

# An Information Theoretic Model of Engineered Cell-Cell Communication

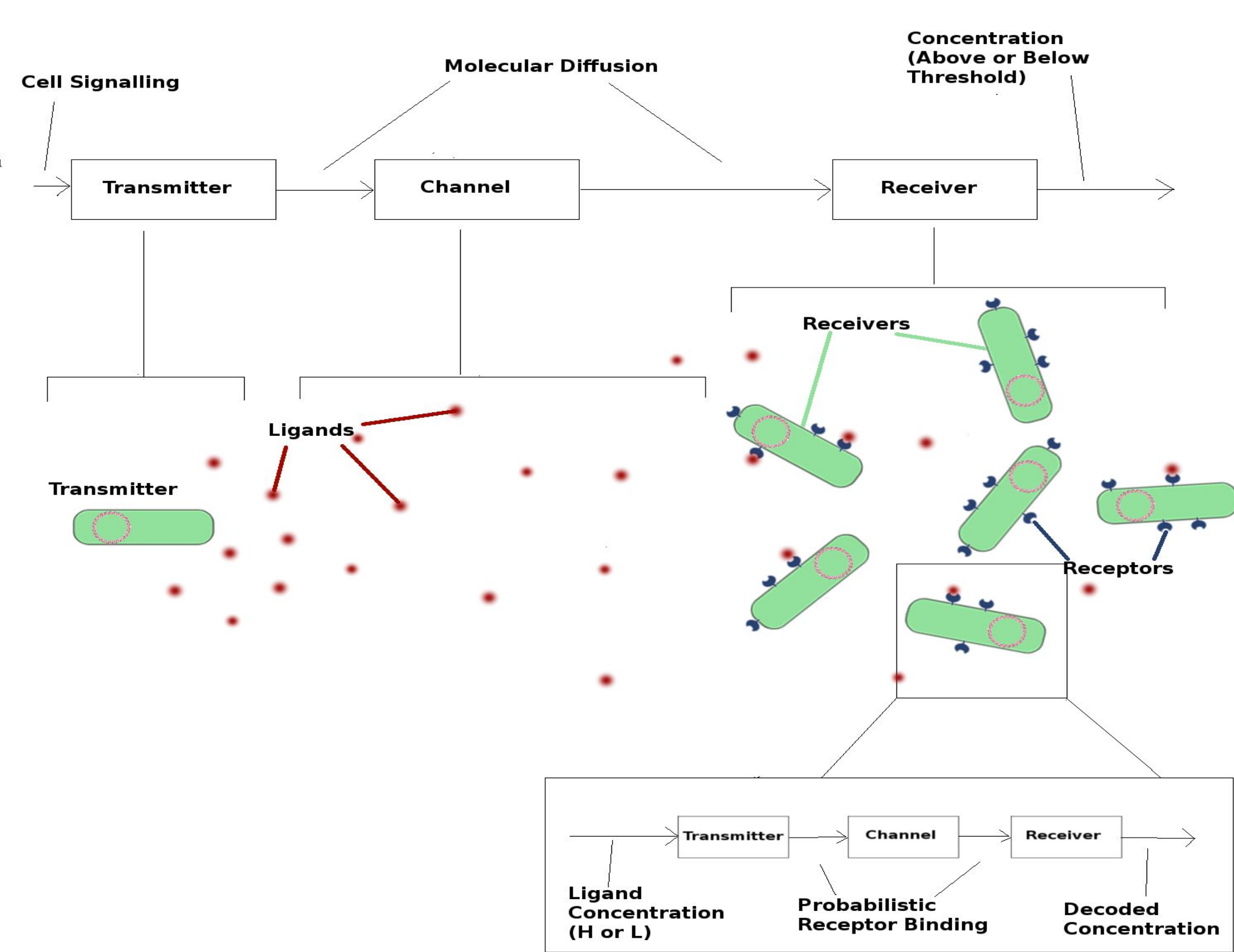
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## 1. Introduction

- The engineering of communicating biological cell swarms is limited by a lack of theoretical design principles
- We develop an information-theoretic model of cell-cell communication to:
  - Determine the information capacity of groups of communicating cells
  - Understand bottlenecks in cellular communication
  - Derive general design principles

