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Swapping in lattice-based cell migration models

Shahzeb Raja Noureen,^{1,*} Jennifer P. Owen,¹ Richard L. Mort,² and Christian A. Yates¹

¹*Centre for Mathematical Biology, University of Bath,
Claverton Down, Bath, BA2 7AY, UK.*

²*Division of Biomedical and Life Sciences,
Faculty of Health and Medicine, Furness Building,
Lancaster University, Bailrigg, Lancaster, LA1 4YG, UK.*

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Abstract

Cell migration is frequently modelled using on-lattice agent-based models (ABMs) that employ the excluded volume interaction. However, cells are also capable of exhibiting more complex cell-cell interactions, such as adhesion, repulsion, pulling, pushing and swapping. Although the first four of these have already been incorporated into mathematical models for cell migration, swapping has not been well studied in this context. In this paper, we develop an ABM for cell movement in which an active agent can ‘swap’ its position with another agent in its neighbourhood with a given swapping probability. We consider a two-species system for which we derive the corresponding macroscopic model and compare it with the average behaviour of the ABM. We see good agreement between the ABM and the macroscopic density. We also analyse the movement of agents at an individual level in the single-species as well as two-species scenarios to quantify the effects of swapping on an agent’s motility.

Keywords: agent-based model, lattice, swapping, cell migration, pattern formation.

* Corresponding author: srn32@bath.ac.uk

23 I. INTRODUCTION

24 Cell migration is an essential biological process required for the correct development of
25 tissues and organs during embryonic development and their proper maintenance, through
26 wound healing and tissue homeostasis, throughout life [1–5]. During embryonic develop-
27 ment, neural crest cells delaminate from the dorsal most aspect of the neural tube and
28 migrate to colonise their target tissues including the gut in the case of enteric ganglia
29 precursors and the skin in the case of melanoblasts the precursors of melanocytes [6].
30 Diseases of the neural crest are known as neurocristopathies for example failure of the
31 enteric ganglia precursors to colonise the developing gut results in Hirschsprung’s disease
32 [7] while failure of melanoblasts to colonise the developing epidermis results in piebaldism
33 [8]. Therefore an in-depth understanding of cell migration is important for identifying
34 the causes of neurocristopathies [9–12] as well as developing new therapeutic targets to
35 prevent metastasis in cancers [9, 13].

36 Traditionally, many biological problems have been modelled using deterministic meth-
37 ods. However, in cell migration, randomness can play a salient role in determining a
38 cell’s trajectory and fate and hence deterministic theory may not be appropriate. Ex-
39 tensive research has gone into modelling cell movement as a stochastic process [14–18].
40 In one widely used approach, cells are modelled as agents whose positions evolve prob-
41 abilistically in space and time according to a predefined set of rules. These models are
42 commonly known as agent-based models (ABMs) or individual-based models (IBMs).
43 The agent-based modelling paradigm can be sub-divided into off-lattice and on-lattice
44 models, both of which have wide applicability to different problems within mathematical
45 biology. Gavagnin and Yates [19] recently reviewed the most commonly used ABMs for
46 cell movement.

47 In this paper, we only concern ourselves with on-lattice models of cell movement. In a
48 lattice-based approach, the domain is divided into a series of compartments in which the
49 cells reside. Cells take up space, preventing other cells from occupying the same space at
50 the same time. For biological plausibility, it is often desirable that mathematical models
51 of cell migration account for the single occupancy of sites. This realism is incorporated in
52 an ABM via the volume-exclusion principle, which states that a cell attempting to move
53 into a neighbouring site successfully moves only if the neighbouring site is not already
54 occupied at the time of moving.

55 Models with volume exclusion at their core have been used to describe the collective

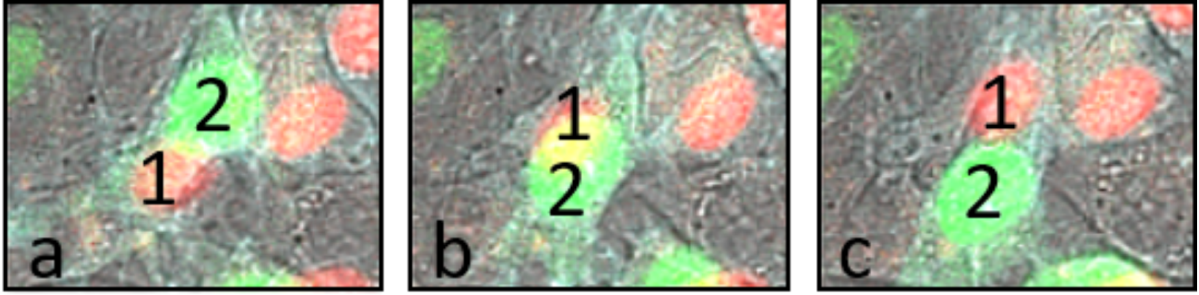


Figure 1. Two *Fucci2a* labelled NIH 3T3 mouse embryonic fibroblasts swapping places with each other in culture [26]. The colour of the cells represents their cell cycle stage (Red = G1, Green = S/G2/M) in this case making it easy to observe the swap. In (a), we show the initial placement of cells: cell 1 (shown in red) is on the bottom-left of cell 2 (shown in green). In (b) a swap starts to take place and in (c) the swap is complete and now cell 1 is on the top-right of cell 2.

56 migration of cells for a wide range of biological applications. Mort *et al.* [8] used an
 57 on-lattice ABM with the exclusion principle to model the invasion of the developing
 58 epidermis by melanoblasts (the embryonic precursors of melanocytes) and investigate
 59 the basis of piebaldism in mice. Lattice-based exclusion models have also been applied
 60 to wound healing [20], migration of breast cancer cells [21], developmental processes on
 61 growing domains [22], cells' responses to chemotaxis [23] and cells exhibiting pushing [24]
 62 and pulling [25] interactions in densely crowded environments.

63 Biologically, although cells are excluded from the space occupied by other cells, their
 64 movement is not completely inhibited by them as typically assumed in volume-exclusion
 65 models. For example melanoblasts are able to move freely between keratinocytes in the
 66 developing epidermis [8]. Experimental data suggests that cells are often able to move
 67 past each other (passing laterally, above or below) exchanging places with one another.
 68 In Figure 1 we show experimental images of two *Fucci2a* labelled NIH-3T3 fibroblasts
 69 exhibiting the swapping behaviour. Swapping has also been observed in blood cells such as
 70 leukocytes, erythrocytes and thrombocytes [27] and in pattern formation for maintaining
 71 sharp boundaries between different groups of cells as part of a cell sorting mechanism
 72 [28]. These examples highlight the importance of incorporating swapping into models of
 73 cell migration. To the best of our knowledge, this has not yet been explored thoroughly
 74 from a mathematical perspective.

75 In this paper, we develop a mathematical model to describe and analyse cell-cell swap-
 76 ping in two species setting. By modifying the movement rules of the traditional volume-
 77 exclusion process, we show that swapping between agents has an effect on the migration

78 of agents at different spatial resolutions. To investigate how swapping manifests itself
79 in the corresponding population-level model (PLM) we derive a set of partial differential
80 equations (PDEs) describing the macroscopic dynamics of the agents. We compare nu-
81 merical solutions of the PDEs with the averaged results from the ABM and comment on
82 the agreement or discrepancy between them. We also analyse the movement of agents at
83 an individual level and derive expressions for the individual-level diffusion coefficient.
84 The remainder of this paper is structured as follows. In Section II A, we develop a
85 model that allows for swapping to take place between pairs of neighbouring agents. In
86 Section II B, we derive the macroscopic PDEs and compare the average behaviour of
87 the ABM to that of the PDEs. In Section III, we analyse the movement of agents at an
88 individual level and derive a relationship between the individual-level diffusion coefficient,
89 swapping probability and background domain density. In Section IV, we give examples
90 to illustrate the applications of swapping and finally in Section V, we conclude the paper
91 with a summary and discussion.

92 II. CELL MIGRATION MODEL WITH SWAPPING

93 We begin by developing an ABM for cell movement with swapping in Section II A and
94 we use this to investigate the effect of swapping on the mobility of agents. In particular,
95 we look at the effect of swapping on mixing of agents in a two-species system. We
96 derive the population-level model and compare this with the average behaviour of the
97 ABM in Section II B. We also analyse the individual-level behaviour in both single and
98 multispecies scenarios in Section III.

99 A. On-lattice agent-based model

100 We model cell migration on a two-dimensional lattice. We discretise the domain into
101 compartments (also known as ‘sites’) such that there are L_x compartments in the ho-
102 rizontal direction and L_y compartments in the vertical direction. We assume that the
103 compartments are square with side length Δ . Supposing that each compartment can
104 contain no more than one agent, Δ can be considered a rough proxy for a cell’s diameter.
105 A site (i, j) for $i = 1, \dots, L_x$ and $j = 1, \dots, L_y$ can be either occupied by a type-A or a
106 type-B agent or unoccupied. Occupancy of a site (i, j) for a type-A (or type-B) agent is
107 defined as a binary indicator, taking a value of 1 if there is a type-A (or type-B) agent

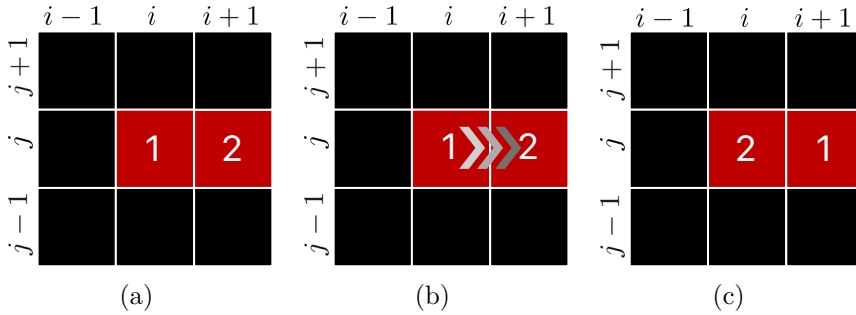


Figure 2. A schematic illustrating the swapping mechanism. Red sites are occupied with agents and black sites are unoccupied. The initial configuration of the lattice before is shown in (a). The agent chosen to move is at site (i, j) and labelled 1. The target site is at position $(i + 1, j)$ and the agent occupying the target site is labelled 2. Agent 1 attempts to move into the target site in (b). The final configuration once the swapping move is complete is shown in (c).

108 at the site (i, j) or 0 if the site is empty.

109 We initialise the lattice with species A and species B agents at densities c_A and c_B ,
 110 respectively, such that $c_A + c_B = c$ and $0 \leq c \leq 1$ where c is the overall domain density.

111 We let the positions of the agents evolve in continuous time according to the Gillespie
 112 algorithm [29]. Let r_A be the rate of movement of a type-A agent and let r_B be the
 113 equivalent for a type-B agent. The rates of movement are defined such that $r_A \delta t$ (and
 114 equivalently $r_B \delta t$) is the probability that a type-A (or type-B) agent attempts to move
 115 during a finite time interval of duration δt . The agent attempts to move into one of
 116 the four sites in its Von Neumann neighbourhood with equal probability. If the chosen
 117 neighbouring site is empty, the focal agent successfully moves and its position is updated.
 118 However, if another agent already occupies the site, the move is aborted [19, 24, 25, 30, 31].
 119 This blocking of the move characterises volume exclusion in our model.

120 Swapping works by modifying the rules of the exclusion process by allowing an exchange
 121 in the positions of a pair of neighbouring agents if the target site is already occupied. We
 122 now introduce the swapping parameter ρ to denote the probability of a successful swap
 123 between a pair of neighbouring agents conditional on one of the agents attempting to
 124 move into the other's position. If $\rho = 0$ then there are no swaps and we arrive back at
 125 the original exclusion process. If $\rho > 0$ then we can have different levels of swapping based
 126 on the value of ρ . For example, for $\rho = 1$ each scenario in which a move is attempted into
 127 an occupied target site will be a successful swap and for $\rho = 0.5$ half of the attempted
 128 moves into occupied sites will be successful.

