

Decoding the epitranscriptomic code in *Plasmodium falciparum*

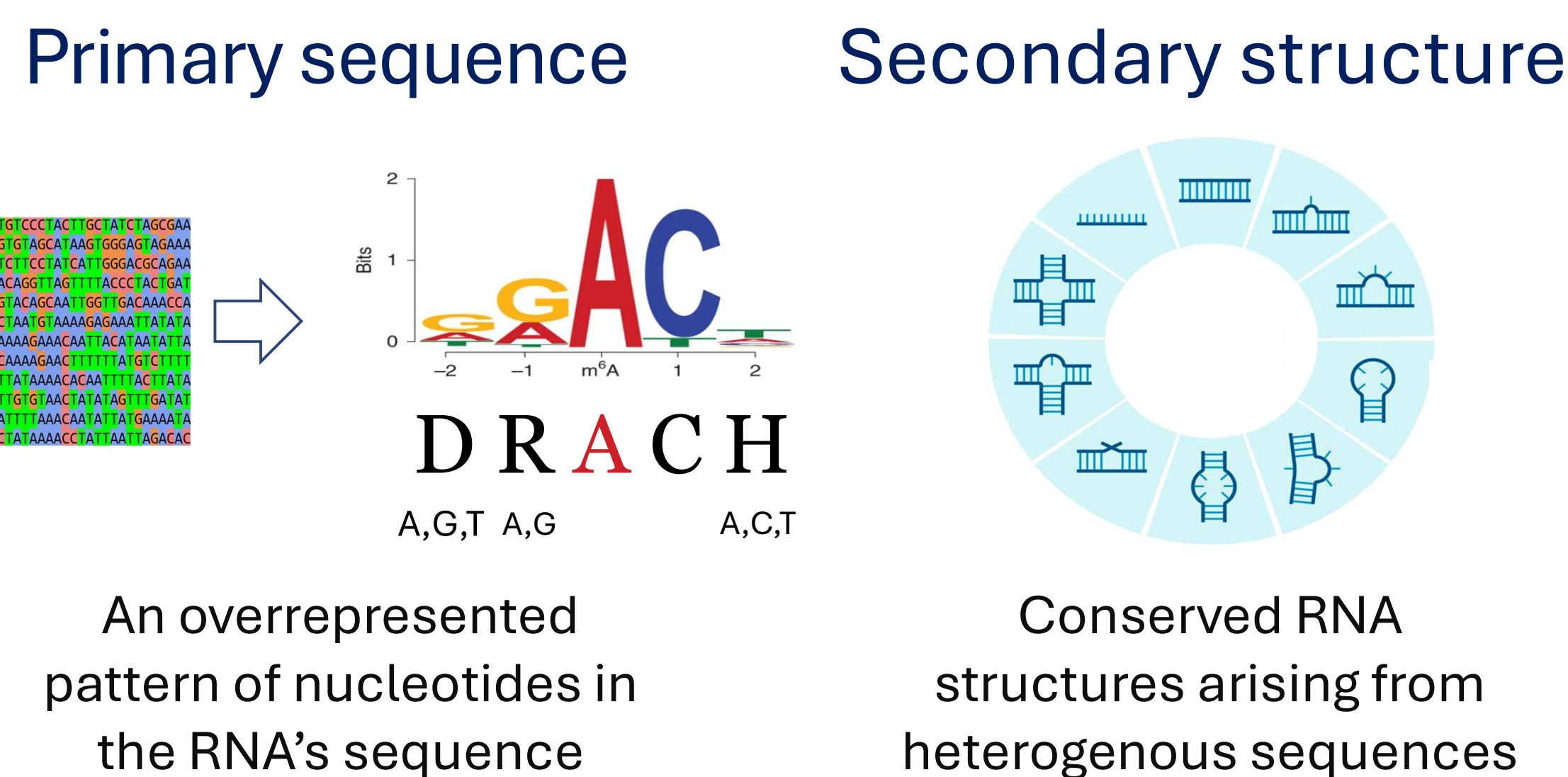
Steven T. Lancashire¹, Joshua Levendis¹, Lakvin Fernando¹, Emma McHugh¹, Stuart A. Ralph¹

