Methanol dynamics in H-ZSM-5 with Si/Al ratio of 25: a quasi-elastic neutron scattering (QENS) study

Santhosh K. Matam1,2 · C. Richard A. Catlow1,2,3 · Ian P. Silverwood1,4 · Alexander J. O'Malley1,5

Accepted: 10 May 2021
© Crown 2021

Abstract
Methanol dynamics in zeolite H-ZSM-5 (Si/Al of 25) with a methanol loading of ~ 30 molecules per unit cell has been studied at 298, 323, 348 and 373 K by incoherent quasi-elastic neutron scattering (QENS). The elastic incoherent structure factor (EISF) reveals that the majority of methanol is immobile, in the range between 70 and 80%, depending on the measurement temperature. At 298 K, ≈ 20% methanol is mobile on the instrumental timescale, exhibiting isotropic rotational dynamics with a rotational diffusion coefficient \(D_R\) of 4.75 \(\times\) 10\(^{-10}\) s\(^{-1}\). Upon increasing the measurement temperature from 298 to 323 K, the nature of the methanol dynamics changes from rotational to translational diffusion dynamics. Similar translational diffusion rates are measured at 348 and 373 K, though with a larger mobile fraction as temperature increases. The translational diffusion is characterised as jump diffusion confined to a sphere with a radius close to that of a ZSM-5 channel. The diffusion coefficients may be calculated using either the Volino–Dianoux (VD) model of diffusion confined to a sphere, or the Chudley–Elliot (CE) jump diffusion model. The VD model gives rise to a self-diffusion co-efficient \(D_s\) of methanol in the range of 7.8–8.4 \(\times\) 10\(^{-10}\) m\(^2\) s\(^{-1}\). The CE model gives a \(D_s\) of around 1.2 (± 0.1) \(\times\) 10\(^{-9}\) m\(^2\) s\(^{-1}\) with a jump distance of 2.8 (either +0.15 or −0.1) Å and a residence time (\(\tau\)) of ~ 10.8 (either +0.1 or −0.2) ps. A correlation between the present and earlier studies that report methanol dynamics in H-ZSM-5 with Si/Al of 36 is made, suggesting that with increasing Si/Al ratio, the mobile fraction of methanol increases while \(D_R\) decreases.

Keywords Zeolite · H-ZSM-5 · Methanol · QENS · Diffusion · Rotational · Translational

1 Introduction
Zeolite ZSM-5 plays a crucial role in many commercial petrochemical and environmental processes due to its unique porous architecture and chemical/catalytic properties [1]. Some prominent examples of these processes include methanol to hydrocarbons, hydrocracking and alkylation [1–6]. In microporous catalytic processes, transportation of reactants and/or products to and from the active Brønsted acid site, located within the zeolite pores with distinct pore architecture, involves adsorption, diffusion and then reaction of molecules at the active site. Therefore, in the case of the widely studied methanol to hydrocarbon (MTH) catalysis, diffusion of methanol and products is one of the key factors for an efficient catalytic process [7]. The knowledge of adsorption geometry, diffusion characteristics and reactivity of diffusing molecules within the zeolite pores will help to design and tailor the porous architecture that favours the processes [7–9].
The diffusion properties of molecules within catalyst pores are measured by different techniques which are broadly classified into (i) macroscopic and (ii) microscopic [10, 11]. Traditionally employed macroscopic techniques include steady-state permeation, sorption-uptake, chromatography and zero length column methods which are often limited by external heat and mass transfer and response time. Microscopic techniques, however, are capable of measuring diffusivity over much shorter time and length scales than is possible with macroscopic techniques. They include pulsed field gradient nuclear magnetic resonance (PFG-NMR) and quasi-elastic neutron scattering (QENS). QENS enables measurement of dynamical processes over a lengthscale of 0.1–100 nm and time scales of 10 fs–100 ns [11, 12]. Whereas, PFG-NMR probes with a length scale of 0.1–1 μm and time scales in the order of 1 ms resolution [11]. Moreover, QENS data can be simulated by molecular dynamics (MD) which sample similar time and length scales [11–13]. Applications of QENS in the field of heterogenous catalysis are limited and only a few QENS studies on methanol diffusion in ZSM-5 (with a comparable Si/Al ratio of 30) have been reported. It is notable that differing conclusions can be found depending on resolution/energy window of the instrument and methanol loading used even in zeolites with comparable Si/Al ratios. [14–17]. For example, one study concluded that methanol diffuses along the ZSM-5 channels with a diffusion coefficient in the order of $10^{-11}$ m$^2$s$^{-1}$ using the high resolution IN10 instrument at ILL, Grenoble [14], while other studies employing lower resolution spectrometers (with wider energy windows) were not able to observe long range methanol diffusion [15–17]. We note that these measurements were conducted on different instruments with varied instrument resolutions [14–17].

