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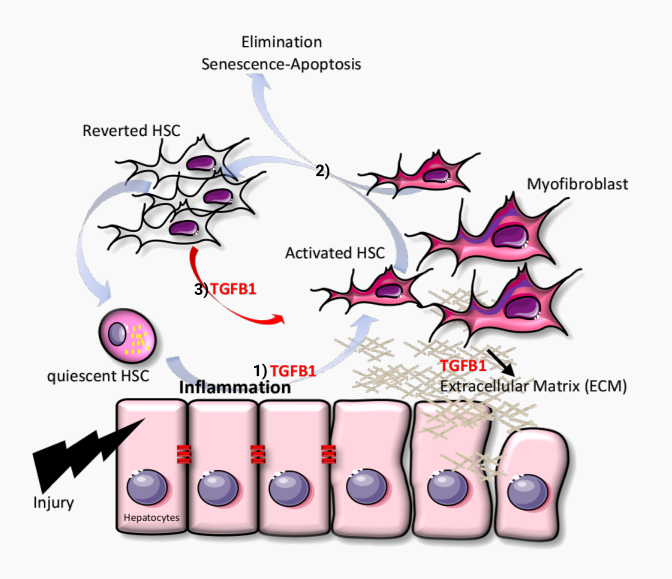
A Kappa model for hepatic stellate cells activation by TGFB1



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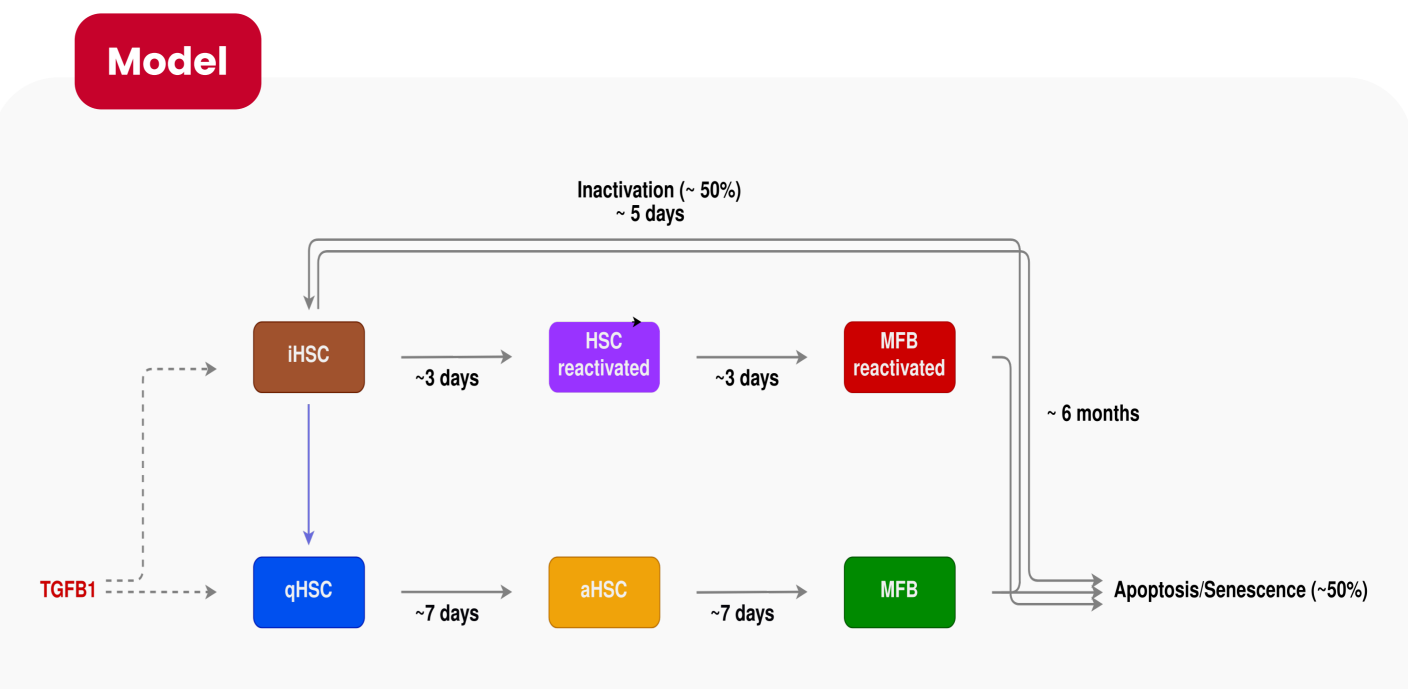
Introduction



Chronic hepatitis is associated with the development of fibrosis, which results in the abnormal deposition of extracellular matrix (**ECM**) and a severe dysfunction of liver functions. The terminal stage of fibrosis is cirrhosis, which constitutes the major risk of occurrence of Hepatocellular Carcinoma (HCC). The matrix microenvironment is therefore the major regulator of events related to the fibrosis-cirrhosis-cancer progression and Hepatic Stellate Cells (**HSC**) are the main actors of the extracellular matrix remodelling.

