

Is Sharing Caring?

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



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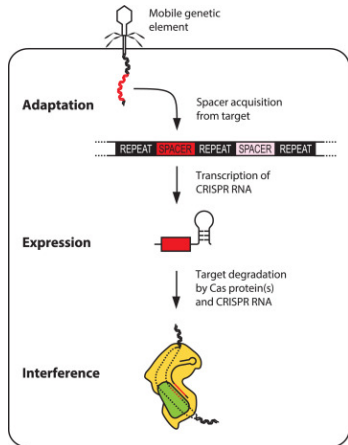
1. CRISPR-Cas systems
2. Horizontal Gene Transfer
3. Phylogenomic Networks
4. Do CRRISPR Systems Affect Horizontal Gene Transfer?
5. My Project

CRISPR-Cas systems

What Are They?

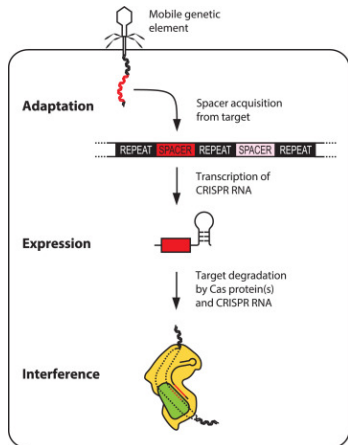
What Are They?

- Adaptive Bacterial Immune System



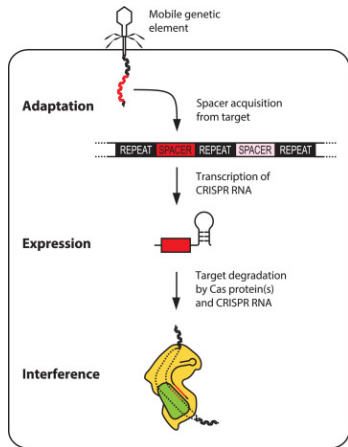
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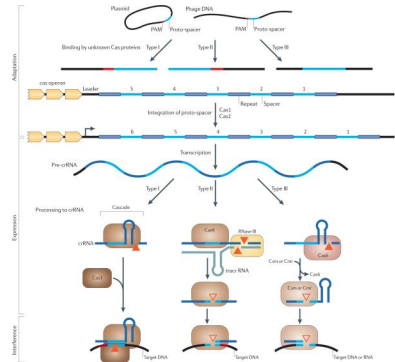
- Adaptive Bacterial Immune System
- Protects against foreign DNA
- Requires Cas proteins and CRISPR loci