The objective of the present work is to utilise QENS (measured employing the instrumentation at ISIS Neutron and Muon Source at Harwell, UK) to study methanol diffusion dynamics in zeolite H-ZSM-5 (Si/Al = 25) as a function of temperature. A comparison with previously reported studies enabled us to assess the effect of Si/Al ratio on methanol diffusion dynamics in ZSM-5. The results show that the methanol mobility is more restricted in the zeolite studied here than that reported previously with a higher Si/Al ratio (Si/Al ≥ 30), especially at room temperature, in line with the higher acid site density of this zeolite.

2 Experimental

2.1 Zeolite H-ZSM-5

The NH$_4$-ZSM-5 zeolite (Si/Al = 25) was procured from Zeolyst International, Inc. The zeolite was calcined in air at 500 °C for 24 h to remove residual template and to obtain H-ZSM-5. Prior to the quasi-elastic neutron scattering (QENS) experiments, the zeolite was dehydrated under vacuum ($10^{-3}$ to $10^{-4}$ mbar) at 350 °C for 12 h and thereafter handled in an argon-filled glovebox. For the methanol loaded samples, methanol (Sigma-Aldrich, ≥ 99.9%) loading was conducted at room temperature by flowing dry N$_2$ (100 mL min$^{-1}$) through a methanol saturator that consisted of a Dreschel bottle at the same temperature [18]. The loading was continued until there was no further uptake by the zeolite (gravimetrically determined) and resulted in a methanol loading of around 30 molecules per unit cell (i.e., ≈ 7.5 molecules per acidic site) [18]. The thin walled aluminium containers of annular geometry were loaded with zeolite under an argon atmosphere in the glovebox.

2.2 Quasi-elastic neutron scattering (QENS)

The zeolite H-ZSM-5 with and without methanol loading were analysed by QENS using the time-of-flight backscattering OSIRIS spectrometer at the ISIS Facility (STFC Rutherford Appleton Laboratory, Chilton, UK) [16, 17 and references therein]. The sample container was placed in a top loading closed cycle refrigerator and a resolution measurement of each undosed (i.e., without methanol) zeolite sample was obtained at a base temperature of around 10 K. QENS measurements were then subsequently conducted at 298, 323, 348 and 373 K on both zeolites with and without the methanol load. The QENS data collected without methanol loading were subtracted from the data obtained with the loaded zeolite. The zeolite with the methanol load measured at base temperature was used as the instrumental resolution function for data fitting. Pyrolytic graphite 002 analyser crystals were used which resulted in an energy resolution of 24.5 μeV with energy transfers measured in a window of ± 0.55 meV and with a Q range of 0.2–1.75 Å$^{-1}$. The data reduction was performed using the MantidPlot software [19]. The data fitting was conducted by the neutron scattering analysis software DAVE (version 2.2) [20].

3 Results and discussion

3.1 Methanol diffusion dynamics at 298 K

The experimental QENS data collected at all studied temperatures were fitted to the combination of a delta function, a single Lorentzian function that accounts for the quasi-elastic broadening of the scattering function and a linear background function convoluted with the resolution data measured at 10 K. Selected QENS data at 298 K are plotted as a function of momentum transfer (Q) in Fig. 1a, along with the Lorentzian function and the total fit. There is a significant elastic component
which is particularly high at low Q values. This intensity decreases with increasing Q, and the intensity of the quasi-elastic component fit by the Lorentzian function increases (becoming particularly significant above 0.9 Å⁻¹) which is typically observed for methanol loaded H-ZSM-5 [17]. In general, methanol mobility over a longer length scale, i.e. translational diffusion, can be expected at lower Q (< 1 Å⁻¹), while higher energy localised methanol dynamics such as rotational motions can be anticipated at higher Q values (> 1 Å⁻¹) [11–13]. The significant decrease in elastic intensity at higher Q indicates the observation of localised methanol dynamics [17, 21]. A large elastic peak intensity at lower Q between 0.24 and 0.9 Å⁻¹ reflects the absence of longer range translational dynamics of methanol (within the instrumental time resolution of ≈ 1–100 ps) [17, 21]. Indeed, at lower Q, the resolution function mainly describes the data, suggesting that the quasi-elastic component, characterised by the Lorentzian function is not significant (see Fig. S1, Supplementary Information (SI)). This observation is in line with previous work that reported no quasi-elastic broadening of the signal of methanol loaded H-ZSM-5 at room temperature (certainly at low Q), though the methanol loading and Si/Al ratio of H-ZSM-5 differ slightly among the reports [14–17, 21].

