

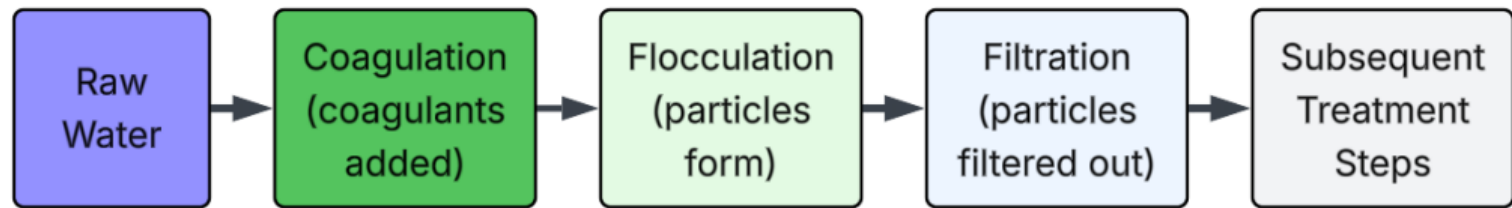
Predicting Water Treatment Plant Chemical Doses

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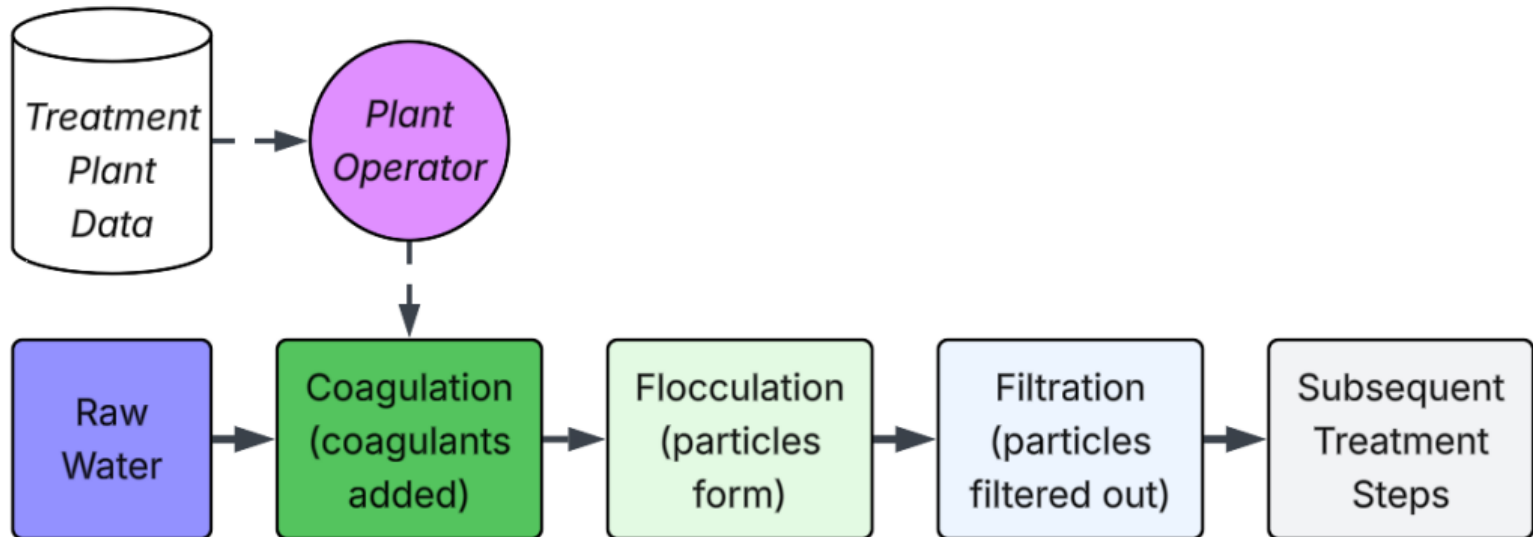
Background and Motivation

The drinking water treatment process takes water from a river, lake, reservoir, or other source, and purifies it to have it reach drinking water standards.



