

Functional Annotation

Lesson 1

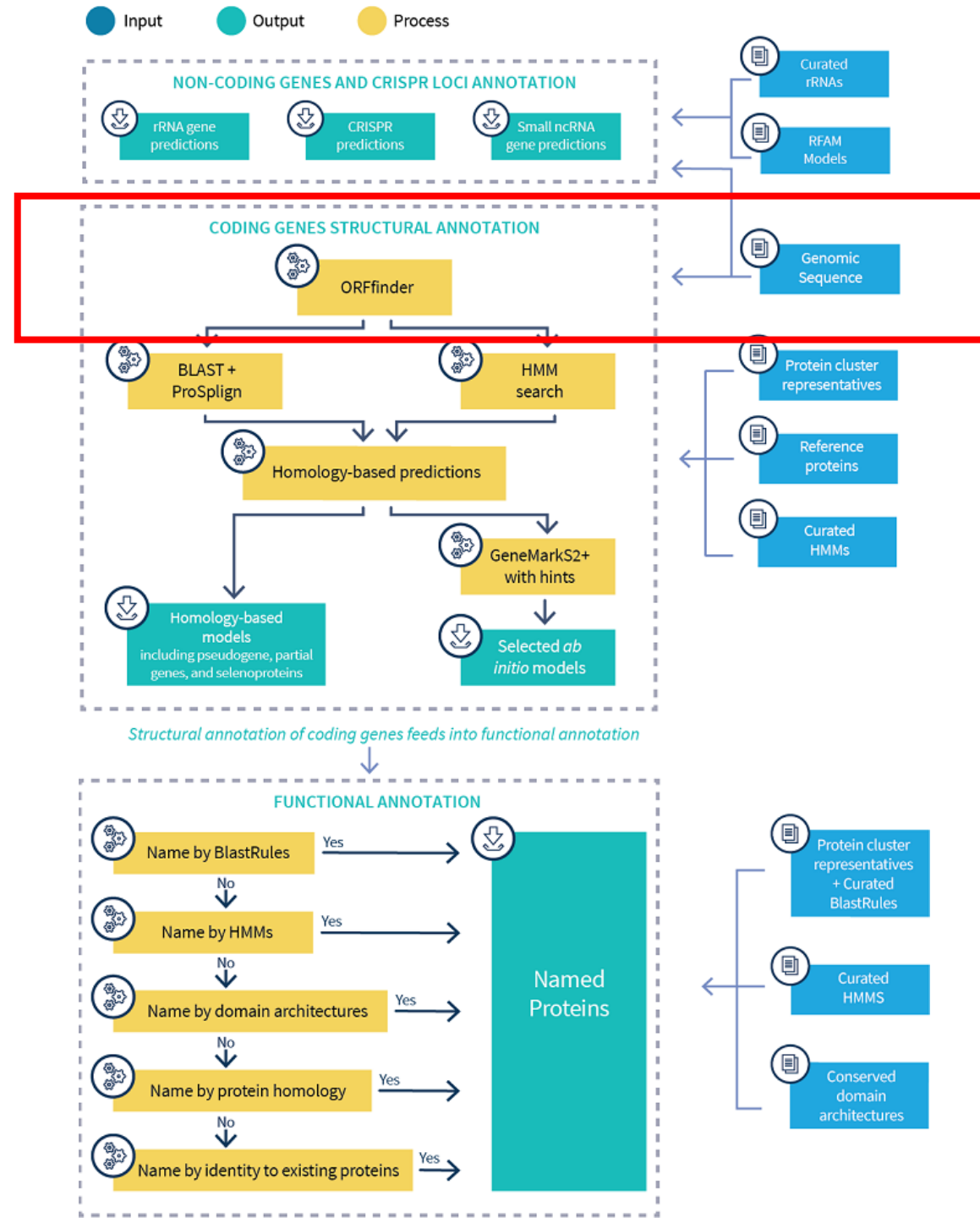
Predicting proteins

- Open reading frames in Bacteria and Archaea
 - Open reading frames in Eukaryotes
 - Translation tables

