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CORRECTION

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Correction: Structural characterization of gas-phase cysteine and cysteine methyl ester complexes with zinc and cadmium dications by infrared multiple photon dissociation spectroscopy

Rebecca A. Coates,^a Christopher P. McNary,^a Georgia C. Boles,^a Giel Berden,^b Jos Oomens^{bc} and P. B. Armentrout ^b *^a

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Correction for 'Structural characterization of gas-phase cysteine and cysteine methyl ester complexes with zinc and cadmium dications by infrared multiple photon dissociation spectroscopy' by Rebecca A. Coates *et al.*, *Phys. Chem. Chem. Phys.*, 2015, **17**, 25799–25808.

Although the overall conclusions of the original article remain unaffected (no experimental or theoretical IR spectra are changed, nor is any calculated thermochemistry at 0 K), the thermal corrections to the Gibbs free energy at 298 K were mistakenly overestimated. Corrected 298 K values for Tables 1 and 2 from the original manuscript are given below. Notably, only very subtle changes are found such that the relative order of all 298 K theoretically determined low-energy species remains the same for the $[Zn(Cys-H)]^+$, $[Cd(Cys-H)]^+$, [Cd(

Table 1 B3LYP,^a B3P86, and MP2(full) relative free energies (kJ mol⁻¹) at 298 K of low-lying conformers of [M(Cys-H)]⁺ and [M(CysOMe-H)]^{+b}

| | | | · · |
|----------------------------|---|--|--|
| Complex | Structure | Zn | Cd |
| [M(Cys-H)] ⁺ | $[N,CO,S^{-}]tgg \\ [N,CO,S^{-}]cgg \\ [CO,S^{-}]cgg \\ [N,S^{-}]tgt \\ [CO^{-},S^{-}]cgg' \\ [N,OH,S^{-}]tgg \\ [CO,S^{-}]ctg \\ [N,CO^{-},S]ggg \\ [N,S^{-}]cgt \\ [N^{-},CO,S]tggg \\ [S^{-}]tgg $ | $\begin{array}{c} 0.0 \ (0.0), \ 0.0, \ 0.0 \\ 24.4 \ (24.0), \ 24.0, \ 23.2 \\ 27.8 \ (32.8), \ 31.5, \ 45.0 \\ 30.8 \ (35.1), \ 37.7, \ 38.0 \\ 36.0 \ (36.9), \ 40.5, \ 42.0 \\ 38.2 \ (36.3), \ 42.1, \ 35.2 \\ 50.2 \ (54.0), \ 54.6, \ 68.5 \\ 60.6 \ (57.7), \ 60.9, \ 58.2 \\ 63.9 \ (67.2), \ 70.2, \ 71.5 \\ 100.5 \ (101.2), \ 101.5, \ 102.8 \\ 143.8 \ (150.7), \ 164.3, \ 179.9 \end{array}$ | $\begin{array}{c} 0.0 \ (0.0), \ 0.0, \ 0.0 \\ 25.1 \ (24.9), \ 24.7, \ 25.9 \\ 23.1 \ (28.5), \ 28.1, \ 33.0 \\ 27.7 \ (34.0), \ 29.8, \ 44.0 \\ 30.5 \ (29.4), \ 34.9, \ 30.9 \\ 41.7 \ (46.0), \ 44.4, \ 58.0 \\ 44.8 \ (46.2), \ 46.1, \ 49.9 \\ 55.9 \ (60.2), \ 60.0, \ 65.7 \\ 77.0 \ (86.0), \ 91.5, \ 98.7 \\ 104.5 \ (108.8), \ 106.6, \ 112.6 \\ 86.3 \ (95.2), \ 100.7, \ 107.9 \end{array}$ |
| [M(CysOMe-H)] ⁺ | [N,CO,S ⁻]tgg [N,CO,S ⁻]cgg [N,S ⁻]tgg [N,S ⁻]tgt [CO,S ⁻]tgg [CO,S ⁻]ctg [N,S ⁻]cgt [N ⁻ ,CO,S]tggg | 0.0 (0.0), 0.0, 0.0 36.2 (33.4), 35.2, 36.2 43.7 (38.9), 46.3, 37.2 36.9 (41.2), 43.4, 46.5 55.5 (61.2), 64.0, 74.1 76.9 (77.6), 83.5, 93.6 84.3 (84.7), 89.5, 92.8 102.3 (102.6), 103.3, 105.9 | $\begin{array}{c} 0.0 \ (0.0), \ 0.0, \ 0.0 \\ 37.2 \ (34.6), \ 36.0, \ 37.6 \\ 36.7 \ (32.8), \ 39.5, \ 33.2 \\ 27.0 \ (32.4), \ 31.5, \ 37.2 \\ 50.7 \ (57.4), \ 57.3, \ 70.0 \\ 72.3 \ (74.2), \ 77.0, \ 90.6 \\ 73.1 \ (74.5), \ 76.3, \ 82.1 \\ 104.4 \ (104.6), \ 106.4, \ 112.7 \end{array}$ |