Diversity & Ubiquity

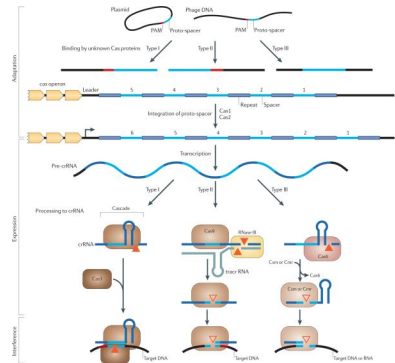
Diversity & Ubiquity

- 45% of bacteria have CRISPR loci ($n = 6782$)²



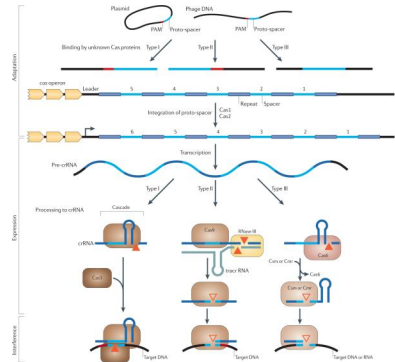
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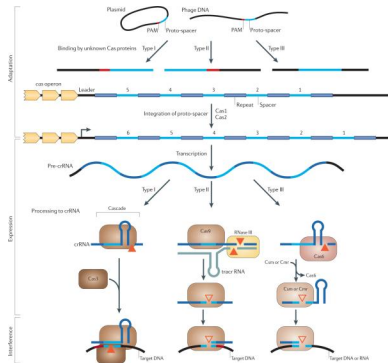
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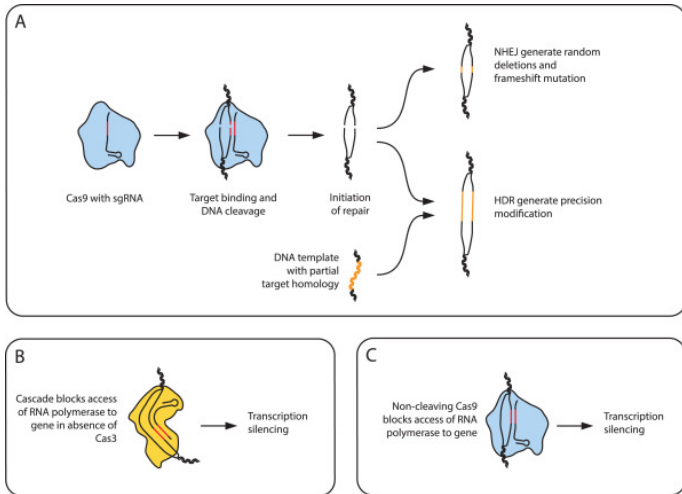
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- 11% – 28% are false or orphaned CRISPR loci⁴



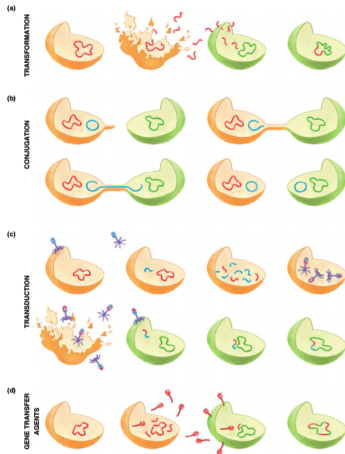
Biotech Application

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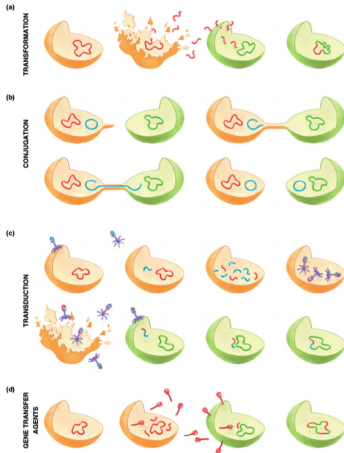


Horizontal Gene Transfer

Mechanisms

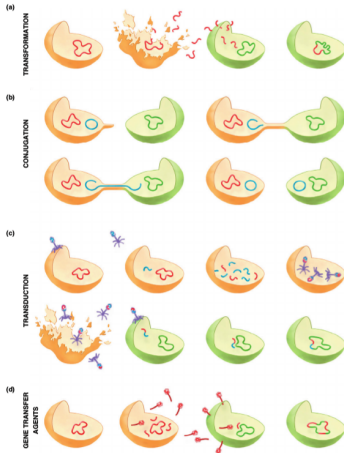


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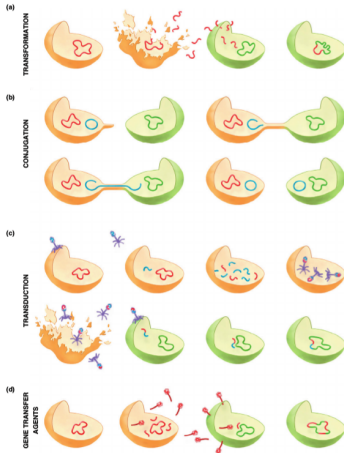
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Mechanisms



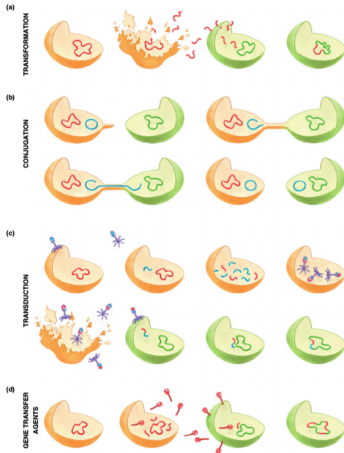
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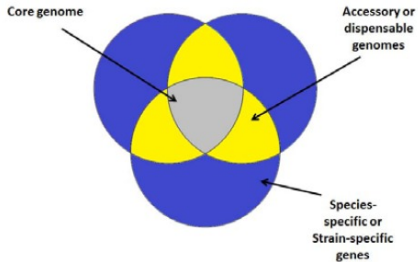
Mechanisms



