

cellular_raza: Agent-based modelling of cellular systems from a clean slate

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Summary

cellular_raza is a library that allows users to define fully-customized cellular agents in order to run numerical simulations. It formulates simulation aspects in the form of rust traits. *TODO CITATION* Cellular agents implement a subset of these simulation aspects and cellular_raza provides generic methods to numerically solve the system and store results. It also comes with predefined building blocks for agents and their physical domain to quickly construct new simulations bottom-up. Furthermore, cellular_raza has been used with the pyo3 and maturin packages to handily create python bindings and act as a numerical backend to a python package.

Statement of need

Agent-based models are common in cellular biology and many tools have been developed so far to asses specific questions in specialized fields (Pleyer & Fleck, 2023). While these tools have proven to be effective for targeted research questions, they often lack the ability to be applied in a more generic context. In order to combat this issue and build up models from first principles without any assumptions on the underlying complexity or abstraction level, we developed cellular_raza.

- TODO CITATIONS*

State of field

Generic agent-based modelling toolkits

There exists a wide variety of many general-purpose agent-based simulation toolkits which are being actively applied in a different fields of study (Abar et al., 2017). - *TODO CITATIONS* These tools are often able to define agents bottom-up and can be a good choice if they allow for the desired cellular representation. However, they lack the explicit forethough to be applied in cellular systems and often implement global rules rather than individual-based ones. - *TODO CITATIONS*.

Cellular agent-based frameworks

In our previous efforts (Pleyer & Fleck, 2023) we have assessed the overall state of modelling toolkits for individual-based cellular simulations. In this mini-review, we focussed on modelling frameworks, which provide a complete workflow. The resulting frameworks are all crafted for specific use-cases and may require a large amount of parameters specific to their domain of usage. These parameters are often not known in practice and are hard to determine

36 experimentally. This creates problems for the extendability of the software and the ability to
37 properly interpret results.

38 We can further reduce the number of modeling frameworks by only considering ones which
39 provide a significant level of flexibility and customizability in their definition of cell-agents.
40 Chaste allows to reuse individual components of their simulation code such as ODE and PDE
41 solvers. - *TODO CITATION* Biocellion has support for different cell shapes such as spheres
42 and cylinders but acknowledges that their current approach lacks flexibility in the subcellular
43 description.

44 ▪ *TODO CITATION*
45 ▪ *TODO check which other frameworks to consider*

46 **Underlying Assumptions and Internals**

47 **List of Simulation Aspects**

Aspect	Description	Depends on
Cellular Agent		
Position	Spatial representation of the cell	Position and Velocity
Velocity	Spatial velocity of the cell	
Mechanics	Calculates the next increment from given force, velocity and position.	
Interaction	Calculates force acting between agents. Also reacts to neighbours.	Position and Velocity
Cycle	Changes core properties of the cell. Responsible for cell-division and death.	
Intracellular	Intracellular representation of the cell.	
Reactions	Intracellular reactions	Intracellular
ReactionsExtra	Couples intra- & extracellular reactions	DomainReactions
ReactionsContact	Models reactions between cells purely by contact	Position, Intracellular
Simulation Domain		
Domain	Represents the physical simulation domain.	
DomainMechanics	Apply boundary conditions to agents.	Position, Velocity
DomainForce	Apply a spatially-dependent force onto the cell.	Mechanics
DomainReactions	Calculate extracellular reactions and effects such as diffusion.	ReactionsExtra
Other		
Controller	Externally apply changes to the cells.	

48 **Spatially Localized Interactions**

49 One of the most fundamental assumptions within `cellular_raza` is that each and every
50 interaction is of finite range. This means that cellular agents only interact with their nearest
51 neighbour and close environment. Any long-ranged interactions must be the result of a
52 collection of short-ranged interactions. This assumption enables us to split the simulation
53 domain into chunks and process them individually although some communication is needed
54 in order to deal with boundary conditions. In practice, this means that any interaction force
55 should be given a cutoff. It also means that any interactions which need to be evaluated
56 between agents should in theory scale linearly with the number of agents $\mathcal{O}(n_{\text{agents}})$.

