

# ML@NOVA DFJ

Predicting Survival Time in Multiple Myeloma Patients

# Team identification

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Final score on the private leaderboard: 2.58777

Leaderboard private ranking: 30

# Task [1.1] - Data preparation and validation pipeline

# What was done in task [1.1]

## 1. Missing Values Analysis

Visualized missing values using multiple methods (bar plot, heatmap, matrix, dendrogram)

Created comprehensive overview of data completeness

Identified patterns in missing data

## 2. Data Cleaning

Dropped rows with missing SurvivalTime values

Removed columns containing missing data (baseline approach)

Excluded censored cases (where Censored == 1)

Retained only complete, uncensored observations

## 3. Feature Exploration

Visualized feature relationships using pairplot

Analyzed correlations between Age, Gender, Stage, TreatmentType, and SurvivalTime

Examined distribution patterns across features

# What was done in task [1.1]

## 4. Data Preparation

Defined feature matrix (X) by dropping target and identifier columns

Isolated target variable (y) as SurvivalTime

Preserved censoring indicator for potential future use

## 5. Validation Strategy Development

Implemented train/validation/test split (64%/16%/20%)

Tested simple split approach with Linear Regression

Implemented 5-fold cross-validation for more robust evaluation

Compared both validation strategies (simple split vs. cross-validation)

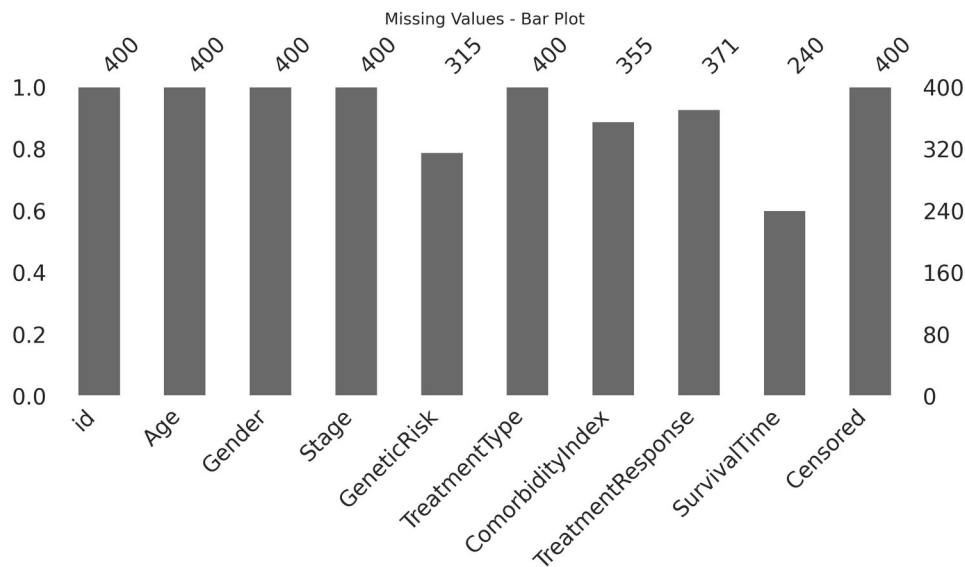
## 6. Performance Evaluation

Calculated MSE (Mean Squared Error) and cMSE (Censored MSE)

