

Data-driven Discovery in the Chemical Sciences

Trinity College, Oxford,
United Kingdom and online

10–12 September 2024



FARADAY DISCUSSIONS

Volume 256, 2025



ROYAL SOCIETY
OF **CHEMISTRY**



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Faraday Discussions (Print ISSN 1359-6640, Electronic ISSN 1364-5498) is published 8 times a year by the Royal Society of Chemistry, Thomas Graham House, Science Park, Milton Road, Cambridge, UK CB4 0WE.

Volume 256 ISBN 978-1-83767-442-8

2025 annual subscription price: print+electronic £1342

US \$2363; electronic only £1279, US \$2250.

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Printed in the UK



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Data-driven Discovery in the Chemical Sciences

Faraday Discussions

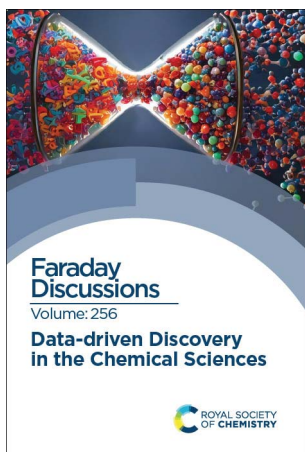
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A General Discussion on Data-driven Discovery in the Chemical Sciences was held in Oxford, UK and online on the 10th, 11th and 12th of September 2024.

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ISSN 1359-6640; ISBN 978-1-83767-442-8



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See Brett Savoie *et al.*, *Faraday Discuss.*, 2025, **256**, 104–119.

Large property models convert design constraints (depicted by letters- left) directly into molecules (emerging from the model - right).

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