

# Comprehensive Evaluation of CAZyme Prediction Tools in Fungal and Bacterial Species

## Introduction

Carbohydrate Active enZymes (CAZymes) are pivotal in pathogen recognition, signalling, structure and energy metabolism. CAZy is the most comprehensive CAZyme database, cataloguing CAZymes into sequence-based CAZy families [1]. The CAZyme prediction tools **dbCAN** [2], **CUPP** [3] and **eCAMI** [4] annotate CAZymes with CAZy families. However, these tools have not been independently evaluated on a common high-quality dataset. Additionally, previous evaluations did not evaluate the **binary classification** of CAZymes/non-CAZymes, and the **multilabel classification** of CAZymes to multiple CAZy families.

## Method

The bioinformatic pipeline **pyrewton** was developed for this independent evaluation (Fig.1).

**GitHub:** <https://github.com/HobnobMancer/pyrewton>

The ground truths were retrieved using **cazy\_webscraper**.

**GitHub:** [https://github.com/HobnobMancer/cazy\\_webscraper](https://github.com/HobnobMancer/cazy_webscraper)

**70 genomic assemblies:**

- 40 Bacteria
- 16 Fungal
- 11 Yeast
- 3 Eukaryotes

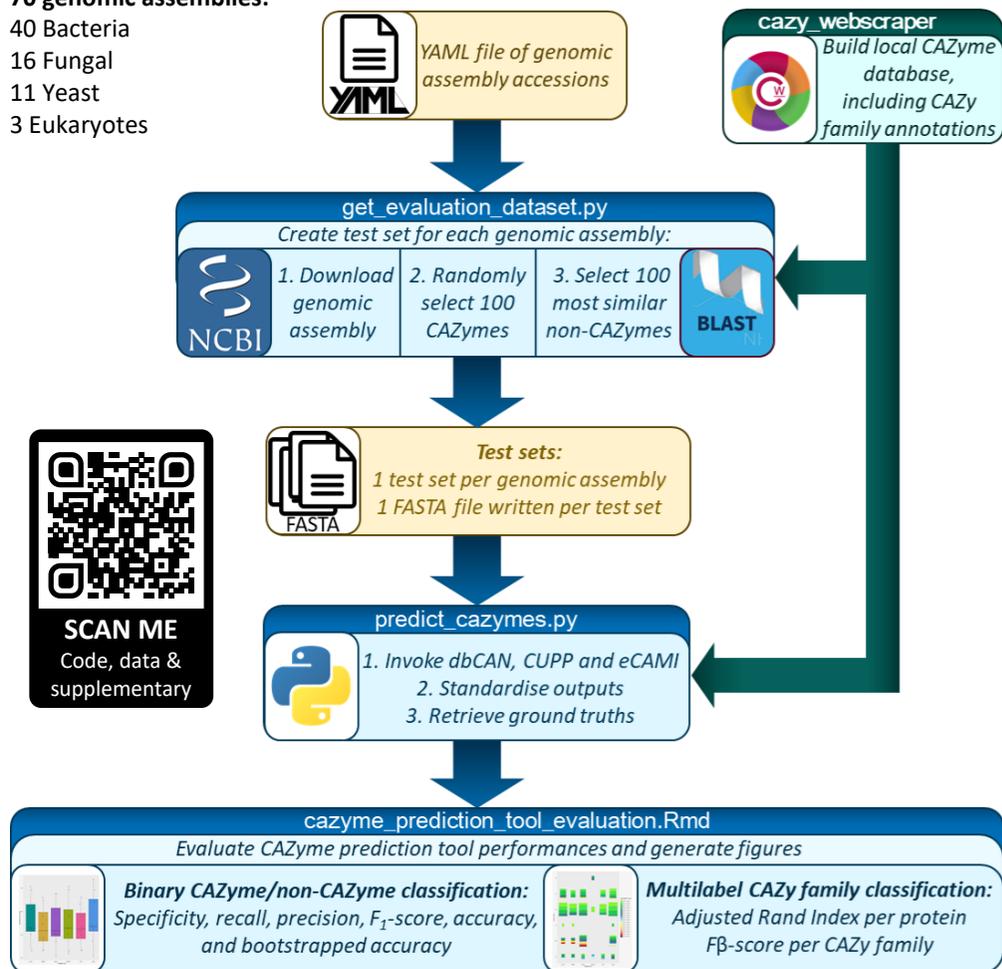


Fig.1 Schematic of the bioinformatic pipeline **pyrewton** for evaluating CAZyme prediction tools

## Results

### Binary CAZyme/non-CAZymes classification evaluation

dbCAN invokes the function prediction tools HMMER, Hotpep and DIAMOND. All prediction tools showed a low probability of misidentifying non-CAZymes as CAZymes, but also showed a tendency to miss identify a small proportion of CAZymes as non-CAZymes (Fig.2).

