

BRAKER2: Incorporating Protein Homology Information into Gene Prediction with GeneMark-EP and AUGUSTUS

Katharina J. Hoff¹, Alexandre Lomsadze², Mario Stanke¹ and Mark Borodovsky²



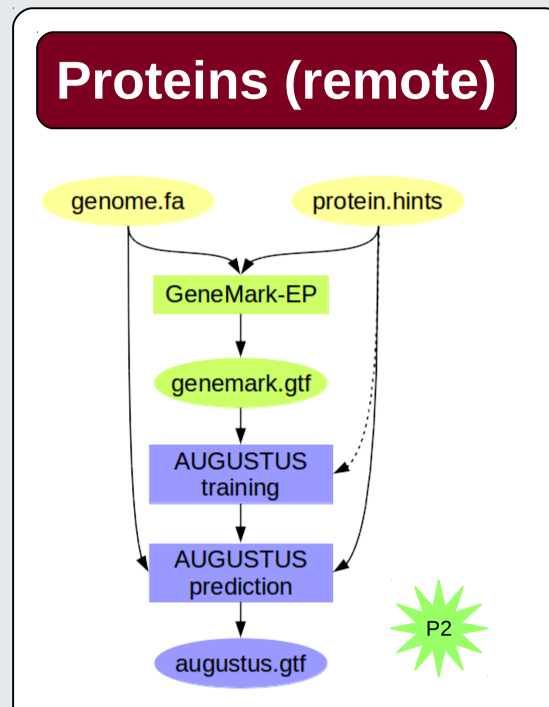
1) Institute for Mathematics and Computer Science, University of Greifswald, Greifswald, GERMANY 2) Joint Georgia Tech and Emory Wallace H Coulter Department of Biomedical Engineering, Georgia Institute of Technology, Atlanta, U.S.A. Contact: katharina.hoff@uni-greifswal.de

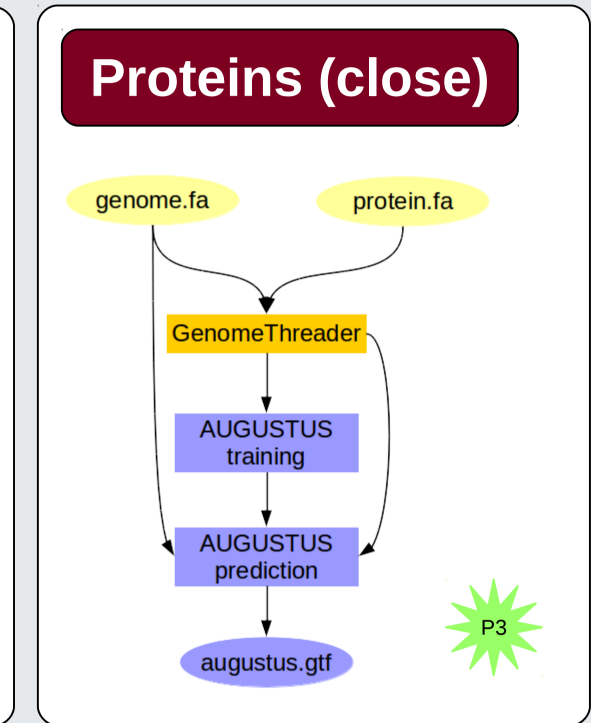
Abstract

The rapidly growing number of sequenced genomes requires fully automated methods for accurate gene structure annotation. With this goal in mind, we have developed BRAKER1 [1], a combination of GeneMark-ET [2] and AUGUSTUS [3], that uses genomic and RNAseq data to automatically generate full gene structure annotations in novel genomes. BRAKER2 is an extension of this earlier developed tool which allows for the integration of protein homology information.

If no annotation of a closely related species is available, a database of protein clusters (that must not contain proteins of very closely related species) can be used instead. In this case, BRAKER2 can execute self-training GeneMark-EP supported by protein alignments generated with ProSplign [4], trains AUGUSTUS on the basis of GeneMark-EP predictions and finds genes with protein homology information with AUGUSTUS. In presence of the proteome of a very closely related species, BRAKER2 can align the proteome with GenomeThreader [6], trains AUGUSTUS on the basis of spliced alignments and predicts genes with AUGUSTUS using protein homology information. BRAKER2 is available for download at http://bioinf.uni-greifswald.de/bioinf/braker and http://exon.gatech.edu.