The significant elastic intensity is attributed to the immobile methanol due either to hydrogen bonding [14–16] or methoxylation [16]. Indeed, complementary diffuse reflectance infrared Fourier transformed spectroscopy (DRIFTS) studies show that both hydrogen bonded methanol and methoxy species are present under these experimental conditions [18, 22]. Besides immobile methanol, the presence of a fraction of locally mobile methanol is also reported previously [14, 17, 21] by analysing the elastic incoherent structure factor (EISF), which is the ratio between the elastic peak intensity and total peak intensity [11–14, 17, 21]. Such localised methanol dynamics can be characterised by different theoretical models that are discussed in detail in references [11–14, 17, 21]. Briefly, the models include whole molecule isotropic rotation [23], translational diffusion confined within a sphere (Volino–Dianoux model) [24] and 3-site jump rotation of the methyl groups [17, 21]. Based on these models, the experimental EISF is analysed. It is important to note that the use of the 3-site jump rotation model can only be applied with a maximum 75% of the protons treated as mobile, as we assume the hydroxyl proton would be anchored to a Brønsted acidic site. The experimental EISF is plotted in Fig. 1B and decreases with increasing Q. The absolute intensity decay as a function of Q is around 20%, which is around 50% lower than those reported previously [17, 21]. It suggests that the localised methanol dynamics are even more restricted in a H-ZSM-5 sample with Si/Al = 25 than in previously reported samples with lower aluminium content (Si/Al > 30). Indeed, simple models with a mobile fraction of 0.75 assigned to 3-site model, as noted earlier, could not fit the experimental EISF, due to the fact that the methanol loaded H-ZSM-5 may show significant interactions with acid sites at room temperature as evident from QENS and also from inelastic neutron scattering (INS) and DRIFTS [16, 18, 22]. Therefore, the effective EISF is considered to include a fraction of immobile methanol in the models as detailed in previous studies [17, 21]. Models, with the static fraction included are now compared with

---

**Fig. 1** A QENS spectra of methanol loaded H-ZSM-5 at 298 K. Experimental data (filled circles), total fit (solid black line) and the Lorentzian function (red). Selected spectra are shown for clarity. B The corresponding experimental elastic incoherent structure factor (EISF) data (green diamond symbols) of methanol in H-ZSM-5 at 298 K are compared with different theoretical models that are discussed in detail in [11–14, 17, 21]. The experimental EISF values at Q between 1.58 and 1.67 are not included in B panel due to contributions from Bragg peaks.
the experimental EISF in Fig. 2A. For the 3-site model, we again assume a mobile fraction of 0.75.

It can be readily seen that the experimental EISF now adequately fits to the isotropic methanol rotation model with a mobile fraction of 21%, which is 50% lower than that reported in H-ZSM-5 with Si/Al of 30 (with mesopores present due to dealumination) and 36 [17, 21]. The 3-site ‘methyl’ jump rotation model fails to fit the data, even when only a fraction of methyl protons is considered as mobile, while translational diffusion of methanol in a confined spherical volume (with a radius of 2.75 Å) model shows a potential agreement with the experimental EISF. However, analysis of quasi-elastic broadening of the spectra by plotting the full width at half maxima (FWHM) of the Lorentzian function with $Q^2$ as in Fig. 2B suggests we are more likely observing isotropic methanol rotation, the model of which is also shown to fit the experimental EISF. The FWHM appear to be $Q^2$ independent, which is characteristic of localised dynamics where the $Q$ resolution is insufficient to distinguish motion in real space, such as isotropic rotation, but not of longer range translational diffusion whether it be Fickian in nature, jump diffusion or confined to a spherical volume [17, 21]. The FWHM give rise to an isotropic methanol rotational diffusion coefficient ($D_R$) of $4.75 \times 10^{10}$ s$^{-1}$ similar to that reported in [21], which is of the same order of magnitude as reported previously [17].