- Upon liver injury, damaged hepatocytes produce signals inducing inflammation that in turn promotes TGFB1-dependent activation of quiescent HSC (qHSC)(1). Activated HSC orchestrate tissue repair and are either eliminated through Senescence and Apoptosis (2) or deactivated towards reverted HSC (iHSC)(2), that can be reactivated more rapidly(3).
- Upon chronic liver injuries, activated HSC(aHSC) progress toward a Myobroblast (MFB) state that escape to control, leading to fibrosis.

The understanding of the dynamics of HSC activation and regulation by TGFB1 is essential to identify markers and therapeutic targets likely to promote the resolution of fibrosis at the expense of its progression. Here we develop a rule-based model to characterize the dynamics of HSC activation and identify the key regulators.



Schematic representation of the model, built using biological data

Kappa

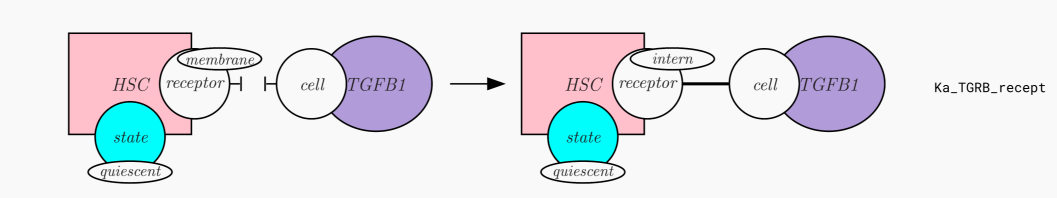
Kappa:

- Rule-based language for modeling systems of interacting agents (https://kappalanguage.org/)
- agents (<u>https://kappalanguage.org/</u>)Entities are graphical structures, rules are graph-rewrite
- directives.
- Rules locally modified the state of a system

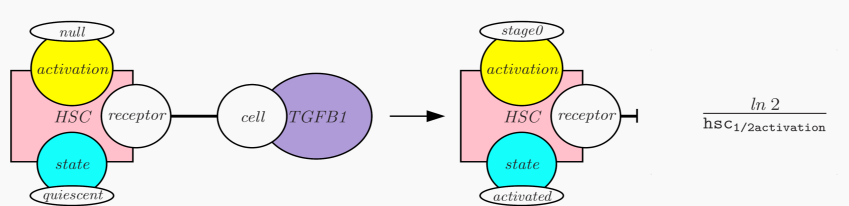
Tools:

- Stochastic simulator **KaSim**
- a. samples trajectory according to their probability density distribution
- b. relies on a representation of the state of the system as a site graph
- c. set of events that may be applied in the current state is computed dynamically
- Modeling plateform

 a. direct simulation
- a. direct simulationb. interactions during the execution of a model



'HSC(state{quiescent} TGFBR1_2{membrane}[.]), TGFB1(cell[.], state{active}) -> HSC(state{quiescent} TGFBR1_2{intern}[1]), TGFB1(cell[1] state{active}) @ 'Ka_TGFB_recep'



'HSC(state{quiescent} TGFBR1_2{intern}[1]), TGFB1(cell[1] state{active}) -> HSC(state{activateds} activation{stage0} TGFBR1_2{intern}[.]), . @ 'ln2' / 'hsc_1/2_activation'

Agent (**HSC** and **TGFB1**) are defined by different sites:

- the site receptor which bind to the site cell of TGFB1
- the site state which have different values:

