

SigHotSpotter: scRNA-seq-based computational tool to control cell subpopulation phenotypes for cellular rejuvenation strategies

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Abstract

Summary: Single-cell RNA-sequencing is increasingly employed to characterize disease or ageing cell subpopulation phenotypes. Despite exponential increase in data generation, systematic identification of key regulatory factors for controlling cellular phenotype to enable cell rejuvenation in disease or ageing remains a challenge. Here, we present SigHotSpotter, a computational tool to predict hotspots of signaling pathways responsible for the stable maintenance of cell subpopulation phenotypes, by integrating signaling and transcriptional networks. Targeted perturbation of these signaling hotspots can enable precise control of cell subpopulation phenotypes. SigHotSpotter correctly predicts the signaling hotspots with known experimental validations in different cellular systems. The tool is simple, user-friendly and is available as web-server or as stand-alone software. We believe SigHotSpotter will serve as a general purpose tool for the systematic prediction of signaling hotspots based on single-cell RNA-seq data, and potentiate novel cell rejuvenation strategies in the context of disease and ageing.

Availability and implementation

SigHotSpotter is at https://SigHotSpotter.lcsb.uni.lu as a web tool. Source code, example datasets and other information are available at https://gitlab.com/srikanth.ravichandran/sighotspotter.

Supplementary information

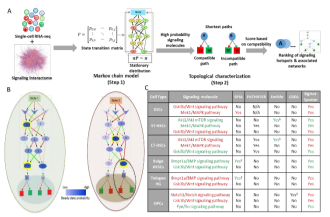
Supplementary data are available at *Bioinformatics* online.

1 Introduction

The ability to control cellular phenotypes offers a great potential for developing novel regenerative medicine strategies. In particular, rejuvenation strategies for counteracting the detrimental effect of the aged or diseased niche that impairs normal cellular functioning are essential (Cheung and Rando 2013; Lane et al. 2014; Neves et al. 2017; Del Sol et al. 2019). Advances in single-cell RNA-seq that allows for profiling of distinct cell subpopulations could aid in this endeavor. However, despite increasing amount of data generation, there is a lack of computational approaches that leverages single-cell omics data to identify specific

factors that can enable cell rejuvenation. Here, we present a general computational tool, SigHotSpotter, which relies on a probabilistic Markov chain model of signal transduction, previously developed by our lab, for the prediction of hotspots (key molecules) of signaling pathways that are constantly activated/inhibited by the niche that maintain neural stem cells (NSC) in a quiescence state (Kalamakis et al. 2019). We define signaling hotspots as those specific molecules that are involved in the sustained transmission of the external niche signals for the stable maintenance of the cell subpopulation phenotypes. Importantly, the tool aims at predicting hotspots, that exhibit highest signal flux through them in a sustained manner, rather than inferring the whole signaling pathways. Functionally, such hotspots are more likely to transmit the constitutive signals from the

niche for phenotype maintenance, in contrast to strong but transient signals which are usually associated with a change in cellular response or phenotype (Wang and Wagers 2011). With the increasing amount of single-cell RNA-seq data being generated, especially in the context of ageing and disease, SigHotSpotter can be of general utility for predicting signaling hotspots that that maintain cell subpopulation phenotypes.



Further, this could aid the development of novel cell rejuvenation strategies that aim to counteract the detrimental effect of the diseased or aged niche.

Fig. 1. SigHotSpotter steps involved and case study application: (A) Series of steps involved in SigHotSpotter. (B) Model of a toy-signaling network in two different phenotypes. The node colors in the network represent the steady state probability of the signal to be in a specific node (signaling molecule). Inhibitory edges are shown in red and activation edges in green. The edge thickness represents the interaction probability of the two molecules inferred from single-cell RNA-seq. The inverted triangle nodes represent receptors/ligands, circles represent intermediate signaling molecules, and the squares represent TFs. Nodes that exhibit both high steady state probability and compatibility with the downstream TF expression are identified as signaling hotspots. The black dotted circle indicates the higher compatibility of node 12 in niche condition 2. (C) Results of SigHotSpotter for the case study applications, a comparison with other signaling pathways/network inference and enrichment methods. Comparison of SigHotSpotter with SPIA, Pathifier and differential expression based enrichment with Enrichr. Yes / No denote if the pathway was found significant, pathway inhibition and activation are represented by red and green color, respectively. If the pathways were not available for certain methods, we looked up similar pathways from KEGG, these are marked in superscript with notation: ¹ PI3K-Akt signaling pathway, ² TGF-beta signaling pathway, ³ Notch4 signaling pathway.