129 To implement swapping, we sample a random number u from the uniform distribution

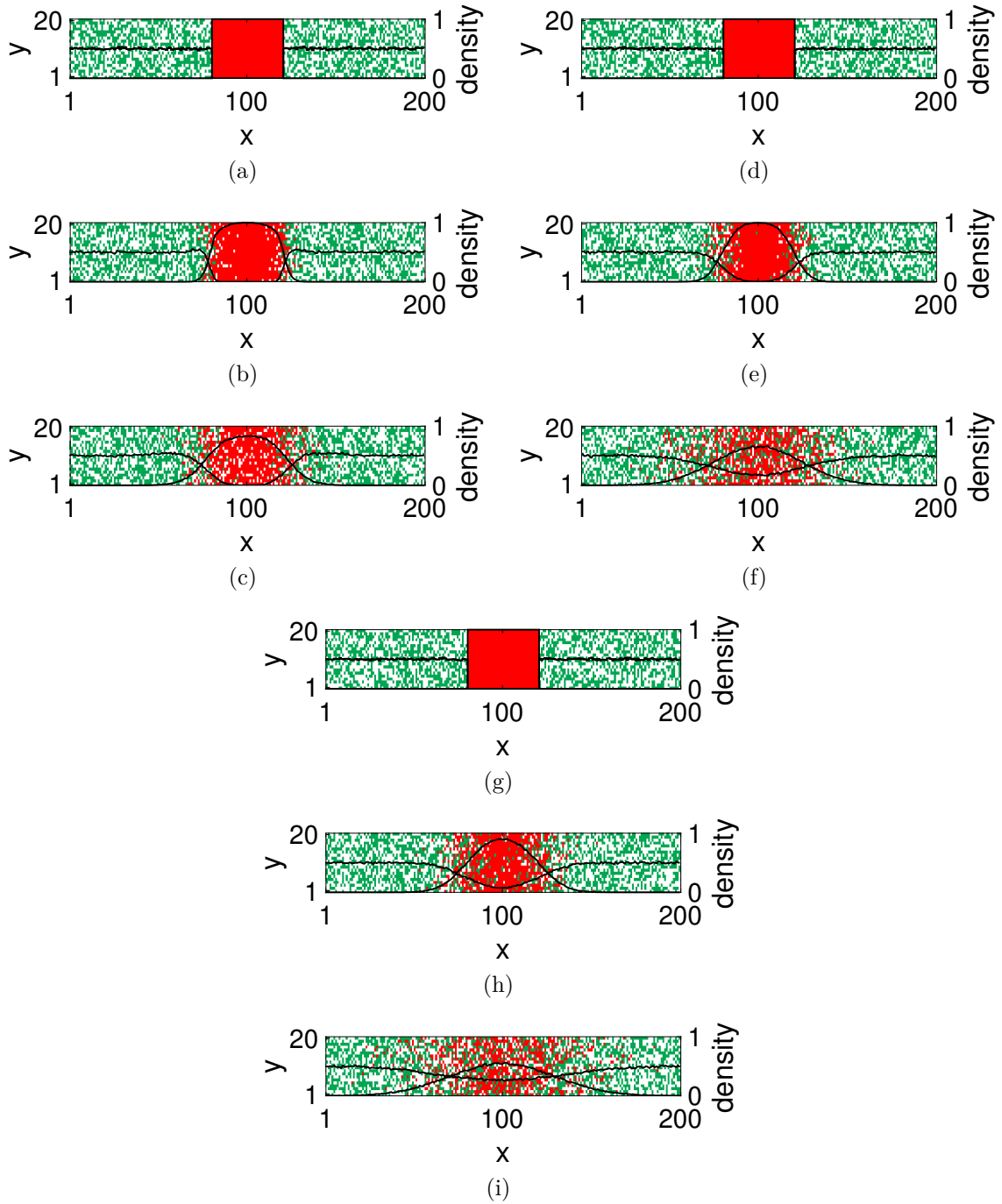


Figure 3. Snapshots of the lattice occupancy of the multi-species swapping model at $t = 0, 100, 1000$ using $r_A = r_B = 1$ for swapping probabilities $\rho = 0$ [(a)-(c)], $\rho = 0.5$ [(d)-(f)] and $\rho = 1$ [(g)-(i)]. Agents are initialised on a domain with dimensions $L_x = 200$ and $L_y = 20$ such that all the sites in the range $81 \leq x \leq 120$ are occupied by type-A agents (red) and the remaining sites are randomly populated with agents of type B (green) at a density of 0.5 [(a),(d),(g)]. Further snapshots of the ABM at $t = 100$ and $t = 1000$ show the dispersal of agents with time. The column-averaged density of the two species over 100 runs of the ABM is also plotted (shown in black). We impose reflective boundary conditions on all four boundaries of the domain.

130 over the unit interval $(0, 1)$. If $u < \rho$, the agent at the site (i, j) swaps with the agent
 131 at the target site (Figure 2(a)-(c)), otherwise the move is aborted. Figure 2 shows a
 132 successful swap between two agents labelled ‘1’ and ‘2’.

133 In this article we present results for the two-species model only unless stated otherwise.
 134 Results for the single-species model can be found in Appendix A.

135 In Figure 3 we present snapshots of lattice occupancy of a 200 by 20 grid occupied
 136 with type-A (red) and type-B (green) agents. The two species move with equal rates,
 137 $r_A = r_B = 1$. We see that a non-zero swapping probability results in faster dispersion
 138 of the agents compared to the $\rho = 0$ case. We also note that increasing the swapping
 139 probability from $\rho = 0.5$ the $\rho = 1$ results in faster dispersion of the agents, as expected.

140 In the next section, we derive the macroscopic PDEs describing the evolution of the mean
 141 lattice occupancy. By analysing the PDEs we generate further insight into the behaviours
 142 observed in Figure 3.

143 B. Continuum model

144 Let $A_{ij}^k(t)$ be the occupancy of site (i, j) at time t on the k th repeat of the ABM such
 145 that $A_{ij}^k(t) = 1$ if the site is occupied by a type-A agent and 0 otherwise. Let $B_{ij}^k(t)$ be
 146 the same for a type-B agent. Then the average density of type-A and type-B agents after
 147 K repeats is given by,

$$\langle A_{ij}(t) \rangle = \frac{1}{K} \sum_{k=1}^K A_{ij}^k(t), \quad \text{and} \quad \langle B_{ij}(t) \rangle = \frac{1}{K} \sum_{k=1}^K B_{ij}^k(t). \quad (1)$$

148 In what follows we will typically drop the notation for time dependence of our species
 149 densities, i.e. $\langle A_{ij}(t) \rangle = \langle A_{ij} \rangle$ and $\langle B_{ij}(t) \rangle = \langle B_{ij} \rangle$, for conciseness. By considering all
 150 the possible ways in which the site (i, j) can gain or lose occupancy of either type-A or
 151 type-B agents during the time step δt , we can write down the corresponding occupancy
 152 master equations at time $t + \delta t$:

$$\begin{aligned} \langle A_{ij}(t + \delta t) \rangle - \langle A_{ij} \rangle &= \frac{r_A}{4} \delta t [(1 - \langle A_{ij} \rangle - \langle B_{ij} \rangle) (\langle A_{i-1,j} \rangle + \langle A_{i+1,j} \rangle + \langle A_{i,j-1} \rangle + \langle A_{i,j+1} \rangle) \\ &\quad - \langle A_{ij} \rangle (4 - \langle A_{i-1,j} \rangle - \langle A_{i+1,j} \rangle - \langle A_{i,j-1} \rangle - \langle A_{i,j+1} \rangle - \langle B_{i-1,j} \rangle \\ &\quad - \langle B_{i+1,j} \rangle - \langle B_{i,j-1} \rangle - \langle B_{i,j+1} \rangle)] \end{aligned}$$

$$\begin{aligned}
& + \frac{(r_A + r_B)}{4} \rho \delta t \langle B_{ij} \rangle (\langle A_{i-1,j} \rangle + \langle A_{i+1,j} \rangle + \langle A_{i,j-1} \rangle + \langle A_{i,j+1} \rangle) \\
& - \frac{(r_A + r_B)}{4} \rho \delta t \langle A_{ij} \rangle (\langle B_{i-1,j} \rangle + \langle B_{i+1,j} \rangle + \langle B_{i,j-1} \rangle + \langle B_{i,j+1} \rangle),
\end{aligned} \tag{2}$$

$$\begin{aligned}
\langle B_{ij}(t + \delta t) \rangle - \langle B_{ij} \rangle &= \frac{r_B}{4} \delta t [(1 - \langle B_{ij} \rangle - \langle A_{ij} \rangle) (\langle B_{i-1,j} \rangle + \langle B_{i+1,j} \rangle + \langle B_{i,j-1} \rangle + \langle B_{i,j+1} \rangle) \\
& - \langle B_{ij} \rangle (4 - \langle B_{i-1,j} \rangle - \langle B_{i+1,j} \rangle - \langle B_{i,j-1} \rangle - \langle B_{i,j+1} \rangle - \langle A_{i-1,j} \rangle \\
& - \langle A_{i+1,j} \rangle - \langle A_{i,j-1} \rangle - \langle A_{i,j+1} \rangle)] \\
& + \frac{(r_A + r_B)}{4} \rho \delta t \langle A_{ij} \rangle (\langle B_{i-1,j} \rangle + \langle B_{i+1,j} \rangle + \langle B_{i,j-1} \rangle + \langle B_{i,j+1} \rangle) \\
& - \frac{(r_A + r_B)}{4} \rho \delta t \langle B_{ij} \rangle (\langle A_{i-1,j} \rangle + \langle A_{i+1,j} \rangle + \langle A_{i,j-1} \rangle + \langle A_{i,j+1} \rangle).
\end{aligned} \tag{3}$$

153 For illustration, we describe the terms in Equation (2). The terms in Equation (3) carry
154 similar interpretation. A site (i, j) can gain occupancy of type A in one of the following
155 three ways:

- 156 1. The site (i, j) is unoccupied and a type-A agent moves in from a neighbouring site
157 (line 1 in Equation (2)).
- 158 2. The site (i, j) is occupied by a type-B agent, which initiates and completes a swap
159 with a type-A agent at a neighbouring site (line 3 in Equation (2)).
- 160 3. The site (i, j) is occupied by a type-B agent and a type-A agent at a neighbouring
161 site initiates a swap to exchange positions with the type-B agent in the site (i, j)
162 (also line 3 in Equation (2)).

163 In all three cases, a type-A agent moves into the site (i, j) . Similarly, there are three ways
164 for a type-A agent to move *out* of the site (i, j) leading to a loss in the corresponding
165 occupancy:

- 166 1. The site is occupied by a type-A agent which jumps out to an unoccupied neigh-
167 bouring site, leaving the site (i, j) empty (line 2 in Equation (2)).
- 168 2. The site (i, j) is occupied by a type-A agent which initiates a swap with a type-B
169 agent in its neighbourhood.

170 3. The site (i, j) is occupied by a type-A agent and a type-B agent in the neighbouring
 171 site initiates a swap to exchange positions with the agent in site (i, j) (line 4 in
 172 Equation (2)).

173 In all three cases, a type-A agent moves out of the site (i, j) .

174 To obtain the continuum model, we Taylor expand the appropriate terms on the RHS
 175 of Equations (2) and (3) around the site (i, j) keeping terms of up to second order. By
 176 letting $\Delta \rightarrow 0$ and $\delta t \rightarrow 0$ such that $\Delta^2/\delta t$ is held constant, we arrive at the coupled
 177 PDEs,

$$\frac{\partial A}{\partial t} = \nabla \cdot [D_1(B)\nabla A + D_2(A)\nabla B], \quad (4)$$

$$\frac{\partial B}{\partial t} = \nabla \cdot [D_3(A)\nabla B + D_4(B)\nabla A], \quad (5)$$

where,

$$D_1(B) = D_A(1 - B) + \rho(D_A + D_B)B, \quad D_2(A) = (D_A - \rho(D_A + D_B))A, \quad (6)$$

$$D_3(A) = D_B(1 - A) + \rho(D_A + D_B)A, \quad D_4(B) = (D_B - \rho(D_A + D_B))B. \quad (7)$$

Here,

$$D_A = \lim_{\Delta \rightarrow 0} \frac{r_A \Delta^2}{4}, \quad \text{and} \quad D_B = \lim_{\Delta \rightarrow 0} \frac{r_B \Delta^2}{4},$$

178 are the macroscopic diffusion coefficients corresponding to species A and B, respectively.

179 Setting $\rho = 0$ in equations (4) and (5) leads to,

$$\frac{\partial A}{\partial t} = D_A \nabla \cdot [(1 - B)\nabla A + A\nabla B], \quad (8)$$

$$\frac{\partial B}{\partial t} = D_B \nabla \cdot [(1 - A)\nabla B + B\nabla A], \quad (9)$$

180 which are the macroscopic equations for the two-species volume-exclusion process [31].

