

Forecasting Ammonia Concentrations and Colour Levels using Machine Learning for Reclaimed Water Treatment Operation and Management

by

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August 2022

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This is to certify that I have examined the above MPhil thesis
and have found that it is complete and satisfactory in all respects,
and that any and all revisions required by
the thesis examination committee have been made.

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TABLE OF CONTENTS

Title Page	i
Authorization Page	ii
Signature Page	iii
Acknowledgments	iv
Table of Contents	v
List of Figures	vii
List of Tables	ix
Abstract	x
Chapter 1 Introduction	1
1.1 Background	1
1.2 Objectives	3
1.3 Organization of the thesis	4
Chapter 2 Literature Review	5
2.1 Introduction to water quality control	5
2.1.1 Automated system for water quality control	5
2.1.2 Artificial Intelligence	7
2.1.3 Machine learning and deep learning	8
2.2 Water quality control with machine learning	9
2.2.1 Wastewater treatment plants	11
2.2.2 Water reclamation system	15
2.3 Tools and techniques for enhancing the performance of machine learning modeling	16
2.3.1 Programming languages	16
2.3.2 Data pre-processing	19
2.3.3 Feature engineering	20
Chapter 3 Methods and Materials	22
3.1 Wastewater treatment plant description	22
3.1.1 Process and data sources in SWHEPP	22
3.2 Data collection and preparation	22
3.2.1 On-line data monitoring and collection	22
3.2.2 Loss function for model evaluation	26
3.2.3 Data cleaning and pre-processing	27
3.2.3.1 Data smoothing with Savitzky-Golay and EWMA filter	28
3.2.3.2 Outlier Removal	30

3.2.3.3	Feature Engineering	32
3.2.4	Data transformation	37
3.2.5	Feature selection	40
3.3	Machine learning models	40
3.3.1	Random Forest	40
3.3.2	Deep Neural Networks	41
3.3.3	Recurrent Neural Network	41
3.3.4	Long Short-term Memory	44
3.3.5	Gate Recurrent Unit	45
3.3.6	Configuration of models	45
Chapter 4	Results and Discussion	49
4.1	Baseline performance of the forecasting models	49
4.2	Improved performance on forecasting models using data pre-processing techniques	52
4.2.1	Models trained by pre-processed datasets	52
4.2.2	The effect of window size of data smoothing filters	60
4.3	Exploit hidden patterns in MBR effluent water quality to enhance model performance	62
4.3.1	Ammonia forecasting models	62
4.3.2	Colour forecasting models	63
4.3.3	Model forecasting results on different forecast horizon	65
Chapter 5	Conclusions and Recommendation	73
5.1	Conclusions	73
5.1.1	Machine learning models vs deep learning models	73
5.1.2	Data pre-processing techniques	73
5.1.3	Feature engineering	74
5.2	Recommendations for future research	75

LIST OF FIGURES

2.1	Proposed framework for control strategy design by Ballhysa et al. (2020).	17
3.1	Sewage treatment process flowchart at SWHEPP (adapted from Drainage Services Department 2020)	23
3.2	Colour levels and ammonia concentration are measured in the effluent container (i.e., on the right of the image.) A water pump transports MBR effluent to the effluent container continuously in real-time. The black vault on the left of the image contains a laptop and a colour spectrophotometer.	23
3.3	Instrument of on-line ammonium monitoring system.	24
3.4	Instruments of on-line colour analysis system.	25
3.5	Schematic diagram of the custom-made on-line colour analysis system.	26
3.6	Ammonia and colour data collected from 23 December 2021 to 22 January 2022.	26
3.7	Machine learning model training steps.	27
3.8	Illustration of the influence of different polynomial degrees in the fitting of SG filter and the weight decay with varied alpha values in EWMA filter.	30
3.9	Illustration of the influence of different polynomial degrees in the fitting of SG filter and the weight decay with varied alpha values in EWMA filter.	31
3.10	Illustration of peak analysis. Four important elements are automatic calculated by the function (MathWorks, 2022b).	32
3.11	Sewer system coverage of SHWEPP. The covered areas (i.e., area circled in red boundary) include Fanling/Sheung-Shui new towns and NENT landfill leachate treatment plant.	33
3.12	Analysis of influent quality composition and the illustration of the positional encoding.	34
3.13	Observed ammonia concentration and colour levels in SHWEPP influent.	35
3.14	Hourly water consumption patterns in households (Abu-Bakar et al., 2021). (a) Cumulative pattern and percentage of hourly consumption for households in the “Evening Peak (EP)” cluster (b) Cumulative pattern and percentage of hourly consumption for households in the “Late Morning Peak Peak (LM)” cluster. (c) Cumulative pattern and percentage of hourly consumption for households in the “Early Morning Peak (EM)” cluster. (d) Cumulative pattern and percentage of hourly consumption for households in the “Multiple Peak (MP)” cluster. Consumption is in (m ³).	36
3.15	The daily patterns of ammonia concentration on 3, 7, 11, 15 January 2022.	36