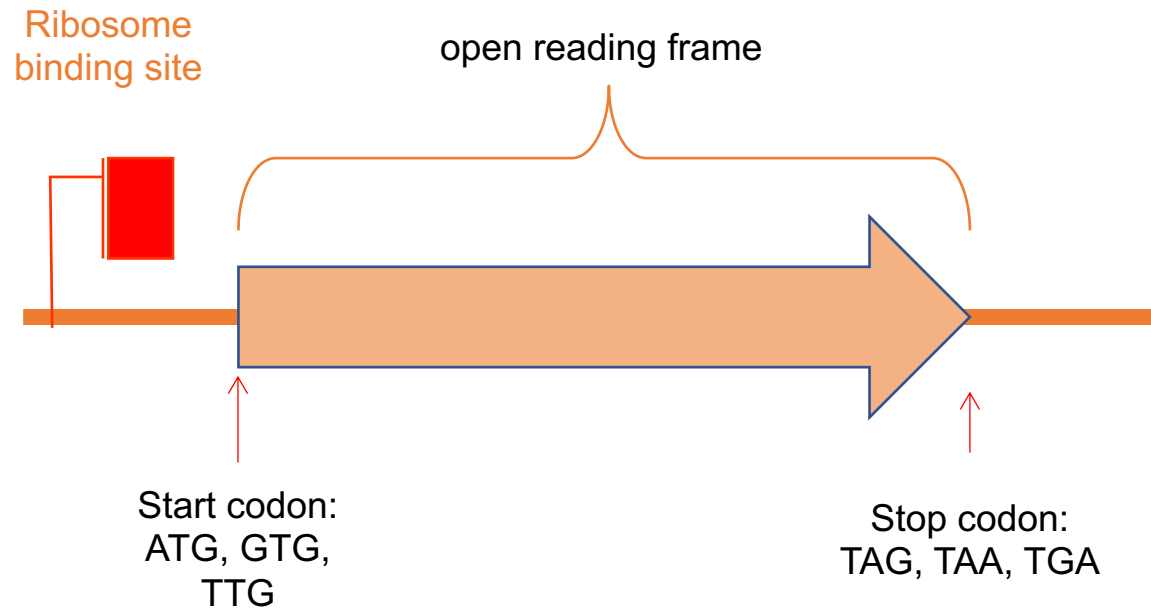


NCBI PGAP

Prokaryotic Genome Annotation Pipeline



Bacteria & Archaea Gene Prediction



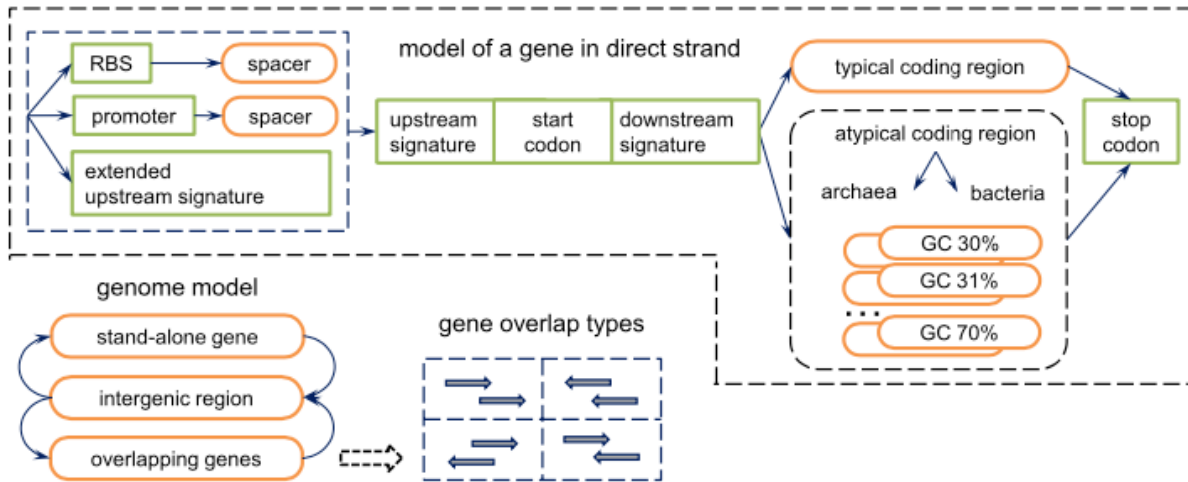
Gene model used by all *ab initio* gene finder

Traits a gene finder is looking for:

- Ribosome-binding site within a predicted distance from a start codon
- One of 3 start codons
- One of 3 stop codons
- No frame interruptions
 - Most gene finders do not handle point mutations that generate missense proteins or superfluous stop codons

Common Gene Prediction Tools

GeneMarkS-2 (2018)



Missed annotated genes (FN)