181 In Figure 4, we compare the column-averaged density of the ABM given by,

$$\bar{A}_i = \frac{1}{L_y} \sum_{j=1}^{L_y} A_{ij}, \quad \bar{B}_i = \frac{1}{L_y} \sum_{j=1}^{L_y} B_{ij},$$

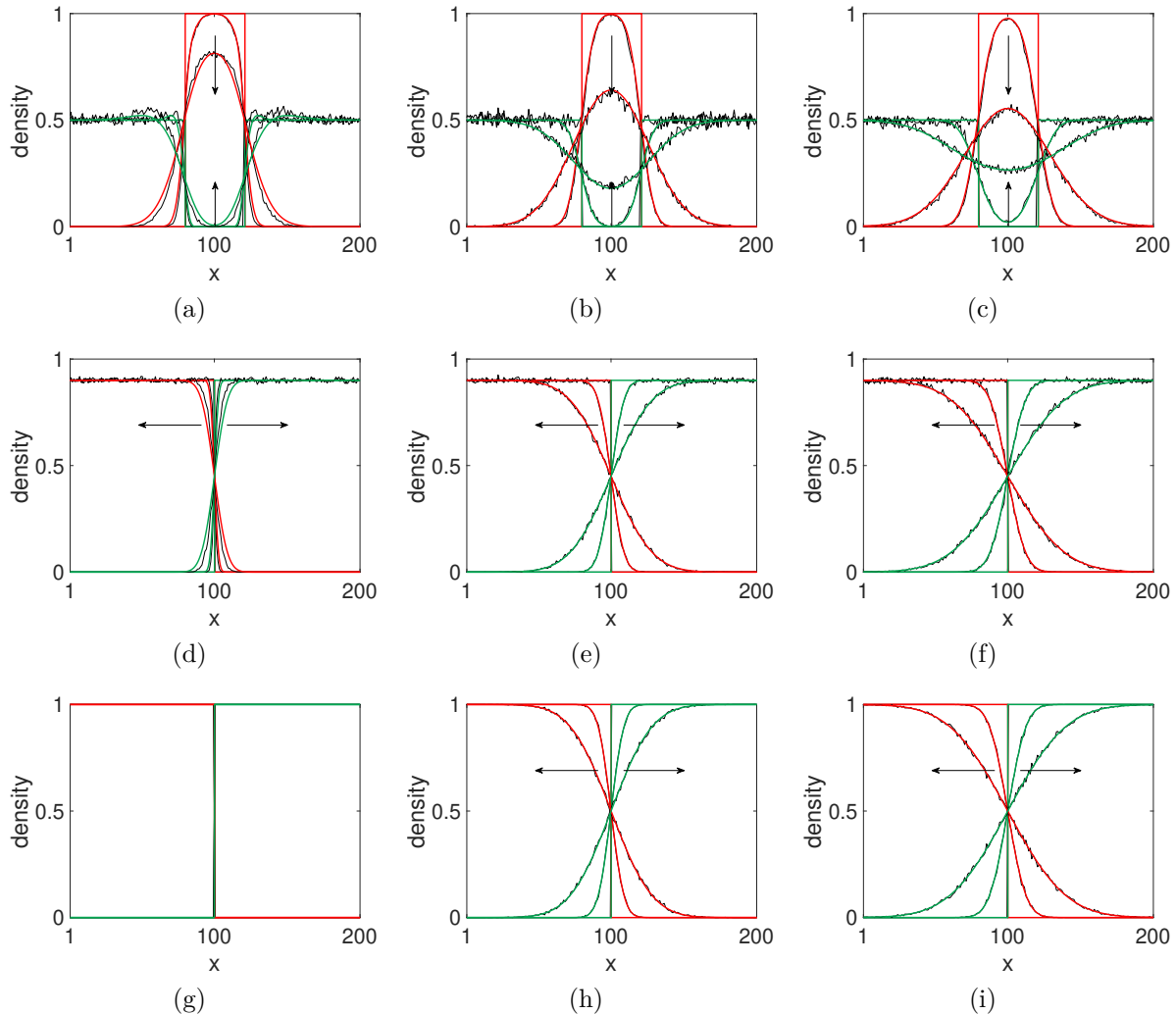


Figure 4. A comparison between the numerical solution of the PDEs (4) and (5) and the averaged behaviour of the ABM for the migration process with swapping with different swapping probabilities: $\rho = 0$ for [(a), (d), (g)]; $\rho = 0.5$ for [(b), (e), (h)] and $\rho = 1$ for [(c), (f), (i)]. The initial conditions for [(a)-(c)] are the same as Figure 3. For [(d)-(f)] we initialised the region $1 \leq x \leq 100$ with agents of type A at a density of 0.9 and the remaining sites with agents of type B also at a density of 0.9. In [(g)-(i)] the lattice was initialised such that all the sites in the region $1 \leq x \leq 100$ are occupied by agents of type A and all the sites in the region $101 \leq x \leq 200$ are occupied by agents of type B. We present solutions at $t = 0$, $t = 100$ and $t = 1000$ in all cases. The averaged column densities \bar{A}_i and \bar{B}_i are shown in black and the approximate PDE solution trajectories are shown in colour (red for agents of type A and green for agents of type B). The black arrows show the direction of increasing time.

182 to the numerical solution of the one-dimensional analogue of Equations (4) and (5) with
 183 reflective boundary conditions by averaging the PDEs over the y direction[32]. In the
 184 $\rho = 0$ case (Figure 4, first column), the PDE solutions and the ABM do not agree well as
 185 evidenced by the disparity between the two profiles. This discrepancy can also be seen in
 186 Simpson *et al.* [31] where the authors devise an ABM for multi-species exclusion processes

187 ($\rho = 0$ case in our model) and compare their ABM with the corresponding continuum
 188 model. We remark that one reason for the discrepancy is that in crowded environments
 189 where the movement of agents is frequently inhibited by other agents, lattice occupancies
 190 cannot be considered independent of each other and spatial correlation are not dissipated
 191 efficiently [33–35]. Independence of lattice sites is a key assumption that is typically made
 192 when deriving the continuum models such as the one above [19, 21, 24, 25, 30, 31, 36].
 193 For non-zero swapping probabilities, the agreement is significantly improved between the
 194 deterministic and stochastic model (Figure 4, second and third columns). Swapping helps
 195 to break down the spatial correlations improving the agreement between the PLM and
 196 the ABM. We also see that in the zero swapping case, crowding of the green agents behind
 197 the red agents leads to profiles for which the maximum density at non-zero time (shown
 198 $t = 100$ and $t = 1000$ in Figure 4(a)) is higher than the initial maximum density ($t = 0$).
 199 The reason for this is that for a multispecies migration process with cross-diffusion terms
 200 there is no maximum principle for the individual species [31, 37, 38]. We note that
 201 the enhanced diffusion which swapping engenders eliminates this effect. However, this
 202 does not necessarily mean that a maximum principle now holds for the systems under
 203 consideration. Investigating this further is beyond the scope of this article.

204 III. INDIVIDUAL-LEVEL ANALYSIS

205 In this section, we analyse the movement of agents at the individual level in the single-
 206 species and two-species case to assess how swapping affects the movement of agents.

207 A. Single-species individual-level analysis

208 The single-species swapping discrete and continuum models are given in Appendix A.
 209 Here, we present the individual-level analysis for the single-species model. Our aim is to
 210 quantify the movement of individually tagged agents by their individual-level diffusion
 211 coefficient. For the analysis that we present next, we neglect any long range temporal cor-
 212 relations in the agents' movement. We first derive the individual-level time-uncorrelated
 213 diffusion coefficient analytically and then we compare it with its ABM approximation.
 214 Let $P_{ij}(t) = P_{ij}$ denote the probability that a focal agent is at position (i, j) at time t .
 215 Defining δt as an infinitesimally small change in time, we can write down the probability
 216 of the agent being at position (i, j) at time $t + \delta t$ as,

$$\begin{aligned}
P_{ij}(t + \delta t) = & \frac{r}{4}P_{i-1,j}[(1 - c) + 2c\rho]\delta t + \frac{r}{4}P_{i+1,j}[(1 - c) + 2c\rho]\delta t + \frac{r}{4}P_{i,j-1}[(1 - c) + 2c\rho]\delta t \\
& + \frac{r}{4}P_{i,j+1}[(1 - c) + 2c\rho]\delta t + P_{ij}[(1 - r\delta t) + rc(1 - \rho)\delta t - rc\rho\delta t] + o(\delta t). \quad (10)
\end{aligned}$$

217 A lattice site is occupied with probability c and empty with probability $1 - c$. The
218 first four terms in Equation (10) are obtained by considering an agent at each of the four
219 lattice sites in the neighbourhood the site (i, j) that has attempted to move into site (i, j)
220 with probability $r/4 \delta t$. If the site (i, j) is vacant, the agent jumps from the neighbouring
221 site to site (i, j) . Otherwise, the agents at the neighbouring site and site (i, j) swap their
222 positions with probability ρ . The 2 in $2\rho c$ corresponds to the two ways in which the
223 position of the agent at site (i, j) can change due to a swap: either the agent at site
224 (i, j) initiates and successfully completes the swap with the agent at the neighbouring
225 site or the agent at the neighbouring site initiates and successfully swaps with the agent
226 at site (i, j) . The last term in Equation (10) gives the probability that the agent already
227 occupying position (i, j) does not attempt to move during the time interval $[t, t + \delta t]$
228 with probability $(1 - r\delta t)$ and the probability that the agent attempts to swap with an
229 agent at an occupied neighbouring site but fails to complete the swap with probability
230 $rc(1 - \rho)\delta t$ and lastly the probability that a neighbouring agent successfully swaps with
231 the agent at site (i, j) with probability $rc\rho\delta t$.

232 By rearranging, dividing both sides of Equation (10) by δt and taking the limit as $\delta t \rightarrow 0$
233 leads to the system of ODEs given by,

$$\frac{dP_{ij}}{dt} = \frac{r}{4}[(1 - c) + 2c\rho](P_{i-1,j} + P_{i+1,j} + P_{i,j-1} + P_{i,j+1}) - rP_{i,j}[(1 - c) + 2c\rho], \quad (11)$$

234 which describe the time evolution of the probability of finding an agent at position (i, j)
235 at time t . From Equation (11) it can be shown that the expected net displacement of
236 an agent is zero. This gives us no information about the statistical fluctuations in the
237 movement of an agent and therefore we use the variance to quantify the net displacement
238 [30, 39]. The equations describing the time-evolution of the variances $\langle i(t)^2 \rangle$ and $\langle j(t)^2 \rangle$
239 are given by,

$$\frac{d\langle i^2 \rangle}{dt} = \frac{d}{dt} \left(\sum_{i=1}^{L_x} i^2 P_{ij} \right) = \frac{r}{2} [(1-c) + 2c\rho], \quad (12)$$

and,

$$\frac{d\langle j^2 \rangle}{dt} = \frac{d}{dt} \left(\sum_{j=1}^{L_y} j^2 P_{ij} \right) = \frac{r}{2} [(1-c) + 2c\rho]. \quad (13)$$

240 Under the initial condition that at time $t = 0$, $\langle i^2 \rangle = \langle j^2 \rangle = 0$, Equations (12) and (13)
 241 solve to give,

$$\langle i(t)^2 \rangle = \langle j(t)^2 \rangle = \frac{r}{2} [(1-c) + 2c\rho]t. \quad (14)$$

242 The time-uncorrelated individual-level diffusion coefficient can be retrieved as,

$$D^*(c, \rho) = \frac{r}{4} [(1-c) + 2c\rho]. \quad (15)$$

243 As briefly discussed above, the master equation neglects temporal correlations in an
 244 agent's position. Consequently, the diffusion coefficient derived from the master equation
 245 will necessarily be inaccurate and fail to represent the true dynamics of the agents in
 246 the system. As a result, we refer to the expression given in Equation (15) as the time-
 247 uncorrelated individual-level diffusion coefficient.

248 To approximate the D^* using the ABM, we initialise a 150 by 150 lattice and randomly
 249 seed it with agents at different background densities $\mathbf{c} = [0, 0.25, 0.5, 0.75, 1]$, where $c = 0$
 250 corresponds to one agent with no agents in the background to interact with and $c =$
 251 1 corresponds to a fully populated lattice. We let the positions of the agents evolve
 252 according to the single-species version of the ABM described in Section II A using the
 253 movement rate $r = 1$ for a range of swapping probabilities $\boldsymbol{\rho} = [0, 0.25, 0.5, 0.75, 1]$. The
 254 positions (X, Y) of agents whose initial position is in the region defined by the central
 255 square $[51, 100] \times [51, 100]$ are recorded over a regular time grid $t_{\text{rec}} = [0, \Delta t, 2\Delta t, \dots, T\Delta t]$
 256 to create a track where Δt is the recording step and T is the track length. Each track
 257 traces the path of an agent over time. We impose periodic boundary conditions on the
 258 domain. The simulation ends at $T_{\text{final}} = 1000$ or when a tracked agent hits a boundary,
 259 whichever happens first.

260 We analyse individual movement of agents using the sum of squared displacement (SSD)
 261 $[30, 36]$,

$$S_t^{(x)} = \sum_{j=1}^{t'} (X(t + j\Delta t) - X(t + (j-1)\Delta t))^2, \quad (16)$$

262 and,

$$S_t^{(y)} = \sum_{j=1}^{t'} (Y(t + j\Delta t) - Y(t + (j-1)\Delta t))^2, \quad t' = 1, \dots, T \quad (17)$$

263 By only taking the difference between successive positions, the SSD neglects temporal
 264 correlations in an agent's position which is consistent with our master equation in Equa-
 265 tion (10). For each value of c and ρ we average the SSD over an ensemble of tracks and
 266 fit a linear model of the form $\hat{S}_t^x = a_x t$ and $\hat{S}_t^y = a_y t$ in each orthogonal direction x and y ,
 267 respectively. The ABM approximation of the time-uncorrelated individual-level diffusion
 268 coefficient can be extracted from the gradient of these linear equations,

$$\hat{D}^*(c, \rho) = \frac{a_x + a_y}{2d}, \quad (18)$$

269 where $d = 2$ is the dimension of the lattice. We put a 'hat' symbol over D^*
 270 to differentiate it from the exact time-uncorrelated diffusion coefficient D^* in Equation
 271 (15).