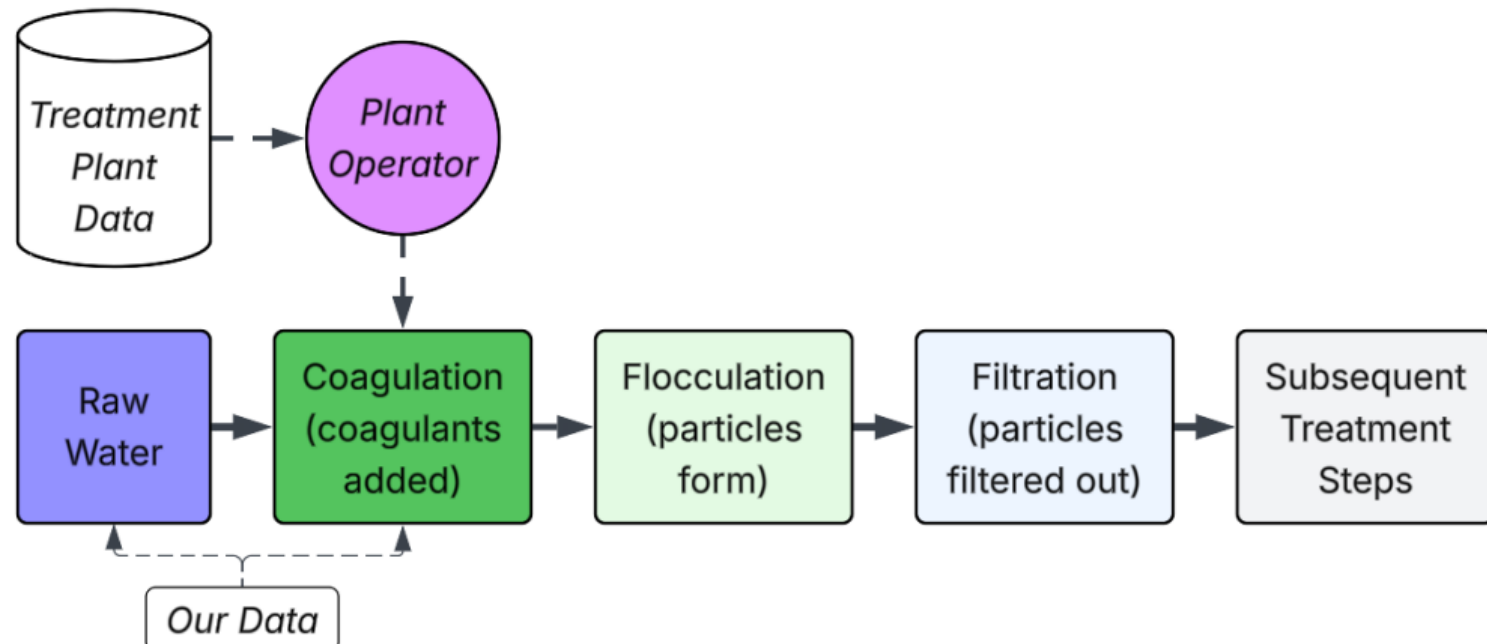
Background and Motivation

One role of an operator in the water treatment process is to look at a wide array of data and make decisions about how to adjust chemical doses.



Background and Motivation

OUR GOAL: Create a model that serves the role of an “operator,” taking raw water data and making predictions of chemical doses.



Data Overview

Three years of data from 2018-2020, provided by a Colorado water treatment plant.

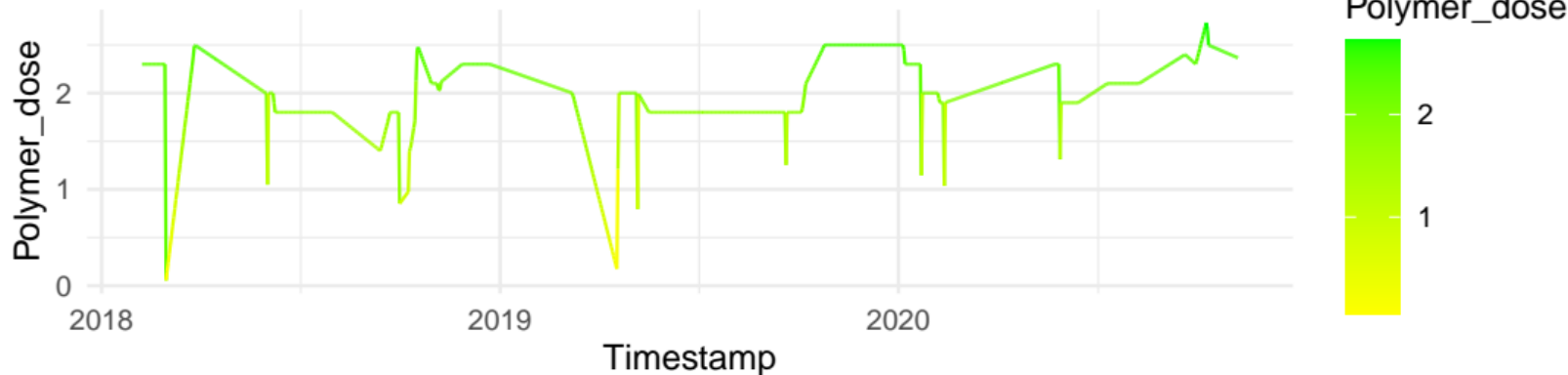
- Raw Water Data
 - pH
 - Temperature (of the water)
 - Turbidity
 - Conductivity
 - Suspended Grain Size
 - Alkalinity
 - Hardness
- Chemical Dosing Data
 - Ferric Chloride Dose: primary additive
 - Cationic Polymer Dose: secondary additive