Evaluated model performance on validation and test sets

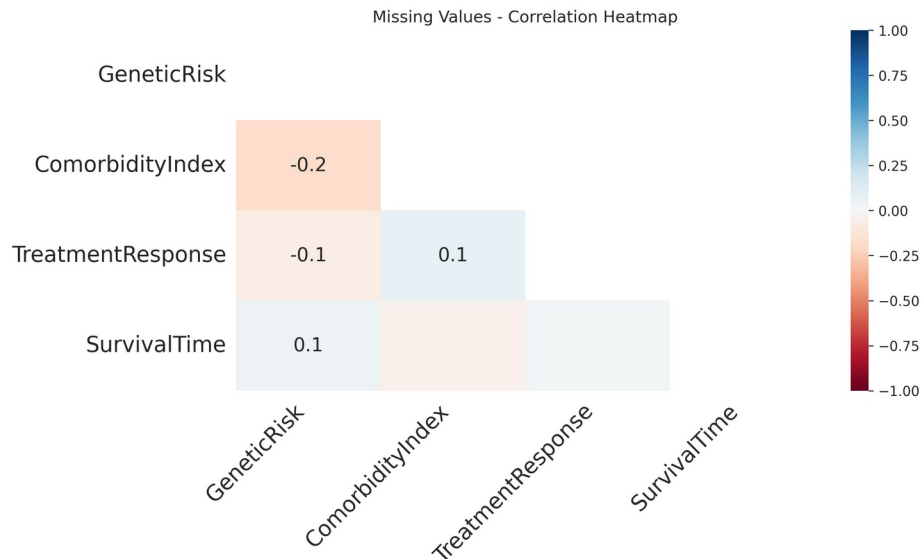
Compared cross-validation results to simple split results

# Results and Analysis from task [1.1]



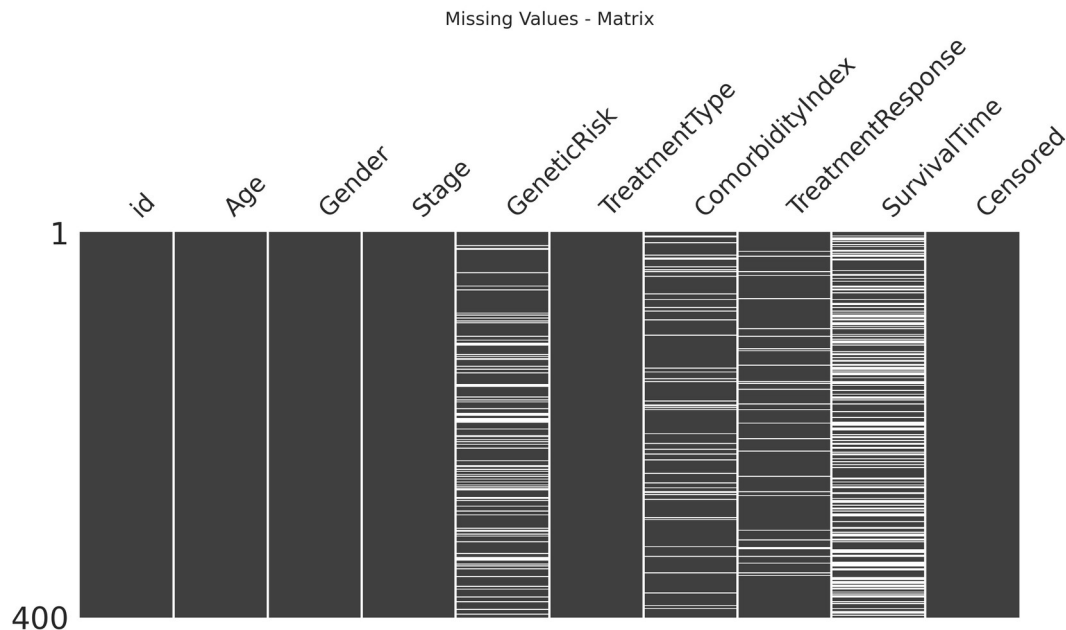
Bar chart showing missing-value counts per variable, with labels indicating how many entries are missing in each column.

# Results and Analysis from task [1.1]



Heatmap showing correlations between missing-value patterns across selected variables, with mostly weak relationships.