272 In Figure 5, we compare the ABM approximation \hat{D}^* (shown in (a)) with the derived
 273 expression D^* (shown in (b)) as a pair of heat maps for the range of values of c and
 274 ρ defined earlier. We also show a line plot that nicely depicts the linear relationship
 275 between D^* and c for different value of ρ (shown in (c)). We can see that there is
 276 excellent agreement between the analytical expression and the simulated results. We
 277 note that in the case of zero background density ($c = 0$) D^* is always 0.25 regardless of
 278 the swapping probability since for a single agent with no other agents to interact with,
 279 the movement of the agent can neither be inhibited by volume exclusion nor enhanced by
 280 swapping. We also note that for swapping probability $\rho = 0.5$, the value of D^* is always
 281 0.25 irrespective of the density since half the number of times a focal agent attempts
 282 to move into an occupied site the moves will be rejected. If this were the only impact
 283 of swapping, we would expect D^* of the focal agent to be reduced. However, just as
 284 often as a focal agent attempts to move into an occupied neighbour's position, an agent
 285 occupying a neighbouring site tries to move into the focal agent's position – achieving
 286 this successfully with probability $\rho = 0.5$. Assuming a well-mixed scenario, this exactly
 287 compensates for the number of aborted moves the focal agent makes, meaning movement

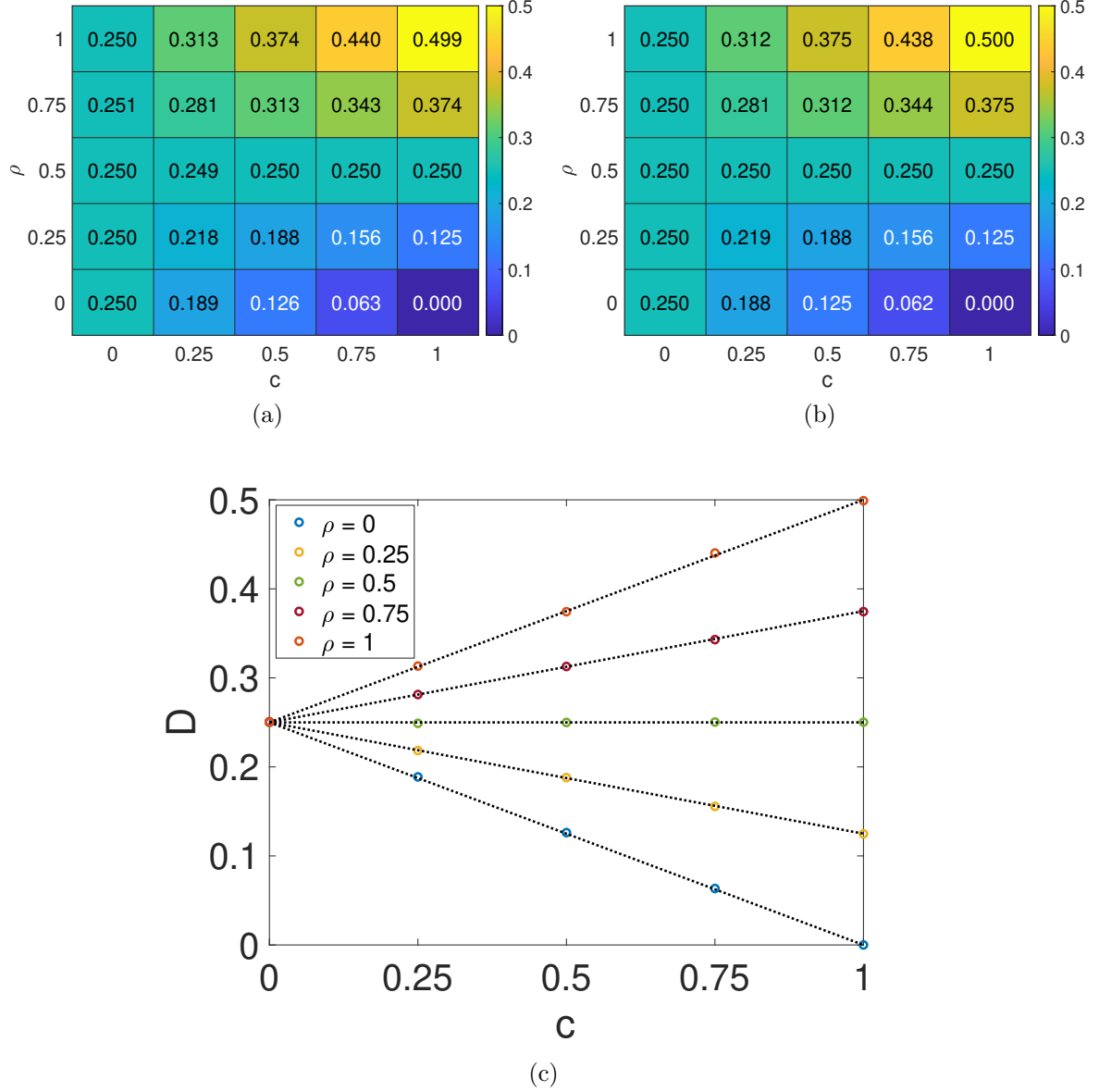


Figure 5. Heat map showing the relationship between the time-uncorrelated individual-level diffusion coefficient, D^* , the domain density, c , and the swapping probability, ρ . For (a), we initialised a 150 by 150 periodic domain with density $c \in \mathbf{c} = [0, 0.25, 0.5, 0.75, 1]$ and for the values of swapping probability $\rho = [0, 0.25, 0.5, 0.75, 1]$ we let the positions of the agents evolve according to the single-species ABM in Section A with $r = 1$. We tracked all agents in the region defined by the square $[51, 100] \times [51, 100]$. Theoretical heatmap in (b) was obtained using Equation (15). In (c) we show the linear relationship between D^* , the domain density, c and the swapping probability, ρ . The circles represent ABM approximations of \hat{D}^* whereas the dotted black lines represent the exact value D^* given in Equation (15).

288 is as if the focal agent were on an unoccupied domain irrespective of density. For $c = 1$
 289 and $\rho = 1$ we note that $D^* = 0.5$ (i.e. twice as large compared to an agent moving on
 290 a domain with zero background density) as every attempted move by the focal agent is
 291 executed successfully and the focal agent is also moved equally often by neighbouring

292 agents swapping into its position.

293 B. Two-species individual-level analysis

294 In this section, we perform the individual-level analysis for a two-species system as set
295 out for the single-species case in Section III A.

296 Let $P_{ij}^A(t) = P_{ij}^A$ be the probability that a focal agent of type-A occupies the position
297 (i, j) and let $P_{ij}^B(t) = P_{ij}^B$ be the equivalent for a type-B focal agent. Recalling δt as a
298 small change in time we can write down the master equations for species A and B at time
299 $t + \delta t$,

$$\begin{aligned} \frac{dP_{ij}^A}{dt} = & \left(\frac{r_A}{4}(1-c) + \frac{r_A}{2}c_A\rho + \frac{(r_A+r_B)}{4}c_B\rho \right) (P_{i-1,j}^A + P_{i+1,j}^A + P_{i,j-1}^A + P_{i,j+1}^A) \\ & + (-r_A(1-c) - 2r_Ac_A\rho - (r_A+r_B)c_B\rho)P_{ij}^A, \end{aligned} \quad (19)$$

$$\begin{aligned} \frac{dP_{ij}^B}{dt} = & \left(\frac{r_B}{4}(1-c) + \frac{r_B}{2}c_B\rho + \frac{(r_A+r_B)}{4}c_A\rho \right) (P_{i-1,j}^B + P_{i+1,j}^B + P_{i,j-1}^B + P_{i,j+1}^B) \\ & + (-r_B(1-c) - 2r_Bc_B\rho - (r_A+r_B)c_A\rho)P_{ij}^B. \end{aligned} \quad (20)$$

300 A full derivation of Equations (19) and (20) can be found in Appendix B accompanying
301 this article.

302 The individual-level time-uncorrelated diffusion coefficients D_A^* and D_B^* for the species A
303 and B, respectively, are given by,

$$D_A^* = \frac{1}{4}(r_A(1-c) + (r_{AC} + r_{ACA} + r_{BCB})\rho), \quad (21)$$

$$D_B^* = \frac{1}{4}(r_B(1-c) + (r_{BC} + r_{BCA} + r_{BCB})\rho). \quad (22)$$

304 In order to investigate the effect of swapping on a two-species system and to verify our
305 theoretical results, we simulate the two-species model, track a set of tagged agents and
306 analyse their movement using the SSD as done in Section III A.

307 In Figure 6 we show the time-uncorrelated diffusion coefficient of species A and species
308 B plotted against background density $c = [0, 0.25, 0.50, 0.75, 1]$ for different values of
309 swapping probability $\rho = [0, 0.25, 0.50, 0.75, 1]$ for the specific case in which $r_A = 1$ and
310 $r_B = 2$. In this figure, both species are present in equal proportions on the domain. We

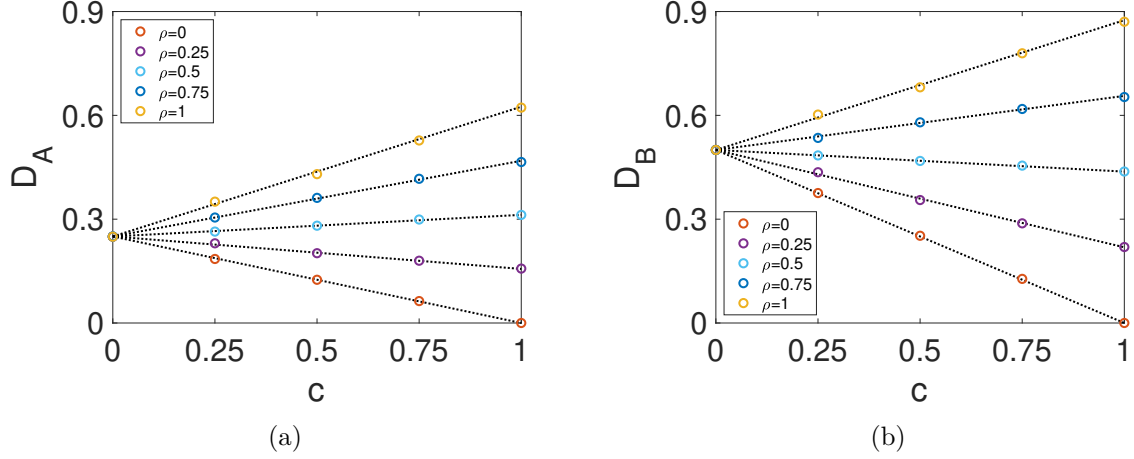


Figure 6. Time-uncorrelated diffusion coefficient of tagged type-A (a) and type-B agents (b) plotted against background densities $c = [0, 0.25, 0.5, 0.75, 1]$ for different swapping probabilities $\rho = [0, 0.25, 0.5, 0.75, 1]$ with fixed movement rate $r_A = 1$ and $r_B = 2$. Both species are present in equal proportions (i.e. $c_A = c_B = 0.5c$). The circles represent the ABM approximations \hat{D}_A^* and \hat{D}_B^* and the dotted lines represent the theoretical values D_A^* and D_B^* from Equations (21) and (22).

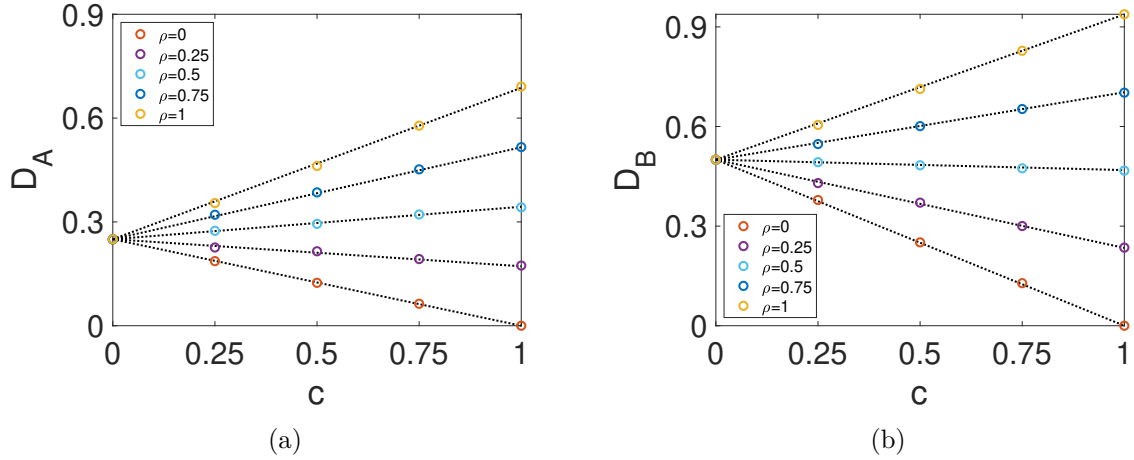


Figure 7. Time-uncorrelated diffusion coefficient of tagged type-A (a) and type-B agents (b) plotted against background densities $c = [0, 0.25, 0.5, 0.75, 1]$ for different swapping probabilities $\rho = [0, 0.25, 0.5, 0.75, 1]$ with fixed movement rate $r_A = 1$ and $r_B = 2$. Type-A agents are present at a density $c_A = 0.25c$ and type-B agents are at $c_B = 0.75c$. The Left panel (a) shows D_A^* and the right-hand panel (b) shows D_B^* .