Data Cleaning

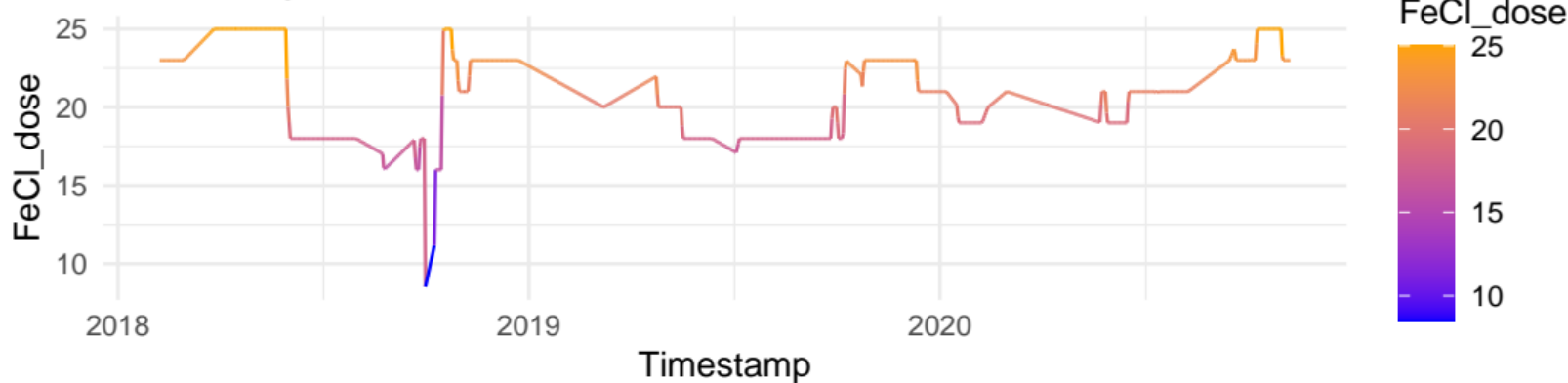
- Dropped Data
 - Suspended grain size was not used as it was highly variable
 - Alkalinity and hardness were not used because of missing data chunks and differing time ranges
- Processing
 - Daily averages were taken for raw water data to match the dosing data's timestep
 - Significant outliers (likely caused by equipment malfunction) were removed or interpolated
 - Dosing and raw water data merged