3.16 Monitored colour levels in MBR effluent and the changes in blending ratio (v/v) of treated leachate effluent to municipal wastewater in the inflow of SWHEPP from December 2021 to January 2022. The date of manually calibration and colour level measured in the laboratory are also provided as a black cross and green dot. The moving average of colour level is calculated by averaging the colour level in the past 24 hours. Note: The colour levels analyzed by the on-line colour monitoring system were compared to the manually measured data obtained from the laboratory, which showed errors of 2.08%, 4.05%, 1.11%, 65.25%, 4.94% and 11.0% in the effluent samples collected 5 Oct, 22 Oct, 3 Nov, 15 Nov, 12 Dec, and 31 Dec 2021, respectively.	37
3.17 Concept of forecasting models (Liu, 2020).	39
3.18 Illustration of feature selections for model training.	40
3.19 Illustration of RF and DNN model structure.	42
3.20 Variant architectures of Recurrent Neural Networks (adapted from Olah (2015)). x_t corresponds to the current input, h_{t-1} to the last hidden state (output), h_t to the current output, tanh is the tangent activation function, σ is the sigmoid activation function, \times is the vector pointwise multiplication, + is the vector pointwise addition.	43
3.21 Illustration of how different step sizes of learning rate reach the minimum loss (Ritchie Ng, 2019).	47
4.1 Baseline performance of ammonia and colour forecasting models.	50
4.2 Visulization of the model forecasting results.	53
4.3 Visulization of the model forecasting results.	54
4.4 Illustration of the heterogeneity and homogeneity between validation and different testing datasets.	57
4.5 Baseline performance of ammonia and colour forecasting models.	61
4.6 Comparisons of the model performance in forecasitng ammonia concentrations.	64
4.7 Comparisons of model performance in forecasting colour levels.	65
4.8 Visualization of ammonia forecasting models at forecast horizon of one.	66
4.9 Visualization of ammonia forecasting models at forecast horizon of two.	67
4.10 Visualization of ammonia forecasting models at forecast horizon of three.	68
4.11 Visualization of colour forecasting models at at forecast horizon of one.	69
4.12 Visualization of colour forecasting models at forecast horizon of two.	70
4.13 Visualization of colour forecasting models at forecast horizon of three.	72

LIST OF TABLES

2.1	Endorsed Reclaimed Water Quality Standards from Water Supply Department.	16
3.1	The selected hyperparameters for SG and EWMA filters.	29
3.2	Final model configuration.	48
4.1	Baseline performance of ammonia forecasting model, evaluated on test dataset from 16 to 22 January 2022 . Loss values are calculated by MSE.	55
4.2	Baseline performance of ammonia forecasting model, evaluated on test dataset from 10 to 16 October 2021 . Loss values are calculated by MSE.	58
4.3	Baseline performance of colour forecasting model, evaluated on test dataset from 16 to 22 January 2022 . Loss values are calculated by MSE.	59