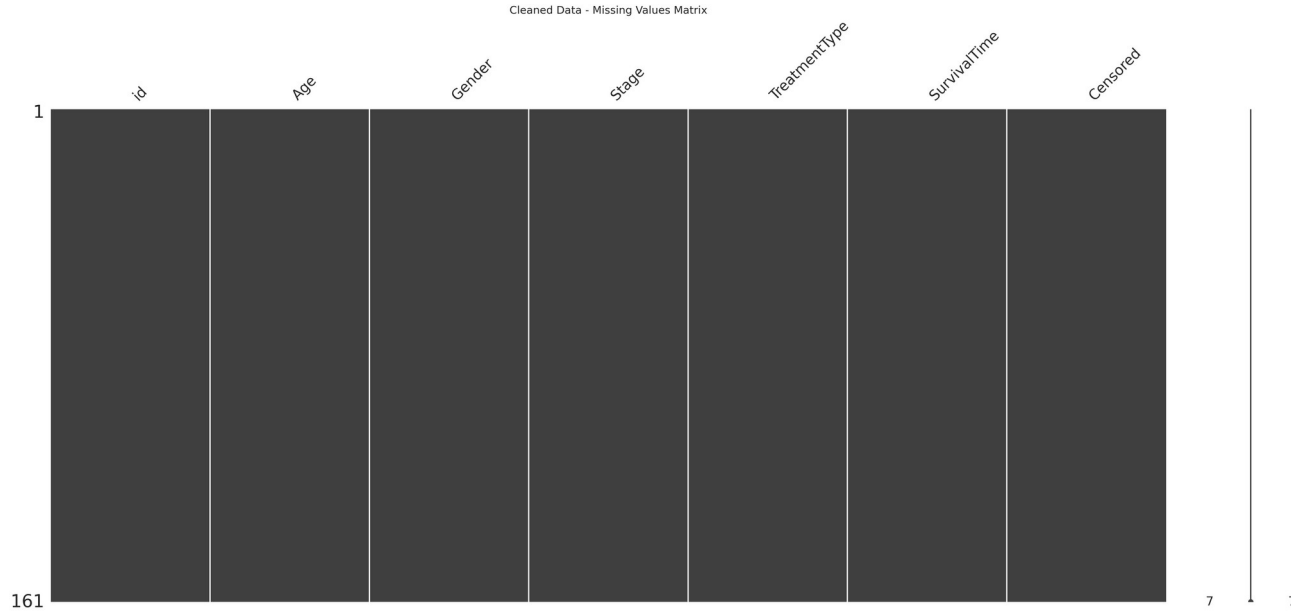
# Results and Analysis from task [1.1]



Matrix plot highlighting which entries are missing across variables, showing the distribution and density of gaps in the dataset.