^{*a*} Values including empirical dispersion corrections are given in parentheses. ^{*b*} Calculations performed at the B3LYP, B3P86, and MP2(full) levels of theory using a 6-311+G(2d,2p) basis set for Zn-containing complexes and def2TZVPP for Cd-containing complexes. Geometries and vibrational frequencies calculated at the B3LYP/6-311+G(d,p) level for Zn-containing complexes and B3LYP/def2TZVP for Cd-containing complexes. ^{*c*} Salt bridge between NH3⁺, CO₂⁻, and S⁻ groups.

^a Department of Chemistry, University of Utah, Salt Lake City, UT 84112, USA. E-mail: armentrout@chem.utah.edu

^b Radboud University, Institute for Molecules and Materials, FELIX Laboratory, Toernooiveld 7, NL-6525ED Nijmegen, The Netherlands

^c van't Hoff Institute for Molecular Sciences, University of Amsterdam, Amsterdam, The Netherlands

Correction

Table 2 Relative free energies (kJ mol⁻¹) at 298 K of low-lying CdCl⁺(CysOMe) conformers^a

| Structure | $B3LYP^b$ | B3P86 | MP2(full) |
|---------------------------|-------------|-------|-----------|
| [N,CO,S]tggg ₊ | 0.0 (0.0) | 0.0 | 0.0 |
| N,CO,S]tggg_ | 1.1 (1.7) | 1.2 | 1.7 |
| $[S^-]tcg^c$ | 17.6 (19.0) | 18.2 | 21.5 |
| [N,CO]tgtg_ | 20.1 (29.3) | 24.9 | 31.1 |
| [N,CO]tggg | 20.7 (29.2) | 25.3 | 30.7 |
| [N,CO] <i>tcgg</i> | 21.2 (30.2) | 25.9 | 31.8 |
| [N,S]tgtg | 23.4 (30.3) | 26.0 | 33.7 |
| $[N,CO]tgtg_+$ | 23.4 (32.8) | 28.3 | 35.0 |
| $[CO,S^-]ttg^c$ | 26.0 (30.1) | 26.7 | 31.0 |
| [N,OMe,S]tggg+ | 27.7 (22.2) | 28.6 | 21.5 |
| [N,OMe,S]tggg_ | 32.2 (27.7) | 33.4 | 26.7 |
| [N,CO,S] <i>cggg</i> | 38.8 (36.1) | 37.9 | 39.2 |
| [CO,S] <i>ttgt</i> | 51.0 (58.8) | 56.8 | 72.6 |
| $[CO,S^{-}]cgg^{c}$ | 66.2 (66.8) | 64.7 | 72.2 |
| [CO,S] <i>ctgt</i> | 82.5 (88.9) | 87.6 | 104.2 |

 a Calculations performed at the stated level of theory using a def2TZVPP basis set. Geometries and vibrational frequencies calculated at the B3LYP/ def2TZVP level of theory. b Values including empirical dispersion corrections are given in parentheses. c Salt bridge between NH3⁺, CO₂⁻, and S⁻ groups.

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