## Matt I J Probert

List of Publications by Year in descending order

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| #  | Article  | IF   | CITATIONS |
|----|--|------|-----------|
| 1  | Materials and Molecular Modeling at the Exascale. Computing in Science and Engineering, 2022, 24, 36-45.   | 1.2  | 7         |
| 2  | Systematic Comparison of Genetic Algorithm and Basin Hopping Approaches to the Global<br>Optimization of Si(111) Surface Reconstructions. Journal of Physical Chemistry A, 2022, 126, 3043-3056.   | 2.5  | 5         |
| 3  | Electron–phonon interaction and superconductivity in hexagonal ternary carbides Nb <sub>2</sub><br>AC (A: Al, S, Ge, As and Sn). Electronic Structure, 2021, 3, 045001.                            | 2.8  | 5         |
| 4  | Effective modelling of the Seebeck coefficient of Fe <sub>2</sub> VAl. Journal of Physics Condensed Matter, 2020, 32, 125401.  | 1.8  | 4         |
| 5  | Anisotropy in antiferromagnets. Journal of Applied Physics, 2020, 128, .   | 2.5  | 19        |
| 6  | The emergence of sequence-dependent structural motifs in stretched, torsionally constrained DNA.<br>Nucleic Acids Research, 2020, 48, 1748-1763.   | 14.5 | 21        |
| 7  | Off-the-shelf DFT-DISPersion methods: Are they now "on-trend―for organic molecular crystals?.<br>Journal of Chemical Physics, 2019, 151, 044106.   | 3.0  | 11        |
| 8  | Simultaneous Prediction of the Magnetic and Crystal Structure of Materials Using a Genetic Algorithm. Crystals, 2019, 9, 439.  | 2.2  | 8         |
| 9  | Huge power factor in p-type half-Heusler alloys NbFeSb and TaFeSb. JPhys Materials, 2019, 2, 035002.   | 4.2  | 33        |
| 10 | Quantum diffusion of H/D on Ni(111)—A partially adiabatic centroid MD study. Journal of Chemical<br>Physics, 2018, 148, 102339.  | 3.0  | 4         |
| 11 | DL_MG: A Parallel Multigrid Poisson and Poisson–Boltzmann Solver for Electronic Structure<br>Calculations in Vacuum and Solution. Journal of Chemical Theory and Computation, 2018, 14, 1412-1432. | 5.3  | 31        |
| 12 | Many-body renormalization of forces in <mml:math<br>xmlns:mml="http://www.w3.org/1998/Math/MathML"&gt;<mml:mi>f</mml:mi> -electron<br/>materials. Physical Review B, 2018, 98, .</mml:math<br>     | 3.2  | 20        |
| 13 | Atomistic dynamics of sulfur-deficient high-symmetry grain boundaries in molybdenum disulfide.<br>Nanoscale, 2017, 9, 10312-10320.   | 5.6  | 18        |
| 14 | Transverse Fluorescence Microscopy with Magnetic and Optical Tweezers. Biophysical Journal, 2017, 112, 299a.   | 0.5  | 0         |
| 15 | Experimental and density functional study of Mn doped Bi2Te3 topological insulator. APL Materials, 2016, 4, .  | 5.1  | 14        |
| 16 | Reproducibility in density functional theory calculations of solids. Science, 2016, 351, aad3000.  | 12.6 | 1,113     |
| 17 | Exploring atomic defects in molybdenum disulphide monolayers. Nature Communications, 2015, 6, 6293.  | 12.8 | 1,124     |
| 18 | Theoretical study of core-loss electron energy-loss spectroscopy at graphene nanoribbon edges.<br>Journal of Physics Condensed Matter, 2015, 27, 305301.   | 1.8  | 5         |

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|----|--|------|-----------|
| 19 | The Effect of Cobalt-Sublattice Disorder on Spin Polarisation in Co2FexMn1â^'xSi Heusler Alloys.<br>Materials, 2014, 7, 1473-1482.   | 2.9  | 9         |
| 20 | Density functional theory in the solid state. Philosophical Transactions Series A, Mathematical,<br>Physical, and Engineering Sciences, 2014, 372, 20130270.                   | 3.4  | 242       |
| 21 | 4th Workshop on Theory, Modelling and Computational Methods for Semiconductors (TMCSIV).<br>Journal of Physics: Conference Series, 2014, 526, 011001.                          | 0.4  | 0         |
| 22 | Quantum simulation of low-temperature metallic liquid hydrogen. Nature Communications, 2013, 4, 2064.  | 12.8 | 75        |
| 23 | Classical and quantum ordering of protons in cold solid hydrogen under megabar pressures. Journal of Physics Condensed Matter, 2013, 25, 085402.                               | 1.8  | 25        |
| 24 | 3rd Workshop on Theory, Modelling and Computational Methods for Semiconductors (TMCSIII).<br>Journal of Physics: Conference Series, 2012, 367, 011001.                         | 0.4  | 0         |
| 25 | Atomistic molecular dynamics simulations of shock compressed quartz. Journal of Chemical Physics, 2011, 135, 044508.   | 3.0  | 9         |
| 26 | Crystal structure prediction for iron as inner core material in heavy terrestrial planets. Earth and Planetary Science Letters, 2011, 312, 237-242.                            | 4.4  | 32        |
| 27 | Electron and vibrational spectroscopies using DFT, plane waves and pseudopotentials: CASTEP implementation. Computational and Theoretical Chemistry, 2010, 954, 22-35.         | 1.5  | 205       |
| 28 | Hydrogen sorption sites in holmium silicide on silicon(1 1 1). Surface Science, 2010, 604, 686-691.  | 1.9  | 0         |
| 29 | An <i>ab initio</i> study of xenon retention in α-quartz. Journal of Physics Condensed Matter, 2010, 22, 025501.   | 1.8  | 19        |
| 30 | Quantum Nature of the Proton in Water-Hydroxyl Overlayers on Metal Surfaces. Physical Review<br>Letters, 2010, 104, 066102.  | 7.8  | 101       |
| 31 | The structure and growth direction of rare earth silicide nanowires on Si(100). Applied Physics<br>Letters, 2010, 96, 241903.  | 3.3  | 13        |
| 32 | Theory, Modelling and Computational methods for Semiconductors. Journal of Physics: Conference Series, 2010, 242, 011001.  | 0.4  | 0         |
| 33 | An experiment on the Purcell effect in a wedge cavity. European Journal of Physics, 2009, 30, S81-S88.   | 0.6  | 3         |
| 34 | Improved real-space genetic algorithm for crystal structure and polymorph prediction. Physical Review B, 2008, 77, .   | 3.2  | 37        |
| 35 | Quantitative LEEDIâ^Vandab initiostudy of theSi(111)â^'3×2â^'Smsurface structure and the missing half-order spots in the3×1diffraction pattern. Physical Review B, 2007, 75, . | 3.2  | 8         |
| 36 | A periodic genetic algorithm with real-space representation for crystal structure and polymorph prediction. Physical Review B, 2006, 73, .                                     | 3.2  | 145       |