GeneMarkS	136	494	434	192	296	1552	
Glimmer3	66	678	1170	341	323	2578	
Prodigal	161	639	417	92	78	1387	
GeneMarkS-2	132	596	370	76	69	1243	
B	Bins (nt)	<150	150–300	300–600	600–900	>900	Total

Currently used by NCBI PGAP

Prodigal (2012)

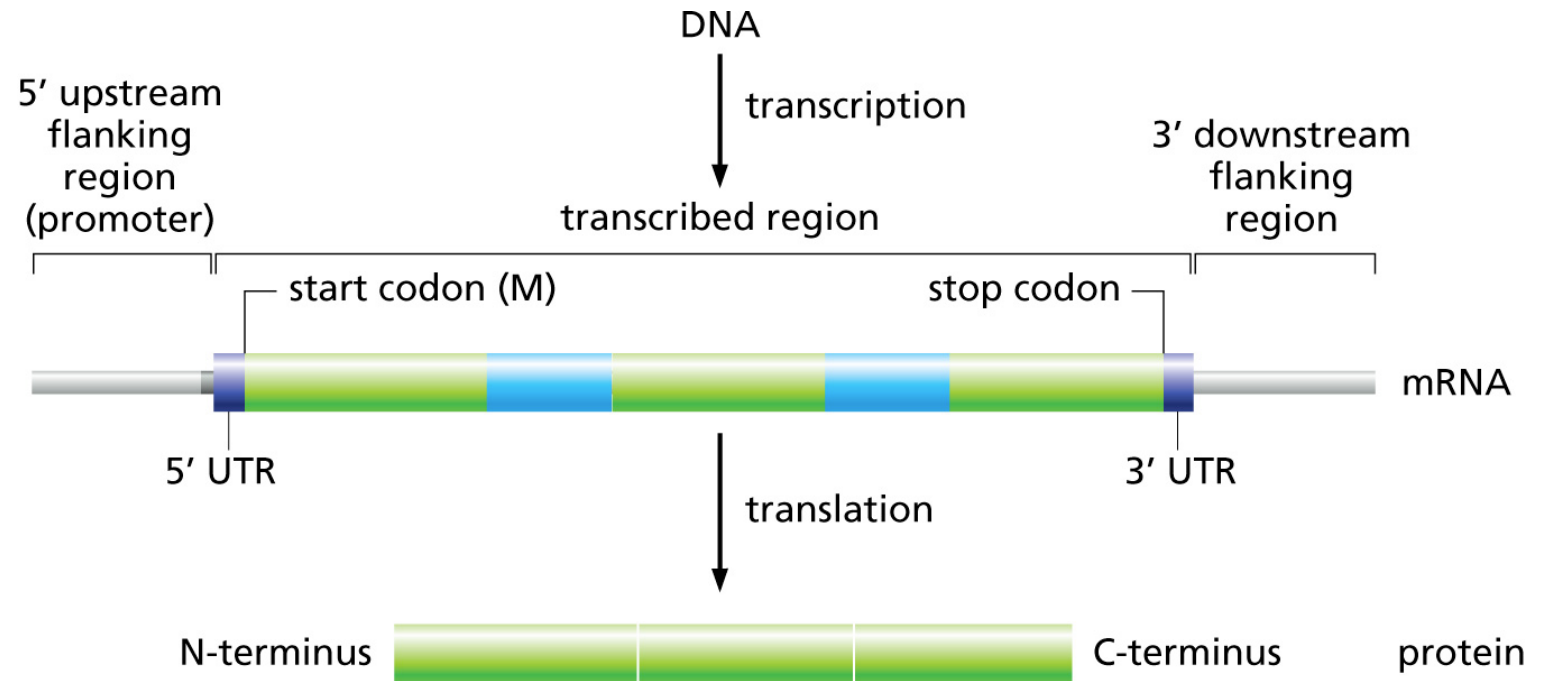
- Predicts protein-coding genes
- Handles draft genomes and metagenomes
- Runs quickly
- Runs unsupervised (unsupervised ML algorithm)
- Handles gaps, scaffolds, and partial genes
- Identifies translation initiation sites
- Outputs detailed summary statistics for each genome

Currently used by JGI

Eukaryotic Gene Prediction

Eukaryotic gene prediction requires consideration of:

- Introns
- Exons
- Untranslated regions (UTRs)
- Splice forms



Common Gene Prediction Tools

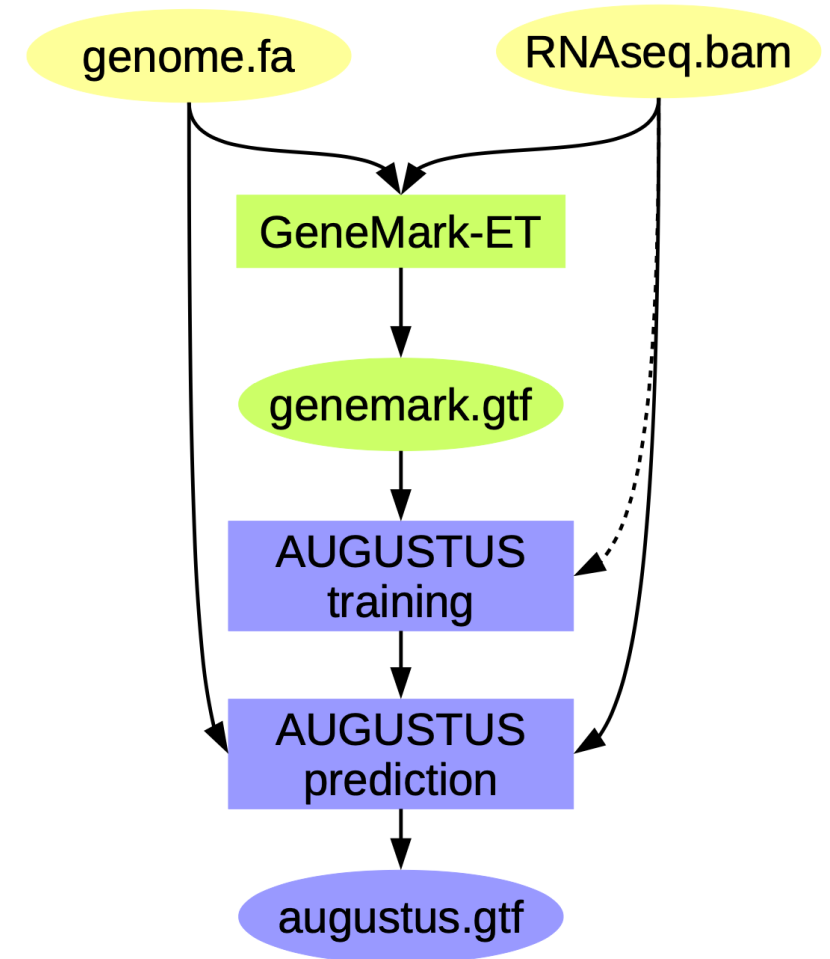
Braker2 (2018)

Preparation steps:

- Create a repeat library and mask your genome with RepeatMasker
- Trim and align RNA-seq to genome → trimgalore
- Align transcripts, ESTs, or long transcript to genome with splice aware aligner → Hisat2
- RUN Braker2

BRAKER1, a combination of GeneMark-ET and AUGUSTUS, that uses genomic and RNA-Seq data to automatically generate full gene structure annotations in novel genome

BRAKER2 is an extension of BRAKER1. BRAKER2 reaches high gene prediction accuracy even in the absence of the annotation of very closely related species and in the absence of RNA-Seq data.