Exploratory Analysis

Polymer Coagulant

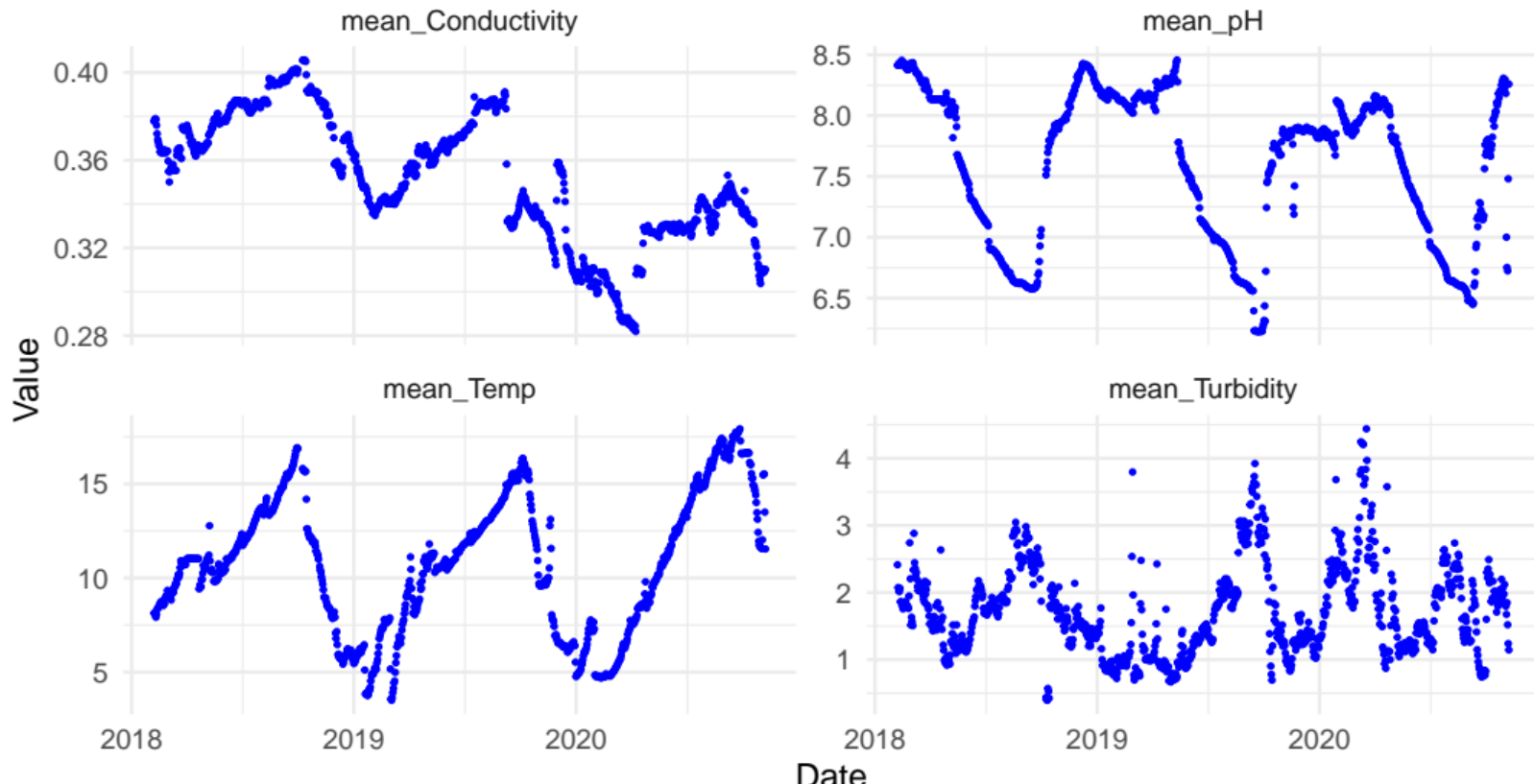


FeCl Coagulant



Exploratory Analysis

Raw Water Characteristics



Predictor Selection

Temperature and pH seem to have the strongest relationship to dose, consistent with feedback from treatment plant operators. Other data were either highly variable or had minimum availability with large gaps. We will focus on the following to predict dose:

- pH
- Temperature
- Conductivity (lots of noise, weaker)
- Turbidity (lots of noise, weaker)

Focus: **FeCl Dose**. Polymer dose is not changed significantly in plants.

Added Variables

Similar to the stock dataset we've seen in labs and homeworks, our time series dataset needs **lagged variables**.

- For each of the main four parameters:
 - Introduce two lagged variables
 - First represents the day before
 - Second represents two days before

We also introduced a **categorical predictor** indicating whether the dose increased, decreased, or stayed the same compared to the day before.

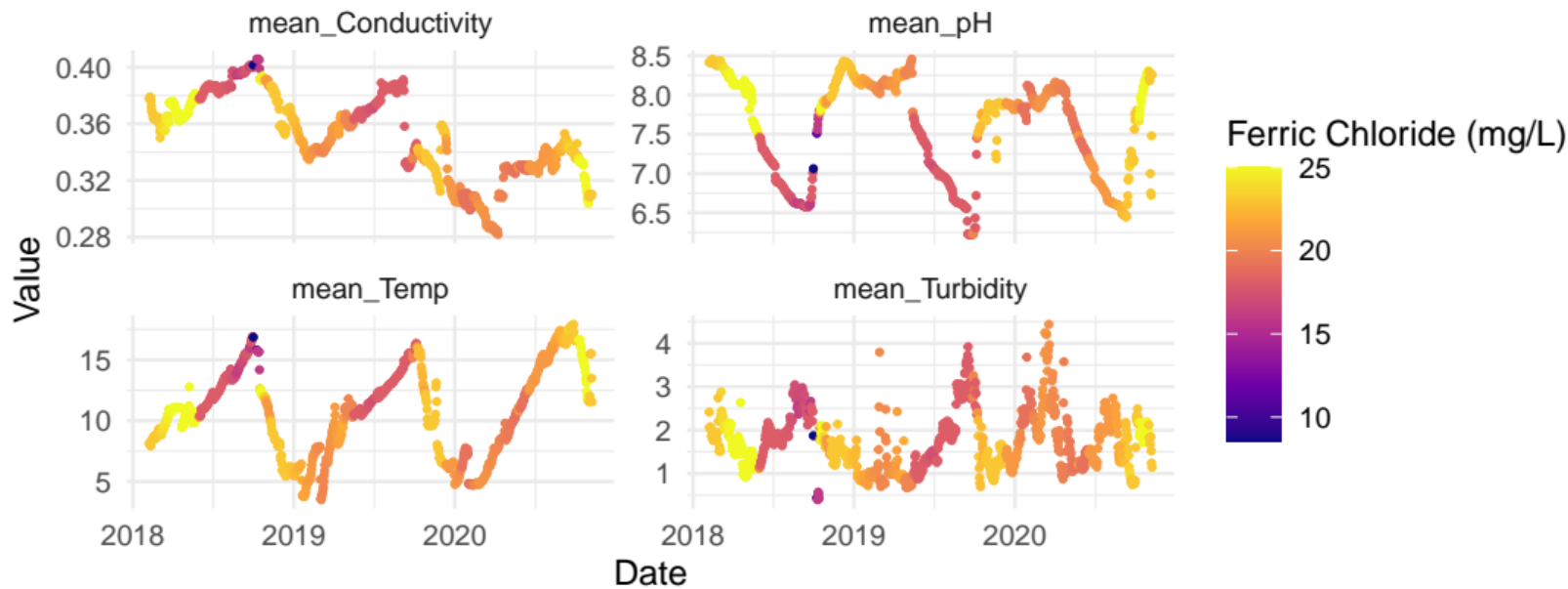
Priorities: Interpretability, Capturing of Nonlinear Behavior

- Training and Testing
 - Training 2018-2019, Testing 2020
 - Approximately 70/30 split
- Coagulant Dose Prediction
 - Linear Regression Models
 - Tree-Based Models
- Coagulant Dose Change Prediction
 - LDA, QDA
 - Multinomial
 - Random Forest

Coagulant Dose Prediction

GOAL: Predict coagulant doses based on raw water characteristics.