### 3.2 Methanol diffusion dynamics at higher temperatures

QENS spectra collected at 323 K are plotted together with the Lorentzian function and the total fit as a function of $Q$ in Fig. 3A (for more clarity see Fig. S2 in SI). The elastic intensity decreases with increasing $Q$, as observed at 298 K (Fig. 1A). As expected, the individual theoretical models alone could not fit the experimental EISF. Therefore, the effective EISF which includes an immobile methanol fraction in the models is considered and compared in Fig. 3B. The isotropic methanol rotation and the 3-site methyl jump rotation models fail to fit the experimental EISF, while the model of translational diffusion confined to a sphere fits the data very well with a spherical radius of 2.75 Å, and a mobile methanol fraction of 22% (Table 1).

It is noteworthy that the methanol dynamics have changed from rotational to translational diffusion on increasing the measurement temperature from 298 to 323 K, which is in line with the previously observed temperature dependent methanol dynamics [17, 21]. Interestingly, the mobile fraction of 22% is similar to that observed at 298 K, although the fraction is derived from two different models (confined diffusion vs rotational model) suggesting that very little extra methanol has become mobile with this temperature increase, but the amount that is mobile, has switched from rotational to translational motions as observed previously [21]. The same Volino–Dianoux (VD) [24] confined diffusion model indicates that the mobile fraction increases with increasing measurement temperature from 323 K to 348 and to 373 K, as expected (Table 1) and the model fits the data very well (Figs. S3 and S4, see Supplementary Information). Upon analysing the half width half maxima (HWHM) of the Lorentzian function as a function of $Q^2$ (Fig. 3C), it is found that the broadenings fit to the Chudley–Elliot (CE) jump diffusion model (the fitting parameters of which will be discussed shortly) [25]. However, it is also observed that the HWHM deviates from this model and plateaus below $Q^2 = 1.3$ Å$^{-2}$. This observation is consistent with jump diffusion confined to a sphere, as has been considered due to contribution from Bragg peaks.

---

**Fig. 2** A Experimental EISF data of methanol in H-ZSM-5 at 298 K, modelled by including a fraction of immobile methanol in different theoretical models. The mobile methanol fraction is shown in parenthesis and the radius of the confined spherical radius of 2.75 Å is considered. The EISF values at $Q$ between 1.58 and 1.67 are not considered due to contribution from Bragg peaks. B The FWHM of the Lorentzian function is plotted as a function of $Q^2$. The FWHM values at 2.52, 2.79 and 3.03 Å$^{-2}$ are discounted due to contribution from Bragg peaks.
Fig. 3  A QENS spectra of methanol in H-ZSM-5 at 323 K. Experimental data (filled circles), total fit (solid black line) and the Lorentzian function (red). Selected spectra are shown for clarity. B Experimental EISF data of methanol in H-ZSM-5 at 323 K, modeled by including a fraction of immobile methanol in different theoretical models. The mobile methanol fraction is shown in the parenthesis of panel (B) and a confined volume radius of 2.75 Å is used. C The Q independent at lower Q (< 1.25 Q$^2$) represents the HWHM of the Lorentzian function that fits reasonably well to the Volino–Dianoux confined diffusion model. The Q dependence of the Lorentzian function HWHM at higher Q > 1 Q$^2$) fits well to the Chudley–Elliot jump diffusion model. The Q$^2$ value corresponding to the 2.75 Å radius of the confined sphere of the Volino–Dianoux model is indicated by the dashed black vertical line. EISF (B) and HWHM (C) values at Q of 1.58 and 1.67 are not considered due to contribution from Bragg peaks.
Table 1 Diffusion coefficients and mobile fractions derived from fitting QENS data with different models, for methanol in H-ZSM-5 with a Si/Al ratio of 25

| T (K) | Mobile fraction (%) | Self-diffusion co-efficient (Ds), m²/s | Residence time (τ), ps |
|-------|---------------------|---------------------------------------|------------------------|
|       | Volino–Dianoux modelb | Chudley–Elliot modelc |
| 298   | 21a                 | –          | –          | –          |
| 323   | 22b                 | 9.2 × 10⁻¹⁰ | 1.15 × 10⁻⁹ | 10.8       |
| 348   | 28b                 | 8.2 × 10⁻¹⁰ | 1.2 × 10⁻⁹  | 10.9       |
| 373   | 32b                 | 9.2 × 10⁻¹⁰ | 1.3 × 10⁻⁹  | 10.6       |

aRotational model
bConfined diffusion Volino–Dianoux model
cChudley–Elliot jump diffusion model with a jump distance of 2.8 (either +0.15 or −0.1) Å depending on temperature

