# Explainable AI for Genomic Tasks with Genome Language Model

**CAP 5610 - Machine Learning** 

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#### **Presented By**

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## Lets Talk DNA



Promoters, silencers, enhancers

(DNA motifs)

Can be very far from binding site, altering chance of protein binding

## **Motivation**



- Advanced Genome Language Models: Transformer-based models such as DNABERT, DNABERT2, Nucleotide Transformers, and HyenaDNA have demonstrated strong performance on various genome-specific classification tasks.
- **Knowledge Gap:** Despite their success, the depth of these models' understanding of genomic concepts remains unclear.
- **Explainability Focus:** This project aims to investigate how well these models can explain their predictions by applying them to a downstream interpretability task.

# Task Description



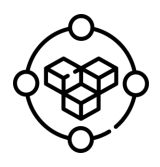
- **Objective:** Use advanced genomic language models (e.g., DNABERT, HyenaDNA) for promoter detection.
- **Focus:** Investigate whether these models truly leverage the presence of TATA boxes for classification.
- **Hypothesis:** TATA boxes are key signals that significantly influence the model's decision-making in identifying promoters.
- **Explainability Goal:** Demonstrate how attention or feature-attribution methods can reveal the role of TATA box motifs in the models' predictions.

## Dataset



Category	Details
Dataset Used	Genome Understanding Evaluation (GUE) for human promoter detection
Sequence Length	300 Nucleotides
Dataset Split	Training Set: 47.4k samples Validation Set: 5.2k samples Test Set: 5.2k samples
Class Distribution	Balanced (Positive:Negative = 50:50)

## Models for Promoter Detection



Model	Purpose
DNABERT	Transformer-based model for DNA sequence analysis
DNABERT2	Improved version of DNABERT with enhanced performance
Nucleotide Transformers (NT)	General-purpose nucleotide sequence transformer
HyenaDNA	A novel architecture designed for genomic data

# Results

Model	Parameters	Architecture	Tokenizer	Accuracy	MCC	F1 Score	FT Time
DNABERT	1.6 M	Transformers	K-mer	0.969	0.909	0.959	230 m
DNABERT2	117 M	Transformers	BPE	0.932	0.863	0.931	150 m
NT	2.5 B	Transformers	K-mer	0.930	0.853	0.927	120 m
HyenaDNA	1.6 M	Implicit Conv.	Single Nucleotid e	0.940	0.862	0.931	35 m

# Model Analysis

- Human-specific pretraining: Models trained on the human genome excel in species-specific tasks.
- Multispecies challenge: Multispecies-pretrained models underperform due to insufficient parameter scaling.
- Scaling advantage: Larger models yield better results.
- Runtime efficiency: Alternatives to transformer architectures offer faster runtimes.

# Tools for Explainability in ML



#### • LIME (Local Interpretable Model-Agnostic Explanations)

Works with any black-box model and provides human-interpretable explanations. Used on text classification(NLP), image classification, and Tabular Data (fraud detection, medical diagnosis)

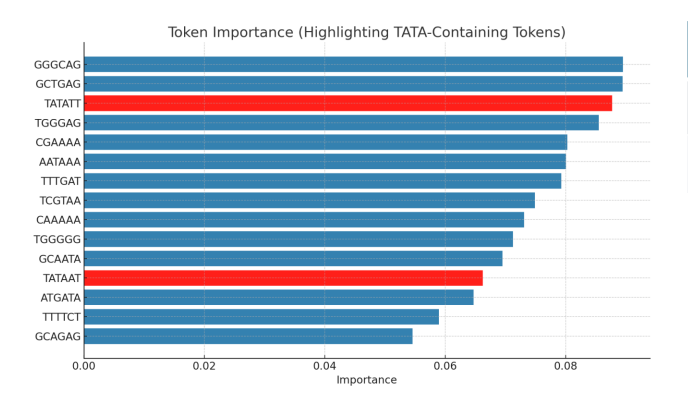
#### • **SHAP** (SHapley Additive Explanations)

Works with various model types (tree-based, deep learning, linear models) and provides both local and global explanations)

