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Correction: Intermolecular network analysis of the liquid and vapor interfaces of pentane and water: microsolvation does not trend with interfacial properties

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Correction for 'Intermolecular network analysis of the liquid and vapor interfaces of pentane and water: microsolvation does not trend with interfacial properties' by Yasaman Ghadar *et al.*, *Phys. Chem. Chem. Phys.*, 2014, **16**, 12475–12487.

Fig. 3(B) is incorrect in the paper. The amended figure is shown below. This error does not affect the main conclusions of the paper.

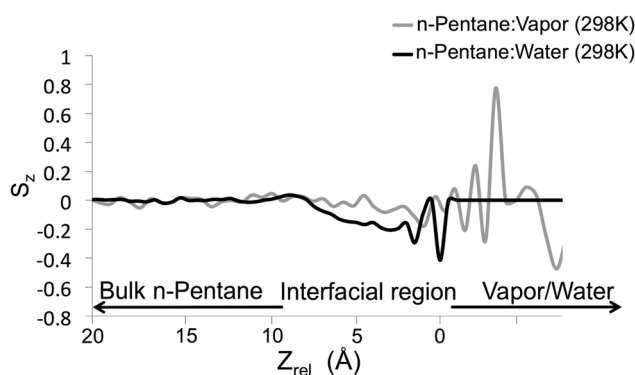


Fig. 3 (B) The *n*-pentane order parameter in *n*-pentane:vapor and water:*n*-pentane at 298 K. A relative *z*-axis (z_{rel}) has been used so as to overlap the spectra for optimal comparisons of behavior.

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