57 **Code Structure**

58 `cellular_raza` consists of multiple crates working in tandem. It was designed to have clear
59 separations between conceptual choices and implementation details. This approach allows us
60 to have a greater amount of modularity and flexibility than regular simulation tools.

61 These crates act on varying levels of abstraction to yield a fully working numerical simulation.
62 Since `cellular_raza` functions on different levels of abstraction, we try to indicate this in the
63 table below.

crate	Abstraction Level	Purpose
<code>cellular_raza</code>	-	Bundle together functionality of all other crates.
<code>concepts</code>	High	Collection of (mainly) traits which need to be implemented to yield a full simulation.
<code>core</code>	Intermediate-High	Contains numerical solvers, storage handlers and more to actually solve a given system.
<code>building_blocks</code>	Intermediate	Predefined components of cell-agents and domains which can be put together to obtain a full simulation.
<code>examples</code>	Application	Showcases and introductions to different simulation approaches.
<code>benchmarks</code>	Application	Performance testing of various configurations.

64 **Backends**

65 To numerically solve a fully specified system, `cellular_raza` provides backends. The function-
66 ality offered by a backend is the most important factor in determining the workflow of the
67 user and how a given simulation is executed. Currently, we provide the default `chili` backend
68 but hope to extend this collection in the future. Backends may choose to purposefully restrict
69 themselves to a subset of simulation aspects or a particular implementation in order to improve
70 performance.

71 **Chili**

72 The `chili` backend is the default choice for any new simulation. It generates source code by
73 extensively using `macros` and `generics`. Afterwards, the generated code is compiled and run.

74 Every backend function is implemented generically by hand. We use [trait bounds](#) to enforce
75 correct usage of every involved type. The generated code is restricted to methods of structs
76 and derivations of their components functionality. To obtain a fully working simulation, the
77 `chili` backend combines these generic methods with user-provided and generated types. The
78 `run_simulation!` macro generates code depending on which type of simulation aspect is
79 activated by the user. By employing this combined scheme of generics and macros, we leverage
80 the strong type-system and Rusts language-specific safety to avoid pitfalls which a purely
81 macro-based approach would yield.

82 Other Backends

83 `cellular_raza` also comes with the `cpu_os_threads` backend which was the first backend
84 created. It is in the midst of being deprecated and only serves for some legacy usecases. In
85 the future, we hope to add a dedicated backend named `cara` to leverage GPU-accelerated
86 (Graphical Processing Unit) algorithms.

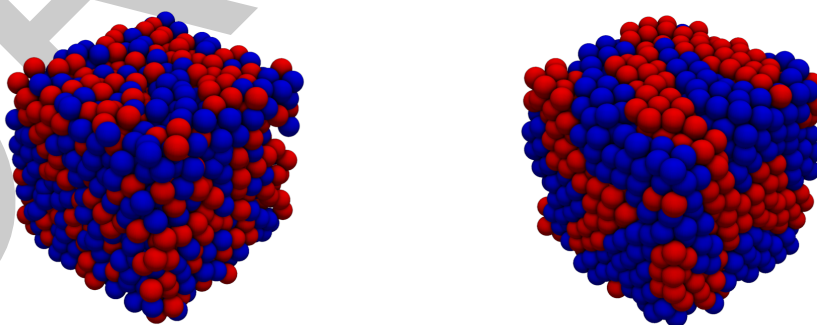
87 Examples

88 All presented examples can be viewed at the [showcase](#) section of the [cellular-raza.com](#) homepage.

89 Cell Sorting

90 Cell Sorting is a naturally occurring phenomenon which drives many biological processes.
91 *TODO CITATION* While the underlying biological reality can be quite complex, it is rather
92 simple to describe such a system in its most basic form. The responsible principle is that
93 the Interaction between cells are specific to their species. In our example, we consider
94 two distinct species represented by soft spheres which physically attract each other at close
95 proximity if their species is identical.