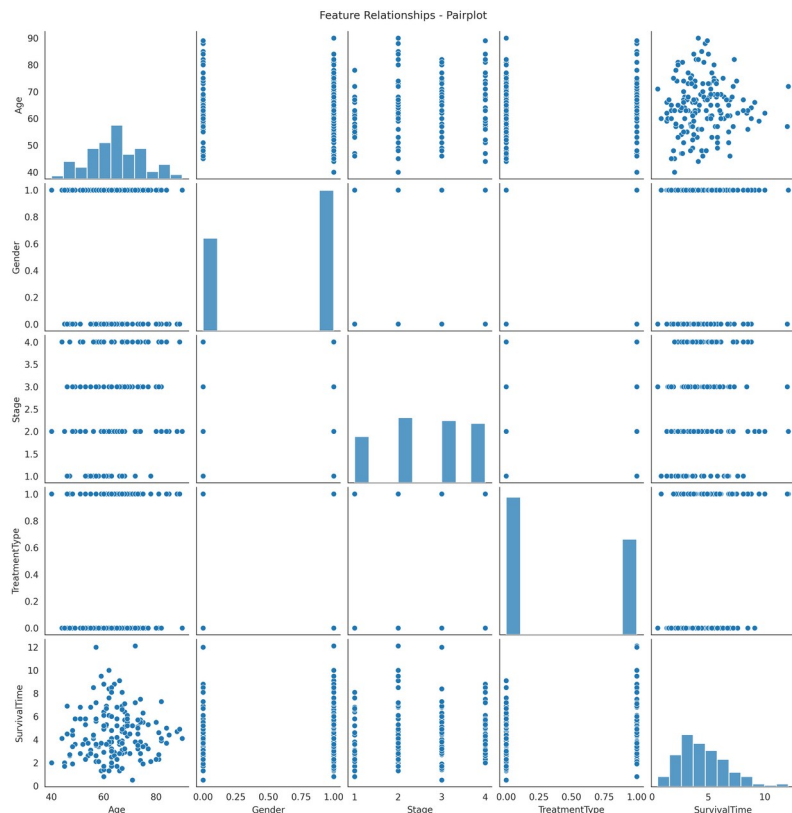


# Results and Analysis from task [1.1]



Matrix plot showing the cleaned dataset, confirming that all variables have no remaining missing values.

# Results and Analysis from task [1.1]



## Age

- Roughly uniform distribution between 40 and 85.
- No clear relationship with Stage, Gender, or TreatmentType.
- Slight indication that lower ages may correspond to higher SurvivalTime, but with large variability.

## Gender

- Binary variable with no meaningful pattern across other features.
- SurvivalTime similarly distributed between genders.

## Stage

- Balanced distribution across stages 1–4.
- Clear trend: higher stages correspond to lower SurvivalTime.
- No noticeable relationship with Age or Gender.

## TreatmentType

- Most individuals fall into one or two treatment categories.
- No strong association with Age or Gender.
- SurvivalTime varies within each treatment group without strong separation.

## SurvivalTime

- Right-skewed distribution, with most values between 1 and 5 years.
- Strongest visible relationship is with Stage (more advanced stages → shorter survival).
- No clear patterns with Gender or TreatmentType.

# Results and Analysis from task [1.1]

	<b>MSE</b>	<b>cMSE</b>
Simple Split	5.0473	5.0473
Cross-Validation	4.0873	4.0873
Pipeline	4.4112	4.4112

The results show that cross-validation improves performance compared to a simple train–test split, reducing the MSE by almost one unit.

The pipeline approach performs slightly worse than cross-validation but still better than the simple split.

## Task [1.2] - Learn the baseline model

# What was done in task [1.2]

## 1. Pipeline Construction

- Built baseline pipeline combining StandardScaler and Linear Regression
- Ensured feature scaling for improved model performance
- Created modular, reusable pipeline structure

## 2. Cross-Validation Training

- Performed 5-fold cross-validation for robust model evaluation
- Calculated CV MSE scores across all folds
- Computed mean and standard deviation of cross-validation performance

## 3. Final Model Training

- Fitted baseline pipeline on entire training dataset
- Generated predictions on training data
- Maximized use of available data for final model