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Abstract

Water scarcity is a global challenge, and one of the promising ways to mitigate the water resource crisis is via wastewater reclamation. Reclaimed water can generate non-potable water to substitute the use of drinking water for irrigation or industrial processes. Water quality and aesthetics are the primary concerns in reclaimed water since under-treated water can pose health risks, and the unpleasant colour is likely to induce public misgiving. Ammoniacal nitrogen ($\text{NH}_3\text{-N}$) and colour substances exist in the reclaimed water and can severely affect the reclaimed water quality in different ways. Chlorine is commonly used for reclaimed water disinfection and requires precise dosing to satisfy endorsed quality standards. However, $\text{NH}_3\text{-N}$ consumes chlorine and affects the chlorine dosing. Colour substances do not consume chlorine, but it requires additional efforts and strategies to remove them from the reclaimed water. Therefore, the on-line monitoring of $\text{NH}_3\text{-N}$ and colour are usually practised in reclaimed water facilities to assist in the removal of both substances. However, the conventional on-line analyzers are wet-chemistry-based, and the measurement takes time. The limitation creates a potential issue: there may not be sufficient time for the downstream chlorine dosing system to respond to sudden surges in colour and ammonia levels. To tackle this challenge, this thesis work developed time-series models based on machine learning to forecast the $\text{NH}_3\text{-N}$ concentrations and colour levels in the reclaimed water three hours into the future. For the training dataset, the $\text{NH}_3\text{-N}$ and colour data were collected by an on-line analyzer and a cus-

tomized auto-sampling spectrophotometer, respectively. Both are installed in a reclaimed water treatment facility in Hong Kong. Baseline models for forecasting ammonia concentrations and colour levels were first developed with five machine learning algorithms. Long Short-Term Memory (LSTM) was found to be the most effective algorithm, with the lowest MSE values of 0.0405 and 0.0148 for ammonia and colour forecasting models, respectively. In the training processes, novel data pre-processing methods and feature engineering techniques were implemented to enhance forecasting model performance. The data pre-processing methods were proved to enhance the quality of training datasets and improve the performance of ammonia and colour forecasting models by reducing the MSE values by 4.2% and 8.1%. The feature engineering results supported that the daily fluctuations in NH₃-N and colour have correlations with the urban water consumption patterns. This finding further enhanced the NH₃-N and colour forecasting model performance by reducing MSE by 8.9% and 28.6% compared to baseline models. The established models can be used to assist the disinfection control strategies based on the model predictions using traditional process control systems. This research offers novel methods and feature engineering processes for NH₃-N concentrations and colour levels forecasting in reclaimed water for treatment optimization.

CHAPTER 1

INTRODUCTION

1.1 Background

Urban water challenge increases as the cities grow larger. The World Bank estimates that the urban population worldwide will double by 2050—with severe implications of escalating water demands in cities by 50–70 percent (TheWorldBank, 2021). Global climate change has primarily affected the amount, distribution, and quality of the available fresh water in the urban water cycle. The report from (UNICEF, 2021) points out that one in four cities is facing challenges in supplying adequate water to inhabitants, and the situation is even worse in cities in the developing world. The rise of urban water usage will generate more wastewater. Thus, converting municipal/industrial wastewater into reusable water has recently drawn much attention. Reuse water increases availability by substituting freshwater for non-potable (drinkable) uses for agricultural irrigation, industrial and urban water reuse, etc. The alternative reuse water can supply many activities and save drinking water for other purposes elsewhere (Adewumi et al., 2010).

The construction of reclaimed water facilities often requires a huge amount of capital investment. Upgrading available wastewater treatment plants with reuse water treatment facilities is an economical solution accompanied by the potential of realizing resource recovery (e.g., nitrogen and phosphorus recovery) (Maryam and Büyükgüngör, 2019; Kehrein et al., 2020). The primary concern of reusing treated wastewater is the potential risks caused to public health. Under unexpected circumstances, the reclaimed water facilities can produce unqualified reclaimed water, which is harmful to the living beings (i.e., as reused water is ingested directly or through irrigated crops) and irrigated soil (Adewumi et al., 2010). In Hong Kong, reclaimed water quality is regulated with up to 10 or more water quality parameters, and any parameters that fail to meet the standard will lead to disqualification. The common practice for controlling the treated water quality is achieved through water quality control strategies. The market controllers have evolved from a simple on-off logic controller called Programmable Logic Controller (PLC),

to a more advanced multi-step response controller called proportional-integral-derivative (PID), and finally to the controller consists of machine learning models.