96 We initially place cells randomly inside a cube with reflective boundary conditions. In the final
97 snapshot, we can clearly see the phase-separation between the different species.



98 **Figure 1:** The initial random placement of cells reorders into a phase-separated spatial pattern.

99 Bacterial Rods

100 Many bacterial species are of elongated shape *TODO CITATION* which grows asymmetrically
101 in the direction of elongation during the growth phase of the cell. To model this behaviour,
102 we describe the physical Mechanics of one cell as a collection of multiple vertices \vec{v}_i which
103 are connected by edges. The edges are modelled as springs and their relative angle at each
104 connecting vertex introduces a stiffening force which is proportional to the angle difference

105 $\alpha - 180^\circ$. The Interaction of two cells is implemented via a force potential which acts
106 between every vertex and the closest point on the other cells edges. The potential that of a
107 soft-sphere with a short-ranged adherent force.

108 In addition, the cell Cycle introduces growth of the bacteria until it reaches a threshold and
109 divides in the middle into two new cells. The growth is downregulated by an increasing number
110 of neighboring cells. This can also be accomplished by the Interaction simulation aspect. It
111 is an phenomenological but effective choice to model the gradual transition into the stationary
112 phase of the bacterial colony.

113 Initially, the cells are placed inside the left-hand side of an elongated box with reflective
114 boundary conditions. The cells are colored continuously from green for fast growth to blue
115 for dormant cells. This setup is reminiscent of a mother machine continuously producing new
116 bacteria. *TODO CITATION*



117
Figure 2: The bacteria extend from the initial placement in the left side towards the right side. Their elongated shape and the confined space favour the orientation facing along the growth direction.

118 **Branching of *Bacillus Subtilis***

119 Spatio-temporal patterns of bacterial growth such as in *Bacillus Subtilis* have been studied
120 for numerous years (Kawasaki et al., 1997; Matsushita et al., 1998). They are typically
121 described by a system of PDEs (Partial Differential Equations) which contain non-spatial and
122 spatial contributions. describing intracellular reactions and cell-cycle and spatial contributions
123 (typically via Diffusion processes) which describe diffusion of nutrients and movement of the
124 cells.

125 With cellular_raza we can clearly distinguish between these simulation aspects. We describe
126 the Mechanics and physical Interaction of the cells as soft spheres. Extracellular reactions
127 (DomainReactions) in the simulation domain are modeled by Diffusion which is coupled via
128 an uptake term (ReactionsExtra) to the cells intracellular Reactions. During its life Cycle,
129 the cell grows continuously and divides upon reaching a threshold.

130 The initial placement of the cells is inside of a centered square. From there, cells start
131 consuming nutrients and growing outwards towards the nutrient-rich area. Cells are colored
132 bright purple while they are actively growing and dividing while dark cells are not subject to
133 growth anymore. The outer domain is colored by the intensity of present nutrients. A lighter
134 color indicates that more nutrients are available while a dark color signifies a lack thereof. The
135 two snapshots show the state after 28% of the total simulation time and at the final simulation
136 step. The diffusivity of the nutrient and the growth rate of the bacteria are the governing
137 criteria for the shape of the pattern.