# What was done in task [1.2]

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4. Performance Metrics
- Calculated Training MSE (Mean Squared Error)
  - Calculated Training cMSE (Censored Mean Squared Error)
  - Established baseline performance benchmarks
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5. Test Predictions & Submission
- Loaded test dataset and prepared features
  - Generated predictions for test samples
  - Created submission file for competition/evaluation
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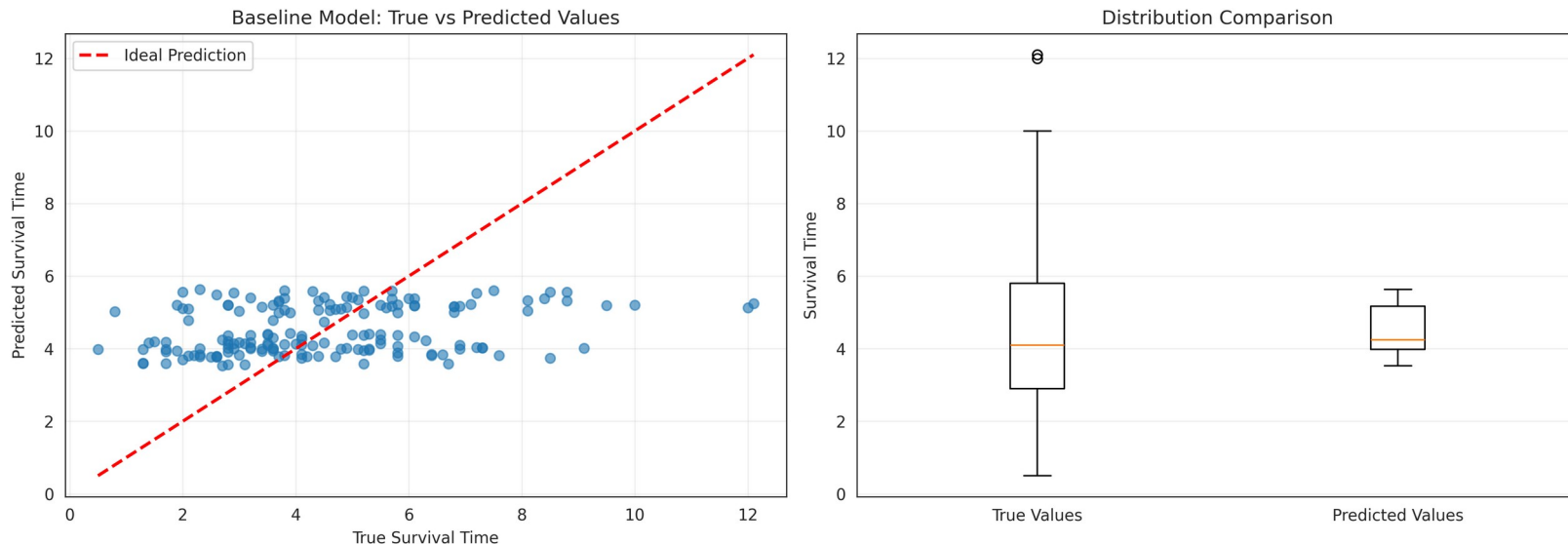
6. Model Visualization
- Created scatter plot comparing true vs predicted survival times
  - Generated boxplot for distribution comparison
  - Visualized model fit quality and prediction patterns
  - Saved individual plots for documentation
-

# Results and Analysis from task [1.2]

- Baseline Model Training with Cross-Validation

MSE	cMSE	Best Kaggle Score
4.0873	4.0873	3.87347

# Results and Analysis from task [1.2]



The scatter plot shows that the model's predictions are squeezed into a narrow band, mostly around 4–6 years and rarely follow the true values across their full range.

The boxplots make this even clearer: the true survival times vary widely, while the predicted ones stay tightly grouped.

Overall, the model isn't capturing the real variability in the data and is underfitting.



# Task [2.1] - Development

# What was done in task [2.1]

## 1. Polynomial Regression Function Development

- Created ``train_polynomial_regression()`` function with hyperparameter search
- Implemented cross-validation for degree selection (testing degrees 1 to `max_degree`)
- Added early stopping mechanism (stops after 2 consecutive iterations without improvement)
- Returned best degree, trained model, and complete CV results dictionary

## 2. k-Nearest Neighbors Function Development

- Created ``train_knn()`` function with hyperparameter search
- Implemented cross-validation for k selection (testing k from 1 to `max_k`)
- Added early stopping mechanism for efficiency
- Returned best k value, trained model, and complete CV results dictionary

## 3. Hyperparameter Selection

- Used 5-fold cross-validation for both models
- Searched polynomial degrees from 1 to 10
- Searched k values from 1 to 20
- Tracked MSE scores with standard deviations for each hyperparameter

# What was done in task [2.1]

## 4. Model Training

- Trained Polynomial Regression with optimal degree on full dataset
- Trained k-NN Regression with optimal k on full dataset
- Generated predictions on training data for both models

## 5. Performance Evaluation

- Calculated training MSE for both models
- Calculated training cMSE for both models
- Compared performance against baseline expectations
- Documented hyperparameter selection results

