## METK Barley SNP-Chip: Linking Genetics to Protein Content

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## Introduction

Our project, utilizing exclusive 2024 data from METK, explores the genetic factors influencing barley's protein content—a key attribute for both nutritional and agricultural purposes. By analyzing the correlation between barley's genetic makeup and its protein levels, we aim to provide insights that could guide future crop breeding efforts and support the growing demand for plant-based protein sources.

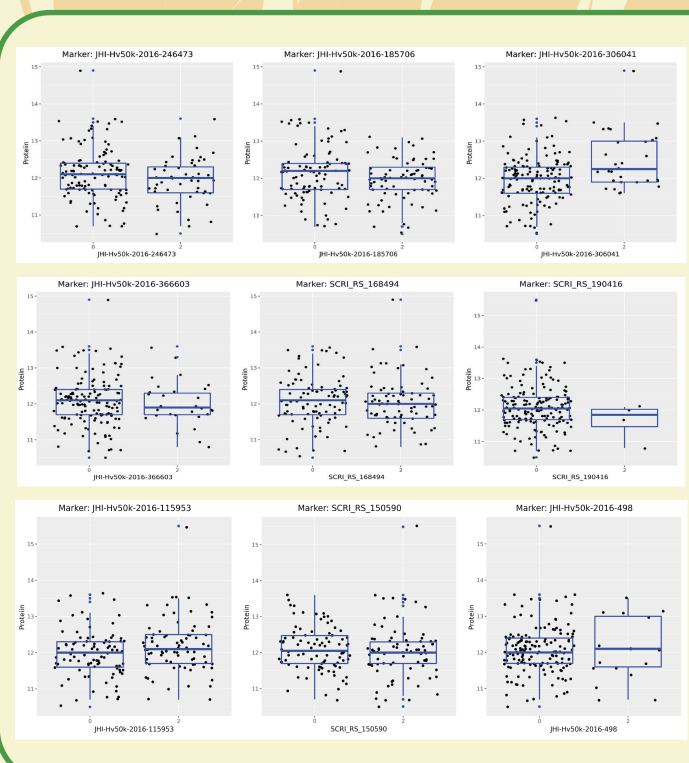
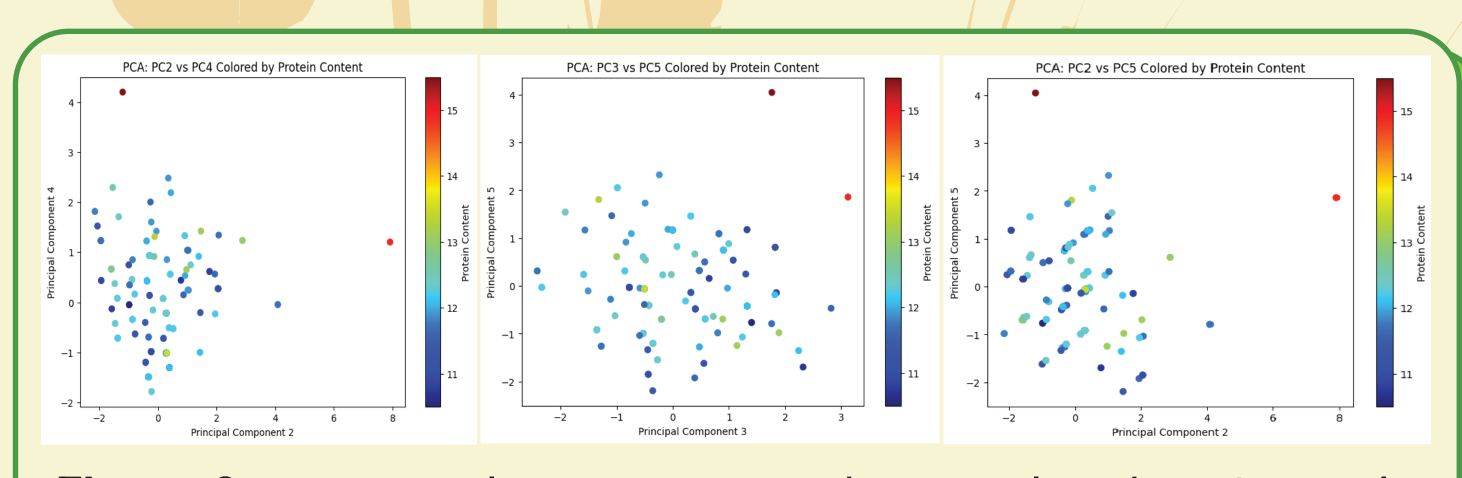


Figure 1: We used scatter and box plots to visualize the variation in protein content across different genotypes for markers with a greater than 2.5% difference in protein content. A total of 9 markers were identified. Two of the most promising markers, JHI-Hv50k-2016-306041 and SCRI\_RS\_190416. Although these markers were not statistically significant, further testing is recommended to explore their potential.