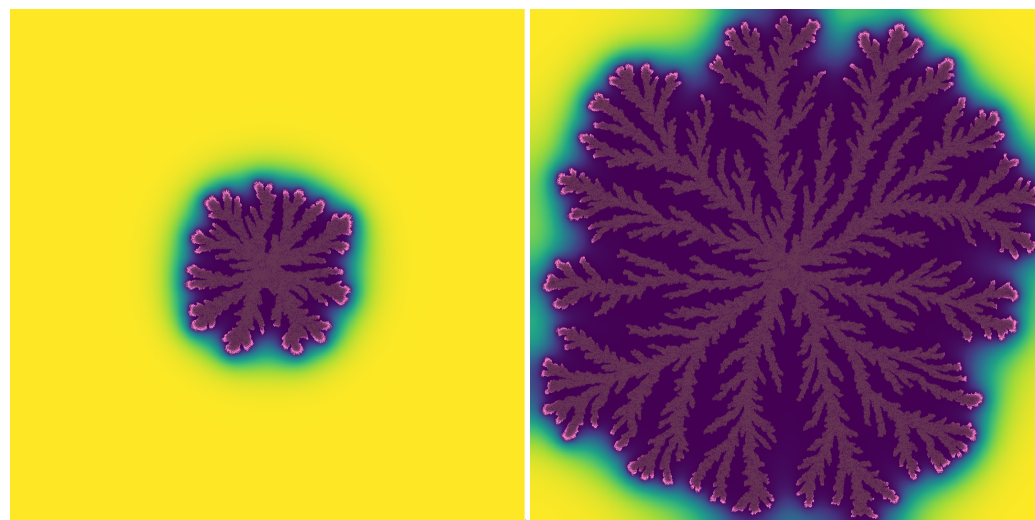


Figure 3: The bacterial colony grows outwards towards the nutrient-rich parts of the domain thus forming branches in the process.

Trichome Patterning in *Arabidopsis Thaliana*

Trichomes are hairs consisting of a single cell which can be seen developing on the surface of leaves of *Arabidopsis Thaliana*. Their specialization from regular epithelial cells is determined by a pattern which can be described by coupling the intracellular reactions of multiple cells to each other. We assume that cells are immobile but exchange the 'trichome-promoting factor TRANSPARENT TESTA GLABRA1' (TTGL) (Bouyer et al., 2008) via their connecting cell wall.

We describe the intracellular gene regulatory network as an Ordinary Differential Equation (ODE) via the Intracellular simulation aspect and couple cells to each other with the ReactionsContact aspect. We restrict the reactions via contact to only neighbouring cells exchange TTGL which is particularly simple to determine for a static tissue. The combined reactions represent a diffusion-driven Turing instability (Turing, 1952) which when given randomized initial values generates peaks that lead to the observed differentiation of the trichome hairs.

There is an ongoing effort (Pleyer, 2024) to use cellular_raza as a simulation backend while generating python bindings with the pyo3 and maturin crates.

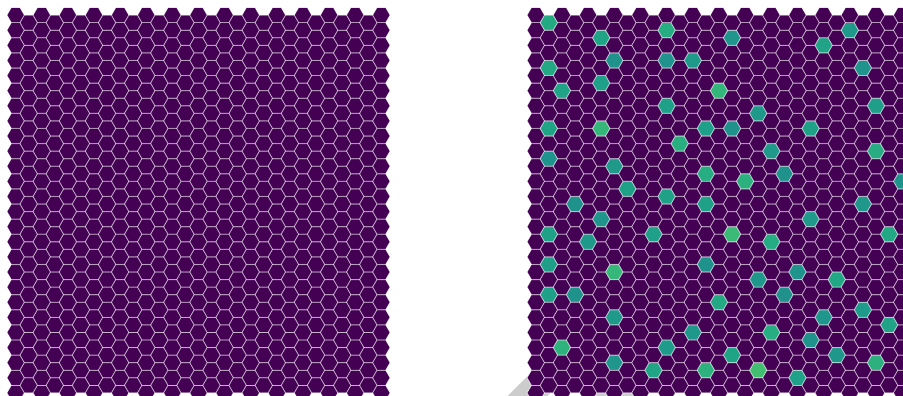


Figure 4: The reaction network produces a pattern of regular peaks that ultimately lead to the differentiation and growth of trichomes.

Performance

Multithreading

One measure of multithreaded performance is to calculate the possible theoretical speedup given by Amdahl's law (Rodgers, 1985)

$$T(n) = T_0 \frac{1}{(1-p) + \frac{p}{n}} \quad (1)$$

where n is the number of used parallel threads and p is the proportion of execution time which benefits from parallelization.