311 see good agreement between the ABM approximations and the theoretical values. We
 312 also see that, as expected and as noted in the single-species system, swapping speeds up
 313 the movement of the agents in the multi-species setting compared to the pure volume-
 314 excluded scenario ($\rho = 0$). Furthermore, since species B has a higher movement rate than

315 species A, species B diffuses faster than species A, apart from the $c = 1$ and $\rho = 0$ trivial
316 case in which D_A^* and D_B^* are both 0 since the agents have nowhere to go.

317 In Figure 7 we present a similar comparison as in Figure 6 but this time the two species are
318 not present in equal proportions. In this particular case, we consider a scenario in which
319 type-A agents make up 25% of the total population and type-B agents 75%. Again, we
320 see good agreement between the ABM and theoretical values and that swapping speeds
321 up the movement of agents.

322 Comparing Figure 6 to Figure 7 we see that for the $\rho = 0$ case D_A^* and D_B^* in Figure 6
323 are equal to their respective values in Figure 7 and hence unaffected by changes in the
324 densities c_A and c_B . Analytically, this can be observed by setting $\rho = 0$ in Equations
325 (21) and (22). In this case, D_A^* and D_B^* depend on the overall background density and
326 not the species proportions. For $\rho > 0$, in the instance where $r_B > r_A$ both species
327 show faster movement in Figure 7 than in Figure 6. Again, this can be checked by
328 referring to the expressions for D_A^* and D_B^* . Changing the proportions of the species
329 (while keeping r_A , r_B and c constant) the term $(r_A c_A + r_B c_B)\rho$ is sensitive to changes
330 in the proportions in both D_A^* and D_B^* . When $r_B > r_A$ and we increase the proportion
331 of the faster moving species, c_B , this results in more swapping events which enhance the
332 movement of both species. On the other hand, if we were to increase the proportion of
333 species A (while simultaneously decreasing the proportion of species B to keep c constant)
334 we would observe reduced movement for both species since an increase in the proportion
335 of the slower moving species (and a decrease in the proportion of faster moving species)
336 would result in fewer swaps.

337 IV. ILLUSTRATIVE EXAMPLES

338 In this section, we show examples of the situations in which swapping has important
339 applications. In Section IV A, we build the swapping mechanism into a cell migration
340 model with proliferation and in Section IV B, we show how the swapping mechanism in
341 conjunction with cell-cell adhesion can facilitate spontaneous pattern formation in densely
342 crowded environments.

343 A. Swapping model with cell proliferation

344 We look at the role of swapping in cell migration with proliferation. For this example,
 345 we concern ourselves with the two-species cell migration model. The movement kinetics
 346 of the agents are the same as the swapping model described in Section II A but in addi-
 347 tion to migrating, agents can attempt to proliferate, placing a daughter at a randomly
 348 chosen neighbouring site if the site is empty, otherwise the division event is aborted. The
 349 proliferation rates per unit time for the two species A and B are denoted by r_p^A and r_p^B ,
 350 respectively.

351 We initialise the domain with $L_x = 100$ sites in the horizontal direction and $L_y = 20$ sites
 352 in the vertical direction. We fill all the sites in the range $41 \leq x \leq 60$ with agents of type
 353 A and all the remaining sites with type-B agents at a density of 0.5. The movement rates
 354 of agents are set to $r_A = r_B = 1$ and the proliferation rates as $r_p^A = 0.01$ and $r_p^B = 0$, i.e.
 355 only the agents of type A divide and the number of type-B agents are held constant. We
 356 let the system evolve according to the specified ABM.

357 In Figure 8 we provide snapshots of the evolving lattice occupancy for $\rho = 0, 0.5, 1$
 358 (columns 1, 2 and 3, respectively) at $t = 0, 500, 1000$ (rows 1, 2 and 3, respectively).
 359 We see, at the same time points, that cells are unsurprisingly more well-mixed in the
 360 case of non-zero swapping (second and third columns in Figure 8) compared to the zero-
 361 swapping situation (first column in Figure 8). We also see faster colonisation of the
 362 domain overall in the non-zero swapping cases than without swapping. This is because
 363 swapping allows the proliferating red agents to disperse more quickly into less dense
 364 regions, which in turn increases the probability of a successful division events for these
 365 agents. Without swapping, it takes longer for proliferative red agents to find the space
 366 to proliferate into. This trend of decreasing colonisation time with increasing swapping
 367 probability is reinforced in Figure 9 where we see that the time to reach the domain's
 368 carrying capacity is a decreasing function of the swapping probability, ρ .

369 The mean-field PDEs describing the approximate population-level dynamics of the agent
 370 are given by,

$$\frac{\partial A}{\partial t} = \nabla \cdot [D_1(B)\nabla A + D_2(A)\nabla B] + r_p^A A(1 - (A + B)), \quad (23)$$

$$\frac{\partial B}{\partial t} = \nabla \cdot [D_3(A)\nabla B + D_4(B)\nabla A], \quad (24)$$

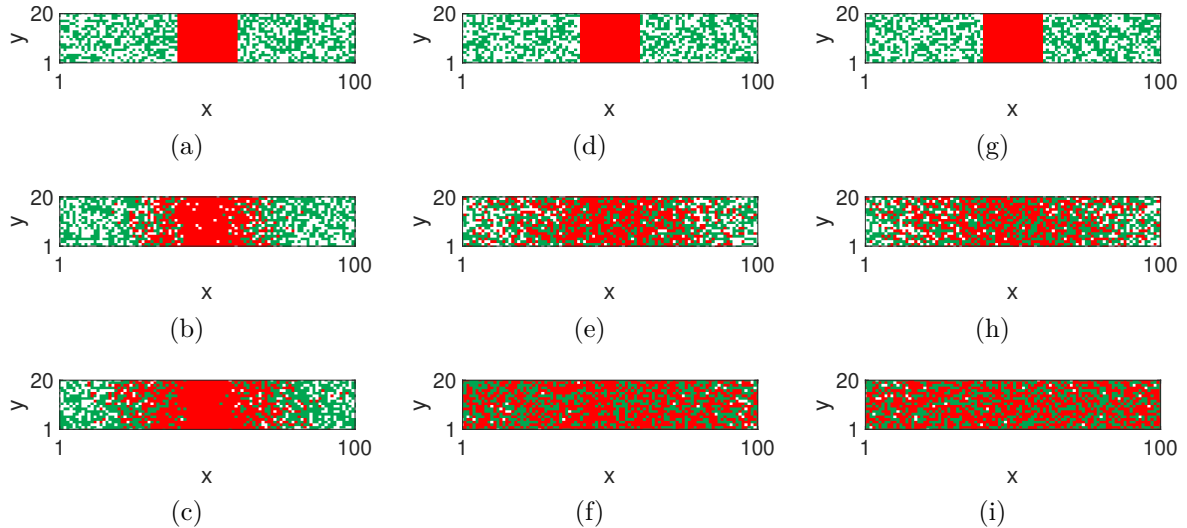


Figure 8. Snapshots of the lattice occupancy with swapping probability $\rho = 0$ in [(a)-(c)], $\rho = 0.5$ in [(d)-(f)] and $\rho = 1$ in [(g)-(i)] at $t = 0, 500, 1000$ for the cell migration process with swapping and proliferation. We initialise the domain as a 20 by 100 lattice where all the sites in the horizontal range 41 to 60 are occupied by the agents of type A (red) and the remaining sites are inhabited by agents of type B at a density of 0.5. Both species diffuse at equal rates $r_A = r_B = 1$. Rates of proliferation are given by $r_p^A = 0.01$ and $r_p^B = 0$.

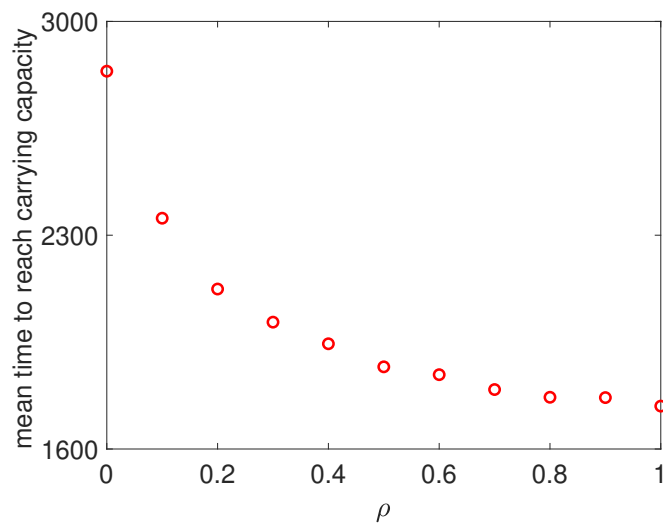


Figure 9. Time to reach the carrying capacity. The red circles show the mean time for the number of agents to reach the carrying capacity of the domain for a range of swapping probabilities (averaged over 100 repeats).

371 where D_1, D_2, D_3 and D_4 are as defined previously. Notice that proliferation of type-A
372 agents gives rise to an additive source term in Equation (23). The derivation of this
373 source is standard and can be found in Plank and Simpson [40], Simpson *et al.* [41], for
374 example.

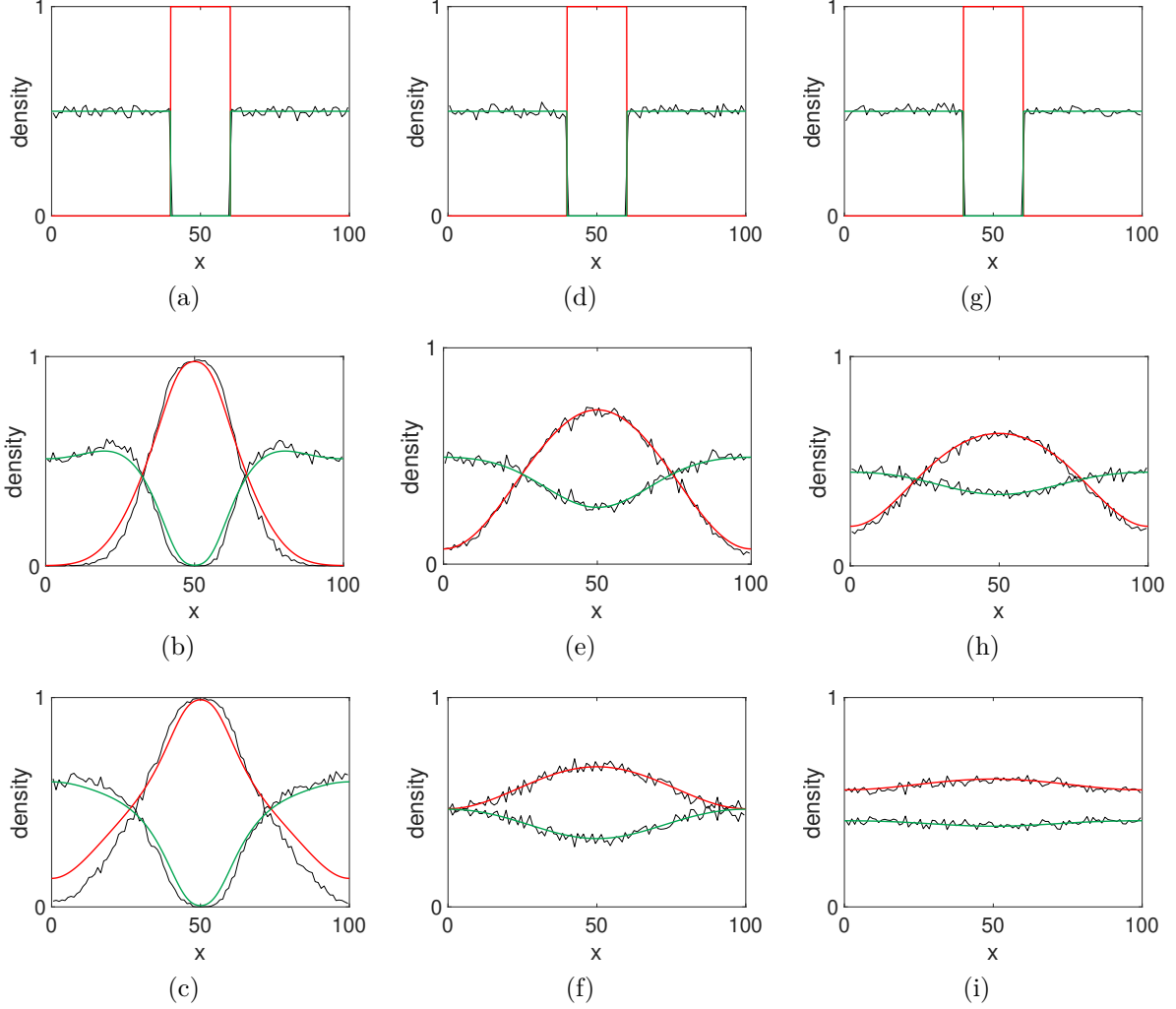


Figure 10. Density profiles for cell migration process with swapping and proliferation. All parameters and initial conditions are the same as Figure 8. Here, we present the column densities at $t = 0, 500, 1000$ for $\rho = 0$ in [(a)-(c)], $\rho = 0.5$ in [(d)-(f)] and $\rho = 1$ in [(g)-(i)] averaged over 100 repeats of the ABM described above. We also plot the corresponding mean-field PDE solutions with $r_p^A = 0.01$ in red for species A and in green for species B .