# Results and Analysis from task [2.1]

- Model performance for polynomial and k-NN Regression

	<b>MSE</b>	<b>cMSE</b>
Polynomial	4.0873	4.0873
k-NN	3.6571	3.6571

# Task [2.2] - Evaluation

# What was done in task [2.2]

- |   |   |
|---|---|
| 1.<br>Comprehensive<br>Model<br>Comparison      | <ul style="list-style-type: none"><li>- Created comparison table with baseline, polynomial regression, and k-NN models</li><li>- Included hyperparameter configurations for each model</li><li>- Displayed min, max, mean, and standard deviation of errors</li><li>- Identified best performing model based on mean cross-validation error</li></ul> |
| 2.<br>Hyperparameter<br>Tuning<br>Visualization | <ul style="list-style-type: none"><li>- Plotted polynomial degree vs MSE with confidence intervals</li><li>- Plotted k-value vs MSE with confidence intervals</li><li>- Marked optimal hyperparameters with vertical lines</li><li>- Showed performance trends across hyperparameter ranges</li></ul>   |
| 3. Model<br>Predictions<br>Comparison           | <ul style="list-style-type: none"><li>- Created scatter plots of true vs predicted values for all three models</li><li>- Displayed MSE on each plot for direct comparison</li><li>- Included ideal prediction line (<math>y=x</math>) as reference</li><li>- Generated combined and individual visualization plots</li></ul>                          |

# What was done in task [2.2]

## 4. Statistical Analysis

- Computed cross-validation statistics for each model
- Analyzed variance in predictions across folds
- Compared model stability through standard deviation metrics
- Evaluated improvement over baseline model

## 5. Test Set Predictions

- Selected best performing model based on CV results
- Generated predictions for test dataset
- Created submission file for evaluation
- Documented model selection rationale

## 6. Results Documentation

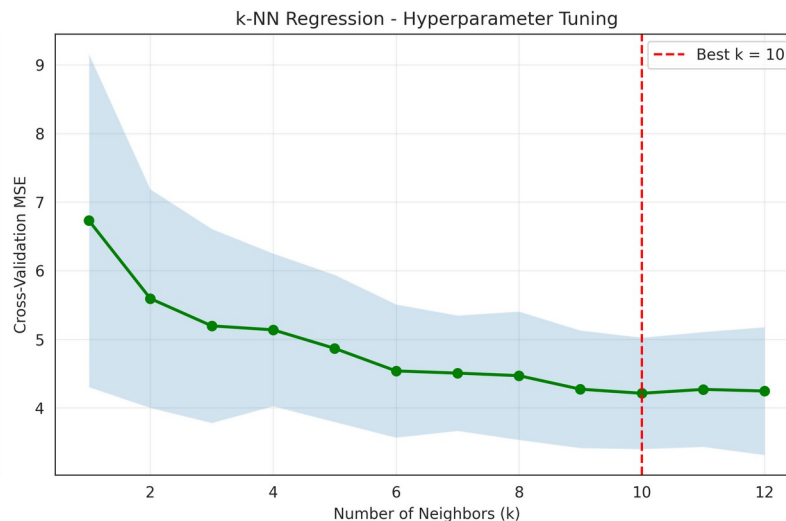
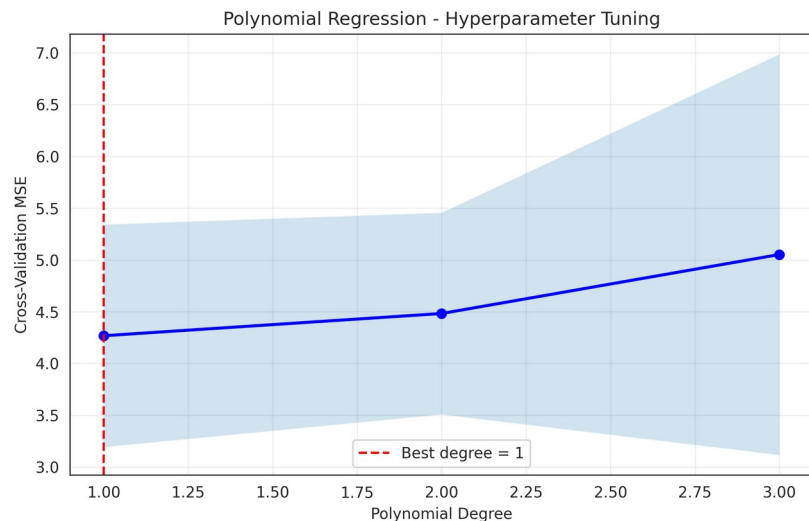
- Saved all comparison plots with task-specific naming
- Generated separate plots for polynomial and k-NN tuning
- Created individual prediction visualizations for each model
- Documented complete evaluation workflow