Measuring the performance of any simulation will be highly dependent on the specific cellular properties and complexity. To measure the performance of `cellular_raza`, we chose the cell sorting example which is the one containing minimal complexity of all the aforementioned example simulations. Any computational overhead which is intrinsic to `cellular_raza` and not related to the chosen example would thus be more likely to manifest in performance results. The total runtime of the simulation is of no relevance since we are only concerned with relative speedup upon using additional resources. In addition, we fixed the frequency of each processor, to circumvent power-dependent behaviour. While it is well known that other aspects such as cache-size and memory latency can have an impact on absolute performance, they should however not introduce any significant deviations in terms of relative performance scaling.

This benchmark was run on three distinct hardware configurations.

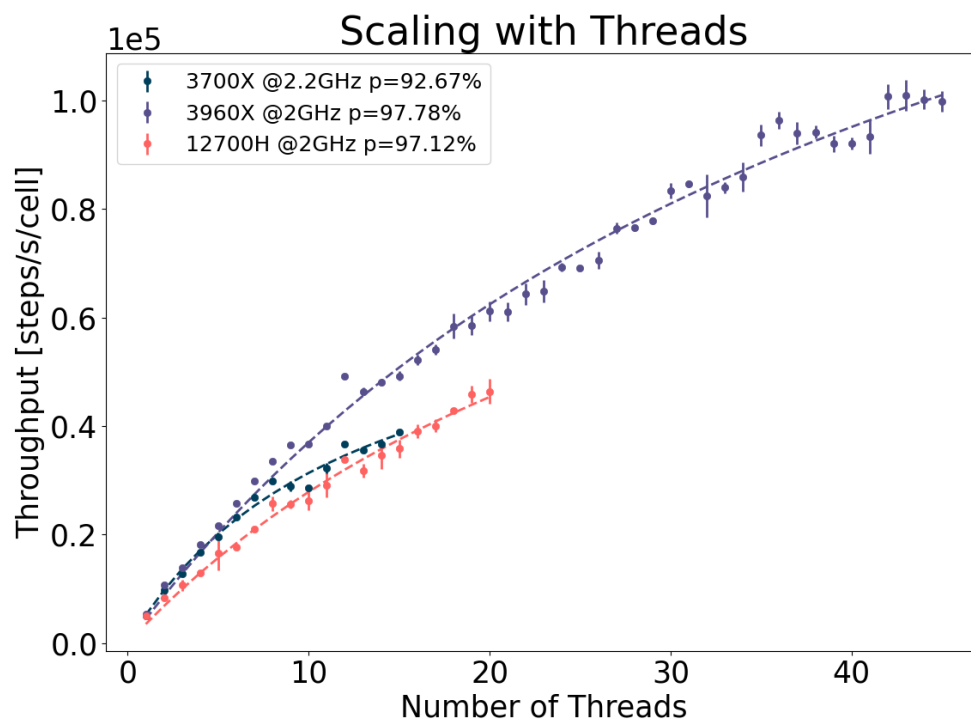


Figure 5: Amdahl's law with increasing amounts of CPU resources.

We fit equation Equation 1 and obtain the parameter p from which the theoretical maximal speedup S can be calculated via

$$S = \frac{1}{1 - p} \quad (2)$$

and thus from figure Figure 5 obtain the values $S_{3700X} = 13.64$, $S_{3960X} = 45.05$ and $S_{12700H} = 34.72$.

