

Anotación de genomas bacterianos

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Anotación

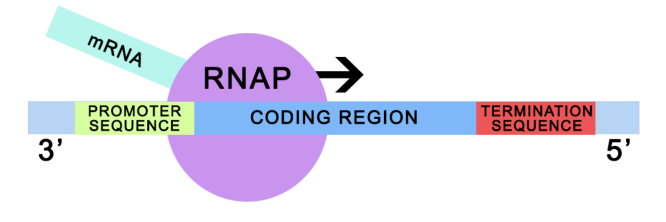
- Identificación de las propiedades funcionales de uno o más genes.
- La mayoría de los softwares de anotación se basan en bases de dato existentes (clase 2).
- Se realiza una comparación entre su secuencia y las presentes en la base de datos con funciones conocidas.



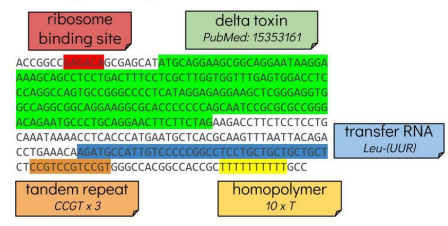


Identificación de secuencias codificantes





Adding biological info to sequences



Blast

1. The query sequence is broken into "words" that will act as seeds in alignments

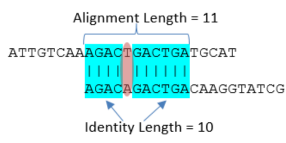
Query Words

2. BLAST searches for matches (or synonyms) in target entries in the database

Word match
Target

If a target entry has two or more matches to "words" from the query, the alignment is extended in both directions looking for additional similarity





Alignment Identity % =
$$\frac{Identity\ Length}{Alignment\ Length} = \frac{10}{11}$$

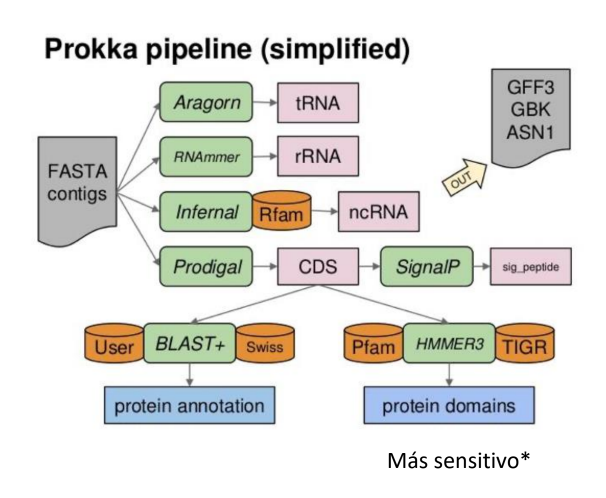
$$Query\ Identity\ \% = \frac{Identity\ Length}{Query\ Length} = \frac{10}{25}$$

Query Coverage
$$\% = \frac{Alignment\ Length}{Query\ Length} = \frac{11}{25}$$

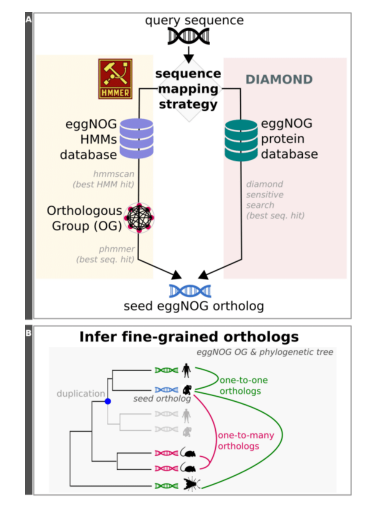
Subject Identity
$$\% = \frac{Identity\ Length}{Subject\ Length} = \frac{10}{21}$$

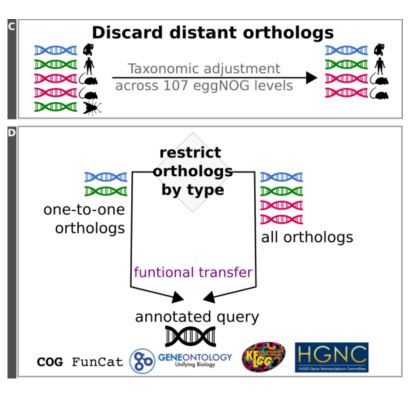
Subject Coverage
$$\% = \frac{Alignment\ Length}{Subject\ Length} = \frac{11}{21}$$