The uses of machine learning models in the water quality controllers for assisting water quality control strategy are ground-breaking applications. Many research papers have proposed various machine learning models for replacing the PLC and PID controllers and demonstrated the benefits of machine learning models. From the study of (Librantz et al., 2018), PID and machine learning-based controllers were deployed to compare the operational costs of dosing the chlorine to the setpoint concentration in a drinking water treatment plant. The results showed that the Artificial Neural Network-based model has a more satisfactory cost reduction in a chlorination dosing control system than the PID controller. Another research finding suggests using a Support Vector Regression (SVR) model as the controller required less time to reach the setpoint concentration of free chlorine residual compared to the PID controller in both simulation and experimental conditions Wang et al. (2020). Incorporating machine learning models in traditional process control systems has also been practiced by Santín et al. (2015) for avoiding violations of total nitrogen in the effluent using the decisions made by Artificial Neural Networks. Long Short-term Model was also used to predict which process control strategy should be selected for eliminating violations of total nitrogen concentrations in the effluent Pisa et al. (2019). Forecasting water quality or predicting future events using machine learning are proved to be effective measures for controlling effluent water quality in wastewater treatment plants, making these approaches to be promising solutions for the reclaimed water treatment operation and management.

The superior performance of machine learning models comes from training high-quality datasets with a good amount of data that can fairly represent the system's dynamics. Most studies have only focused on evaluating the model performance by comparing the test loss values between models and the improvements over PID controllers without considering the collected dataset's quality. The noises in the data and the number of features (i.e., inputs or variables) are the two critical factors affecting machine learning models' accuracy and robustness. Many data pre-processing techniques are proposed and applied to enhance the dataset's quality by removing the data noise. For instance, some papers discussed pre-processed data for removing the noise in raw datasets using data smoothing filters (Cheng et al., 2020), or creating new features in addition to the original ones (Mamandipoor et al., 2020) to achieve data augmentation. Despite the efforts being made, the influences of the

proposed data pre-processing techniques on the model performance have not yet been established.

Machine learning models for water quality control have two main types of algorithms, regression and classification. The former provides forecasting results of specific values, while the latter offers a decision of yes or no (i.e., 1 or 0). The regression model is also called the forecasting model, which plays a vital role in water quality control in drinking water treatment plants (DWTPs) and wastewater treatment plants (WWTPs). The forecasted results can be effectively used to provide critical information for the water quality control strategy. The need to use forecasting models is to cope with the unpredictable nature of water quality and to plan control strategies ahead. Without future information, the treatment operations are less likely to guarantee the production of effluent quality satisfying the government regulation Chen et al. (2003) regardless of how the influent water quality may vary daily. In the reclaimed water system in Shek Wu Hui Effluent Polish Plant (SWHEPP), forecasting models are recommended for effluent treatment management and operation. From the available datasets, we noticed SHWEPP effluent contains ammonia concentrations and colour levels which exceed the reclaimed water standard. To generate non-potable reuse water, it is critical to use on-line data to assist water quality control strategy. Currently, the available on-line sensors on-site are limited. Although the model can only train on limited data, it is still possible to train forecasting models with one feature, known as the univariate forecasting model. In this study, we will attempt to install one more on-line sensor and build machine learning models for forecasting ammonia concentrations and colour levels in the reclaimed water system. Meanwhile, data pre-processing and feature engineering techniques will be proposed and evaluated to address the research gaps of insufficient understanding of data pre-processing in building forecasting models in the reclaimed water system.

1.2 Objectives

The specific objectives of this thesis work are:

- (1) To build baseline univariate forecasting models using machine learning and deep learning models.
- (2) To develop data pre-processing techniques for removing data noise for enhancing model performance.

- (3) To extract relevant information from reclaimed water system using domain knowledge for feature engineering.
- (4) To create new features for augmenting dataset's quality for further improving forecasting model performance.

1.3 Organization of the thesis

In Chapter 1, “Introduction,” the background information, objectives, and organization of the thesis were presented.

Chapter 2, “Literature Review”, provides an overview of water quality control strategies in water treatment plants, wastewater treatment plants, and reclaimed water systems.

In Chapter 3, “Materials and Methods,” the instruments for data collection of ammonia concentrations and colour levels, computer programming environment, and data preparation techniques were summarized. The processes of formulating extra features for training forecasting models were illustrated.

In Chapter 4, “Results and discussion,” the performance of machine learning and deep learning models were compared. Forecasting models trained by different data pre-processing techniques and the influences of feature engineering on model performances were compared with the baseline model performances in test loss.