375 In Figure 10 we compare the average column density of the ABM with the numerical
376 solution of the one-dimensional analogue of the mean-field PDEs obtained by averaging
377 over the y direction with $r_p^A = 0.01$. We chose this value for the proliferation rate as
378 we wanted to keep the ratio $r_A/r_p^A \ll 1$ [41, 42]. There are two reasons for this: firstly,
379 a modelling choice to prevent agents clustering into proliferation-induced patches and
380 secondly, it is biological realism that given the parameters of the model we could ex-
381 pect real biological cells will attempt proliferation events less frequently than movement
382 events. We see good agreement between the two profiles for non-zero swapping probabil-
383 ity. However, when the swapping probability is set to 0, we see discrepancies arising that

384 are amplified as time increases. Recall that the disparity between the PDE and ABM
385 profiles can be also observed in Figure 3 due to spatial correlations that are not accoun-
386 ted for by the mean-field PDEs. The addition of proliferation into the model increases
387 the spatial correlations between site occupancies, leading to greater disparity. One way
388 to partially rectify this problem is by modelling the higher order moments in the PDE
389 description [33–35] however for the purposes of this work, we simply note that allowing
390 swapping breaks up the correlations more effectively than in its absence leading to better
391 agreement between the ABM and the population-level densities.

392 **B. Swapping model with cell-cell adhesion**

393 Another interesting application of swapping is the formation of patterns in densely
394 crowded environments. In this section, we use a cell-cell adhesion model with swap-
395 ping to investigate how biologically plausible patterns can form starting from a randomly
396 seeded domain. Our model is based on the similar cell-cell adhesion model studied pre-
397 viously by [20, 21, 23] who consider adhesion between identical agents. Here, we extend
398 the model to incorporate two types of agents with swapping to facilitate the movement
399 events.

400 For the purpose of this paper, we assume adhesion between two species, A and B, on a fully
401 populated domain. For a simple exclusion-based ABM the agents on the fully populated
402 domain would not successfully move at all. This is since the exclusion principle forbids
403 cells from occupying the already occupied lattice sites. In our model, the movement of the
404 agents and the formation of patterns will be facilitated by the swapping mechanism. In
405 an on-lattice adhesion model, agents can adhere to other agents in their neighbourhood,
406 making them less likely to successfully complete the movement event. As well as the
407 number of agents in the neighbourhood, the strength of adhesion determines how likely
408 an agent is to successfully move. In a simple model with species A and B, $0 \leq p \leq 1$
409 characterises the strength of adhesion between two type-A agents and $0 \leq q \leq 1$ the
410 strength of adhesion between two type-B agents.

411 We assume cell movement in a densely crowded domain where the underlying lattice is
412 fully populated with type-A and B agents, their positions chosen uniformly at random.
413 When an agent is chosen to move into an occupied neighbouring site, we check the feasibil-
414 ity of swapping by sampling a random number u from the standard uniform distribution
415 and comparing it with the swapping probability, ρ . A swapping move breaks existing

416 interactions between the two swapping agents and their neighbours and makes new con-
417 nections following a successful swap. Since the movement of the focal agent and the target
418 agent depends on their respective neighbours, the success of a swapping move depends on
419 whether the respective neighbouring sites are occupied by type-A or type-B agents. Sup-
420 pose the focal agent is at site (i, j) and let $Z_{ij} = \{(i-1, j), (i+1, j), (i, j-1), (i, j+1)\}$ be
421 the set containing the positions its neighbouring sites on the two-dimensional lattice. If
422 the focal agent is a type-A agent then the probability of it breaking existing connections
423 with its neighbours is given by,

$$p_{\text{break}}^{\text{agent}} = (1 - p)^{\sum_{z \in Z} A_z}. \quad (25)$$

424 However, if the focal agent is a type-B agent then,

$$p_{\text{break}}^{\text{agent}} = (1 - q)^{\sum_{z \in Z} B_z}. \quad (26)$$

425 Here, A_z are binary taking a value of unity if the site with position $z \in Z_{ij}$ is occupied
426 by a type-A agent or 0 otherwise. Therefore, $\sum_{z \in Z} A_z$ is the sum of occupancies of
427 the sites in Z_{ij} that are occupied by type-A agents. Likewise, $\sum_{z \in Z} B_z$ is the sum of
428 occupancies of the sites in Z_{ij} that are occupied by type-B agents. Similarly, for a type-A
429 agent occupying the target site the probability that it breaks existing connections with
430 its neighbours is given by,

$$p_{\text{break}}^{\text{targ}} = (1 - p)^{\sum_{y \in Y_z} A_y}. \quad (27)$$

431 However, if the agent occupying the target site is a type-B agent then,

$$p_{\text{break}}^{\text{targ}} = (1 - q)^{\sum_{y \in Y_z} B_y}. \quad (28)$$

432 Here, the set Y_z contains the positions of sites in the neighbourhood of the target site, z .
433 Therefore, $\sum_{y \in Y_z} A_y$ is the sum of occupancies of all the sites in Y_z that are occupied by
434 a type-A agent and $\sum_{y \in Y_z} B_y$ denotes the sum of occupancies of all the sites in Y_z that
435 are occupied by a type-B agent.

436 The probability of a successful swap given a movement event has been attempted is
437 therefore a product of the swapping probability, the probability of the focal agent at site
438 (i, j) breaking links with its neighbours in order to move out and the probability of the
439 agent at the target site breaking links with its neighbours in order to move in, i.e.,

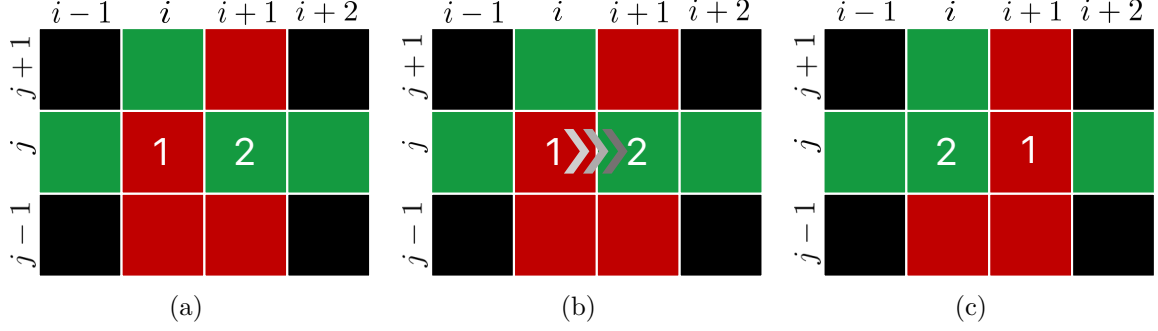


Figure 11. A schematic illustrating swapping in the cell-cell adhesion model. A red site represents a type-A agent and a green site represent a type-B agent. Agent 1 at site (i, j) attempts to swap with agent 2 at site $(i + 1, j)$. We have coloured in black the neighbouring sites that are unimportant in this context (i.e. occupancy of these sites does not affect the probability of a successful swap happening between agent 1 and agent 2). The initial configuration of the lattice is shown in (a). Agent 1 attempts to swap with agent 2 (b) and the state of the lattice after the successful swap is shown in (c).

$$p_{\text{swap}} = \rho p_{\text{break}}^{\text{agent}} p_{\text{break}}^{\text{targ}}. \quad (29)$$

440 Here we are considering that the link between the two swapping agents has to be broken in
441 order for the swap to take place. As an example, consider the case in which the focal agent
442 at site (i, j) attempts to jump into the neighbouring site $(i + 1, j)$ (Figure 11). The agent
443 occupying the focal site is a type-A agent, shown in red and labelled as 1 and the target
444 site $(i + 1, j)$ is occupied by a type-B agent, shown in green and labelled as 2. Since the
445 focal agent is red, it is adhesive to other red agents in its Von Neumann neighbourhood.
446 There is only one site (with position $(i, j - 1)$) neighbouring agent 1 that is occupied by
447 a red agent and therefore, $\sum_{z \in Z_{ij}} A_z = 1$. Since agent 2 (target site) is a green agent, it
448 is adhesive to the other green agents in its Von Neumann neighbourhood. There is only
449 one green agent in the neighbourhood (position $(i + 2, j)$) and hence, $\sum_{y \in Y_{i+1, j}} B_y = 1$.
450 For an arbitrary ρ , p and q , the probability of swap in this situation given a neighbouring
451 site to move into has already been chosen can be written as $p_{\text{swap}} = \rho(1 - p)(1 - q)$.

452 In Figure 12, we present some results that demonstrate the importance and impact of
453 swapping in a model of adhesion-mediated pattern formation. For this, we consider a
454 square lattice with $L = 20$ sites in both the horizontal and vertical direction. As before,
455 one compartment can accommodate no more than a single agent at a time. We impose
456 periodic boundary conditions. We seed the lattice with type-A and type-B agents at a
457 density of 0.5 each, where the initial positions of the agents on the lattice are assigned

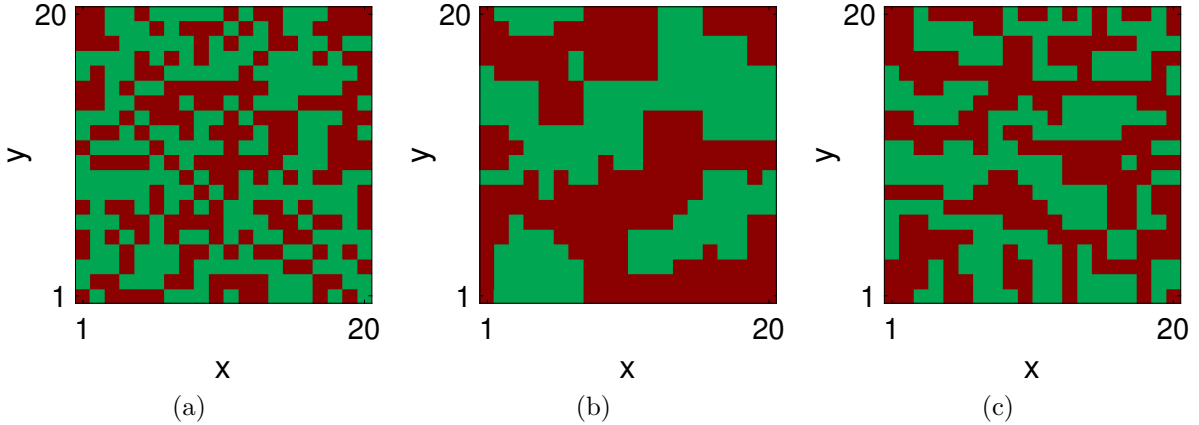


Figure 12. Pattern formation in a crowded environment. The domain is initialised as a square lattice with $L = 20$ sites in the horizontal and vertical directions, respectively. Initially, the domain is fully populated by type-A and type-B agents at a density of 0.5 each, where the positions of the agents are assigned uniformly at random (as shown in (a)). We present the state of the lattice at $t = 15,000$ for two different sets of adhesion strengths: $p = 0.90$ and $q = 0.70$ (b) and $p = q = 0.98$ (c). In both cases, we let $\rho = 1$ and the movement rates of the two species $r_A = r_B = 1$.

458 uniformly at random (Figure 12(a)). We let positions of the agents evolve according to
 459 the kinetics described above using the movement rate $r_A = r_B = 1$ and $\rho = 1$.
 460 Snapshots of the evolving lattice occupancy at $t = 15,000$ for two different sets of p and
 461 q values are shown in Figure 12(b) and (c). We can see self-organisation of agents into
 462 clusters of like type agents. The characteristic size of the aggregates is sensitive to the
 463 magnitude of the adhesion strengths. The clusters are bigger where self-adhesion within
 464 one species is stronger than within the other ($p = 0.9$ and $q = 0.7$) whereas we see
 465 more labyrinthine patterns when both the adhesion strengths are very strong and equal
 466 ($p = q = 0.98$). It is evident from these figures that in densely crowded domains, swapping
 467 plays a vital role in allowing agents to organise themselves to form patterns. Without
 468 swapping, no agent movement and hence no pattern formation would be possible.