Raw Water Characteristics Over Time Colored by Dosing



Coagulant Dose Prediction: Linear Regression

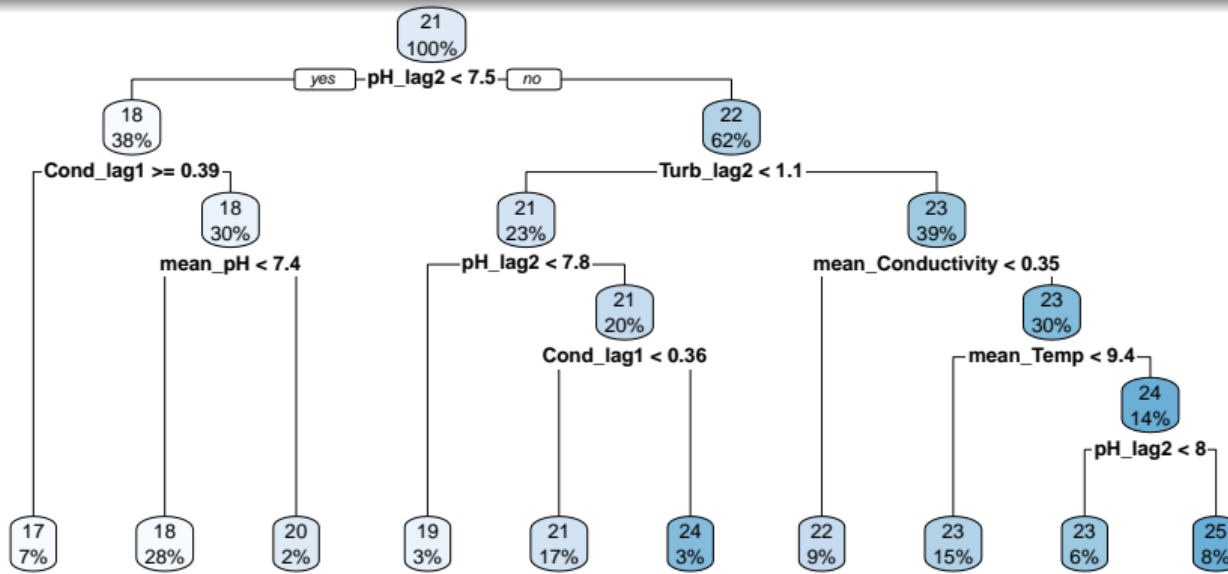
- Created 6 models with different predictors and combinations of interaction terms
- Used lag variables for temp, pH, conductivity, turbidity
- Dropped Turbidity as it was not a good predictor for FeCL dosing
- Difficult to determine what factors of the dataset are driving the predictions
- RMSE and MAE vary widely depending which predictors are selected and the interactions

Coagulant Dose Prediction: Linear Regression

Model Performance Summary

| Model | RMSE | MAE | Description |
|--------------|-------------|------------|--|
| Model 4 | 1.672 | 1.421 | Cond \times pH + Temp \times Cond interactions |
| Model 3 | 1.891 | 1.516 | Temp \times Cond interaction |
| Model 5 | 2.350 | 1.941 | Only Temp and pH |
| Model 0 | 2.971 | 2.580 | Main effects only |
| Model 2 | 3.201 | 2.784 | Temp \times pH + Cond \times Turb interactions |
| Model 1 | 3.483 | 2.938 | Temp \times pH interaction |