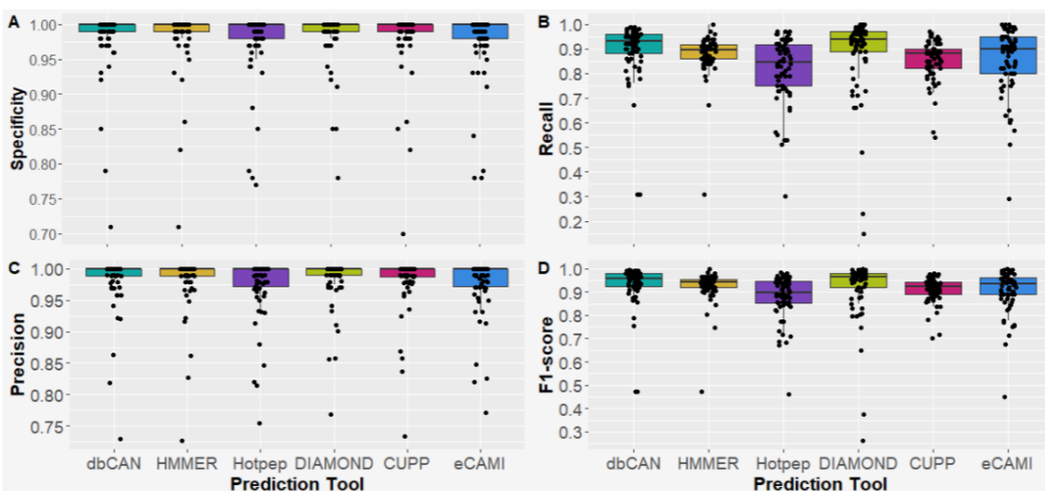


Fig.2 Evaluation of CAZyme/non-CAZyme differentiation performance.

One-dimensional scatterplots overlaying boxplots for [A] specificity, [B] recall (sensitivity), [C] precision and [D] F1-score.

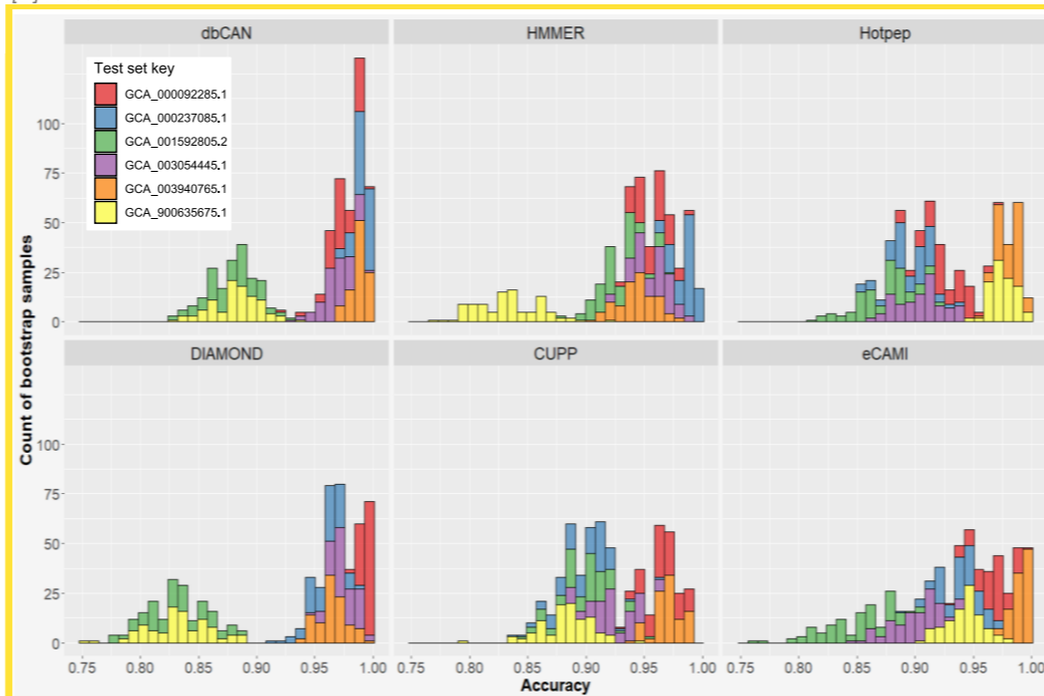


Fig.3 Stacked histograms of the range of performances of CAZyme prediction tools

6 test sets (identified by their genomic accessions) were randomly selected. The predictions for each prediction tool were bootstrap resampled 100 for each test set, and the accuracy for each bootstrap sample calculated.