## 2. Model



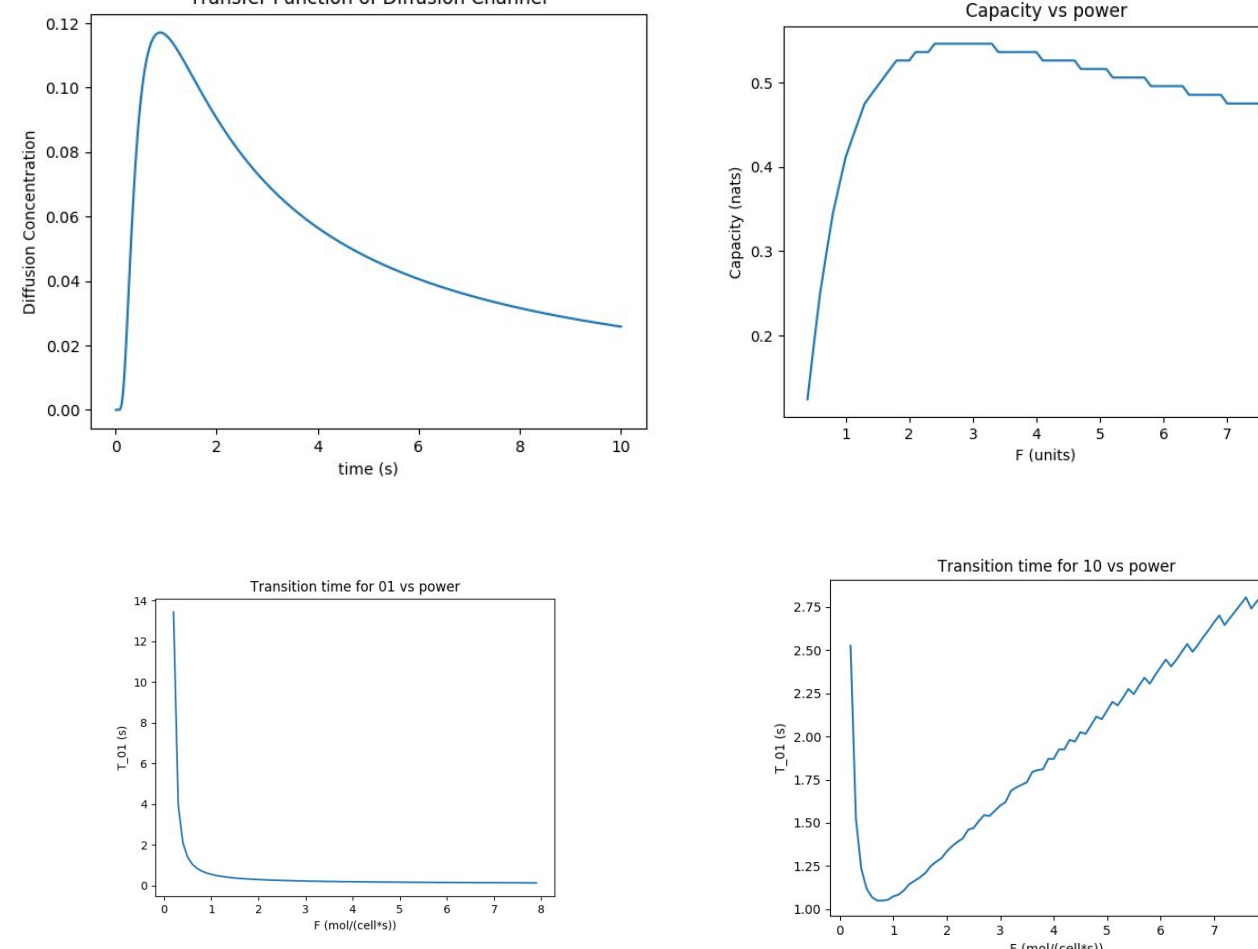
## 4. Analysis

### Diffusion Channel

The diffusion channel is a memory-channel with a transfer function given by Fick's second law:

$$\frac{\partial c(x, t)}{\partial t} = D \nabla^2 c(x, t) + r(x, t)$$

The input is encoded as low or high concentrations (1 bit). Transition time is shown for 0->1 and 1->0 signal sequences [2].

