Analysis and classification of long terminal repeat sequences from plant LTR-retrotransposons

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Long Terminal Repeats (LTRs) are DNA sequences common in various organisms. By understanding their structure we may be able to understand the evolutionary processes of the organisms in which they are located.

These sequences are, however, difficult to detect using common methods in bioinformatics.

This thesis aims to analyze their inner structure, as well as train machine learning models which are able to recognize LTR sequences. Different models of different complexities are used for this approach in order to find the best one.

The findings of this work may be used by scientists studying evolution of viruses in plants.