### Multilabel CAZy family classification evaluation

Multilabel classification arises from the ability of a CAZyme to be assigned multiple CAZy families. The Adjusted Rand Index (ARI) was calculated per protein (Fig.4[A]). The  $F\beta$ -score ( $\beta=1$ ) was calculated for each CAZy family, true negative non-CAZyme predictions were excluded (Fig.4[B]).

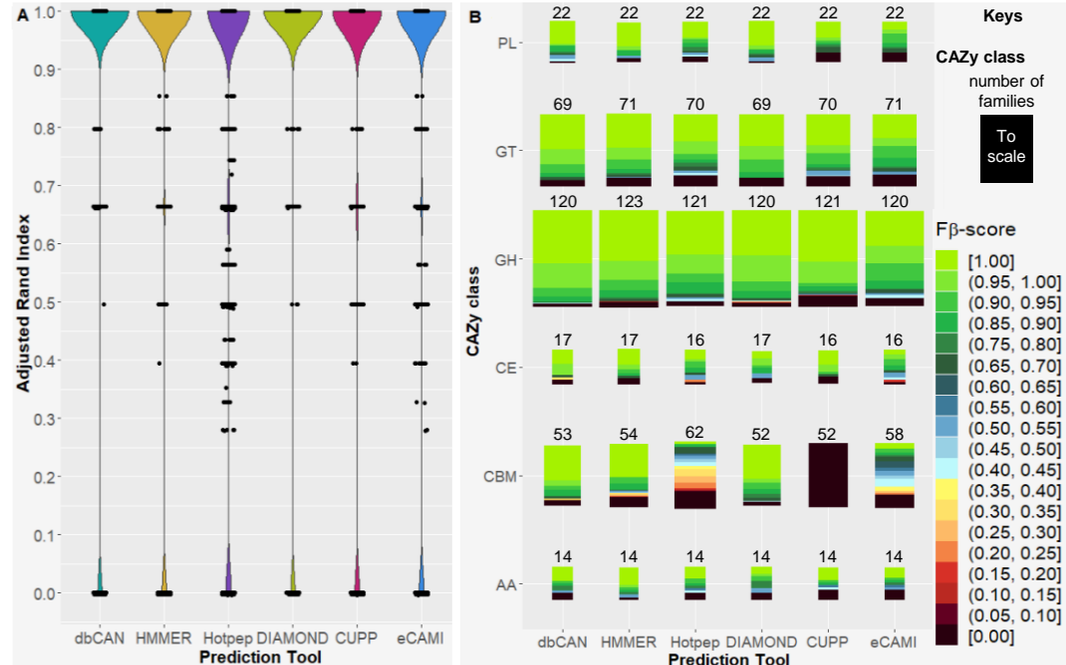


Fig.4 Evaluation of CAZy family multilabel classification

[A] Adjusted Rand Index per protein sequence. [B] Proportional area plot of CAZy classes sized by the number of families analysed, and coloured by the proportion of CAZy family  $F\beta$ -scores within each range of the scale, ( $\beta=1$ ).

## Conclusions

- Created a bioinformatic pipeline for reproducible evaluation of CAZyme prediction tools
- Benchmarked dbCAN, CUPP and eCAMI against a high quality test set
- Evaluated the binary and multilabel classification of CAZymes
- Statistically evaluated the expected range of performance
- dbCAN performed most strongly overall, and Hotpep (a component of dbCAN) was the weakest
- Better dbCAN performance may be achieved by replacing Hotpep with CUPP and/or eCAMI

## References

- Lombard, V. *et al.* (2014) 'The carbohydrate-active enzymes database (CAZy) in 2013', *Nucleic Acids Research*, 42, pp.D490–D495
- Zhang *et al.* (2018) 'dbCAN2: a meta server for automated carbohydrate-active enzyme annotation', *Nucleic Acids Research*, 46, W1, pp. W95–W101
- Barrett, K., Lange, L. (2019) 'Peptide-based functional annotation of carbohydrate active enzymes by conserved unique peptide patterns (CUPP)', *Biotechnology for biofuels*, 12, 102
- Xu *et al.* (2020) 'eCAMI: simultaneous classification and motif identification for enzyme annotation', *Bioinformatics*, 36, 7, pp.2068–2075

## Acknowledgements

We would like to thank the EASTBIO Doctoral Training Partnership (BBSCR) for funding our research.