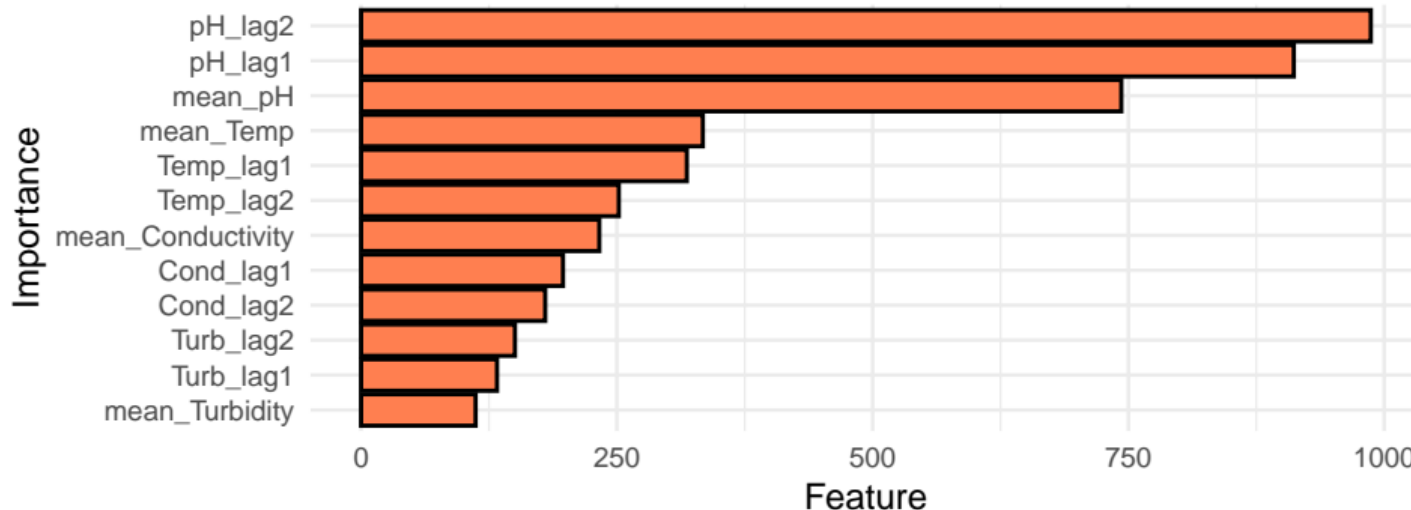
Coagulant Dose Prediction: Decision Tree



| Metric | Value |
|-----------|-------|
| RMSE | 2.756 |
| R-Squared | 0.012 |
| MAE | 2.484 |

Coagulant Dose Prediction: Random Forest

Feature Importance (Random Forest)

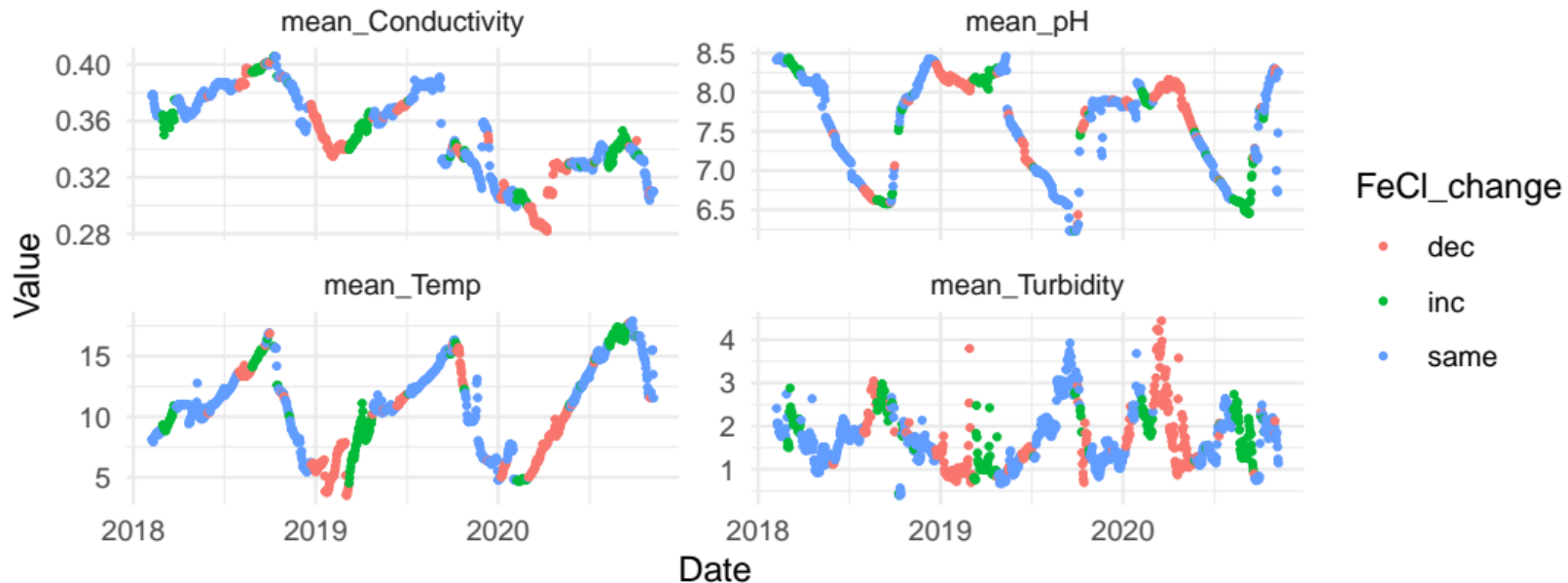


| Metric | Value |
|-----------|-------|
| RMSE | 2.811 |
| R-Squared | 0.021 |
| MAE | 2.283 |

Coagulant Dose Change Prediction

GOAL: Predict whether coagulant dose increased, decreased, or stayed the same based on raw water characteristics.