469 V. DISCUSSION AND CONCLUSION

470 Cell movement is often modelled as a volume-exclusion process. However, in reality cell
 471 movement is not completely inhibited volume exclusion. In this paper, motivated by
 472 real-life examples, we developed an ABM that allows a pair of neighbouring agents to
 473 swap positions with each other. Our model maintains the important carrying capacity
 474 component of volume-exclusion models but allows the flexibility of movement observed

475 even amongst some densely packed configurations. We considered a two-species system
476 and allowed it to evolve according to the dynamics of our model. We found that swapping
477 enhances the movement of agents by allowing agents to mix more compared to the pure
478 volume-exclusion model. Comparing the ABM to the population-level model we found
479 excellent agreement between the two as long as the swapping probability was sufficiently
480 large.

481 To understand how swapping affects agent movement at an individual level, we analysed
482 simulated agent tracks to determine the time-uncorrelated individual-level diffusion coef-
483 ficient. We found that swapping enhanced the movement of agents in all cases compared
484 to the volume-exclusion model. Using the probability master equation, we were able to
485 analytically derive an expression for the diffusion coefficient that confirms the relationship
486 obtained via the simulated tracks.

487 In Section IV, we demonstrated the importance of swapping via a couple of examples. In
488 the first application, we considered a cell migration model with proliferation. We found
489 that swapping accelerates the proliferation process by allowing the agents to disperse
490 more and breaking spatial correlations. We found that the time to reach the domain's
491 carrying capacity varies inversely with the swapping probability. Deriving the PLM and
492 comparing it to the average density of the ABM, we showed that there is a good agreement
493 between the two profiles for sufficiently large swapping probability. For the $\rho = 0$ case, we
494 note discrepancies that are caused by the build up of spatial correlations between sites.

495 For the second example, by incorporating swapping into a cell-cell adhesion model, we
496 showed that agents can spontaneously rearrange themselves into clusters to form patterns
497 even on densely populated domains. We stress that without swapping these patterns
498 would not be realised. Biological patterns involving cell-cell adhesion have been observed
499 amply in experimental work. Honda *et al.* [43] reported chequered patterns in Japan-
500 ese quail oviduct epithelium which were later modelled mathematically by Glazier and
501 Graner [14] using a cellular Potts model. Armstrong [44] observed sorting of neural and
502 pigmented retinal epithelial cells in chick embryo where neural cells completely engulfed
503 the pigmented cells from an initially disordered arrangement. Engulfment was modelled
504 mathematically using a cellular Potts model [14, 15] and by Armstrong *et al.* [45] using
505 a continuous approach.

506 Models incorporating swapping may prove useful when analysing multi-species systems
507 in which cells need to migrate in a crowded epithelium or tissue parenchyma. These sys-

508 tems are often interrogated experimentally using the generation of chimaeras or mosaics,
509 analysing the resulting patterns generated by cell mixing [46–50]. For example the gen-
510 eration of corneal epithelial stripes in X-inactivation mosaic female mice hemizygous for
511 an X-linked copy of the LacZ gene, expressing the enzyme β -galactosidase [51]. Here, a
512 mixture of β -gal positive and β -gal negative limbal epithelial cells are specified as limbal
513 stem cells whose progeny then migrate centripetally to generate β -gal positive and β -gal
514 negative corneal epithelial stripes [52]. Experiments suggest that the degree of mixing of
515 the limbal progenitors will generate stripes of different width composed of multiple clones
516 of the same colour [51]. Swapping would be useful in understanding how cell mixing at
517 the limbus impacts the final stripe pattern generated.

518 In conclusion, motivating our study by real-life examples, we have developed a cell migra-
519 tion model that incorporates swapping as a viable movement process. As well as adding
520 biological realism, our model has the added benefit of better agreement between the
521 corresponding continuum description and the ABM compared to the classical volume-
522 exclusion process. We also saw that the ABM with swapping and cell-cell adhesion,
523 when applied to cells in densely crowded environments, leads to pattern formation. We
524 once again stress that the patterns would be unattainable under the traditional volume-
525 exclusion model. The results in this paper hint that swapping is an important and
526 overlooked mechanism in the context of modelling real biological scenarios and merits
527 further explorations in conjunction with experimental data.

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540 **Appendix A: Single-species PDE vs. ABM results**

541 Below we summarise the single-species swapping model and present results comparing
542 the discrete and continuum models.

543 The single-species model is a simplified version of the two-species model in the sense
544 that there is only species on the domain of interest at density c and hence only a single
545 movement rate. We summarise the the model here.

546 We let r be the rate of movement of an agent such that $r\delta t$ is the probability that the agent
547 attempts to move during a finite time interval of duration δt . The agent attempts to move
548 into one of its four neighbouring sites with equal probability. If the chosen neighbouring
549 site is empty, the focal agent successfully moves and its position is updated. If another
550 agent already occupies the site, the agent at site (i, j) attempts to swap positions with the
551 neighbouring agent with probability ρ . If the swap is successful, the two agents exchange
552 positions with each other. Otherwise, the move is aborted.

553 In Figure 13 we present snapshots of the lattice occupancy for the single-species model
554 at $t = 0, 100, 1000$ and for swapping probabilities $\rho = 0, 0.5, 1$ with $r = 1$ with reflective
555 boundary conditions. We see that swapping seems to have no effect on the dispersion of
556 the agents at a macroscopic scale as the density profiles are indistinguishable regardless
557 of the swapping probability. If two agents are identical to each other and unlabelled then
558 exchanging positions by swapping is equivalent to an aborted movement attempt in the
559 volume-exclusion process and produces no change in the state of the system.

560 In the next section, we derive the macroscopic PDE describing the evolution of the mean
561 lattice occupancy in the single-species model.

562 **1. Single-species continuum model**

563 Let $C_{ij}^k(t)$ be the occupancy of site (i, j) on the k th repeat at time t , where $C_{ij}^k(t) = 1$ if
564 the site (i, j) is occupied and $C_{ij}^k(t) = 0$ otherwise. The average occupancy of site (i, j)
565 at time t after K runs is then given by,

$$\langle C_{ij}(t) \rangle = \frac{1}{K} \sum_{k=1}^K C_{ij}^k(t). \quad (\text{A1})$$

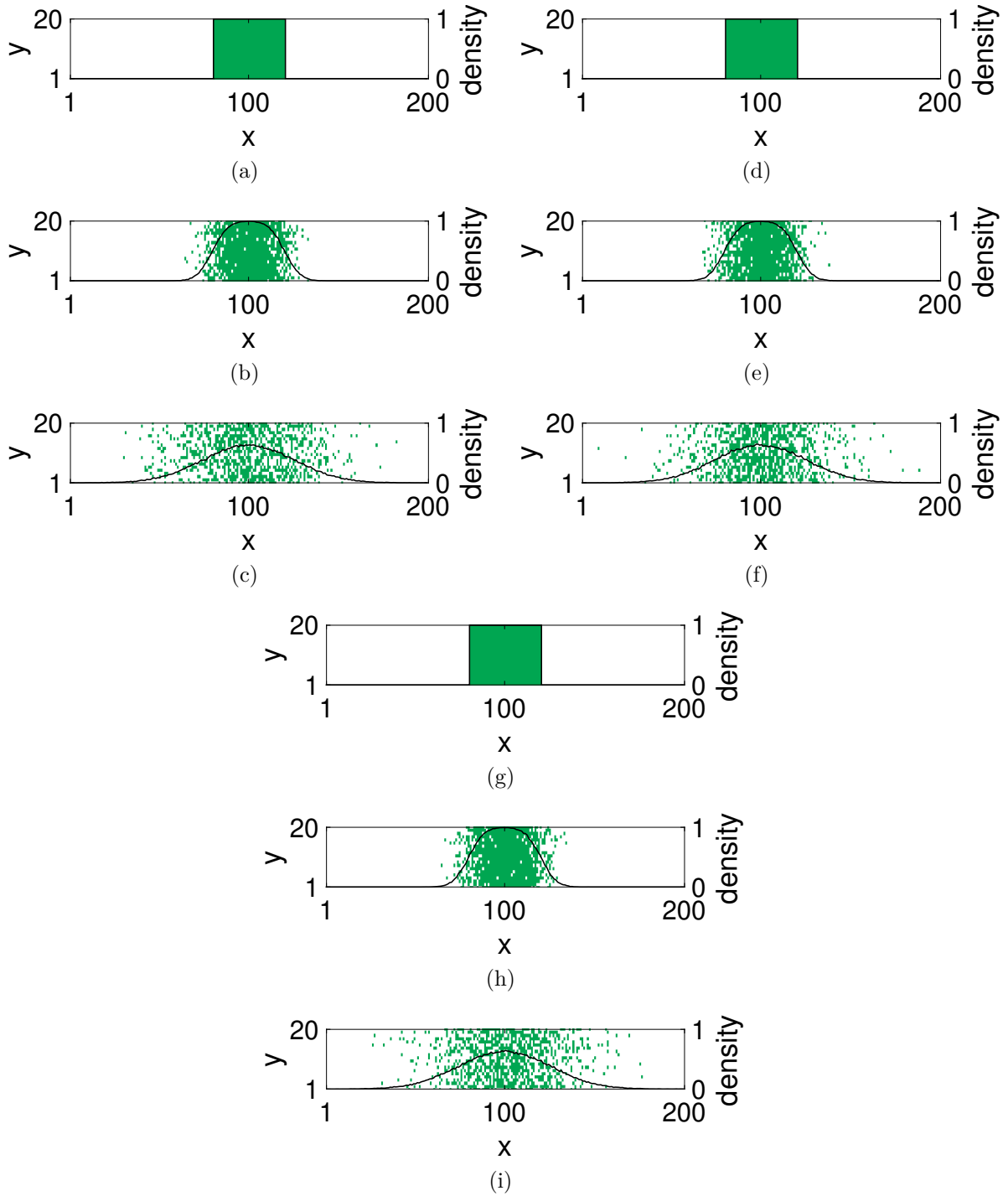


Figure 13. Snapshots of lattice occupancy for the single-species swapping model at $t = 0, 100, 1000$ with $r = 1$ for swapping probabilities $\rho = 0$ [(a)-(c)], $\rho = 0.5$ [(d)-(f)] and $\rho = 1$ [(g)-(i)]. Agents are initialised on a domain with dimensions $L_x = 200$ and $L_y = 20$ such that all the sites in the range $81 \leq x \leq 120$ are occupied with agents (green) [(a),(d),(g)]. Further snapshots of the IBM at $t = 100$ and $t = 1000$ show the dispersal of agents with time. The column-averaged density of agents over 100 runs of the ABM is also plotted (shown in black). We impose reflective boundary conditions on all four boundaries of the domain.

566 Let $C_{ij}(t) = C_{ij}$ for conciseness. By considering the possible movement events of the
 567 agent at the site (i, j) during the small time step δt [24, 25], we can write down the
 568 master equation for the occupancy of the site at time $t + \delta t$,

$$\begin{aligned}
 \langle C_{ij}(t + \delta t) \rangle - \langle C_{ij} \rangle &= \frac{r}{4} \delta t [(1 - \langle C_{ij} \rangle) (\langle C_{i-1,j} \rangle + \langle C_{i+1,j} \rangle + \langle C_{i,j-1} \rangle + \langle C_{i,j+1} \rangle) \\
 &\quad - \langle C_{ij} \rangle (4 - \langle C_{i-1,j} \rangle - \langle C_{i+1,j} \rangle - \langle C_{i,j-1} \rangle - \langle C_{i,j+1} \rangle)] \\
 &\quad + \frac{r}{4} \rho \delta t \langle C_{ij} \rangle (\langle C_{i-1,j} \rangle + \langle C_{i+1,j} \rangle + \langle C_{i,j-1} \rangle + \langle C_{i,j+1} \rangle) \\
 &\quad + \frac{r}{4} \rho \delta t \langle C_{ij} \rangle (\langle C_{i-1,j} \rangle + \langle C_{i+1,j} \rangle + \langle C_{i,j-1} \rangle + \langle C_{i,j+1} \rangle) \\
 &\quad - \frac{r}{4} \rho \delta t \langle C_{ij} \rangle (\langle C_{i-1,j} \rangle + \langle C_{i+1,j} \rangle + \langle C_{i,j-1} \rangle + \langle C_{i,j+1} \rangle) \\
 &\quad - \frac{r}{4} \rho \delta t \langle C_{ij} \rangle (\langle C_{i-1,j} \rangle + \langle C_{i+1,j} \rangle + \langle C_{i,j-1} \rangle + \langle C_{i,j+1} \rangle). \quad (\text{A2})
 \end{aligned}$$