¹. The Department of Biochemistry and Pharmacology, Bio 21 Molecular Science and Biotechnology Institute, The University of Melbourne, Victoria 3010

Background

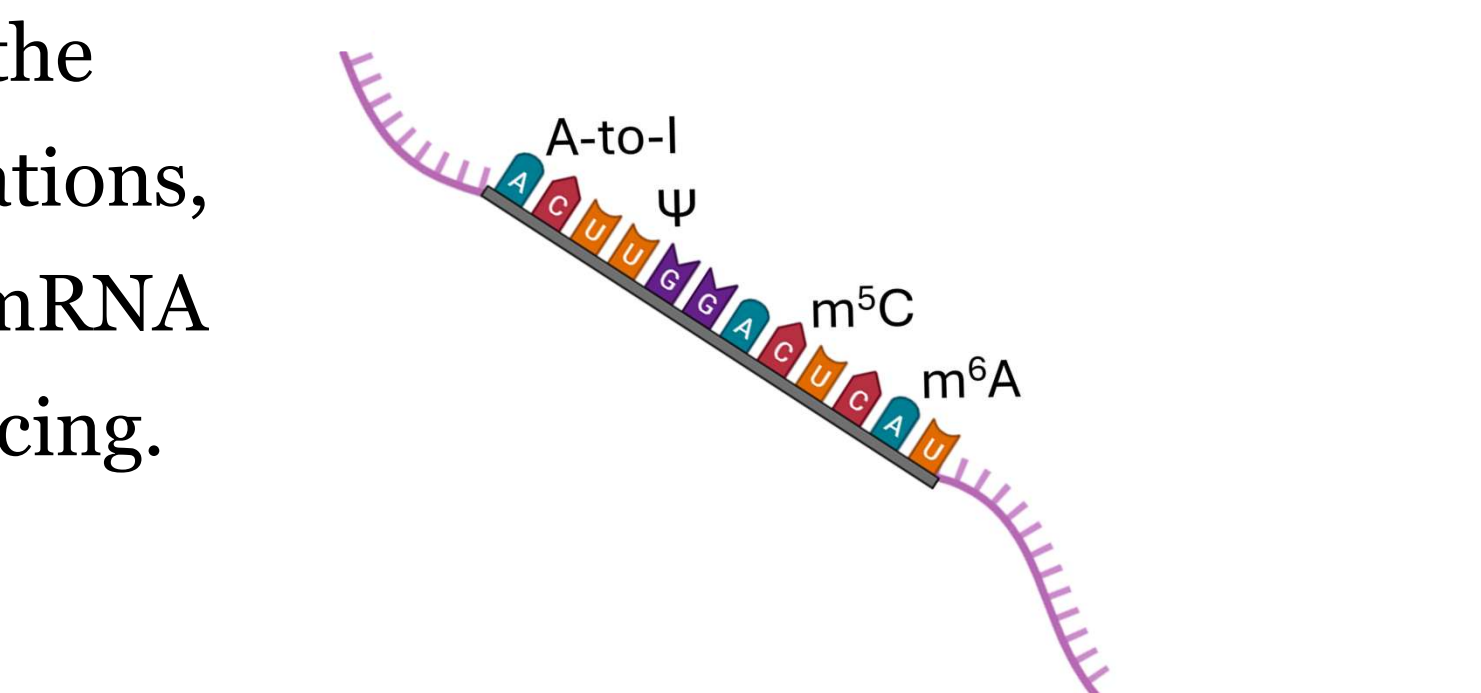
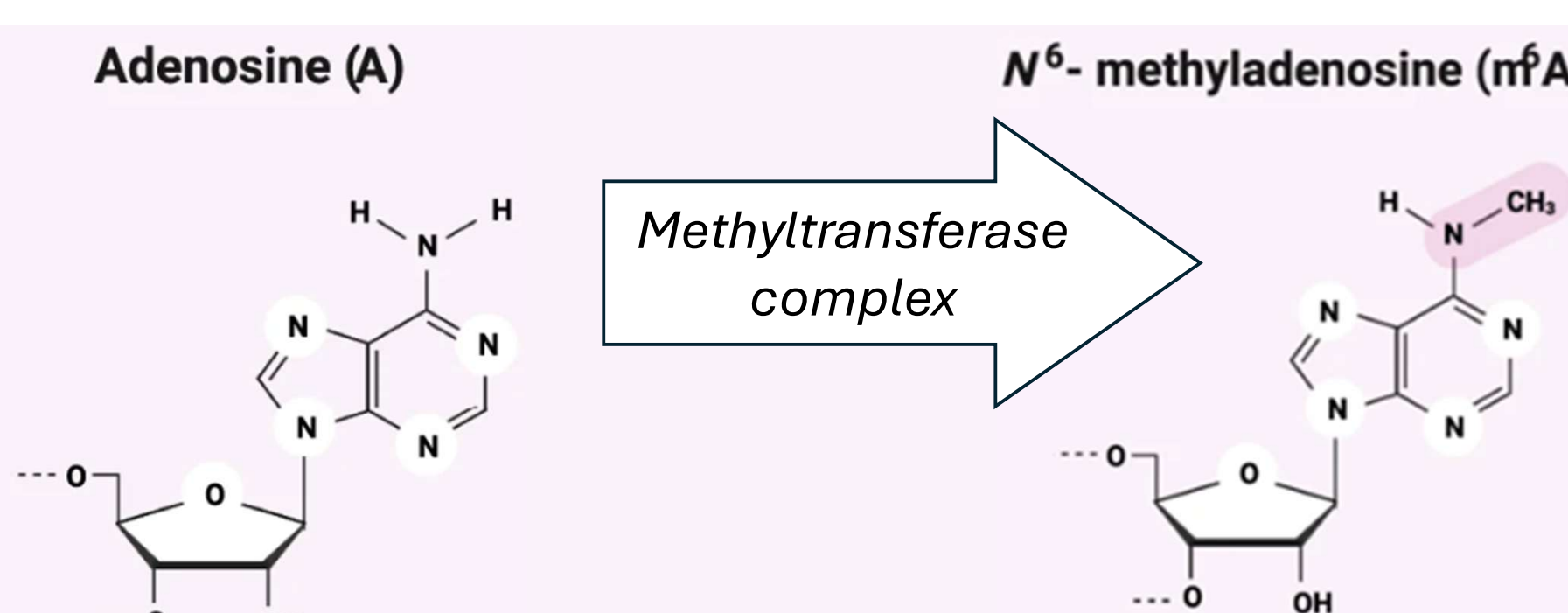
Plasmodium falciparum, the infectious parasite that causes malaria in humans has an mRNA transcriptome that undergoes extensive post-transcriptional chemical modifications. These modifications are implicated in the transcriptional regulation of the parasite's complex lifecycle. The most prevalent modification is adenosine to N6-methyl-adenosine (m⁶A). These modifications to mRNA occur at DRACH sequence motifs, however less than 10% of DRACH sites are modified. This bioinformatics masters project aims to identify the other primary sequence and secondary structural motifs required for m⁶A modification.

