On Possible Indicators of Negative Selection in Germinal Centers

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What is a Germinal Center?

GC: Keystone of Adaptive Immunity



Figure: Young (2021)

GC: Small Scale Evolutionary Optimization Alg.



Figure: Young (2021)

- Body gets hit by pathogen.
- Anitgen gets presented to B-Cells in Germinal Center.
- Speedy evolution for Affinity (binding ability).
- Get high quality antibodies (yay!)

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GCs are Neat (For Theory)

- Fast & small scale evolution.
- Contrained system.
- Known objective function.

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Question: How *Does* **Selection Take Place?**

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Traditional population growth model:

New cell rate = (Cell "Fitness") * (Number of Cells)

Traditional population growth model:

New cell rate = (birth rate – death rate) * (Number of Cells)

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Normal models: Single Parameter

by Edelstein-Resnet (1900).

1.1 Continuous Growth Models

Single species models are of relevance to laboratory studies in particular but, in the real world, can reflect a telescoping of effects which influence the population dynamics. Let N(t) be the population of the species at time t, then the rate of change

$$\frac{dN}{dt} = \text{births} - \text{deaths} + \text{migration} , \qquad (1.1)$$

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is a conservation equation for the population. The form of the various terms on the right hand side of (1.1) necessitates modelling the situation that we are concerned with. The simplest model has no migration and the birth and death terms are proportional to N. That is

$$\frac{dN}{dt} = bN - dN \quad \Rightarrow \quad N(t) = N_0 e^{(b-d)t}$$

where b, d are positive constants and the initial population $N(0) = N_0$. Thus if b > d the population grows exponentially while if b < d it dies out. This

Figure: Murray (1993)

Birth and Death Are Not The Same

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Birth and Death Are Not The Same

Birth Selection/Positive Selection: High fitness = divide faster





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Birth and Death Are Not The Same

Death Selection/Negative Selection: High fitness = die slower





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Previous Work: The Moran Model

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Previous Work: The Moran Model

Definition

The Moran Model is a simple model of cancer development.

- Looks at how mutants strains spread through tissues.
- Perserves tissue structure.
- Sloooooow mutation rate.
- Evolves with simple birth-death rules.

Moran Model(s)



Key Metric

Definition

The **Partial Takeover Time** measures how long it takes for a novel mutant to take over X% of the tissue.

Fixation Times Distinguish Birth and Death



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Intuition:

- Birth Selection starts fast, slows down near total fixation.
- Death Selection starts slow, but fixates fast.
- Truncation skip slow regime for Birth Selection, changing the qualitative shape.

Can we find a convenient signature of Death Selection in Germinal Centers?

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We want a model that...

- 1 Recreates the basic dynamics of GCs, and
- 2 Includes are few parts as possible.

We want a model that will produce the strongest possible signal of selection, with a minimum of confounding variables.

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Tools

Differential Equations

Equations that model how a set of variables change over time.

Stochastic Processes

• The study of how random events accumulate over time.

Toy-Model Numerics

 Building and running simple simulations to check if our pencil-and-paper calculations make sense.

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Only two affinities, High (H) and Low (L) for maximum disparity.

h = fraction of population which is high affinity.

GC has three potential selection mechanisms:

- Birth Selection (r_B) ,
- Death Selection (r_D) ,
- Mutational Selection (??).

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Typical Selection Curve



- Mutations in GC occur 10⁶ times more often than default.
- GCs occasionally have Clonal Bursts, where one cell line repeatedly divides.
- When bursting, the line has a **notably lower** mutation rate than normal for the GC.
- This opens the possibility of a **tunable mutation rate**.

Pure Mutational Selection



Let $\ell = 1 - h$, then the average selection dynamics are given by:

$$\frac{dh}{dt} = \frac{2\rho_L \eta_L}{1 + n/N} \frac{\ell}{r_B h + \ell} + \frac{1 - 2\rho_H \eta_H}{1 + n/N} \frac{r_B h}{r_B h + \ell} - \frac{n/N}{1 + n/N} \frac{h/r_D}{h/r_D + \ell}$$

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Skew Hypothesis

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Skew Hypothesis



Figure: Tatsuya (2022)

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Skew Hypothesis

Since L and H have different levels of mutational activity, would they seperate?



