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## Correction: Structure and solvation of confined water and water-ethanol clusters within microporous Brønsted acids and their effects on ethanol dehydration catalysis

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Correction for 'Structure and solvation of confined water and water–ethanol clusters within microporous Brønsted acids and their effects on ethanol dehydration catalysis' by Jason S. Bates *et al.*, *Chem. Sci.*, 2020, **11**, 7102–7122, DOI: 10.1039/D0SC02589E.

The authors regret that in the original article there is an error in Fig. 7(a) as it is missing the data set for H-Al-MFI (closed circles). The correct version of Fig. 7 (in which the closed circle data series appears in panel (a)) is presented below.

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**Fig. 7** (a) Apparent first-order bimolecular ethanol dehydration rate constant (per H<sup>+</sup>, 373 K) and (b) activity coefficient ratio ( $\chi = \gamma_{EW_p}/\gamma_{\pm}$ ) from eqn (7), as a function of H<sub>2</sub>O pressure on H-Al-Beta-F(2.0) ( $\blacksquare$ ), H-Al-TON ( $\blacktriangle$ ), H-Al-FAU ( $\diamondsuit$ ), H-Al-MFI ( $\bigcirc$ ), H-Al-AEI ( $\square$ ), H-Al-CHA ( $\bigcirc$ ), and HPW/Si-MCM-41 ( $\bullet$ ). Solid lines in (a) reflect regression of measured rate constants to eqn (7). Inset (a): dependence of  $k_3K_2K_4^{-1}$  values on the largest included sphere diameter of zeolite pores ( $d_{LC}$ ). Inset (b): slopes of the data sets in (b) in the high water pressure limit, as a function of the channel undulation parameter ( $d_{PL}/d_{LC}$ ). Solid lines in both insets are to guide the eye.

The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.