- Conjugation: Transfer of DNA through cell-cell connections⁶
- Transformation: Incorporation of free-floating DNA into the genome⁶
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- **CRISPR-Cas directly affects Transduction and Transformation⁶**

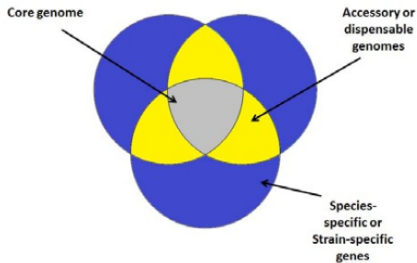
Pan-Genomes

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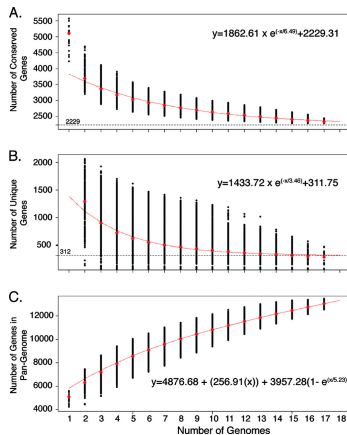


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Pan-Genomes



7



8

Rate Influencing Factors

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- Amount of exogenous DNA/cell density/phage density

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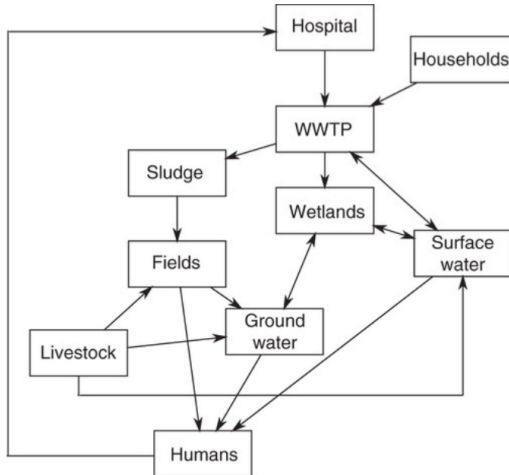
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Applications

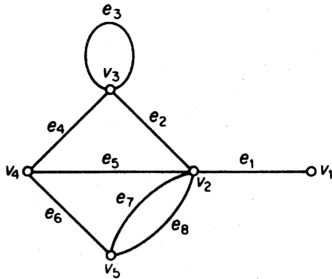
Applications



Phylogenomic Networks

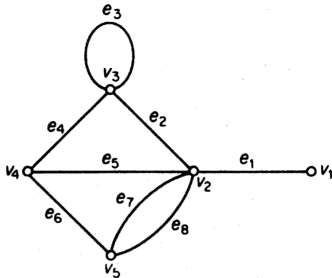
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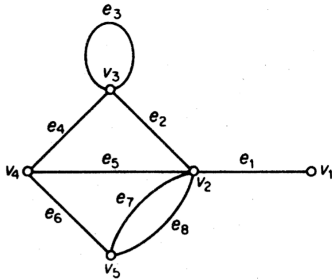
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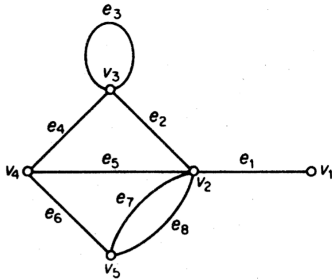
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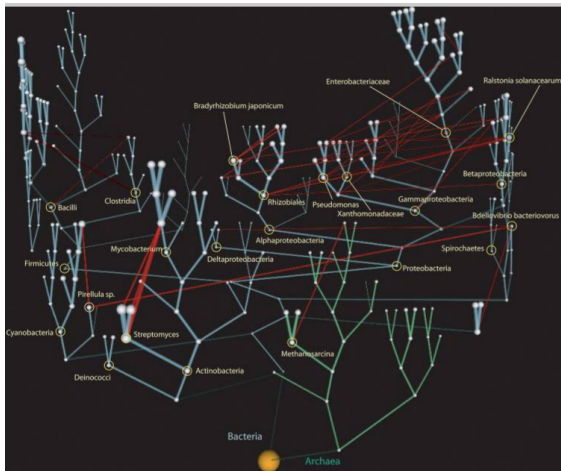
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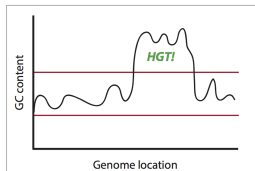
Prokaryotic “Net of Life”

