

# <sup>1</sup> dKMC: Delocalised kinetic Monte Carlo for simulating fundamental transport processes involving partially delocalised carriers in disordered materials

<sup>4</sup> **Daniel Balzer**  <sup>1</sup> and **Ivan Kassal**  

<sup>5</sup> <sup>1</sup> School of Chemistry, University of Sydney, NSW 2006, Australia  Corresponding author

DOI: [10.xxxxxx/draft](https://doi.org/10.xxxxxx/draft)

## Software

- [Review](#) 
- [Repository](#) 
- [Archive](#) 

Editor: [Open Journals](#) 

Reviewers:

- [@openjournals](#)

Submitted: 01 January 1970

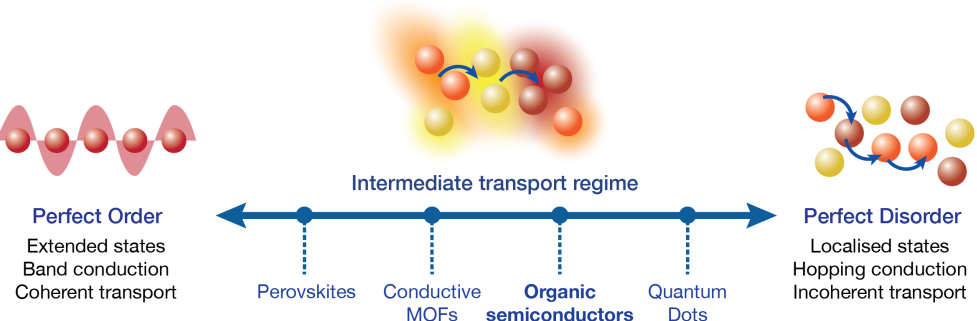
Published: unpublished

## License

Authors of papers retain copyright and release the work under a Creative Commons Attribution 4.0 International License ([CC BY 4.0](#)).<sup>17</sup><sup>18</sup><sup>19</sup><sup>20</sup><sup>21</sup>

## <sup>6</sup> Summary

<sup>7</sup> The movement of charge and energy is a fundamental process in materials science, underpinning <sup>8</sup> technologies such as solar cells, light-emitting diodes, batteries, and electronics. Transport <sup>9</sup> is well understood in both highly ordered materials (band conduction) and highly disordered <sup>10</sup> ones (hopping conduction). However, in moderately disordered materials—including many <sup>11</sup> organic semiconductors—transport lies in the intermediate transport regime between these <sup>12</sup> well-understood extremes. Accurately modelling intermediate-regime conduction is difficult <sup>13</sup> because describing wavefunction delocalisation requires a fully quantum-mechanical treatment, <sup>14</sup> which is challenging in disordered materials that lack periodicity. We describe delocalised <sup>15</sup> kinetic Monte Carlo (dKMC), the first theoretical approach to treat, in three dimensions, all the <sup>16</sup> processes crucial in organic semiconductors: disorder, delocalisation, and polaron formation. As <sup>17</sup> a result, it can treat the intermediate transport regime between band and hopping conduction. <sup>18</sup> dKMC reveals that the fundamental physics of transport in moderately disordered materials is <sup>19</sup> that of charges and excitons hopping between partially delocalised electronic states. In this <sup>20</sup> work, we release the dKMC.jl package, which contains modules for simulating the fundamental <sup>21</sup> processes of charge and exciton transport as well as charge separation and generation.



**Figure 1:** Figure 1: While transport mechanisms are well understood in the extremes, coherent band conduction through extended states and incoherent hopping through localised states, they remain poorly understood in the intermediate regime where many organic semiconductors lie. Figure adapted with permission from (Balzer et al., 2021).

## <sup>22</sup> Statement of need

<sup>23</sup> The need for dKMC is demonstrated by its ability to solve two important problems.

