; 1024 scans; recycle delay of 4 s; evolution delay time, $1/2J_{CH}$, equal to 3.4 ms; the value of the echo time, τ , was chosen to be $1/2J_{\rm CH}$, where $J_{\rm CH}$ is the spin-spin coupling constant for a ${}^{13}{\rm C}{-}^{1}{\rm H}$ group and is approximately 145 Hz. All the ¹³C NMR spectra were acquired relative to the ¹³C resonance of tetramethylsilane (TMS).

Fig. 1 shows the evolution of the ¹³C DEPT-45 spectrum as a function of time. The large peak at approximately 130 ppm arise from the overlapping NMR signals coming from the aromatic rings of both benzyl alcohol (reactant) and benzaldehyde (product), hence such peak cannot be used to monitor unambiguously the reaction. However, the unambiguous and clearly distinguishable signals of the -CH₂ - group of benzyl alcohol, at approximately 60 ppm, and that of the -CHO of the benzaldehyde, at approximately 200 ppm, can be used to monitor both kinetics, conversion and selectivity of the reaction. The formation of benzaldehyde, the main oxidation product, is evident by the rising peak at 200 ppm, whereas the benzyl alcohol consumption can be observed by the decreasing signal at approximately 60 ppm. Spectra of similar quality were observed for benzyl alcohol dissolved in aliphatic alcohols. Again, for all solvents studied, the resonances of the -CH2- group of benzyl alcohol and that the -CHO of the benzaldehyde were clearly of



Fig. 1 Evolution of 13 C DEPT-45 NMR spectra during the aerobic catalytic oxidation of benzyl alcohol over Pd/Al₂O₃ catalyst. The formation of benzaldehyde is clearly evident by the rising NMR signal at 200 ppm, which belongs to the –CHO group of benzaldehyde. Simultaneously, the conversion of the benzyl alcohol is shown by the decrease of the signal at approximately 60 ppm, which belongs to the –CH₂– group of benzyl alcohol.

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distinguishable. No oxidation of the alcohol solvent was observed at the operating conditions chosen for the reaction. It is noted that Pd/Al_2O_3 catalysts have been reported to be highly selective to convert benzyl alcohol to benzaldehyde with almost 100% selectivity⁸ and indeed in our case benzaldehyde was the only oxidation product.

In order to quantify chemical composition using the spectra reported in Fig. 1 several important points need to be considered. The DEPT ¹³C NMR signal is influenced by a number of factors that, if not accounted for, will make the acquired signal intensity non-quantitative. These influences are: (a) non-uniform longitudinal relaxation (characterised by T_1); (b) non-uniform transverse relaxation (characterised by T_2); (c) effects arising from the pulse-sequence itself that leads to differential enhancement of different CH_n resonances. Non-uniform longitudinal relaxation may be avoided by ensuring a long recycle delay whereas effects arising from differential enhancement of CH_n resonances can be corrected by considering that the intensity of the peaks of a ¹³C DEPT NMR spectrum is described by:⁹

$$I(CH_n) = n \sin \theta \cos^{n-1} \theta$$

where n = 1, 2 or 3 and θ is the pulse angle of the final ¹H pulse in the DEPT sequence. Addressing effects arising from non-uniform transverse relaxation (T_2) is a more difficult task. Such effects are likely to be present especially when dealing with liquids in porous catalysts, the latter affecting significantly relaxation times of the confined liquids.¹⁰ In principle, the effect of transverse relaxation rate on peak intensity may be predicted by knowing the T_2 value of the peak that needs to be corrected. However, DEPT ¹³C NMR involves multiple quantum coherence (mqc) transverse relaxation. Measurements of mqc relaxation rates for pure liquids have been reported by Mlynarik;¹¹ however, attempts to extend these measurements to heterogeneous systems have not been successful.¹²

In the present work, quantification of DEPT ¹³C NMR spectra, in terms of benzaldehyde composition on a solvent-free basis, was carried out using a calibration procedure. Samples of known benzaldehyde composition in benzyl alcohol imbibed in Pd/Al_2O_3 catalyst pellets, were prepared and DEPT-45 ¹³C NMR spectra acquired, keeping the same parameters used to probe reactions. Spectra were acquired at atmospheric pressure and a check to verify that no reaction occurred during the acquisition time of each spectrum was also carried out. No reaction occurred at ambient conditions during the acquisition time. The actual molar composition was compared with the molar composition calculated from NMR spectra by considering the areas of the $-CH_2$ - peak of benzyl alcohol at 60 ppm and that of the -CHO peak of benzaldehyde at 200 ppm.

Results of the calibration are shown in Fig. 2. It can be clearly seen that the calibration plots are in some cases significantly different from the ideal plot (dotted line). To further investigate the trend in Fig. 2 we carried out T_2 ¹H





Fig. 2 Calibration plots of ¹³C DEPT-45 NMR spectra of benzaldehyde in different benzyl alcohol/solvent mixtures, within Pd/Al_2O_3 catalyst. Composition of benzaldehyde is solvent-free. Error bars are calculated based on signal-to-noise ratio.

NMR relaxation measurements of benzyl alcohol and benzaldehyde in the absence and presence of solvents with the Pd/Al₂O₃ catalyst in order to assess any possible effect of non-uniform T2 relaxation on the DEPT ¹³C NMR spectra. Here, it is noted that DEPT ¹³C NMR relies on T_2 relaxation of ¹H nuclei and not of ¹³C nuclei, which is due to the nature of the pulse sequence (*i.e.*, ${}^{1}H^{-13}C$ polarisation transfer). We found that whilst the T_2 of benzaldehyde was approximately the same in all cases (~45 ms), that of benzyl alcohol was significantly shorter (<25 ms) and especially non-uniform across the various solvents. In particular, we observed that the shorter the T_2 of benzyl alcohol the further the calibration curve deviates from the ideal plot. This is because shorter T_2 values of the benzyl alcohol peak lead to a larger drop of its NMR signal, hence making the benzaldehyde composition appearing larger than the actual composition. This implies that there is a significant effect on the spectra due to nonuniform T_2 relaxation; hence quantification through a calibration procedure is needed to correctly quantify intraparticle composition during reaction.