569 The site (i, j) can gain occupancy if it is unoccupied at time t and the agent from a
 570 neighbouring site moves in (first line on the RHS of Equation (A2)). Similarly, the site
 571 (i, j) can lose occupancy if the site is occupied at time t and the residing agent jumps
 572 out to a neighbouring compartment, leaving the site (i, j) empty (second line in Equation
 573 (A2)). Movement of agents due to a successful swapping event is captured by the lines 3-6
 574 in Equation (A2). However, since the agents are assumed to be identical and unlabelled
 575 lines 3-6 cancel each other out, eliminating the effect of swapping. Consequently, we are
 576 left with lines 1 and 2 only. Taylor expanding these remaining terms around the site
 577 (i, j) up to second-order and taking the limit as $\Delta \rightarrow 0$ gives the diffusion equation, as
 578 expected:

$$\frac{\partial C}{\partial t} = D \nabla^2 C. \quad (\text{A3})$$

579 Here,

$$D = \lim_{\Delta \rightarrow 0} \frac{r \Delta^2}{4},$$

580 is the diffusion coefficient given that $\Delta^2/\delta t$ is held constant in the diffusive limit. Equation
 581 (A3) describes the evolution of the lattice occupancy over time. It is well-known that
 582 for the simple exclusion process, the occupancy is described by the diffusion equation
 583 [19, 31]. It makes sense therefore that in Figure 13 swapping made no difference to the

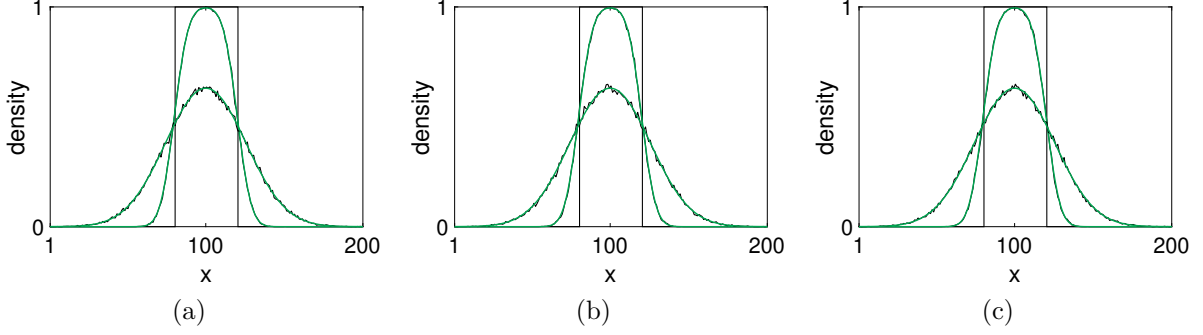


Figure 14. A comparison between the numerical solution of the one-dimensional version of the PDE (A3) and the averaged behaviour of the ABM with $r = 1$ for $\rho = 0$ (a), $\rho = 0.5$ (b) and $\rho = 1$ (c). The averaged densities are shown in black and the corresponding PDE approximation is shown in green. We present solutions at $t = 0$, $t = 100$ and $t = 1000$. The black arrows show the direction of increasing time.

584 overall occupancy of the lattice. In Figure 14 we compare the average column density of
 585 the ABM which is given by,

$$\bar{C}_i(t) = \frac{1}{L_y} \sum_{j=1}^{L_y} C_{ij}(t),$$

586 to the solution of the one-dimensional analogue of Equation (A3) with reflective boundary
 587 conditions by averaging the PDEs over the y direction.

588 As expected, we see an excellent agreement between the two density profiles. We also
 589 see that there is perfect agreement even in the zero-swapping case which we did not see
 590 in the two-species example in Section II B. This is because in a single-species system
 591 there is no way of distinguishing between a successful swapping of the position of a pair
 592 of neighbouring agents and an aborted movement event due to the volume-exclusion
 593 principle. This leads to identical profiles for different ρ regardless of its magnitude.

594 In contrast to the two-species case, here there is perfect agreement between the ABM
 595 and the PDE profiles since in the single-species case the agents are indistinguishable from
 596 each other. In the two-species scenario, because the agents are competing for space, in the
 597 no-swapping situation one species affects the occupancy of the other, leading to spatial
 598 correlation and discrepancy between the ABM and PDE descriptions. Swapping serves
 599 to break up the correlation and improve agreement between the discrete and continuum
 600 models.

601 **Appendix B: Two-species individual-level diffusion coefficients**

602 Let $P_{ij}^A(t) = P_{ij}$ be the probability that a type-A agent occupies the site (i, j) and let
 603 $P_{ij}^B(t) = P_{ij}^B$ be the equivalent for species B. We can write down the master equation for
 604 species A and species B at time $t + \delta t$ where δt is a small change in time,

$$\begin{aligned}
 P_{ij}^A(t + \delta t) &= \frac{r_A}{4} \delta t (1 - c) \left[P_{i-1,j}^A + P_{i+1,j}^A + P_{i,j-1}^A + P_{i,j+1}^A \right] + \frac{r_A}{4} c_A \rho \delta t \left[2P_{i-1,j}^A + 2P_{i+1,j}^A \right. \\
 &\quad \left. + 2P_{i,j-1}^A + 2P_{i,j+1}^A \right] + \frac{r_A}{4} c_B \rho \delta t \left[P_{i-1,j}^A + P_{i+1,j}^A + P_{i,j-1}^A + P_{i,j+1}^A \right] \\
 &\quad + \frac{r_B}{4} c_B \rho \delta t \left[P_{i-1,j}^A + P_{i+1,j}^A + P_{i,j-1}^A + P_{i,j+1}^A \right] + P_{ij}^A \left[1 - r_A (1 - c) \delta t \right. \\
 &\quad \left. - 2r_A \delta t c_A \rho - r_A \delta t c_B \rho - r_B \delta t c_B \rho \right] \tag{B1} \\
 &= \left(\frac{r_A}{4} (1 - c) + \frac{r_A}{2} c_A \rho + \frac{(r_{ACB} + r_{BCB})}{4} \rho \right) \delta t (P_{i-1,j}^A + P_{i+1,j}^A + P_{i,j-1}^A + P_{i,j+1}^A) \\
 &\quad + P_{ij}^A + (-r_A (1 - c) - 2r_{AC} \rho - (r_{BCB} + r_{ACB}) \rho) \delta t P_{ij}^A
 \end{aligned}$$

$$\begin{aligned}
 P_{ij}^B(t + \delta t) &= \frac{r_B}{4} \delta t (1 - c) \left[P_{i-1,j}^B + P_{i+1,j}^B + P_{i,j-1}^B + P_{i,j+1}^B \right] + \frac{r_B}{4} c_B \rho \delta t \left[2P_{i-1,j}^B + 2P_{i+1,j}^B \right. \\
 &\quad \left. + 2P_{i,j-1}^B + 2P_{i,j+1}^B \right] + \frac{r_B}{4} c_A \rho \delta t \left[P_{i-1,j}^B + P_{i+1,j}^B + P_{i,j-1}^B + P_{i,j+1}^B \right] \\
 &\quad + \frac{r_A}{4} c_A \rho \delta t \left[P_{i-1,j}^B + P_{i+1,j}^B + P_{i,j-1}^B + P_{i,j+1}^B \right] + P_{ij}^B \left[1 - r_B (1 - c) \delta t \right. \\
 &\quad \left. - 2r_B \delta t c_B \rho - r_B \delta t c_A \rho - r_A \delta t c_A \rho \right] \tag{B2} \\
 &= \left(\frac{r_B}{4} (1 - c) + \frac{r_B}{2} c_B \rho + \frac{(r_{BCA} + r_{ACA})}{4} \rho \right) \delta t (P_{i-1,j}^B + P_{i+1,j}^B + P_{i,j-1}^B + P_{i,j+1}^B) \\
 &\quad + P_{ij}^B + (-r_B (1 - c) - 2r_{BC} \rho - (r_{ACA} + r_{BCA}) \rho) \delta t P_{ij}^B
 \end{aligned}$$

605 We explain the meaning of the terms in the discrete-time master Equation (B1). Equation
 606 (B2) can be interpreted similarly. The equations describe the evolution of the probability
 607 that a focal agent of type A is occupying site (i, j) by considering possible movement
 608 events of the focal agent at site (i, j) and agents at neighbouring sites. Firstly, the terms
 609 which correspond movements that increase the probability that the focal agent sits at

610 position (i, j) are as follows:

- 611 1. The focal type-A agent residing at a neighbouring site moves into the empty site
612 (i, j) (line 1, term 1).
- 613 2. The focal type-A agent at a neighbouring site initiates and successfully completes
614 a swap with a type-A agent at site (i, j) or, alternatively a type-A agent at site
615 (i, j) initiates the swap and exchanges position with the focal type-A agent at the
616 neighbouring site (hence the multiplier 2 for the two equally likely probabilities)
617 (term 2 on line 1 through line 2 up to first closing square bracket).
- 618 3. The focal type-A agent at a neighbouring site initiates and successfully swaps with
619 a type-B agent at site (i, j) (term 2 on line 2).
- 620 4. A type-B agent at site (i, j) initiates and successfully swaps positions with the focal
621 type-A agent at a neighbouring site (term 1 on line 3).

622 Secondly the terms which correspond to the site already being occupied and no event
623 occurring to change that state correspond to the terms inside the pair of square brackets
624 which spans lines 3 and 4. Recalling that the probability of nothing happening in the
625 time interval $[t, t + \delta t)$ is unity minus the probability that something happens, the terms
626 in square brackets can be described as follows:

- 627 1. The unit corresponds to the probability that the focal agent occupied site (i, j) at
628 time t .
- 629 2. The second term correspond to the focal type-A agent at site (i, j) moving to an
630 unoccupied neighbouring site.
- 631 3. The third term corresponds to the probability that the focal agent of type A at
632 site (i, j) initiates and successfully swaps with a type-A agent at a neighbouring
633 site and the probability that a type-A agent at a neighbouring site initiates and
634 successfully swaps with the focal agent of type A at site (i, j) (hence the factor of
635 2 for these equally probably events).
- 636 4. The fourth term corresponds to the probability that the focal type-A agent at
637 position (i, j) initiates and successfully swaps with a neighbouring agent of type B.

638 5. Finally, the fifth term in the square brackets corresponds to a type-B agent at
639 a neighbouring site successfully swapping positions with the type-A focal agent
640 occupying site (i, j) .

641 Subtracting P_{ij}^A from both side and dividing by δt and taking the limit as $\delta t \rightarrow 0$ gives,

$$\begin{aligned} \frac{dP_{ij}^A}{dt} = & \left(\frac{r_A}{4}(1-c) + \frac{r_A}{2}c_{A\rho} + \frac{(r_A+r_B)}{4}c_{B\rho} \right) (P_{i-1,j}^A + P_{i+1,j}^A + P_{i,j-1}^A + P_{i,j+1}^A) \\ & + (-r_A(1-c) - 2r_Ac_{A\rho} - (r_A+r_B)c_{B\rho})P_{ij}^A, \end{aligned} \quad (\text{B3})$$

$$\begin{aligned} \frac{dP_{ij}^B}{dt} = & \left(\frac{r_B}{4}(1-c) + \frac{r_B}{2}c_{B\rho} + \frac{(r_B+r_A)}{4}c_{A\rho} \right) (P_{i-1,j}^B + P_{i+1,j}^B + P_{i,j-1}^B + P_{i,j+1}^B) \\ & + (-r_B(1-c) - 2r_Bc_{B\rho} - (r_A+r_B)c_{A\rho})P_{ij}^B. \end{aligned} \quad (\text{B4})$$

642 After using the definitions in Equation (12) and (13), and suitably transforming the
643 indices i and j , after simplifying it can be shown that the equations describing the
644 evolution of the second moment of the position i and j of species A and B are given
645 by,

$$\frac{d(\langle i^2 \rangle + \langle j^2 \rangle)_A}{dt} = r_A(1-c) + 2r_Ac_{A\rho} + (r_A+r_B)c_{B\rho}, \quad (\text{B5})$$

$$\frac{d(\langle i^2 \rangle + \langle j^2 \rangle)_B}{dt} = r_B(1-c) + 2r_Bc_{B\rho} + (r_A+r_B)c_{A\rho}. \quad (\text{B6})$$

646 Given that $\langle i(0) \rangle = \langle j(0) \rangle = 0$,

$$(\langle i^2 \rangle + \langle j^2 \rangle)_A = r_A(1-c) + 2r_Ac_{A\rho} + (r_A+r_B)c_{B\rho}t,$$

$$(\langle i^2 \rangle + \langle j^2 \rangle)_B = r_B(1-c) + 2r_Bc_{B\rho} + (r_A+r_B)c_{A\rho}t.$$

647 Thus,

$$D_A^* = \frac{1}{4}(r_A(1 - c) + (r_{AC} + r_{ACA} + r_{BCB})\rho), \quad (\text{B7})$$

$$D_B^* = \frac{1}{4}(r_B(1 - c) + (r_{BC} + r_{ACA} + r_{BCB})\rho). \quad (\text{B8})$$

648 are the individual-level time-uncorrelated diffusion coefficients of species A and B, re-
649 spectively.

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