Reducing noise in protein multialignments

Introduction

Multialignments are noisy. Homologous proteins can contain regions that are not inherited and should therefore not be aligned, and other regions may have evolved so fast that the correct multialignment is impossible to infer. In order to get rid of such problematic regions in subsequential analysis, noisy columns are removed. In this project, a column is considered noisy if

- there are indels.
- at least 50% of amino acids are unique

To validate whether these reduction is useful or not, I had to infer tree from the given alignments and the noise reduced alignments of alignment set having different mutation rate. Then the reference tree of particular alignments was compared with noise-reduced tree and tree from given alignment.

Results and Discussion

In case of symmetric data, frequency of recovery of the references tree improve a bit but in case of asymmetric data it doesn't make any difference. So it's not worthwhile to use multialignment noise reduction based on above specified criterion.

Frequency of reference tree being recovered:

Data Group	Before noise reduction	After noise reduction
asymmetric_0.5	1	1
asymmetric_1.0	0	0
asymmetric_2.0	0	0
symmetric_0.5	21	27
symmetric_1.0	16	18
symmetric_2.0	6	12

Average symmetric difference:

Data Group	Before noise reduction	After noise reduction
asymmetric_0.5	7.33	7.29
asymmetric_1.0	9.4	8.88
asymmetric_2.0	12.51	11.39
symmetric_0.5	4.07	4.06
symmetric_1.0	5.07	4.81
symmetric_2.0	7.06	6.28