Raw Water Characteristics Colored by Dose Change



Coagulant Dose Change Prediction: LDA

- Experimented with predictor sets and found that pH and temperature (plus the associated lagged variables) were the main effective predictors.
- Turbidity and conductivity did not improve prediction.

LDA ACCURACY: 43.7%

Table 3: LDA Confusion Matrix

| True | dec | inc | same |
|------|-----|-----|------|
| dec | 26 | 2 | 74 |
| inc | 22 | 6 | 44 |
| same | 17 | 16 | 104 |

Coagulant Dose Change Prediction: QDA

- Experimented with predictor sets and found that pH and temperature (plus the associated lagged variables) were the main effective predictors.
- Turbidity and conductivity did not improve prediction.

QDA ACCURACY: 35.7%

Table 4: QDA Confusion Matrix

| True | dec | inc | same |
|------|-----|-----|------|
| dec | 71 | 2 | 29 |
| inc | 57 | 5 | 10 |
| same | 86 | 16 | 35 |

Coagulant Dose Change Prediction: Multinomial Classification

- Experimented with predictor sets and found that pH and temperature (plus the associated lagged variables) were the main effective predictors.
- Turbidity and conductivity did not improve prediction.

MULTINOMIAL ACCURACY: 46.3%

Table 5: Multinomial Regression Confusion Matrix

| True | dec | inc | same |
|------|-----|-----|------|
| dec | 31 | 2 | 69 |
| inc | 22 | 2 | 48 |
| same | 21 | 5 | 111 |

Coagulant Dose Change Prediction: Random Forest

- Experimented with predictor sets and found that pH and temperature (plus the associated lagged variables) were the main effective predictors.
- Turbidity and conductivity did not improve prediction.