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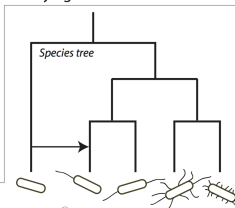


Construction

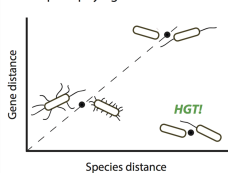
1. Parametric methods



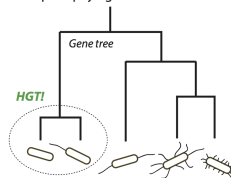
2. Phylogenetic methods



2a. Implicit phylogenetic methods



2b. Explicit phylogenetic methods



Do CRRISPR Systems Affect Horizontal Gene Transfer?

Yes

CRISPR Cost Complexity

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- Cost tradeoff factors:

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 - Metabolic maintenance¹

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 - Prophage abundance¹⁵

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- Selective CRISPR inactivation¹
- CRISPR can enhance transduction-mediated HGT¹⁵

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 - Assume all singletons arose from HGT
 - Used GC% to identify HGT
- Contradicted by a former thesis student
 - Can see inhibitory effects of CRISPR on HGT over short evolutionary time scales
 - Higher gene indel rates for CRISPR containing genera than non-CRISPR containing outgroups

My Project

Hypothesis

Null Hypothesis

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 containing strains, compare the node statistics of CRISPR-containing strain to non-CRISPR-containing strains.

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Between Network Comparisons

For genera with no CRISPR containing strains, compare the network statistics of mixed to non-CRISPR-containing networks.

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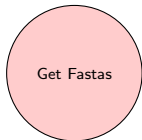
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For genera with no CRISPR containing strains, compare the network statistics of mixed to non-CRISPR-containing networks.