**Figure 2:** A scatter plot was generated to visualize the PCA results, with points colored according to protein content. In some of the plots, two sample points with higest protein content (Kornelija and Ilma) were visually clustered separately. However, no distinct separation was observed between the low and medium protein content groups.

## Data analysis

Data Cleaning: By filtering out SNP markers with insufficient variation (where one genotype or NaN values dominated), we focused only on markers with useful genetic diversity.

Protein Content Variation: The analysis showed which SNP markers had significant variation in protein content between different genotypes. Some SNP markers exhibited a large range of protein content differences, indicating that the genotype at these markers could have a notable effect on protein expression. (Figure 1)

PCA: We performed a series of steps to analyze SNP (Single Nucleotide Polymorphism) data in relation to protein content, with the goal of identifying any potential patterns or clusters in the data. (Figure 2)

Dendrogram: We used Hamming distance to quantify the similarity between SNP samples, which is appropriate for binary or categorical data. Missing data was handled using mean imputation, ensuring that incomplete rows wouldn't disrupt the analysis. Applied hierarchical clustering to group similar samples based on their SNP profiles, helping to identify patterns or clusters of genetic similarity. (Figure 3)

**Prediction Models:** For protein content prediction different models were used: RandomForestRegressor, DecisionTreeRegressor, Simple Linear, Lasso, and Ridge Regression. Ridge Regression had the best R2 score (0.089) and MSE (0.535) out of the selection. Unfortunately, these models are probably not sophisticated enough and a more specialised software such as MEGA should be used.

## Results

We found 2 markers that have a stronger correlation to protein: JHI-Hv50k-2016-306041 and SCRI\_RS\_190416. Both are located on chromosome 5H. 2 varieties (Kornelija, Ilma) are genetically very different from the rest, having more unique DNA as well as a higher protein content. Although current findings do not point towards any significant correlation between protein content and SNP markers, this is a solid base for future research.

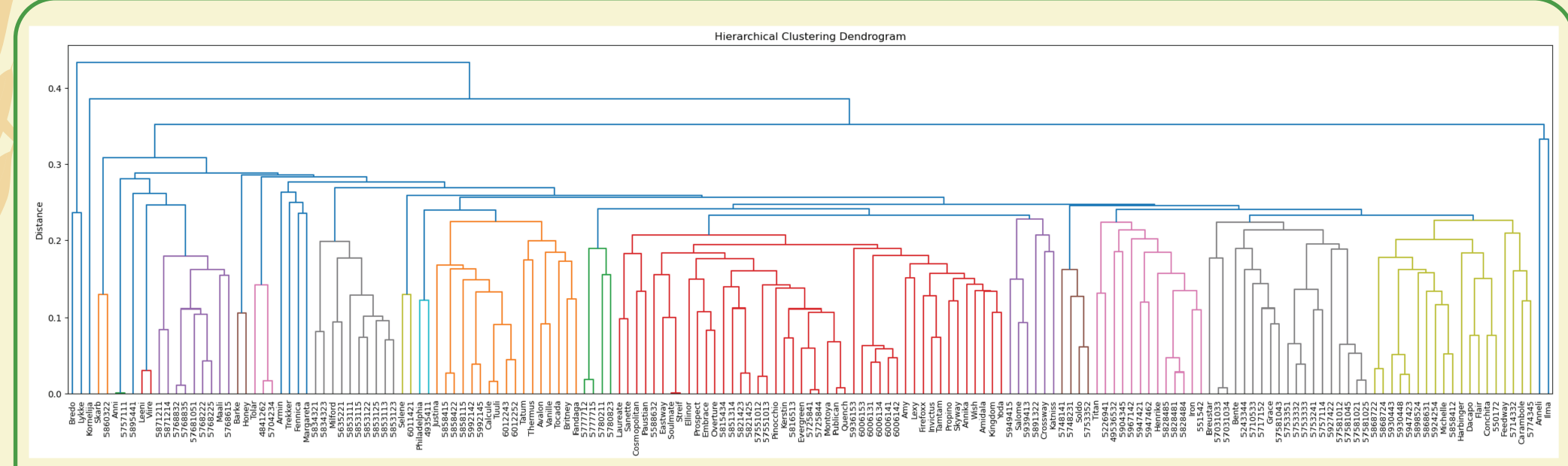


Figure 3: Dendrogram highlighted sample relationships and distinct clusters.