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|----|--|-----|-----------|
| 37 | Ab Initio Path Integral Molecular Dynamics Simulation of Hydrogen in Silicon. AIP Conference<br>Proceedings, 2006, , .   | 0.4 | 0         |
| 38 | STM andab initiostudy of holmium nanowires on a Ge(111) surface. Physical Review B, 2006, 74, .  | 3.2 | 6         |
| 39 | Orientational effects of twisted light on twisted nematic liquid crystals. Journal of Physics B:<br>Atomic, Molecular and Optical Physics, 2006, 39, S523-S528.                                    | 1.5 | 2         |
| 40 | Constant pressure Langevin dynamics: theory and application. Computer Physics Communications, 2005, 169, 322-325.  | 7.5 | 13        |
| 41 | Phase behavior of a three-dimensional core-softened model system. Physical Review E, 2005, 71, 065701.   | 2.1 | 16        |
| 42 | Progression of phase behavior for a sequence of model core-softened potentials. Physical Review E, 2005, 72, 061202.   | 2.1 | 4         |
| 43 | First principles methods using CASTEP. Zeitschrift Fur Kristallographie - Crystalline Materials, 2005, 220, .  | 0.8 | 9,458     |
| 44 | Stiffness and thermal expansion of ZrB2: anab initiostudy. Journal of Physics Condensed Matter, 2005, 17, 2233-2241.   | 1.8 | 85        |
| 45 | Langevin dynamics in constant pressure extended systems. Journal of Chemical Physics, 2004, 120, 11432-11441.  | 3.0 | 116       |
| 46 | Improved algorithm for geometry optimisation using damped molecular dynamics. Journal of Computational Physics, 2003, 191, 130-146.  | 3.8 | 15        |
| 47 | Non-equilibrium electron transport in degenerate nitride heterostructures—dynamic screening<br>effects. Physica E: Low-Dimensional Systems and Nanostructures, 2003, 17, 272-275.                  | 2.7 | 5         |
| 48 | Improving the convergence of defect calculations in supercells: Anab initiostudy of the neutral silicon vacancy. Physical Review B, 2003, 67, .  | 3.2 | 163       |
| 49 | Molecular dynamics studies of liquids using a Beowulf computer. Contemporary Physics, 2003, 44, 435-450.   | 1.8 | 1         |
| 50 | Comment on `Checking the influence of numerically induced chaos in the computational study of intramolecular dynamics using trajectory equivalence'. Chemical Physics Letters, 2002, 354, 529-531. | 2.6 | 1         |
| 51 | First-principles simulation: ideas, illustrations and the CASTEP code. Journal of Physics Condensed Matter, 2002, 14, 2717-2744.   | 1.8 | 8,382     |
| 52 | Comment on "Algorithm for normal random numbers― Physical Review E, 2001, 63, 058701; author<br>reply 058702.  | 2.1 | 3         |
| 53 | Anab initiostudy of muons in ethanal. Journal of Physics Condensed Matter, 1997, 9, 3241-3257.   | 1.8 | 6         |
| 54 | Ab initiostudies of magnetism in the organic radicalp-NPNN. Journal of Physics Condensed Matter, 1997, 9, 3635-3645.   | 1.8 | 4         |

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|----|--|-----|-----------|
| 55 | Potential for a novel μSR experiment — the results of an ab initio study. Chemical Physics Letters, 1996,<br>259, 271-275.   | 2.6 | 7         |
| 56 | Theoretical studies of implanted muons in organic magnets. Materials Science and Engineering B:<br>Solid-State Materials for Advanced Technology, 1996, 37, 247-250. | 3.5 | 3         |
| 57 | Computer simulations of flux motion in high-temperature superconductors. Physica B: Condensed Matter, 1995, 205, 180-182.  | 2.7 | 0         |
| 58 | Nature of the Irreversibility Line in High-Temperature Superconductors. Physical Review Letters, 1995, 75, 1835-1838.  | 7.8 | 14        |