Based on the comparison with previous studies on H-ZSM-5 with Si/Al of 30 [17] and 36 [21], a correlation between the methanol mobile fraction (%) and rotational diffusion coefficient (D₉) as a function of Si/Al ratio is plotted in Fig. 4. It is evident that the mobile fraction increases and isotropic methanol rotational diffusion coefficient (D₉) decreases with increasing Si/Al ratio of H-ZSM-5 from 25 [present study] to 30 [17] and to 36 [21] (Fig. 4), which is attributed to the acid site density of the zeolites; the higher the acid site density the lower the mobile fraction. However, the comparison between zeolites with Si/Al ratio of 25 and 36 with that of 30 should be treated with caution due to the formation of a fraction of larger pores due to dealumination of H-ZSM-5 during the MTH process and measurement temperature. Nonetheless, the comparison presents a clear composition dependent methanol rotational dynamics in zeolite H-ZSM-5. In line with this, further increase in Si/Al of the zeolite from 36 to 135 is shown in previous work to have led to a switch from D₉ to translational diffusion dynamics of methanol (Ds) (at a similar methanol loading and measurement temperature) [21].

Interestingly, the Ds obtained within a confined spherical volume appears to be similar in magnitude in zeolites with Si/Al ratio of 25 (Table 1), 36 and 135 [21]; for example, Ds of 8.8–9.2 × 10⁻¹⁰ m² s⁻¹ observed at 373 K. Computational studies are in progress to assess the relationship between Si/Al ratio of H-ZSM-5 and the longer range translational diffusion of methanol.

4 Summary and conclusions

Methanol diffusion dynamics in zeolite H-ZSM-5 (Si/Al of 25) with a methanol loading of ~30 molecules per unit cell have been studied by incoherent quasi elastic neutron scattering (QENS) at 298, 323, 348 and 373 K. The experimental QENS data fit to a delta function convoluted with the resolution data collected at 10 K, a Lorentzian function that accounts for the quasi-elastic broadening and a background function. Based on the fitting, the following main conclusions can be drawn:

(i) Methanol diffusion dynamics are, as expected, temperature dependent: the higher the temperature, the greater the methanol mobility. However, the majority of the methanol in H-ZSM-5 remains immobile, in the range of 70–80%, depending on the measurement temperature, as shown from the effective elastic incoherent structure factor (EISF). At 298 K, around 20% methanol is mobile and shows isotropic rotational dynamics with a rotational diffusion coefficient (D₉) of 4.75 × 10¹⁰ s⁻¹. The D₉ decreases while the mobile fraction of methanol increases with increasing Si/Al.
ratio of H-ZSM-5 from 25 to 36 as evident from a correlation between the present and earlier studies.

(ii) The nature of methanol dynamics switches from rotational to translational diffusion on increasing the measurement temperature from 298 to 323 K. Translational diffusion dynamics is observed at similar rates, but with higher mobile populations at 348 and 373 K. Even at 373 K however, the majority of methanol is still immobile (≈ 70%) on the instrumental timescale.

(iii) The translational motion is characterised as jump diffusion confined to a sphere with a radius of ≈ 2.75 Å, which matches with the ZSM-5 channel size of 5.5 Å. The self-diffusion co-efficient ($D_s$) may be quantified using either the VD model of diffusion confined to a sphere or the CE jump diffusion model. Using the VD confined diffusion model to fit the plateaued broadenings at low $Q$, a $D_s$ of 8.7 $(± 0.5) \times 10^{-10}$ m$^2$ s$^{-1}$ is obtained. The CE jump diffusion model gives a $D_s$ of around 1.2 $(± 0.1) \times 10^{-9}$ m$^2$ s$^{-1}$ after fitting a jump distance of around 2.8 (either +0.15 or −0.1) Å depending on temperature and a residence time ($τ$) of around 10.8 (either +0.1 or −0.2) ps at temperatures between 323 and 373 K.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s11244-021-01450-z.