2 Methods

key steps involved in SigHotSpotter is represented in Fig. 1A. Detailed description of the method is provided in the Supplementary Material. As an input, SigHotSpotter requires single-cell RNA-seq data. Further, it relies on a signaling interactome network, constructed by combining Reactome, Omnipath databases and transcriptional interactions from Metacore (Clarivate Analytics) obtained from Zaffaroni et al (Wu et al. 2010; Turei et al. 2016; Zaffaroni et al. 2019). In the first step, a state transition matrix is constructed based on the signaling interactome and the input single-cell RNA-seg data. The signal transduction process from the niche to intracellular signaling pathways is modeled as a finite discrete time-homogenous Markov chain (Kalamakis et al. 2019). The stationary distribution of this Markov chain enables shortlisting those signaling molecules (defined as receptors, ligands, kinases. phosphatases) that exhibit high steady probability, which reflects the high signal flux through

them for a given phenotype (Fig. 1). This information alone is not sufficient to infer whether these molecules are in an active or inactive state to maintain the phenotype. In the second step, SigHotSpotter attempts to delineate the potential regulatory activity status of the high probability signaling molecules by employing a topological characterization of their compatibility with differential expression status of the downstream transcription factors (TFs) (Fig. 1) (Supplementary Fig. S1 & S2). A compatibility score is calculated for each high probability signaling molecule based on their net effect on the differentially expressed downstream TFs, via all the shortest paths in the network. This score relies on the steady state probabilities from the Markov chain model and serves to both classify and rank, the active and inactive signaling hotspots by their importance to maintain the corresponding phenotype (Fig. 1). Finally, we use Igraph (Gabor Csardi and Nepusz 2006) implementation of Dijkstra's shortest paths algorithm (Dijkstra 1959) to extract the subnetwork controlled by the predicted hotspots. A screenshot of SigHotSpotter web tool is shown in Supplementary Figure S3.

3 Results

We demonstrate SigHotSpotter as a case study in four different cellular systems based on single-cell RNA-seq data from embryonic stem cells (ESCs) (Kolodziejczyk et al. 2015), hematopoietic stem cells (HSCs) (Kowalczyk et al. 2015), hair-follicle stem cells (HFSc) (Yang et al. 2017) and oligodendrocyte progenitor cells (OPCs) (Marques et al. 2016). The list of computational predictions and associated literature support is listed in Supplementary Table S1. Embryonic stem cells (ESCs) maintained under in vitro culture conditions serve as good model system to initially assess the performance of SigHotSpotter, since these cells can be stably maintained in different defined culture conditions such as 2i or LIF, and also exhibit condition dependent differences in their phenotypes (Ying et al. 2008). The culture conditions employing 2i (inhibition of Gsk3b and Mek) is known to maintain the mESCs in a naive pluripotency state, whereas, the LIF alone maintain the mESCs in a relatively primed/metastable pluripotency state (Ying et al. 2008). Importantly, sustained inhibition of Gsk3b and Mek is required for stable maintenance of naive pluripotency. In the example of mESCs (Kolodziejczyk et al. 2015), where exact signaling molecules that are constantly inhibited by the niche (culture conditions in this case) are clearly known, SigHotSpotter could correctly predict the inhibition (i.e. as inactive) of Gsk3b and Mek (Map2k1) under 2i conditions (Supplementary Table S1, Fig S1). Although, to our knowledge no general method currently exists for the task of identifying signaling hotspots that control cellular phenotypes, we compared the performance of SigHotSpotter with other general methods for signaling pathway/network inference and enrichment analysis that rely on only differential expression or network topology characterization (Tarca et al. 2009; Drier et al. 2013; Kuleshov et al. 2016). Notably, these methods were not able identify either Wnt signaling or Map kinase signaling along with their deregulation status for mESCs phenotype control (Fig. 1). Other case study applications (Fig. 1, Supplementary Table S2) and the comparison of SigHotSpotter with 4 other pathways/network inference and enrichment methods, namely, SPIA (Marques et al. 2016), GSEA (Subramanian et al. 2005), EnrichR (Wu et al. 2010) and Pathifier (Drier et al. 2013) are described in the Supplementary Material.

4 Discussion and Conclusion

Computational methods that combine molecular interaction databases and genomics data have been very useful for the inference of dysregulated signaling pathways and networks, especially in the context of cancer (Leiserson et al. 2015). However, these methods are not specifically built for the prediction of key molecules or hotspots that constantly mediate cell-extrinsic niche cues for the stable maintenance of the cellular phenotype. Furthermore, this is a challenge, since signal transduction involves several post translational modifications, and is not a deterministic linear cascade of biochemical interactions (as often depicted in pathway diagrams), but rather a probabilistic process involving multiple proteinprotein interactions (Ladbury and Arold 2012). In this regard, although SigHotSpotter is based on transcriptomics data, it benefits from the heterogeneity of single-cell gene expression, and attempts to overcome some of these challenges by relying on a probabilistic model to infer signaling hotspots that most likely transmit the sustained niche-induced signals, rather than inferring the entire dysregulated signaling pathways. Hence, SigHotSpotter is qualitatively different from the plethora of methods for pathway enrichment or inference of functional signaling networks (Amadoz et al. 2018), as it predicts specific signaling molecules and their regulatory effect on the cellular phenotype. In addition, the ranking of the hotspots along with their associated network controlling the downstream TFs will serve as a guide experimentalists to prioritize the predicted targets for further study.

In summary, SigHotSpotter employs single-cell RNA-seq data to serve as a general purpose tool for predicting signaling hotspots that control cell subpopulation phenotypes. Importantly, this can enable the development of cell rejuvenation strategies for counteracting the detrimental effect of the niche due to disease or ageing, where endogenous stem cells lose their activation potential.

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Conflict of Interest: none declared.

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