







# Investigating the Histone Binding Potential of the ZZ and SANT Domains of Ada2

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# In silico Histone N-terminal Interrogation





Peptide Library

• Developed novel computational algorithm, Seq2PTM, to generate library of post-translationally modified Histone H3 and H4 N-terminal tail pdb files

Seq2PTM generates code to fabricate the peptides with PTMs in PyMol.

• Seq2PTM can spawn a library of peptides with PTMs rather than the one-



methodology to simulate docking between peptides and Ada2 domains

# **Future Work**

In silico

 Determine best program & methodology to dock N-terminal tails with Ada2 Broaden functionality of Seq2PTM by adding support for more PTMs

## In vitro

## In vivo

• Compare levels of PTMs between wild type and mutated versions of Ada2

# **Author Contribution Statement**

SS: Contributed to the design of experiments, designed the poster, generated all data; EM, CC, & HR: Contributed to the design of *in vitro* experiments; BS: Oversaw the project design and execution

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