RNA-Seq GeneMark-ET genemark.gtf AUGUSTUS training AUGUSTUS prediction





Conclusions

- BRAKER2 trained on protein data employing GeneMark-EP (P2) achieves high accuracy even in absence of close relatives in the protein database.
- Spliced alignments of proteins against genome with GenomeThreader loose sensivity with increasing distance between target and reference species.
- If the proteome of a very closely related species is available, BRAKER2 (P3) trained on spliced alignments from GenomeThreader is superior to BRAKER1 (P1) and BRAKER2 (P2) with GeneMark-EP.

Results

Table: Accuracy of BRAKER in *D. melanogaster***.** Expert trained AUGUSTUS (AUG) ab initio, self-trained GeneMark-ES (GM-ES) ab initio, BRAKER1 GeneMark-ET (GM-ET) and AUGUSTUS, BRAKER2 with inNog except for all *Drosophila* species GeneMark-EP (GM-EP) and AUGUSTUS, BRAKER2 with inNog with D. grimshawi, D. virilis and D. willistoni and without all other Drosohpila species (w_gvw), BRAKER2 with inNog with D. grimshawi, D. virilis, D. willistoni, D. pseudoobscura and D. ananassae and without all other *Drosohpila* species (w gvwpa).

| | | BRAKER1 (P1) | | | BRAKER2 (P2) | | BRAKER2 (P2) | | BRAKER2 (P2) | |
|-------------------|----------|--------------|--------|------|--------------|------|--------------|------|--------------|------|
| | ab inito | ab initio | RNAseq | | noDrosophila | | w_gvw | | w_gvwpa | |
| | AUG | GM-ES | GM-ET | AUG | GM-EP | AUG | GM-EP | AUG | GM-EP | AUG |
| Gene Sens. | 54.7 | 45.1 | 48.9 | 62.8 | 49.2 | 55.2 | 49.7 | 60.7 | 49.7 | 60.6 |
| Gene Spec. | 60.5 | 62.2 | 49.2 | 62.4 | 49.9 | 58.4 | 49.8 | 62.7 | 49.8 | 62.3 |
| Trans. Sens. | 37.7 | 31.4 | 34.0 | 44.7 | 34.3 | 39.7 | 34.5 | 43.1 | 34.6 | 42.9 |
| Trans. Spec. | 60.5 | 46.2 | 49.2 | 59.2 | 49.9 | 54.5 | 49.8 | 57.3 | 49.8 | 55.8 |
| Exon Sens. | 71.2 | 66.1 | 67.6 | 77.3 | 67.2 | 69.3 | 67.6 | 73.2 | 67.7 | 73.8 |
| Exon Spec. | 83.0 | 70.9 | 74.0 | 80.6 | 75.2 | 81.3 | 74.8 | 82.3 | 74.8 | 81.2 |