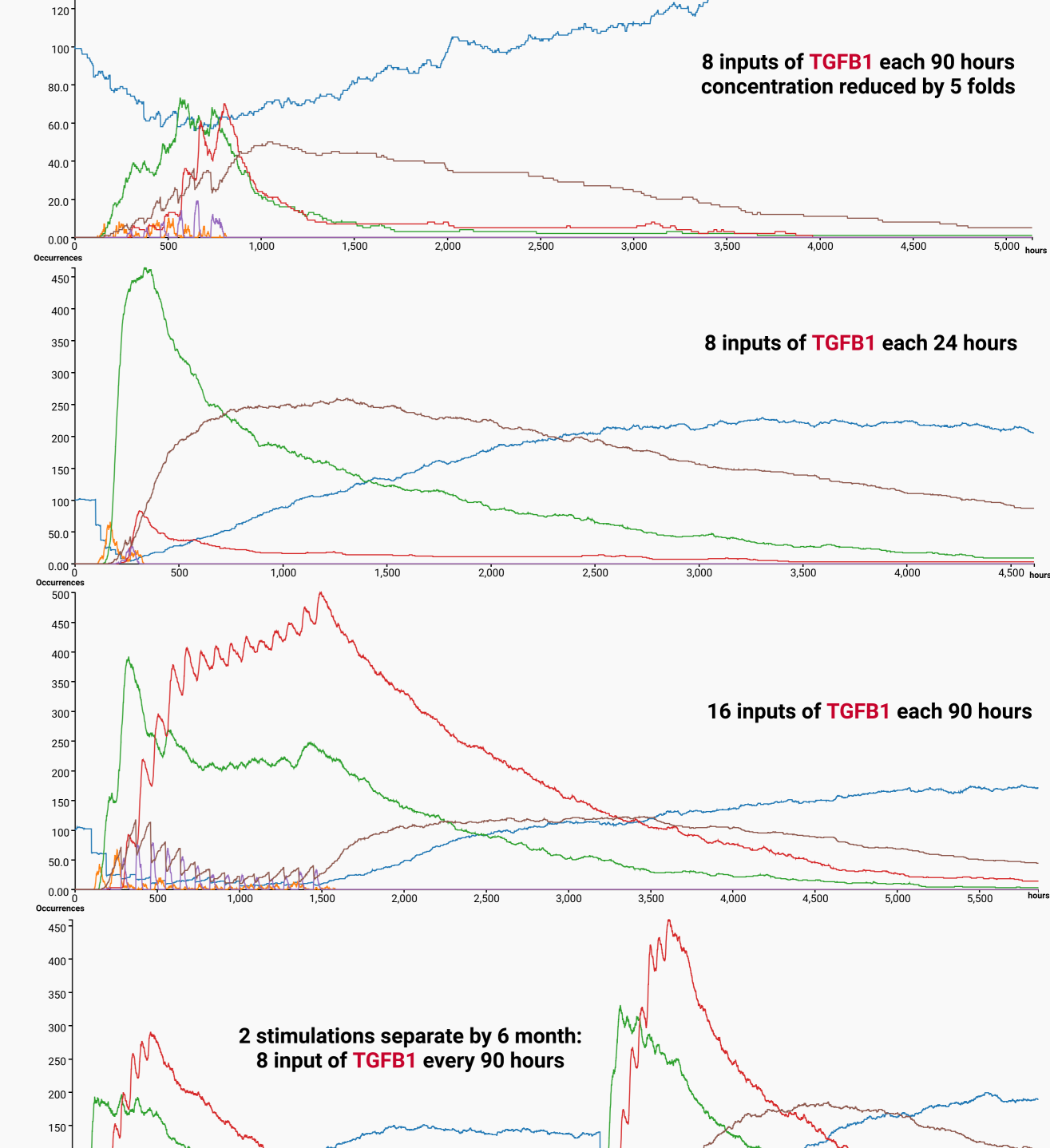
quiescent or activated

M Bouguéon, P Boutillier, J Feret, OHazard and N Theret. 2021 A Kappa model for hepatic stellate cells activation by TGFB, WILEY. Systems Biology Modelling and Analysis: Formal Bioinformatics Methods and Tools.

QHSC aHSC MFB iHSC reactivated reactivated

8 input of TGFB1 every 90 hours

hours



Conclusion / Perspectives

Conclusion

- Reverted HSC (iHSC) are key regulators
- TGFB1 is necessary to induce HSC activation but chronic stimulation by TGFB1 doesn't allow to reach a fibrotic state characterised by persistent high level of MFB

Perspectives

Because ECM deposition induced by **activated HSC** and **MFB** lead to an inscrease of stifness, which may also contribute to activation of **HSC (4)**. The futur challenges will be the integration of a new agent **ECM** to take into account this information.