The composition of benzaldehyde within the Pd/Al_2O_3 catalyst during reaction in different solvents was obtained using the calibration plots of Fig. 2; this was achieved by using the benzaldehyde composition calculated from the NMR peak area measured in a specific solvent (or in solventfree conditions), hence the vertical axis values of Fig. 2, which were then converted into actual composition, horizontal axis of Fig. 2, using the calibration plot for the solvent (or solvent-free) under investigation. The results are reported in Fig. 3. It can be noted as the benzaldehyde production increases more rapidly in the first 5 h of reaction and then reaches a plateau. Further investigation revealed that such a plateau was due to oxygen depletion. Since air



Fig. 3 Evolution of benzaldehyde composition during the course of the oxidation reaction of benzyl alcohol over Pd/Al_2O_3 catalyst. Composition of benzaldehyde is on a solvent-free basis. Solid lines are a guide to the eye. Error bars are calculated based on signal-to-noise ratio.

(approximately 20% oxygen and 80% nitrogen) is used as the oxidant gas, the oxygen consumed is being replaced by air, which eventually leads to accumulation of inert nitrogen and consequent oxygen depletion, stopping the reaction. Indeed, after reaching the plateau, when the reactor was depressurised (purged of used air) and pressurised again with fresh air, the conversion started increasing again, confirming our hypothesis.

From Fig. 3 it is possible to observe differences in reaction rates for different solvents. In particular the initial reaction rate of benzyl alcohol follows the trend:

methanol (5.4
$$h^{-1}$$
) > ethanol (3.5 h^{-1})
> 1 - propanol (3.0 h^{-1})

In order to obtain further insights into such solvent effects, we carried out diffusion measurements of the various reactant/solvent/catalyst systems using ¹H pulsed-field gradient (PFG) NMR. A typical data set of such measurements is shown in Fig. 4, which shows the ¹H PFG NMR signal attenuation for an equimolar mixture of benzyl alcohol/1-propanol within the Pd/AL₂O₃ catalyst. The spectrum shows that the peak assignment is unambiguous as it is possible to distinguish the resonances of 1-propanol (0.93, 1.6 and 3.6 ppm) from those of benzyl alcohols are not present in the spectrum due to T_2 -weighting of the PFG NMR signal. From such spectra it is possible to quantify the diffusion decay of the benzyl alcohol peak in the various solvents and calculate its diffusion coefficient.

In addition to transport properties, we also estimated solvent affinities with the Pd/Al₂O₃ catalyst by calculation of the T_1/T_2 ratio of methanol, ethanol and 1-propanol adsorbed in Pd/Al₂O₃. Such a ratio has been previously used to assess solvent inhibition in heterogeneous catalysis¹³ and gives a good indication of surface adsorption strength.¹⁰ Solvents with a higher T_1/T_2 ratio exhibit stronger interaction with the surface,^{14,15} and hence may inhibit a favourable adsorption of the reactant decreasing reaction rate at the surface.

A summary of initial reaction rates, NMR diffusion coefficients and relaxation time measurements is reported in Table 1. The typical relative error for diffusion and T_1/T_2 measurements is approximately 2 and 4%, respectively. It is possible to see a trend in diffusion rate of the reactant in different solvents as well as in T_1/T_2 of the solvent. In particular, solvents with higher reaction rates are associated



Fig. 4 PFG NMR spectra of an equimolar mixture of benzyl alcohol/1-propanol in Pd/Al₂O₃ catalyst as a function of the gradient strength.

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Table 1 Initial rate, reactant diffusion and 1H NMR T_1/T_2 for different solvents in Pd/Al_2O_3 catalyst

Solvent	Rate [h ⁻¹]	$D_{\text{Reactant}} \left[\text{m}^2 \text{ s}^{-1} \right]$	$T_1/T_2 \begin{bmatrix} - \end{bmatrix}$
Methanol	5.4 ± 0.2	$\begin{array}{c} 1.85 \pm 0.04 \\ 1.69 \pm 0.03 \\ 1.52 \pm 0.03 \end{array}$	20.6 ± 0.8
Ethanol	3.5 ± 0.1		29.8 ± 1.2
1-Propapol	3.0 ± 0.1		35.0 ± 1.4

to a faster diffusion rate of the reactant and to a lower T_1/T_2 of the solvent, indicating improved transport and a lower degree of solvent inhibition, which leads to favourable adsorption conditions for the reactant over the surface. Such results strongly suggest that both transport and adsorption play an important role in determining catalyst performances. It is interesting to note as a similar trend in T_1/T_2 for primary alcohols in porous oxide catalysts has been reported by Robinson *et al.*¹⁶ in a study involving both NMR and DFT calculations.

In summary, we have reported the first *in situ* ¹H and ¹³C NMR measurements to study aerobic, high-pressure oxidation reactions occurring inside heterogeneous catalysts. The subsequent analysis of the NMR data shows that it is possible to monitor, unambiguously, the evolution of intra-particle composition during the oxidation of benzyl alcohol to benzaldehyde inside Pd/Al₂O₃ catalyst pellets using air as the the combination of ¹³C Moreover, NMR oxidant. measurements, (to extract kinetic information) with measurements of NMR diffusion coefficients and relaxation times gives a more detailed understanding of the physicochemical process of the catalytic system under different working conditions.

Conflicts of interest

There are no conflicts to declare.

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