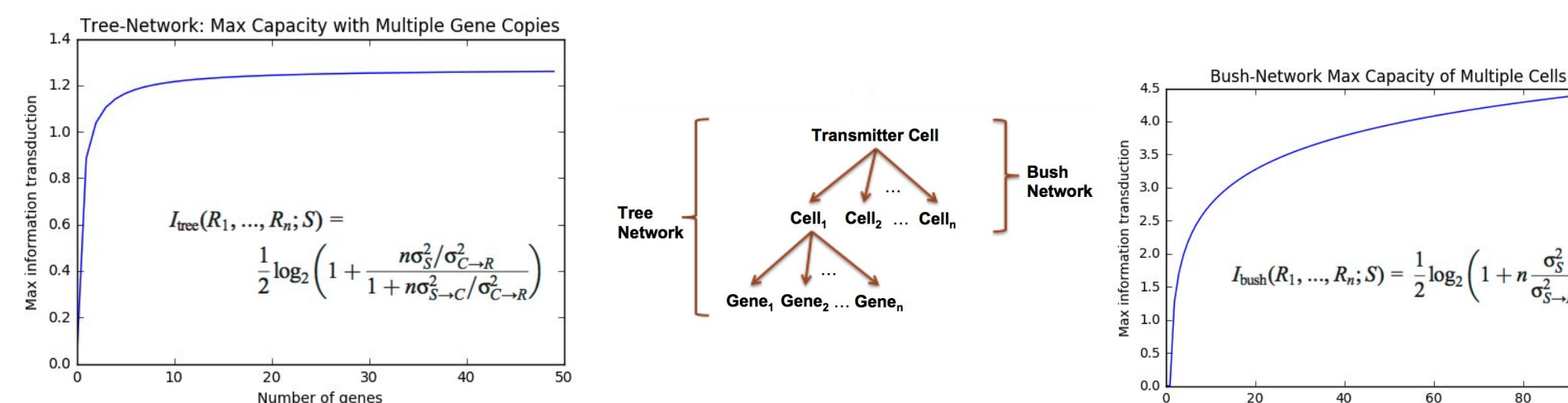


### Multiple receptors

$$0 \xrightleftharpoons[\tau\beta]{n\tau\alpha_{H/L}} 1 \xrightleftharpoons[2\tau\beta]{(n-1)\tau\alpha_{H/L}} 2 \dots \xrightleftharpoons[(k+1)\tau\beta]{(n-k)\tau\alpha_{H/L}} k+1 \dots \xrightleftharpoons[n\tau\beta]{\tau\alpha_{H/L}} n$$

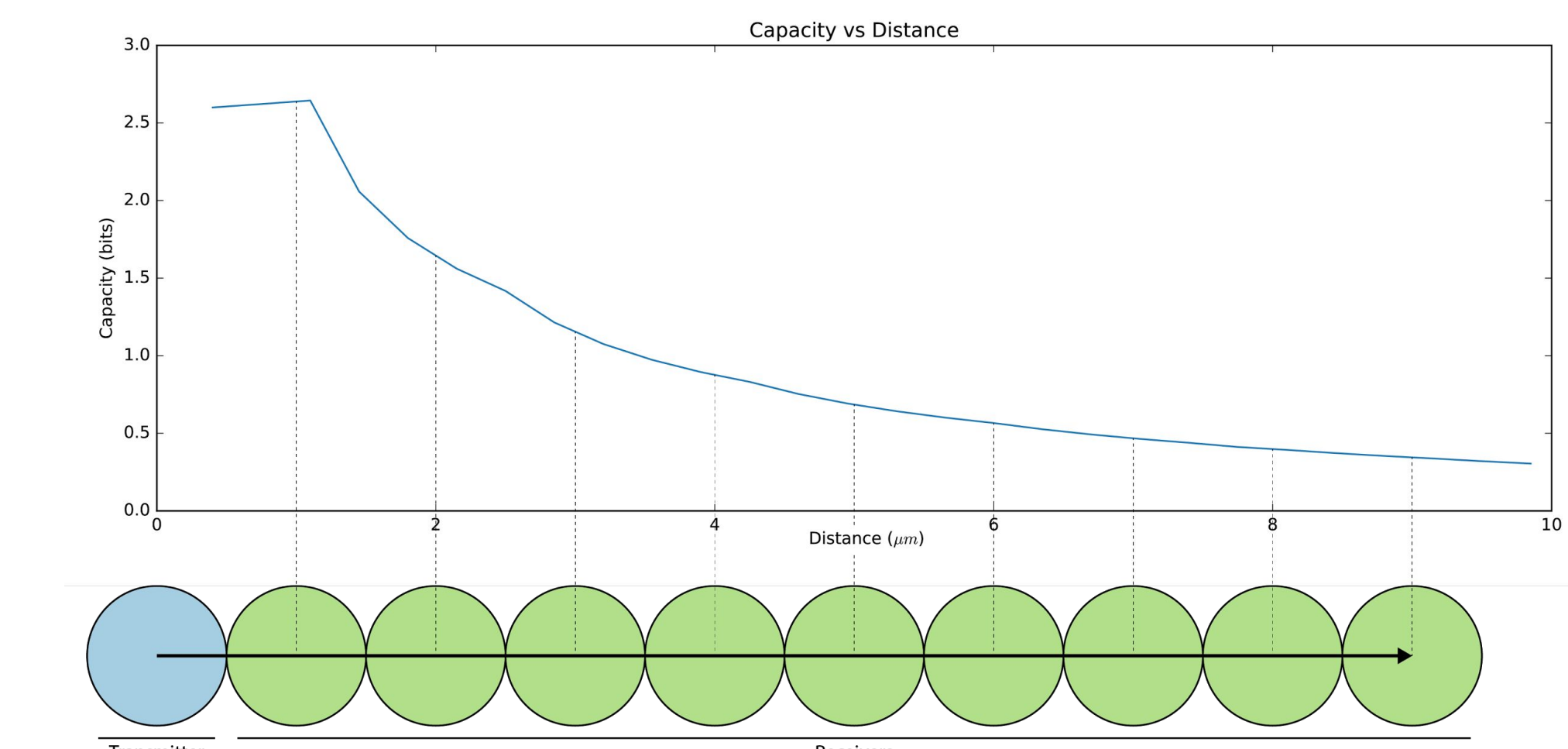
A cell with  $n$  receptors can be in any of  $n+1$  states, where state  $k$  means any  $k$  receptors are bound. The transition diagram is shown above. Capacity for receiving cells with  $n$  independent receptors is  $n$ -times greater than cells with single receptors, and IID input maximizes capacity (left) [1].