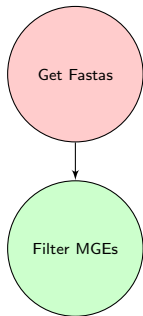
Gene Indel Rates vs. Network Statistics

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

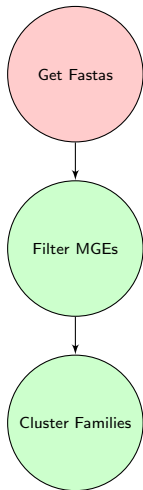
Workflow



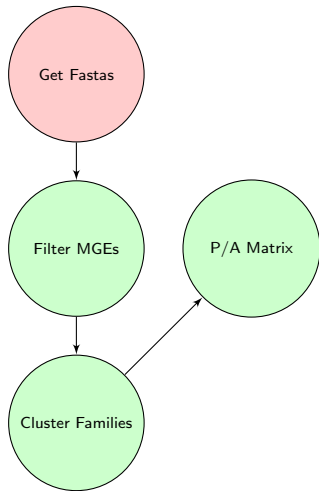
Workflow



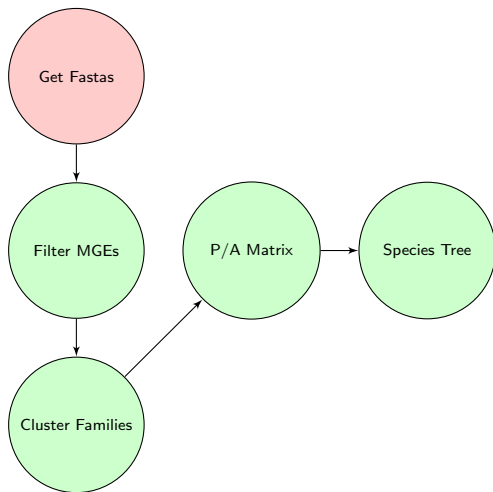
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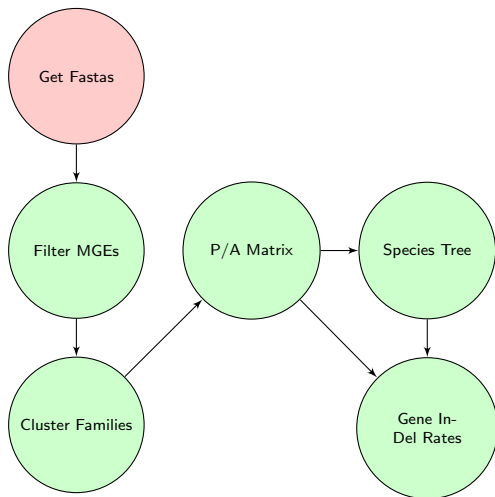
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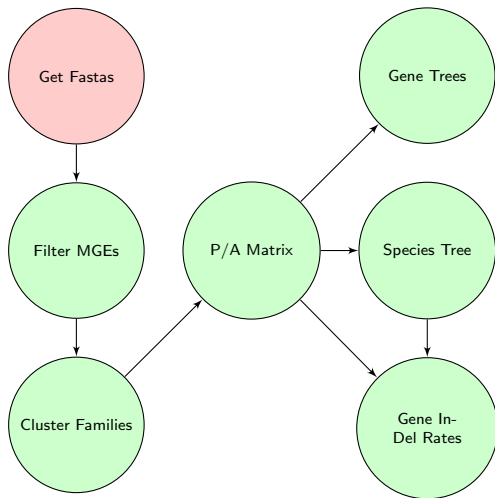
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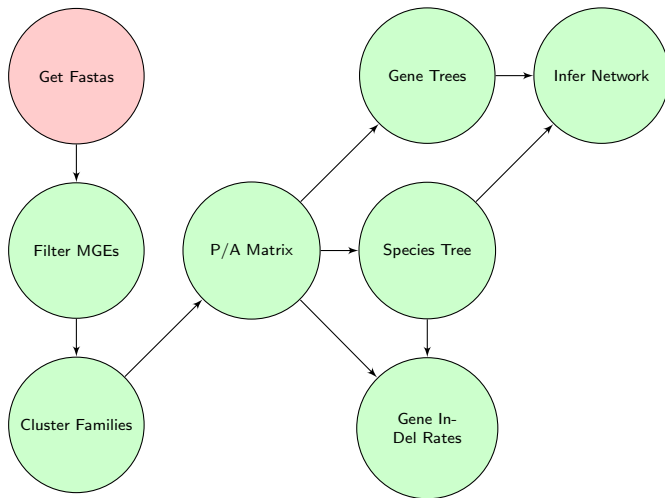
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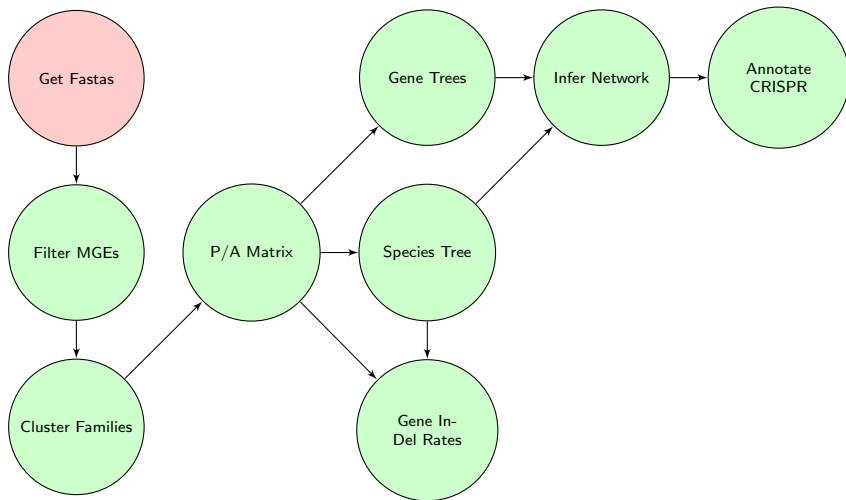
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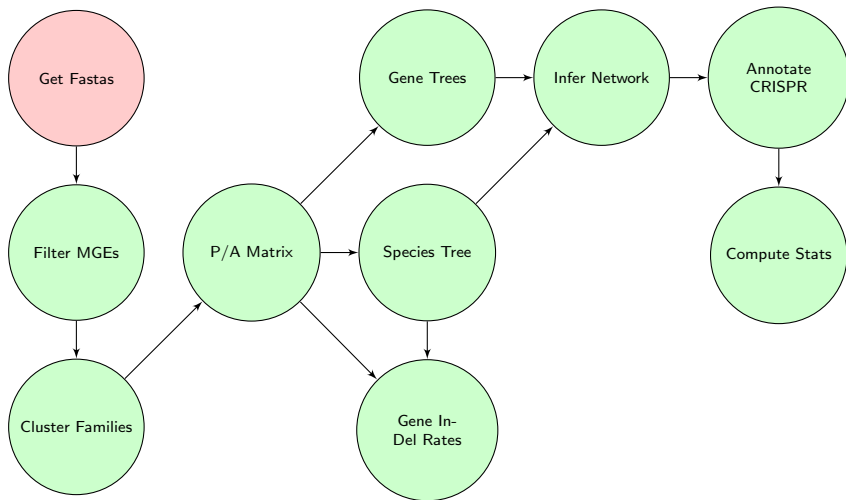
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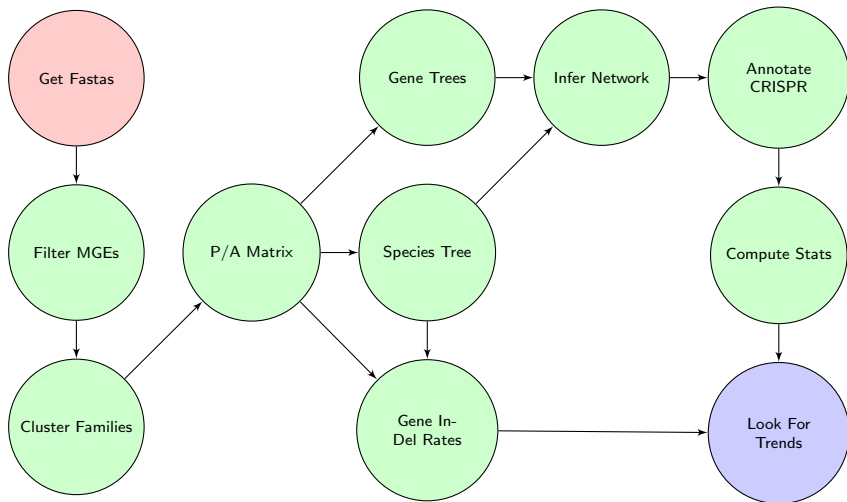
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

