


 Cite this: *Chem. Commun.*, 2018, **54**, 8136

DOI: 10.1039/c8cc90298d

rsc.li/chemcomm

## Correction: Model molecules to classify CH...O hydrogen-bonds

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 Correction for 'Model molecules to classify CH...O hydrogen-bonds' by Amol M. Vibhute et al., *Chem. Commun.*, 2018, **54**, 4629–4632.

The editorial office regrets that the bond distance data in the sentence beginning “The short C...O distances of...” in the left hand column of page 4630 are incorrect in the original article due to a production error. The corrected sentence is shown below.

The short C...O distances of 2.82–3.02 Å (much below the van der Waals distance of 3.22 Å for O and C) and H...O distances of 2.33–2.49 Å (much below the van der Waals distance of 2.72 Å for O and H) as well as CH...O bond angles of 110°–114° in **5–12** suggest the possibility of an intramolecular CH...O H-bond within them (Table 1).

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

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