What are RNA motifs?



What is the epitranscriptome?

The epitranscriptome is the set of chemical alterations to the nucleotides of RNA. There are 170+ known RNA modifications, with m⁶A the most abundant in mRNA. Modifications to mRNA influence translational efficiency, mRNA stability and splicing.



Methylation of a DRACH motif's central adenosine to m⁶A is performed by a methyl-transferase heterodimer that is conserved in eukaryotes.

Less than
10%
of
D R A C H
motifs carry
m⁶A

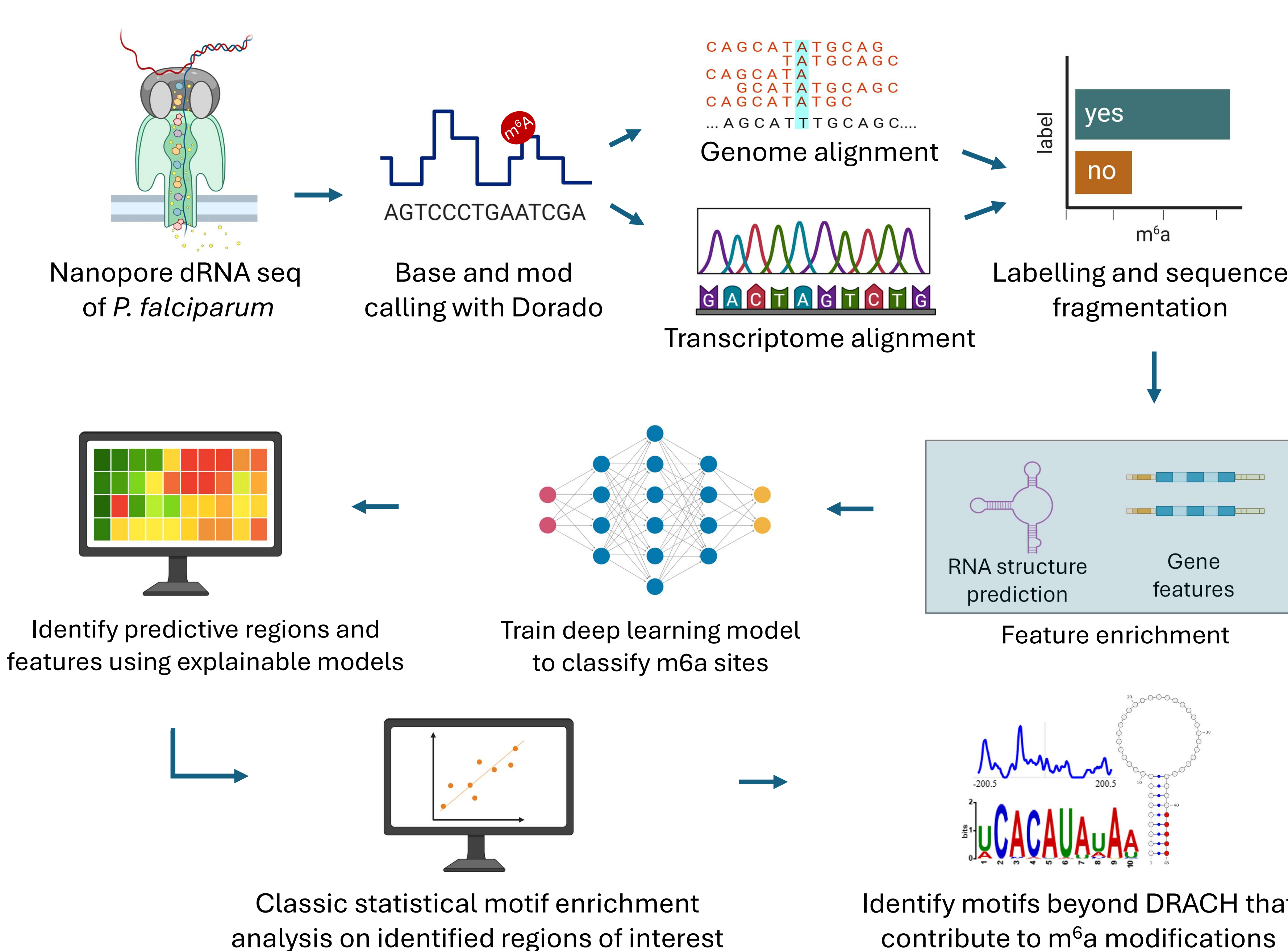
m⁶A impacts translational plasticity across *P. falciparum*'s lifecycle

- Modification reduces and delays translation, resulting in parasite lifecycle supporting just-in-time translation
- The m⁶A reader, PfYTH2, represses protein translation and is essential for parasite survival
- m⁶A in 3' UTRs facilitates 3' cleavage of the mRNA transcript

What RNA sequence and structural motifs do m⁶A modifications require?