24 First, dKMC is the first computational technique that is able to include all of the processes  
25 crucial in organic semiconductors while remaining computationally tractable enough to treat  
26 realistic, three-dimensional systems on mesoscopic time and length scales (Balzer et al., 2021).  
27 Before dKMC, the difficulty in modelling transport in the intermediate regime had prevented  
28 the development of such a theory. Instead, prior approaches either needed to exclude one  
29 of the key ingredients (disorder, delocalisation, and polaron formation), which could lead  
30 to inaccurate results, or included them all but restricted the application to smaller systems  
31 (dimension, time, and length scales). dKMC solves these problems, striking a balance between  
32 the level of approximation and the size of the system it can simulate. Therefore, dKMC provides  
33 a simulation tool that can both accurately and efficiently simulate fundamental transport  
34 processes involving partially delocalised carriers.

35 Second, dKMC explains the often confusing behaviour of organic electronics, including organic  
36 photovoltaics (OPVs). Most models of transport in disordered organic semiconductors assume  
37 hopping transport, where charge carriers or excitons are localised onto individual molecules and  
38 move via thermally assisted hops from one molecule to the next. However, hopping transport  
39 fails to explain how charges and excitons move as fast as they do, or how charges in OPVs  
40 overcome their strong Coulomb attraction and separate from CT states as efficiently as they  
41 do, or how charges are generated so efficiently even in OPV devices with little to no energetic  
42 offsets. Hopping transport fails because, in many organic semiconductors, the charges and  
43 excitons remain delocalised across multiple molecules. By going beyond hopping, dKMC reveals  
44 that delocalisation improves each of the four fundamental transport processes in OPVs: charge  
45 transport (Balzer et al., 2021), exciton transport (Balzer & Kassal, 2023), charge separation  
46 (Balzer & Kassal, 2022), and charge generation (Balzer & Kassal, 2024). Delocalisation  
47 improves all of these important transport processes in essentially the same way, by enabling  
48 carriers to hop further and faster, explaining the failure of classical theories. Therefore, dKMC is  
49 a tool for explaining the otherwise unpredictable behaviour observed in devices.

50 dKMC was developed for organic semiconductors, but it can be more generally applied to other  
51 materials that lie in the intermediate regime. In particular, the first application of dKMC was to  
52 OPVs and therefore the package contains a module for each of the four fundamental processes in  
53 OPVs: charge transport, exciton transport, charge separation, and charge generation. However,  
54 the transport modules can be more generally applied to organic semiconductor materials used  
55 in other devices, such as organic light-emitting diodes or organic field-effect transistors.

## 56 Acknowledgements

57 We were supported by the Westpac Scholars Trust (Research Fellowship and Future Leaders  
58 Scholarship), by the Australian Research Council (DP220103584), by the Australian Government  
59 Research Training Program, and by the University of Sydney Nano Institute Grand Challenge  
60 Computational Materials Discovery. We were also supported by computational resources from  
61 the National Computational Infrastructure (Gadi) and by the University of Sydney Informatics  
62 Hub (Artemis).

## 63 References

- 64 Balzer, D., & Kassal, I. (2022). Even a little delocalization produces large kinetic enhancements  
65 of charge-separation efficiency in organic photovoltaics. *Science Advances*, 8(32), eabl9692.  
66 <https://doi.org/10.1126/sciadv.abl9692>
- 67 Balzer, D., & Kassal, I. (2023). Mechanism of delocalization-enhanced exciton transport  
68 in disordered organic semiconductors. *J. Phys. Chem. Lett.*, 14(8), 2155–2162. <https://doi.org/10.1021/acs.jpclett.2c03886>
- 70 Balzer, D., & Kassal, I. (2024). Delocalisation enables efficient charge generation in organic

<sup>71</sup> photovoltaics, even with little to no energetic offset. *Chemical Science*, 15, 4779–4789.  
<sup>72</sup> <https://doi.org/10.1039/D3SC05409H>

<sup>73</sup> Balzer, D., Smolders, T. J. A. M., Blyth, D., Hood, S. N., & Kassal, I. (2021). Delocalised  
<sup>74</sup> kinetic Monte Carlo for simulating delocalisation-enhanced charge and exciton transport  
<sup>75</sup> in disordered materials. *Chemical Science*, 12, 2276–2285. <https://doi.org/10.1039/D0SC04116E>  
<sup>76</sup>

DRAFT