# Is Sharing Caring?

Elucidating the Effects of the Presence of CRISPR-Cas Systems on Rates of Horizontal Gene Transfer Using Network Analysis



Siddharth Reed MolBiol 4C12 Thesis

> Golding Lab, Biology Department, McMaster University

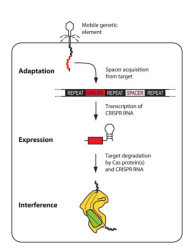
March 29, 2019

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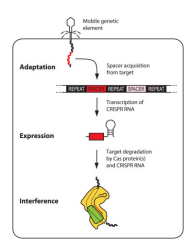
- 1. CRISPR-Cas systems
- 2. Horizontal Gene Transfer
- 3. Phylogenomic Networks
- 4. Do CRRISPR Systems Affect Horizontal Gene Transfer?
- 5. My Project

# CRISPR-Cas systems

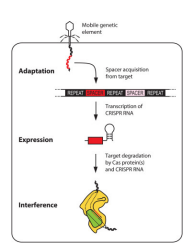
 Adaptive Bacterial Immune System



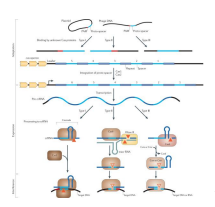
- Adaptive Bacterial Immune System
- Protects against foreign DNA



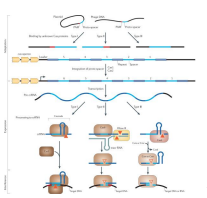
- Adaptive Bacterial Immune System
- Protects against foreign DNA
- Requires Cas proteins and CRISPR loci



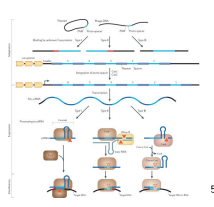
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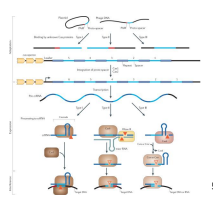
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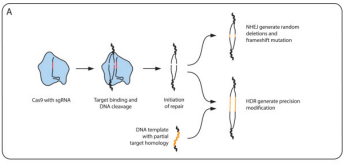


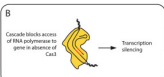
- 45% of bacteria have CRISPR loci  $(n = 6782)^2$
- 3 Main Types, multiple subtypes<sup>3</sup>
- CRISPR arrays represent unique life history of an organism
- 11% 28% are false or orphaned CRISPR loci<sup>4</sup>

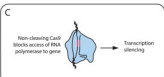


# Biotech Application

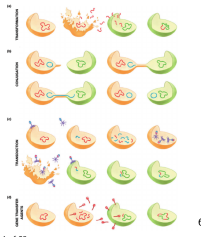
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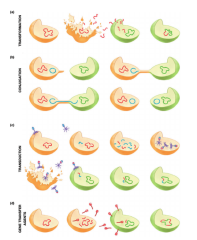




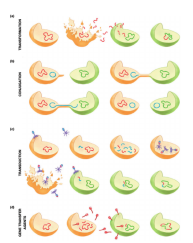


# Horizontal Gene Transfer

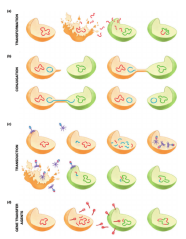




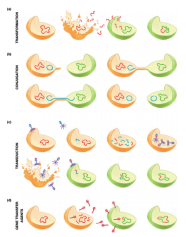
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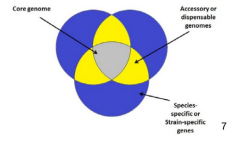
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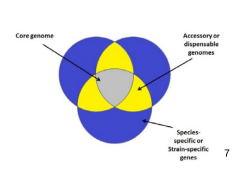
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- Transformation: Incorportaion of free-floating DNA into the genome<sup>6</sup>
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- CRISPR-Cas directly affects Transduction and Transformation<sup>6</sup>

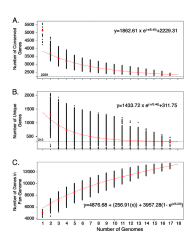
# Pan-Genomes

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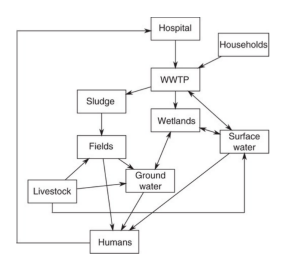
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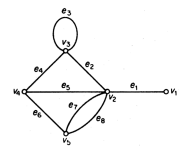
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- Metabolic costs
- Sequence compatibility

# Applications

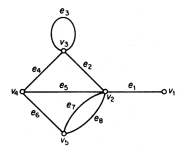
# **Applications**



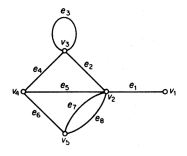
# Phylogenomic Networks



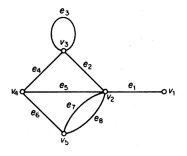
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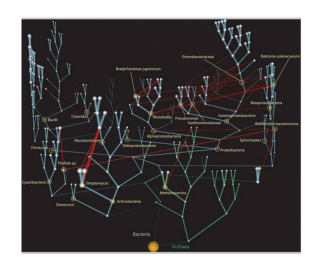
- Useful mathematical abstraction of real world system
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- Directed or Undirected Edges