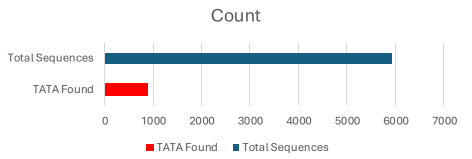
#### • ANCHOR (High-Precision Model-Agnostic Explanations)

More stable model-agnostic tool that provides high-confidence explanations. Used for Text classification, Image classification, Fraud detection, and healthcare)

## LIME - Results

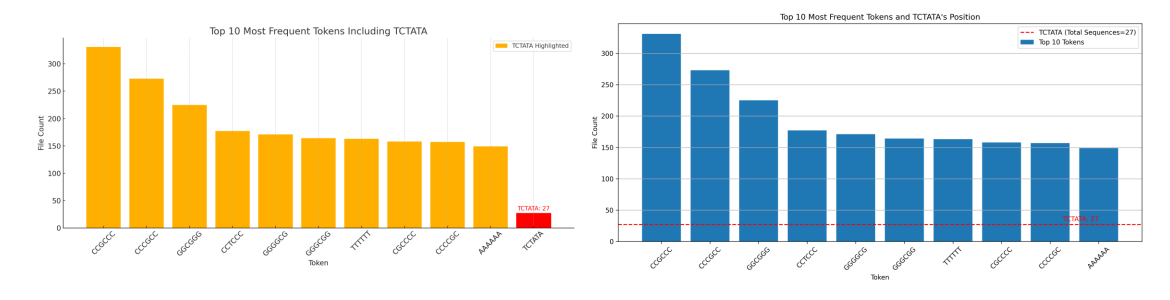


Importance/Label	0	1
Positive	54	99
Negative	<mark>639</mark>	84



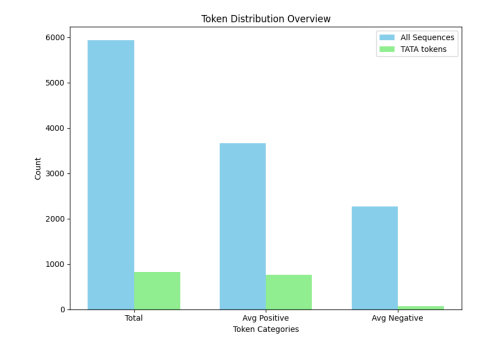
## LIME – Results – cont.

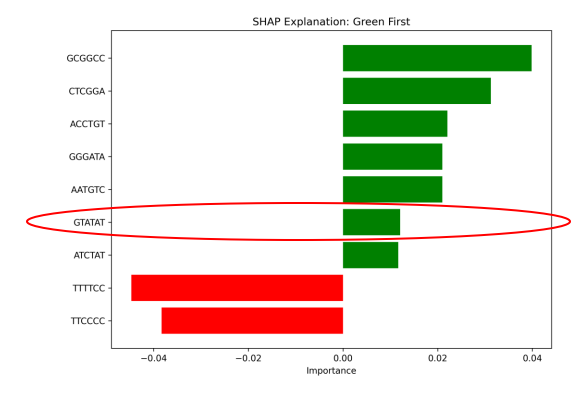
**TCTATA**, the most common **TATA-containing token**, appears in **27** sequences **(20%)**, whereas the top tokens average around **150** appearances—indicating TCTATA's moderate presence.



# SHAP (SHapley Additive exPlanations)

- Model-agnostic : Can explain any machine learning model
- > Unlike LIME, SHAP utilizes Global understanding
- > Creates set of features to find importance of each





- ☐ Total sequence = 5930
- ☐ Tokens analyzed per sequence = 10 ☐
  - Total sequences = 5930
- ☐ Total TATA tokens = 828