Motif discovery

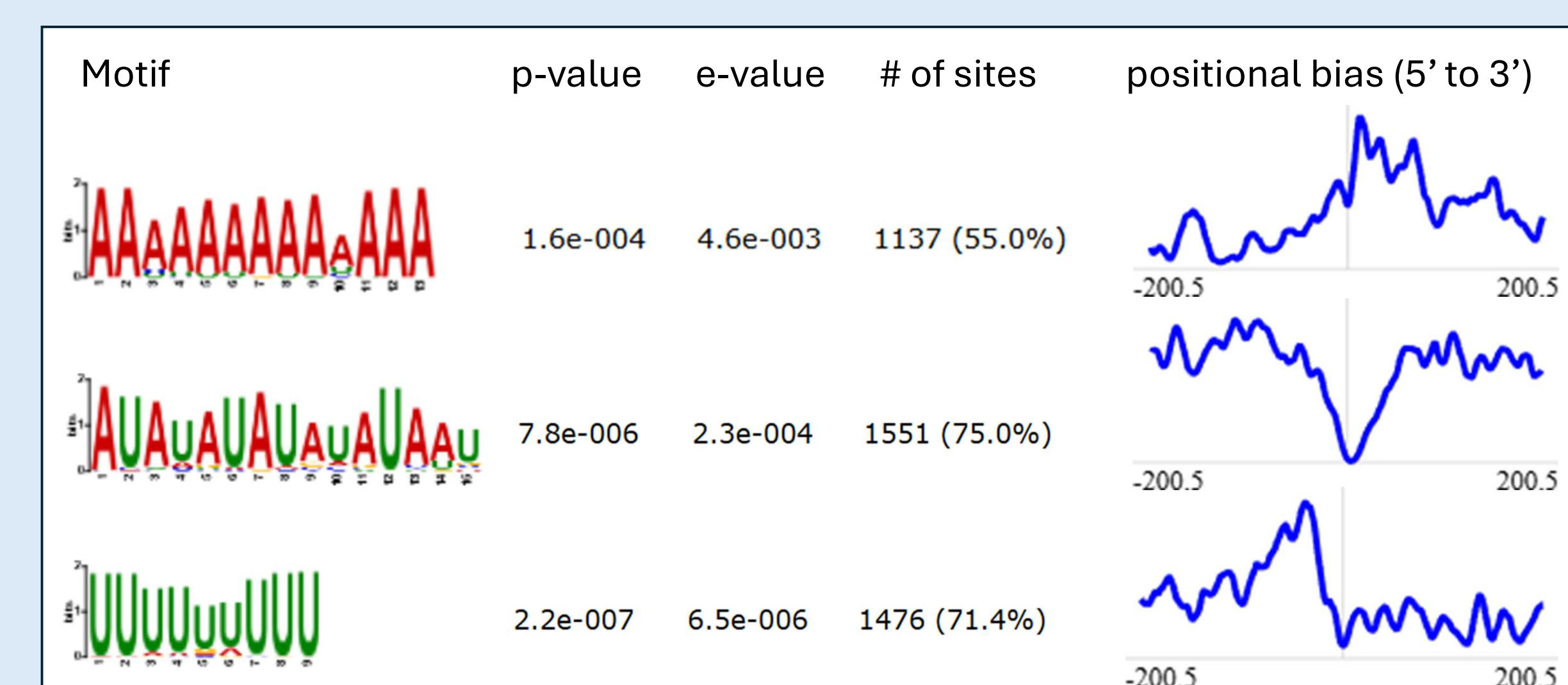
Plasmodium falciparum long read dRNA-seq data are combined with RNA structure and gene features. Deep learning models are used to identify RNA regions predictive of m⁶A. Statistical motif enrichment analysis is then applied to these regions to discover motifs.