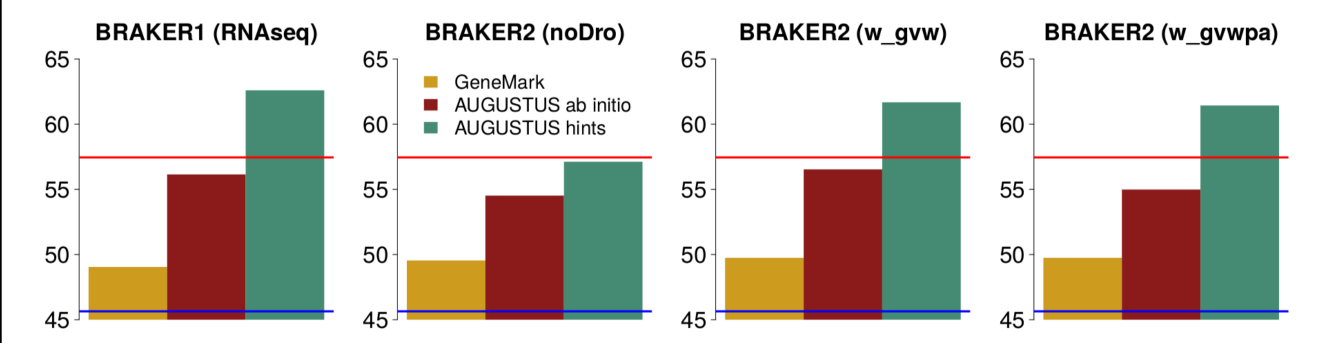
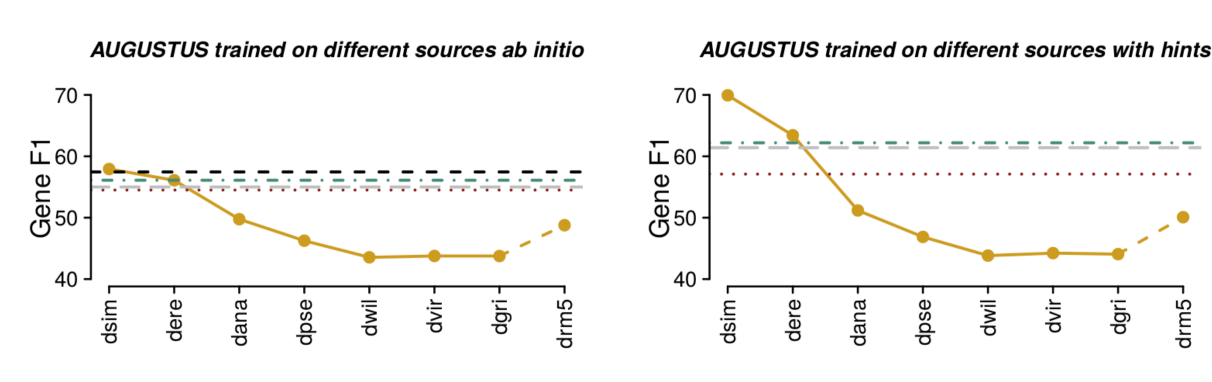


Figure 1: Gene F1 (P1/P2) of sensitivity and specificity from GeneMark ab initio predictions, AUGUSTUS ab initio predictions, **AUGUSTUS** predictions with hints; GeneMark-ES (blue), AUGUSTUS expert trained (red).

Figure 2: GenomeThreader trained AUGUSTUS (P3) vs. BRAKER1 with RNAseq (P1) & BRAKER2 trained on ProSplign alignments against inNog with GeneMark-EP (P2).



expert training BRAKER1 RNAseq (P1) BRAKER2 no Drosophila (P2) BRAKER2 w_gvwpa (P2) GenomeThreader spliced alignments (P3)

Data: A case study on *D. melanogaster*

- **Genome:** dmel-all-chromosome-r6.18.fasta
- RNAseq (P1): SRR023505, SRR023546, SRR023608, SRR026433, SRR027108 aligned against softmasked Genome with STAR [6]
- Reference annotation: dmel-no-analysis-r6.18.gff
- Proteomes for running BRAKER2 (P2): EggNog (inNog)n excluding
 - I. all *Drosophila* species (noDro)
 - II. *Drosophila* species except for
 - D. grimshawi, D. virilis, D. willistoni (w_gvw)
 - III. *Drosophila* species except for
 - D. grimshawi, D. virilis, D. willistoni,
 - D. pseudoobscura, D. ananassae (w_gvwpa) aligned with ProSplign [4].

dpse

Image: S. König, L. Romoth, M. Stanke (2018)

Comparative Genome Annotation

1.0

- Proteomes for BRAKER2 with spliced Alignments of close relatives (P3): Flybase proteomes of

dsim D. simulans dsim dere dere D. erecta danaD. ananassae dpse D. pseudodana obscura dwil D. willistoni /er 0.8 dvir D. virilis dgri D.grimshawi 0.6 0.4 Average Mutations per Genomic Site drm5dana, dpse,

aligned with GenomeThreader.

dwil, dvir, dgri

References

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