Acknowledgements The UK Catalysis Hub is kindly thanked for resources and support provided via our membership of the UK Catalysis Hub Consortium and funded by EPSRC Grant: EP/R026939/1, EP/R026815/1, EP/R026645/1, EP/R027129/1. AJOM acknowledges Roger and Sue Whorrod for the funding of the Whorrod Fellowship. The authors acknowledge the ISIS Neutron and Muon Source at the STFC Rutherford Appleton Laboratory for granting access to OSIRIS instrument and the beamline facilities. The raw experimental data resulted from our beamtime RB1920441 can be found at https://doi.org/10.5286/ISIS.E.RB1920441.

Declarations

Conflict of interest There are no conflict of interest.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. van Speybroeck V, Hemelsoet K, Joos L, Waroquier M, Bell RG, Catlow CRA (2015) Chem Soc Rev 44:7044
2. Chang CD, Silvestri AJ (1977) J Catal 47:249
3. Hunter R, Hutchings GJ (1987) Chem Commun 5:377
4. Svelle S, Visur M, Olsbye U, Saeuprahman BM (2011) Topics Catal 54:897
5. Van der Mynsbrugge J, Moors SLC, De Wispelaere K, Van Speybroeck V (2014) ChemCatChem 6:1906

 Springer
6. Minova IB, Matam SK, Greenaway A, Catlow CRA, Frogley MD, Cinque G, Wright PA, Howe RF (2019) ACS Catal 9:6564
7. Jost W (1960) Diffusion in Solids, Liquids and Gases. Academic Press, New York
8. Hoogschagen J (1955) Ind Eng Chem 47:906
9. Weisz PB (1973) Chem Tech 3:498
10. Karger J, Ruthven DM (1989) Zeolites 9:267
11. Jobic H, Theodorou DN (2007) Micro Meso Mater 102:21
12. O’Malley AJ, Catlow CRA (2017) Sorbate dynamics in zeolite catalysts. In: Fernandez-Alonso F, Price DL (eds) Experimental Methods in the Physical Sciences; Neutron Scattering - Applications in Biology, Chemistry, and Materials Science. Academic Press, Boca Raton, pp 349–401
13. O’Malley AJ, Parker SF, Catlow CRA (2017) Chem Commun 53:12164
14. Jobic H, Renouprez A, Bee M, Poignignon C (1986) J Phys Chem 90:1059
15. Mitra S, Kamble VS, Tripathi AK, Gupta NM, Mukhopadhuyay R (2004) Pramana J Phys 63:443
16. O’Malley AJ, Parker SF, Chutia A, Farrow MR, Silverwood IP, Sakai VG, Catlow CRA (2016) Chem Commun 52:2897
17. Matam SK, Omalley AJ, Catlow CRA, Suwardiyanto CP, Hawkins AP, Zachariou A, Lennon D, Silverwood IP, Parker SF, Howe RF (2018) Catal Sci Technol 8:3304
18. Matam SK, Nastae SAF, Logsdail AJ, Catlow CRA (2020) Chem Sci 11:6805
19. Arnold O, Bilheux JC, Borreguero JM, Buts A, Campbell SI, Chapon L, Doucet M, Draper N, Leal RF, Gigg MA, Lynch VE, Markvardsen A, Mikkelsen DJ, Mikkelson RL, Miller R, Palmen K, Parker P, Passos G, Perrig TG, Peterson PF, Ren S, Reuter MA, Savici AT, Taylor JW, Taylor RJ, Tolchenov R, Zhou W, Zikovsky J (2014) Instrum Methods Phys Res Sect A 764:156
20. Azuah RT, Kneller LR, Qiu Y, Tregenna-Piggott PLW, Brown CM, Copley JRD, Dimeo RM (2009) J Res Natl Inst Stand Technol 114(6):341
21. Omojola T, Silverwood IP, O’Malley AJ (2020) Catal Sci Technol 10:4305
22. Matam SK, Howe RF, Thetford A, Catlow CRA (2018) Chem Commun 54:12875
23. Sears VF (1966) Can J Phys 44(6):1279
24. Volino F, Dianoux AJ (1980) Mol Phys 41(2):271
25. Chudley CT, Elliott RJ (1961) Proc Phys Soc Lond 77(2):353
26. O’Malley AJ, Sarwar M, Armstrong J, Catlow CRA, Silverwood IP, York APE, Hitchcock I (2018) Phys Chem Chem Phys 20(17):11976

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.