LTR Parser Results

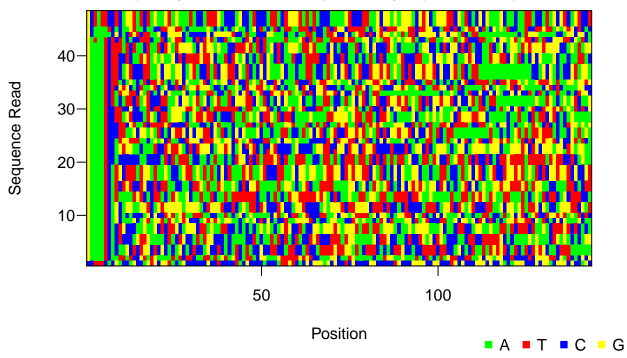
Kevin McCormick

20 April, 2020

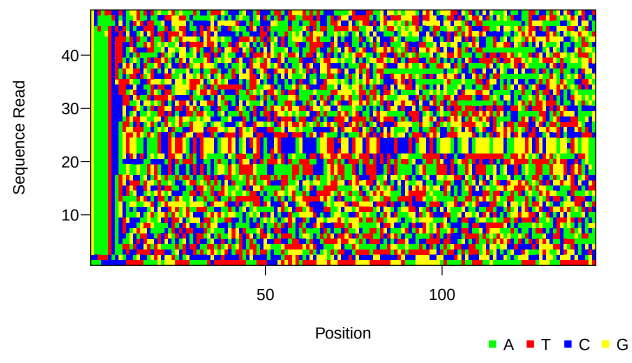
Overview

Total number of reads in sample	48
Number of distinct reads in sample	48
Number of copies of most common read	1

Visualization of 48 sequencing reads, selected randomly and arranged by nucleotide sequence:



Visualization of 48 distinct sequencing reads (after removing duplicates), selected randomly and arrangedy by nucleotide sequence:



No primer LTR pairs appeared in more than 5% of total reads.