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- **Node Associativity:** $\frac{j(j+1)(\bar{k}-\mu_q)}{2E\sigma_q^2}$ where j is the excess degree of the node and \bar{k} is the average excess degree of the node's neighbors and μ_q and σ_q are the mean and standard variation of the excess degree distribution.
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Thanks




Thank you to

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



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-  GRissa, I. and Drevet, C. and Couvin, D. *CRISPRdb*. <http://crispr.i2bc.paris-saclay.fr/>. Online; accessed 22 October 2018. 2017.



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-  Quan Zhang and Yuzhen Ye. “Not all predicted CRISPR–Cas systems are equal: isolated cas genes and classes of CRISPR like elements”. In: *BMC Bioinformatics* 18.1 (Feb. 2017), p. 92. ISSN: 1471-2105. DOI: [10.1186/s12859-017-1512-4](https://doi.org/10.1186/s12859-017-1512-4). URL: <https://doi.org/10.1186/s12859-017-1512-4>.
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



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-  Ovidiu Popa and Tal Dagan. “Trends and barriers to lateral gene transfer in prokaryotes”. In: *Current Opinion in Microbiology* 14.5 (2011). Antimicrobials/Genomics, pp. 615–623. ISSN: 1369-5274. DOI: <https://doi.org/10.1016/j.mib.2011.07.027>. URL: <http://www.sciencedirect.com/science/article/pii/S1369527411001111>.
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

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