Prokka



eggNOG-mapper





Herramientas especializadas

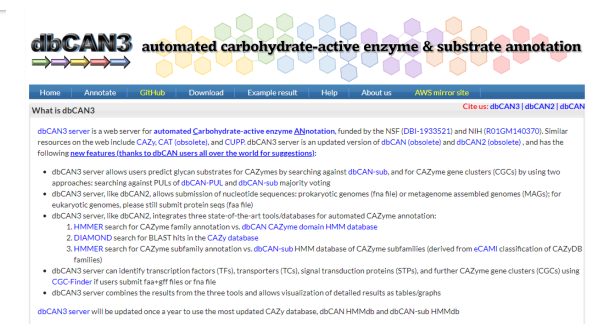
JOURNAL ARTICLE

NCycDB: a curated integrative database for fast and accurate metagenomic profiling of nitrogen cycling genes

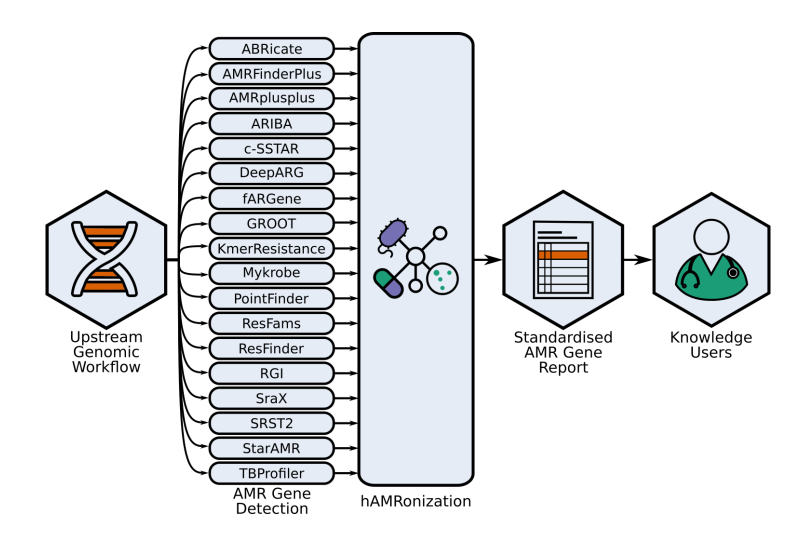
Qichao Tu 巫, Lu Lin 巫, Lei Cheng, Ye Deng, Zhili He

Bioinformatics, Volume 35, Issue 6, March 2019, Pages 1040–1048, https://doi.org/10.1093/bioinformatics/bty741

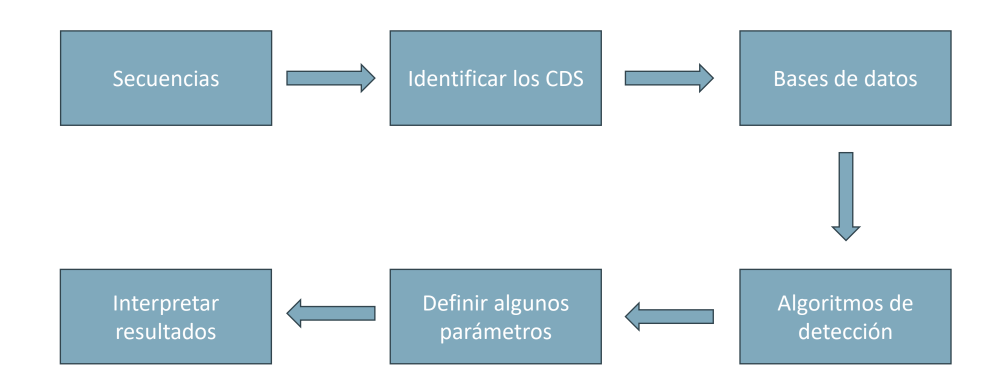
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Genes de resistencia



Qué necesitamos para un anotación?



Preguntas?