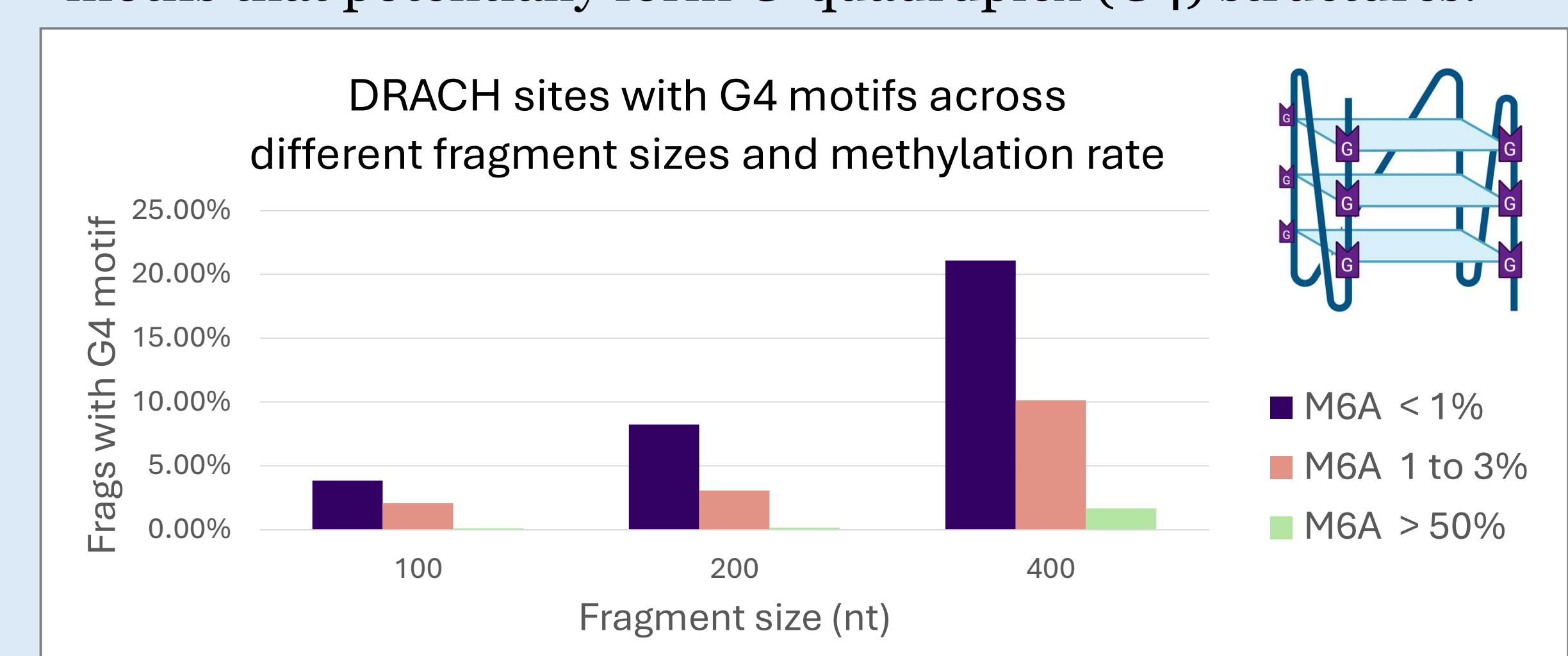
Preliminary results

m⁶A is selectively enriched and depleted in different motif contexts

Result 1: Enriched homopolymer motifs show directional bias around m⁶a modified DRACH sites.



Result 2: DRACH sites are less likely to be modified near motifs that potentially form G-quadruplex (G4) structures.



References

- Sinha, A. et al. Functional Characterization of the m6A-Dependent Translational Modulator PfYTH2 in the Human Malaria Parasite. *mbio* **12**, 10.1128/mbio.00661-21 (2021).
- Baumgarten, S. et al. Transcriptome-wide dynamics of extensive m6A mRNA methylation during *Plasmodium falciparum* blood-stage development. *Nat Microbiol* **4**, 2246–2259 (2019).
- Bailey, T. L. STREME: accurate and versatile sequence motif discovery. *Bioinformatics* **37**, 2834–2840 (2021).

Acknowledgements

Stuart Ralph, Joshua Levendis, Lakvin Fernando, Emma McHugh, Sophie Collier, Long Huynh, Haowen Deng, Zoe Tregloan-Dunn

stevelan.github.io/m6a-motifs