Skew Hypothesis: Pilot Data



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How is Skew Measured?

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Definition

The *k*th **Moment** of *N* numbers is:

$$\langle x^k \rangle = \frac{x_1^k + x_2^k + \ldots + x_N^k}{N}$$

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Definition

The shape of a distribution with mean μ is determined by its **Central Moments**:

$$C(x^{k}) = \frac{(x_{1} - \mu)^{k} + (x_{2} - \mu)^{k} + \ldots + (x_{N} - \mu)^{k}}{N}$$

The second central moment measures the width of the distribution (the Variance).

Definition

The **Skew** of a distribution is given by:

Skew(x) =
$$\frac{C(x^3)}{C(x^2)^{3/2}}$$

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Calculation Outline: Mutation Dynamics

$$h_m = \frac{\text{number of H cells with m mutations}}{\text{number of cells}}$$

$$\ell_m = \frac{\text{number of L cells with m mutations}}{\text{number of cells}}$$

$$\langle m^k \rangle =$$
 k'th moment of mutation distribution
= $\sum_m (h_m + \ell_m) m^k$

Calculation Outline: Linear Structure

$$\partial_t \langle m_H^k \rangle = A_H \langle m_H^k \rangle + B_H \sum_{w=0}^k \binom{k}{w} \langle m_H^k \rangle + C_H \sum_{w=0}^k \binom{k}{w} \langle m_L^k \rangle$$
$$\partial_t \langle m_L^k \rangle = A_L \langle m_L^k \rangle + B_L \sum_{w=0}^k \binom{k}{w} \langle m_H^k \rangle + C_L \sum_{w=0}^k \binom{k}{w} \langle m_L^k \rangle$$

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Calculation Outline: Back of Envelope

$$\mathsf{Skew} = \frac{C(m^3)}{C(m^2)^{3/2}}$$

$$egin{aligned} C(m^3) &pprox \langle m^3
angle &= \mathbb{O}(t^3) \ C(m^2) &pprox \langle m^2
angle &= \mathbb{O}(t^2) \end{aligned}$$

Therefore Skew
$$ightarrow$$
 Constant

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NOPE



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Calculation Outline: Using Careful Methods

$$\mathsf{Skew} = \frac{C(m^3)}{C(m^2)^{3/2}}$$

where

$$C(m^3) = \langle m^3 \rangle - 3 \langle m^2 \rangle \langle m \rangle + 2 \langle m \rangle^3 = \mathbb{O}(t)$$

 $C(m^2) = \langle m^2 \rangle - 2 \langle m \rangle^2 = \mathbb{O}(t)$

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Mutations: The Truth



Mutations: The Truth

$$\mathsf{Skew} = \mathbb{O}(t^{-1/2}) \to 0$$



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Dynamical Footprint of Selection



Dynamical Footprint of Selection



Ancestry Hypothesis

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Ancestry Hypothesis



Figure: Tatsuya (2022)

Image: A mathematical states and a mathem

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$$F_{H} = \frac{\text{number of H cells with H ancestors}}{\text{number of H cells}}$$
$$F_{L} = \frac{\text{number of L cells with H ancestors}}{\text{number of L cells}}$$

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Ancestry Hypothesis: False

Notably, F_H and F_L only depend on the combined $r_B r_D$.



Ancestry Hypothesis: Alternates



Is there anything static that distiguishes r_D and r_B ?

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Fraction of cells which are High affinity =
$$\frac{1}{1 + r_B g(r_B r_D)}$$

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Static Signals of Selection Scheme?

Criteria	Status
Mutation Distribution Shape	Nope
Mutation Dynamics	Sure
Preferential Ancestry	Not Really
Overall Selection Strength	Yep

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If we combine two metrics, we can solve for fitness, e.g.

$$r_B = \left(\frac{L \to H \text{ mutation rate}}{H \to H \text{ mutation rate}}\right) \left(\frac{F_H}{1 - F_H}\right) \left(\frac{1 - h}{h}\right)$$

... but we want qualitative indicators, not just quantitative ones.

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Talk (will be) available at: ottinoloffler.com

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