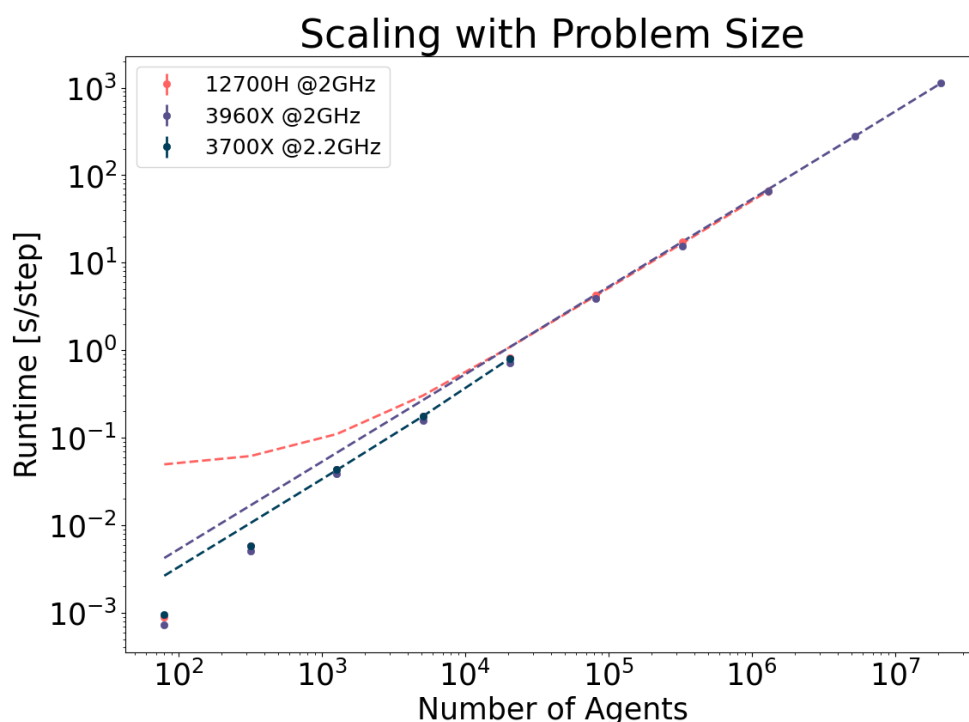


Figure 6: Scaling of the total simulation size.

Discussion

Acknowledgements

References

- Abar, S., Theodoropoulos, G. K., Lemariner, P., & O'Hare, G. M. P. (2017). Agent based modelling and simulation tools: A review of the state-of-art software. *Computer Science Review*, 24, 13–33. <https://doi.org/10.1016/j.cosrev.2017.03.001>
- Bouyer, D., Geier, F., Kragler, F., Schnittger, A., Pesch, M., Wester, K., Balkunde, R., Timmer, J., Fleck, C., & Hülkamp, M. (2008). Two-dimensional patterning by a trapping/depletion mechanism: The role of TTG1 and GL3 in arabidopsis trichome formation. *PLoS Biology*, 6(6), e141. <https://doi.org/10.1371/journal.pbio.0060141>
- Kawasaki, K., Mochizuki, A., Matsushita, M., Umeda, T., & Shigesada, N. (1997). *Modeling Spatio-Temporal Patterns Generated by Bacillus subtilis*. <https://doi.org/10.1006/jtbi.1997.0462>
- Matsushita, M., Wakita, J., Itoh, H., Ràfols, I., Matsuyama, T., Sakaguchi, H., & Mimura, M. (1998). *Interface growth and pattern formation in bacterial colonies*. [https://doi.org/10.1016/S0378-4371\(97\)00511-6](https://doi.org/10.1016/S0378-4371(97)00511-6)
- Pleyer, J. (2024). *Jonaspleyer/cr_trichome*. https://github.com/jonaspleyer/cr_trichome
- Pleyer, J., & Fleck, C. (2023). Agent-based models in cellular systems. *Frontiers in Physics*,

- 196 10. <https://doi.org/10.3389/fphy.2022.968409>
- 197 Rodgers, D. P. (1985). Improvements in multiprocessor system design. *ACM SIGARCH*
198 *Computer Architecture News*, 13(3), 225–231. <https://doi.org/10.1145/327070.327215>
- 199 Turing, A. M. (1952). The chemical basis of morphogenesis. *Philosophical Transactions*
200 *of the Royal Society of London. Series B, Biological Sciences*, 237(641), 37–72. <https://doi.org/10.1098/rstb.1952.0012>
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