## Results and Analysis from task [2.2]

- Best submission for non-linear in Kaggle: **3.29914**



# Results and Analysis from task [2.2]

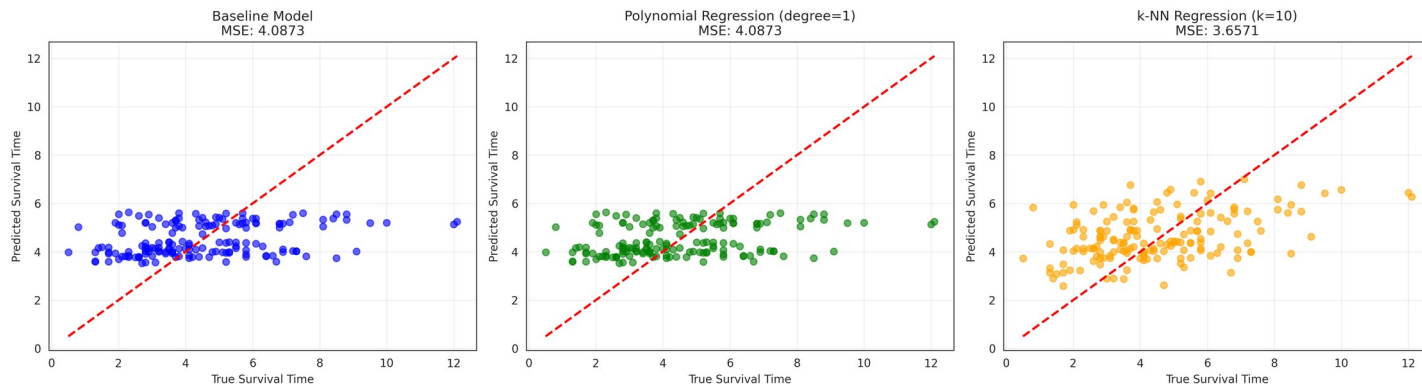


The tuning results show that polynomial regression performs best at degree 1, meaning that adding polynomial terms does not improve the model and only increases error.

In contrast, k-NN regression benefits from increasing the number of neighbors: the error decreases steadily and reaches its minimum around  $k = 10$ .

This indicates that a smoother, more averaged prediction strategy fits the data better than more complex polynomial relationships.

# Results and Analysis from task [2.2]



The baseline and polynomial regression models behave almost identically: both compress predictions into a narrow band and fail to follow the true survival times across their full range. The k-NN model improves this pattern. It spreads predictions more realistically and gets closer to the ideal diagonal, which is reflected in its lower MSE. Even so, the improvement is modest, and the model still underestimates higher survival times.

# Task [3.1] - Missing data imputation

# What was done in task [3.1]

## 1. Data Preparation

- Load original dataset with missing values
- Analyze missing value patterns
- Prepare feature matrix (X) and target variable (y) with missing data intact

