## **PCCP**



## CORRECTION

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## Correction: Trajectory analysis of single molecules exhibiting non-Brownian motion

Lindsay C. C. Elliott, Moussa Barhoum, Joel M. Harris\* and Paul W. Bohn\*c

Correction for 'Trajectory analysis of single molecules exhibiting non-Brownian motion' by Lindsay C. C. Elliott *et al., Phys. Chem. Chem. Phys.*, 2011, **13**, 4326–4334.

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It has been brought to our attention that an error exists in our 2011 paper, 'Trajectory analysis of single molecules exhibiting non-Brownian motion'.

The problem is in the definition of the eigenvalues, and just after eqn (2) on p. 4328 (3rd page of the article) should read: "... where  $R_1^2$  and  $R_2^2$  are the major and minor eigenvalues, respectively, of the radius of gyration tensor, T." This definition permits correct calculation of the radius of gyration,  $R_g$ , and allows the units to work out as you would expect, with  $R_g$  in meters.

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

a Department of Chemistry, University of Illinois at Urbana-Champaign, 600 S. Mathews Ave., Urbana, IL 61801, USA. E-mail: pbohn@nd.edu, harrisj@chem.utah.edu

<sup>&</sup>lt;sup>b</sup> Department of Chemistry, University of Utah, 315 South 1400 East, Salt Lake City, UT 84112, USA

<sup>&</sup>lt;sup>c</sup> Department of Chemical and Biomolecular Engineering and Department of Chemistry and Biochemistry, University of Notre Dame, Notre Dame, IN 46556, USA