Appendix B: Statement of Authorship

This declaration concerns the article entitled:											
Atomistic insights into the order-disorder transition in Cu2ZnSnS4 solar cells from Monte Carlo simulations											
Pages 89 - 99 of thesis											
Publication status (tick one)											
draft manuscript		Submitted		In review		Accept	ted		Published	Х	
Publication details (reference)	Atomistic insights into the order-disorder transition in Cu2ZnSnS4 solar cells from Monte Carlo simulations SK Wallace, JM Frost, A Walsh Journal of Materials Chemistry A, 2019, DOI: 10.1039/C8TA04812F										
Candidate's contribution to the paper (detailed, and also given as a percentage).	Form J. Fr prov analy Wall throw through the simulation of the sim	Formulation of ideas (50%): The original code used in this work was provided by J. Frost and the idea to investigate Cu-Zn disorder in Cu2ZnSnS4 (CZTS) was provided by A. Walsh. Ideas for subsequent developments of the code and analysis of the data were mostly formulated and implemented by myself (S. Wallace) with support and discussions provided by J. Frost and A. Walsh throughout this study. **Design of methodology (50%): The Bespoke Monte Carlo model, Eris, (doi: 10.5281/zenodo.1248445) developed for this publication is based on the Starry Night code (doi: 10.5281/zenodo.10543) written by J. Frost to simulate dipole-dipole interactions and ferroelectric domains in a hybrid organic-inorganic perovskite solar cell. Eris was originally adapted by J. Frost from Starry Night to simulate thermodynamic Cu-Zn disorder in CZTS. Developments of the Eris code by S. Wallace include writing routines for: **Lattice initialisation methods to ensure a stoichiometric CZTS crystal** **Testing the convergence in the electrostatic summations for calculating the change in lattice energy when performing a Monte Carlo move* **Developing and testing methods to check that the equilibrium disordered configuration was achieved for each simulation temperature* **Additional outputs from the code including: data to compute an order parameter, tests for convergence when computing on-site electrostatic potentials and outputting lattice on-site electrostatic potentials in various forms for further analysis*									
Statement from Candidate	This paper reports on original research I conducted during the period of my Higher Degree by Research candidature.										
Signed		5.6	Wa	u			Date	0	5/12/18		