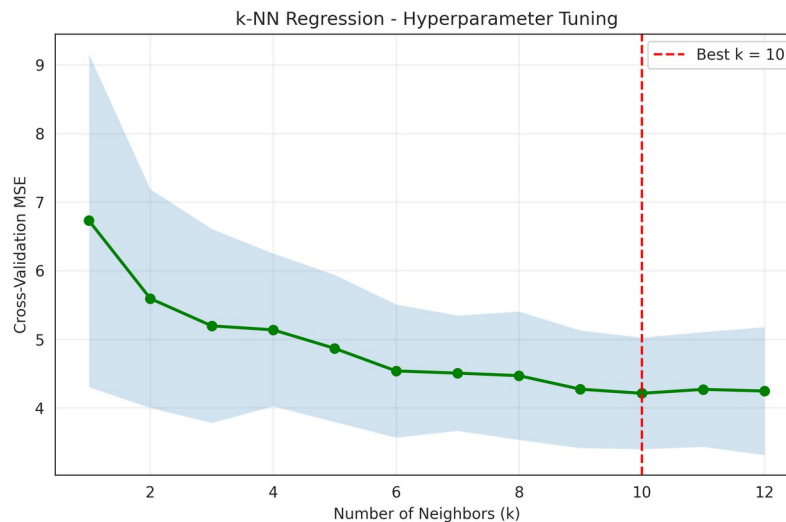
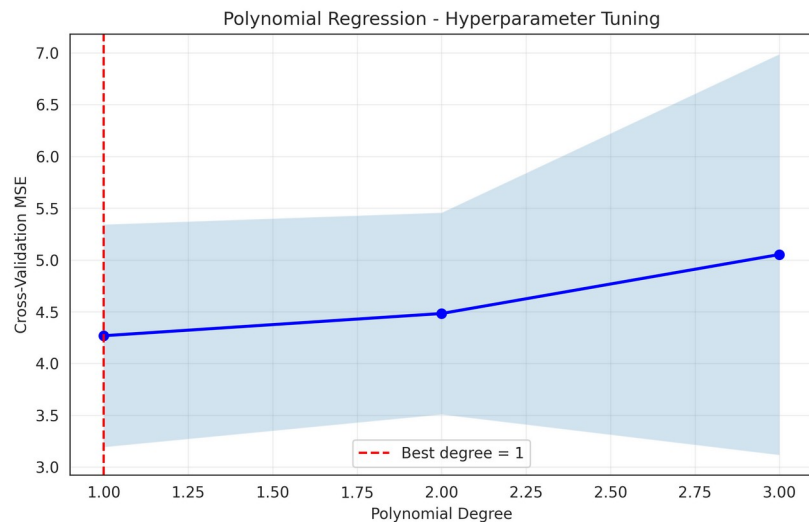
## 2. Imputation Strategies

- Mean Imputation: Replace missing values with column means
- KNN Imputation: Use k-nearest neighbors to estimate missing values
- Iterative Imputation: Use Bayesian Ridge regression for multivariate imputation

## 3. Model Evaluation

- Train baseline Linear Regression model on each imputed dataset
- Evaluate using both train/test split and cross-validation approaches
- Compare performance using cMSE (Censored Mean Squared Error)
- Test with KNN Regression model for comparison

# Results and Analysis from task [3.1]



Task [3.2] - Train models that do not require imputation

What was done in task [3.2]

## Results and Analysis from task [3.2]



# Task [3.3] - Evaluation

# What was done in task [3.3]

## 1. Comparison Analysis

- Built comparison table with all strategies: baseline, imputation methods (Mean, KNN, Iterative), and models handling missing data (Decision Tree, HistGradientBoosting, CatBoost AFT)
- Displayed MSE and cMSE metrics for all approaches
- Created y vs y-hat scatter plots for visual comparison of model performance

## 2. Combined Approach Testing

- Selected best imputation strategy from Task 3.1: Mean Imputation (cMSE: 1.7645)
- Combined with best model from Task 3.2: CatBoost AFT
- Trained CatBoost AFT on mean-imputed data
- Evaluated performance: cMSE of 3.1236 (worse than native missing handling)

## What was done in task [3.3]

### 3. Best Model Selection

- Compared all strategies including combined approach
- Identified CatBoost AFT with native missing support as best performer (cMSE: 1.7339)

### 4. Test Predictions & Submission

- Generated predictions on test data using best model (CatBoost AFT)
- Created Kaggle submission file: `handle-missing-submission-xx.csv`

## Results and Analysis from task [3.3]

# Code Demo

# Overall assessment

# What went wrong

What went great