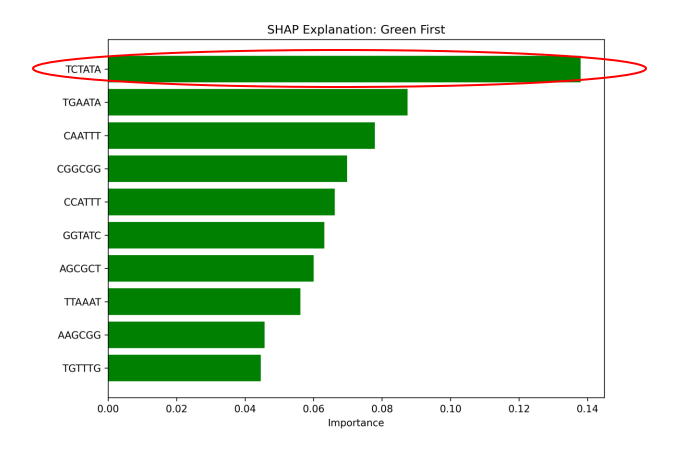
- ☐ Avg Positive tokens = 3662
  - Avg Negative tokens = 2268
- ☐ Positive TATA = 763
  - Negative TATA = 65
- Accurate TATA = 277
- Non-accurate TATA=486

## **SHAP**

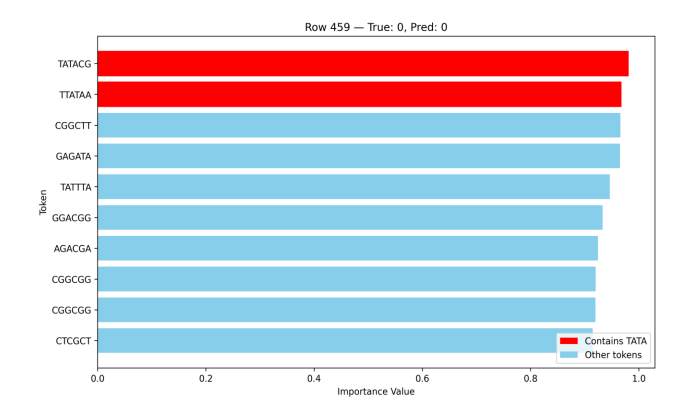
#### > Top 10 tokens with positive importance

TCTATA	118
СТСССС	78
GGCGGG	78
GGTGGA	65
GCGGCG	65
GCCTGG	65
TGAATA	65
CCAGAG	65
CCTCCC	65
TATTTT	52

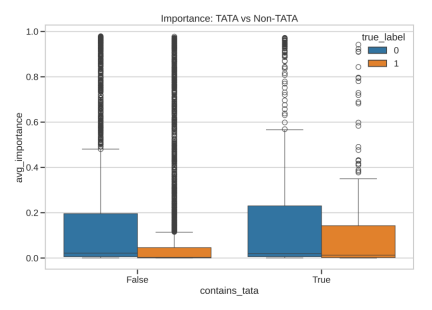
- **➤** Model Used : Nucleotide Transformers
- $\triangleright$  Token size = 6



## **ANCHOR**



Category	Count
Contains TATA	554 (9.34%)
Lable 0 with TATA	407
Label 1 with TATA	147
TATA in Label 0, Negative	0
TATA in Label 0, Positive	407
TATA in Label 1, Negative	0
TATA in Label 1, Positive	147



## Limitations

- ☐ Resource
  - > GPU
  - Job Distributor
  - Worker Process (for running the programs independently on 16 GPUs)
- ☐ Time
  - > Analyzing 1 sequence with SHAP tool takes around 500 seconds.
  - > Total 5930 sequences take 5930\*500 = 824 Hours = 34 Days
  - > Had to use parallel processing to complete these tasks within timeline
- □ Model Characteristics
  - Not all the model used for promoter detection are suitable for explainabilty
  - > HyenaDNA model cannot be used for our project

### Conclusion

- > TATA box is a short, specific DNA sequence of about 25-35 base pairs long (TATAAA, or something similar) located at its distance from the transcription initiation site which is part of many promoters.
- It is biologically proved that, "TATA" sequence will be present in the DNA sequence if there is promoter.
- Even ML models, are to some extent dependent on this TATA sequence to detect promoter in DNA





https://github.com/roufunr/explainable-AI-for-genomic-tasks

What??
You Want
More??

