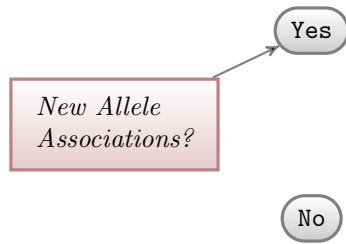


*Use New and Existing
Allele Associations*



*Use Epitope Prediction
To Discover Properties
of Beneficial Alleles*

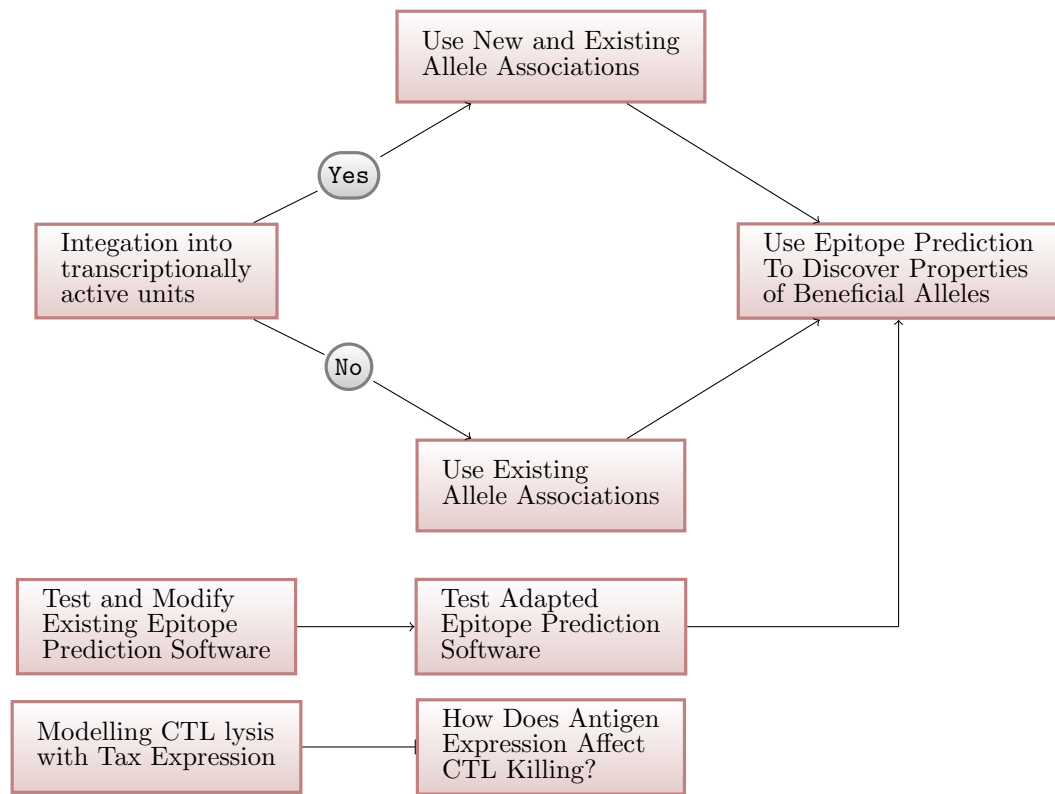
*Use Existing
Allele Associations*

*Test and Modify
Existing Epitope
Prediction Software*

*Test Adapted
Epitope Prediction
Software*

*Modelling CTL lysis
with Tax Expression*

*How Does Antigen
Expression Affect
CTL Killing?*



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