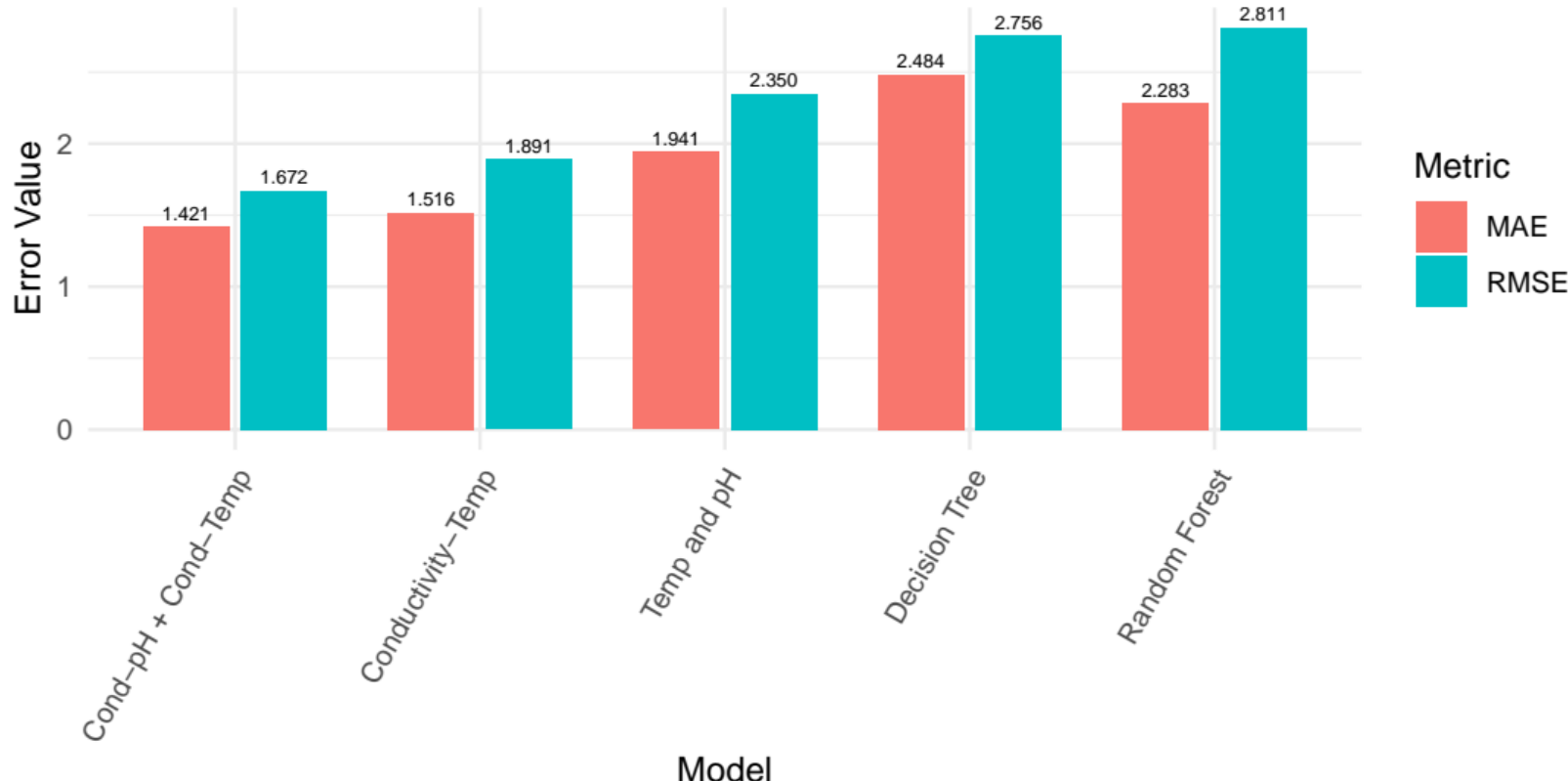
RANDOM FOREST ACCURACY: 39.2%

Table 6: Random Forest Confusion Matrix

| True | dec | inc | same |
|------|-----|-----|------|
| dec | 37 | 6 | 59 |
| inc | 17 | 7 | 48 |
| same | 37 | 22 | 78 |

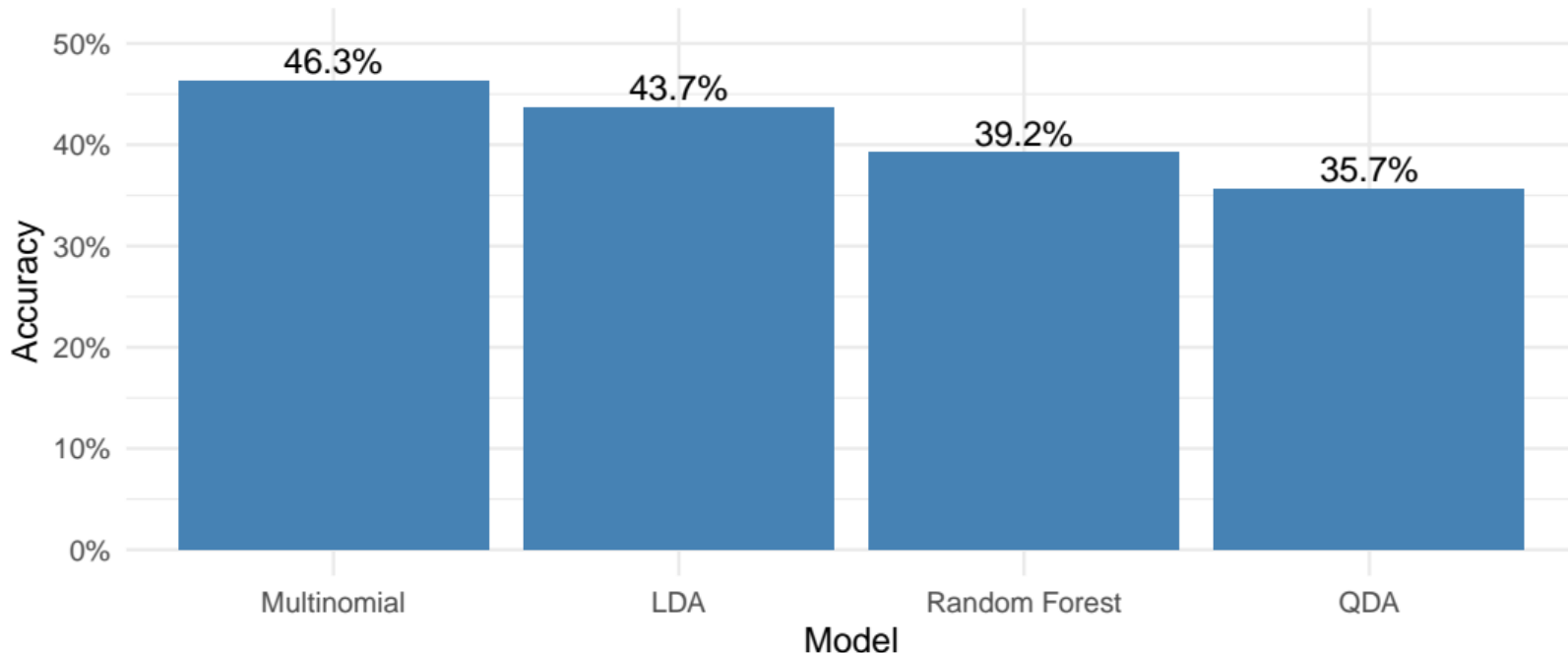
Summary of Dose Prediction Models

Dose Prediction Model Comparison (RMSE and MAE)



Summary of Dose Change Prediction Models

Overall ineffective, and prone to overfitting (especially random forest).
Model predicting no dose change would be more accurate.



Conclusion and Next Steps

- Predicting chemical doses based on these raw water characteristics proved to be ineffective
- Most likely reasons:
 - Limited size of dataset
 - Limited predictor set
 - Operator-dependent
 - Complexity of chemical relationships
 - Other factors (example: reservoir turnover)
- Operators make decisions based on variety of other factors: chemical waste production, filter performance, lab tests, etc.
- Incorporate more varieties and time ranges of data in future modeling efforts