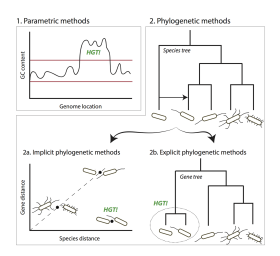
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- Directed or Undirected Edges
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# Prokaryotic "Net of Life"

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#### Construction



# Do CRRISPR Systems Affect Horizontal Gene Transfer?

# Yes

• Cost tradeoff factors:

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- CRISPR can enhance transduction-mediated HGT<sup>15</sup>

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  - Can see inhibitory effects of CRIPSR on HGT over short evolutionary time scales
  - Higher gene indel rates for CRISPR containing genera than non-CRISPR containing outgroups

# My Project

#### **Hypothesis**

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Bacterial strains or genera with known CRISPR systems will show no significant differences in network statistics compared to those strains or genera without known CRISPR systems.

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#### Alternative Hypothesis

Bacterial strains or genera with known CRISPR systems will show a significant difference in at least 1 network statistic compared to those strains or genera without known CRISPR systems.

# Objectives

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#### Within Network Comparisons

For genera with CRISPR conatining strains, compare the node statistics of CRIPSR-containing strain to non-CRISPR-containing strains.

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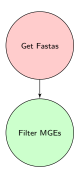
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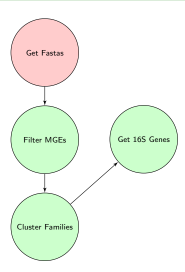
#### Gene Indel Rates vs. Network Statistics

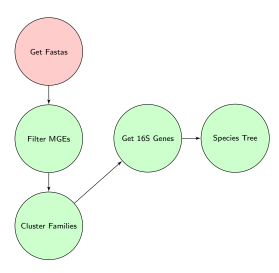
Compare gene InDel rates to node/network statistics for CRISPR-containing and non-CRISPR-containing strains/genera.

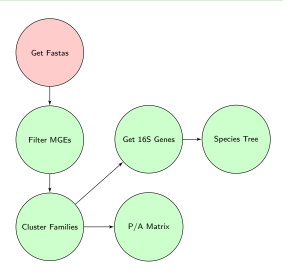


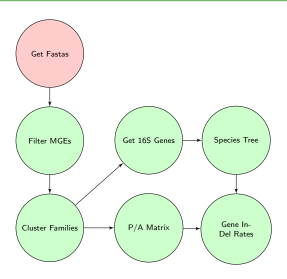


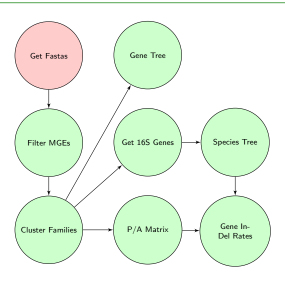


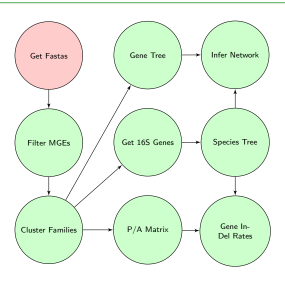


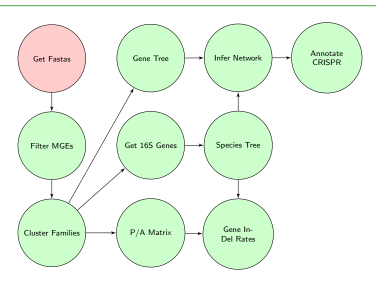


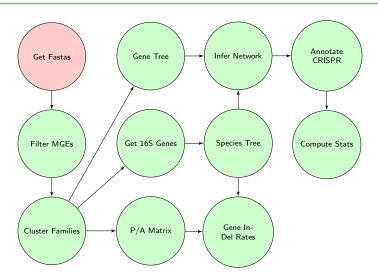


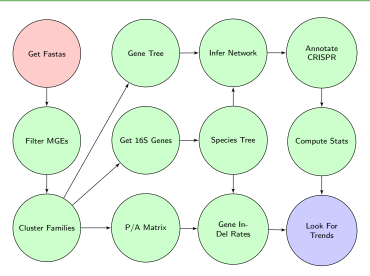












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- Network Modularity:  $Q = \frac{1}{2m} \sum_{uv}^{W} [W_{uv} \frac{k_u k_v}{2m}] \delta(u, v)$  where m is the total weight of alledges,  $k_u$  is the degree of u and  $\delta(u, v)$  is 1 if u and v both have or do not have CRISPR systems and 0 otherwise.  $Q \in [-1, 1]^{21}$

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- Taxonomic Mistakes: Inconsistencies in taxonomic labelling can result in ignored or misplaced OTUs.
- Multifurcation Error: Some species trees contained multifurcations, which were resolved randomly to generate a bifurcating tree. Estimating this error by examining variance over different resolutions is possible.

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- Continuous CRISPR activity: Labelling nodes by estimated CRISPR activity (array length, transciptomic data, etc.)
- Considering bacterial ecology and environments: Consider geographically close OTUs or differences between networks due to environmental factors

### **Thanks**

### Thank you to

- Dr. G. Brian Golding
- Dr. Ben Evans
- The Golding lab
  - Caitlin Simopoulos
  - Daniella Lato
  - Zachery Dickson
  - Sam Long
  - Geoge Long
  - Lucy Zhang
  - Brianne Laverty
  - Nicole Zhang
- Everyone here for listening

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