### Multiple receiver cells & gene copies

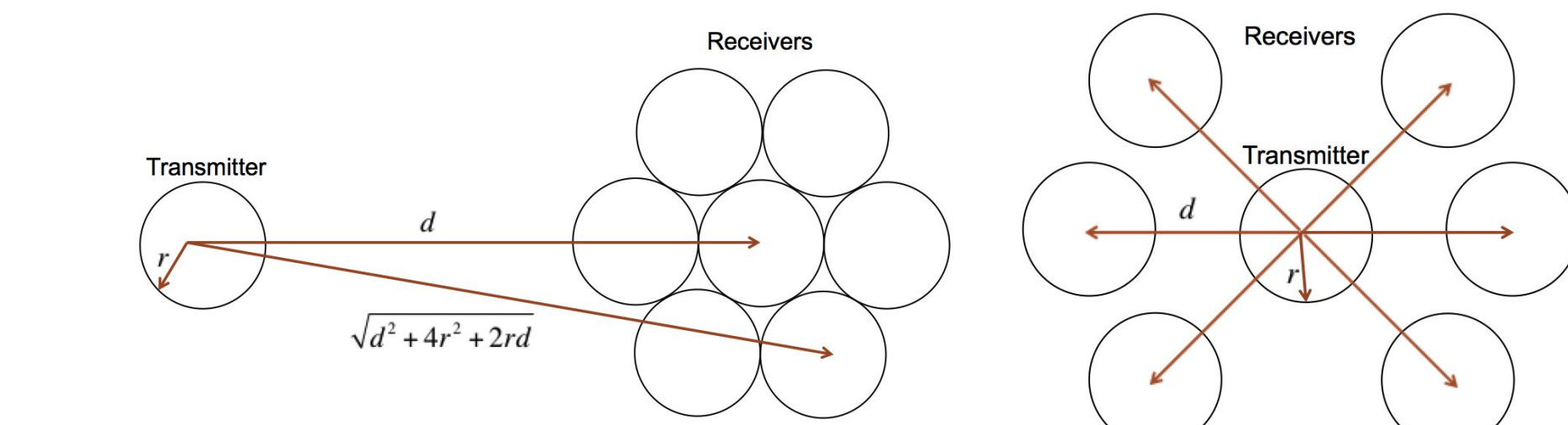


Information rate in signaling pathways such as TNF-a is limited by receiver receptors [3]. We model multiple receiving cells as a bush network to evaluate maximal improvements in capacity with multiple cells. Multiple copies of genes involved with intracellular signaling gives diminishing capacity gains because of the tree network structure.

## 5. Results



The capacity of receiver cells decreases with increasing distance from the transmitter. This is due to greater saturation of the memory-channel with greater distance.



We calculate the average information rate possible from one cell to a group of distant cells to be 0.88 bits/cell on average based on distance = 4 μm, and from one cell to a surrounding group of six cells to be 3.9 bits total using a fixed distance of 1 μm and the bush network model. We approximate the maximum possible information rate of a cell with a biologically relevant number of receptors (10,000 receptors for E. coli) to be 25,778 bits—this capacity is not limiting. Finally, we calculate the theoretical maximal information flux across a 1 mm colony of 150,000 E. coli, each cell receiving independent inputs from 6 neighbors, to be 271,103 bits. This synthesis provides a basis for designing simple cell-cell communication systems.

## 6. References

- [1] P. J. Thomas and A. W. Eckford. Shannon Capacity of Signal Transduction for Multiple Independent Receptors. *Proc. IEEE ISIT*. 2016-April.
- [2] A. Einolghozati, M. Sardari, A. Beirami, and F. Fekri. *t. Proc. IEEE ISIT*. 723-727, 2011-Aug.
- [3] Cheong, Raymond, et al. "Information transduction capacity of noisy biochemical signaling networks." *science* 334.6054 (2011): 354-358.