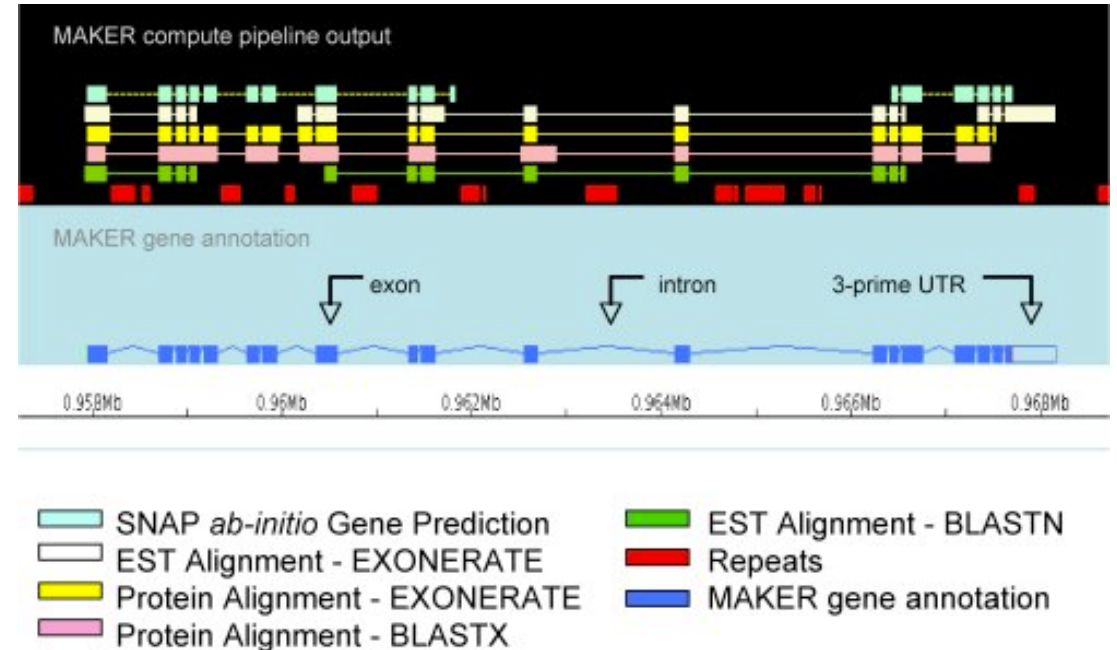
Common Gene Prediction Tools

MAKER (v3 2020)

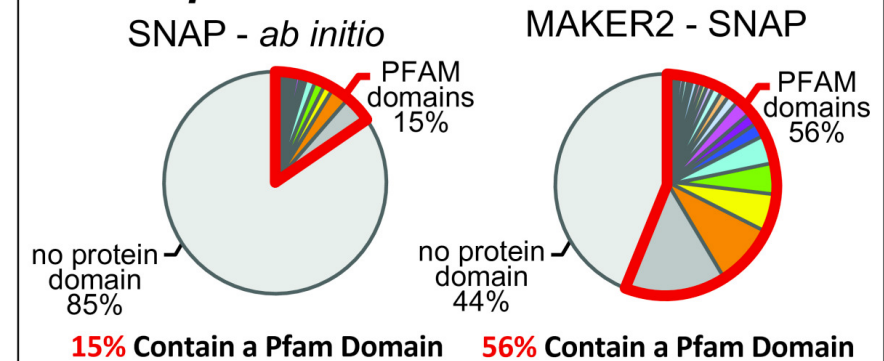
Steps:

- Mask out repeats
- Align EST to genome
- Align proteins to genome
- Produce ab initio predictions
- Synthesize into final predictions

MAKER is an annotation pipeline, not a gene predictor. MAKER does not predict genes, rather MAKER leverages existing software tools (some of which are gene predictors) and integrates their output to produce what MAKER finds to be the best possible gene model for a given location based on evidence alignments.



(b) *Linepithema humile*



Translation tables

- [1. The Standard Code](#)
- [2. The Vertebrate Mitochondrial Code](#)
- [3. The Yeast Mitochondrial Code](#)
- [4. The Mold, Protozoan, and Coelenterate Mitochondrial Code and the Mycoplasma/Spiroplasma Code](#)
- [5. The Invertebrate Mitochondrial Code](#)
- [6. The Ciliate, Dasycladacean and Hexamita Nuclear Code](#)
- [9. The Echinoderm and Flatworm Mitochondrial Code](#)
- [10. The Euplotid Nuclear Code](#)
- [11. The Bacterial, Archaeal and Plant Plastid Code](#)
- [12. The Alternative Yeast Nuclear Code](#)
- [13. The Ascidian Mitochondrial Code](#)
- [14. The Alternative Flatworm Mitochondrial Code](#)
- [16. Chlorophycean Mitochondrial Code](#)
- [21. Trematode Mitochondrial Code](#)
- [22. Scenedesmus obliquus Mitochondrial Code](#)
- [23. Thraustochytrium Mitochondrial Code](#)
- [24. Pterobranchia Mitochondrial Code](#)
- [25. Candidate Division SR1 and Gracilibacteria Code](#)
- [26. Pachysolen tannophilus Nuclear Code](#)
- [27. Karyorelict Nuclear Code](#)
- [28. Condyllostoma Nuclear Code](#)
- [29. Mesodinium Nuclear Code](#)
- [30. Peritrich Nuclear Code](#)
- [31. Blastocrithidia Nuclear Code](#)
- [33. Cephalodiscidae Mitochondrial UAA-Tyr Code](#)

33 translation tables – organism dependent

<https://www.ncbi.nlm.nih.gov/Taxonomy/Utils/wprintgc.cgi>