In Chapter 5, “Conclusions and Recommendations,” the findings obtained from this thesis work were summarized, and possible future studies were recommended.

CHAPTER 2

LITERATURE REVIEW

2.1 Introduction to water quality control

2.1.1 Automated system for water quality control

A programmable logic controller (PLC) is an industrial computer system designed for any process requiring a series of devices and equipment to operate cohesively to achieve multiple purposes in manufacturing or treatment processes. The main components of PLC include a central process unit (CPU), input modules, and output modules (I/O). CPU is responsible for processing digital signals from input modules and sending commands through output modules based on the control logic programmed on the PLC. For chemical dosing control in water treatment plants (WTPs), the PLC system receives readings from turbidity and pH sensors and uses pumps to dose aluminum solution automatically (Andhare and Palkar, 2014). The PLC system with the capability of producing real-time output commands in response to the input signals also makes it widely used in wastewater treatment plants (WWTPs). For oxygen concentration control in the aeration tank, the PLC system receives signals from dissolved oxygen (DO) detectors and transmits signals to open or close the electric butterfly valves to alter the DO concentration (Zhu and Qiu, 2017). Although PLC systems are the most used systems across industries for their easy programming and reliable control, PLC system is merely a device that can be programmed to control operative devices with on-off logic (i.e., a logic control with two states). The straightforward implementation of the PLC system compromised its ability to perform complex tasks in a more dynamic water treatment environment. In reality, many WTPs or WWTPs require precise control of the treatment processes. Being aware of the limitations of the PLC systems, a more advanced controller called proportional–integral–derivative (PID) controller for receiving analog signals was developed to obtain more sophisticated controls over the operative devices.

To react to rapidly-changing environments in wastewater treatment plants, a PID controller generates an output value based on the continuous calculation of an error value

$e(t)$, which is the difference between the desired setpoint (SP) and a measured process variable. Then, the controller applies a correction based on proportional, integral, and derivative terms in the control functions. The use of the "P," "I," and "D" allows the system to quickly reach steady-states with feedback control systems (i.e., the system output is returned to the system input, which is included in the decision-making process of PID controller). Generally speaking, a PID controller is a technology (i.e., a specialist algorithm) for controlling a single device with more complex logic, while a PLC system is a physical system consisting of different modules capable of controlling dozens of devices only with two-state logic. In addition, A PID controller can be designed to operate on a PLC device and provide a more specific control strategy to a designated device. In WWTPs, a single-variable feedback analog control loop in PID can be used to control the temperature in the activated sludge treatment by stabilizing the system temperature in a shorter time (Bados and Morejon, 2020). The feedback control scheme is also applied in DWTPs to adjust the addition of chlorine dosage (i.e., also known as the disinfection process, chlorination, or post-chlorination) to reach the target concentration of free chlorine residual (FRC) (Wang and Xiang, 2019). The disinfection process is carried out in a chlorine contact tank, which provides sufficient time for the chlorine to disinfect pollutants. Since the chlorine added by the dosing device requires time to travel from the entry to the exit, the system output can only reflect the changes in water quality in a delayed time of 30 minutes. In the case of chlorination, the time lag makes feedback control difficult (Kobylinski et al., 2006) as the system is delayed in responding to any sudden surge of the pollutants when it can only receive output at the end of the disinfection process. PID controllers in WWTPs also encounter similar challenges as the increasing complexity of water quality and stricter regulations on the discharged water quality.

Many control strategies are proposed to address the challenges encountered in the process control system. For instance, feed forward-feedback control, linearized and optimal control, model-predictive control, fuzzy control (Demir and Woo, 2014), etc. Among the algorithms used in control strategies, Artificial Intelligence (AI) modeling has gained the most attention in recent years compared to modeling based on mathematical or empirical formulas. In DWTPs or WWTPs, fully understanding the treatment plants' physical, biological, and chemical interactions is very difficult. The unpredictable behaviors during the water treatment can be the significant changes in influent flow rate, water quality fluctuations, the complexity of the biological treatment process, and the large time delay

between control variables and the process inputs, etc. Therefore, AI modeling shows great potential in dealing with the highly complex conditions in the treatment process (Li et al., 2021). The next sections will discuss the applications of different AI modeling methods.

2.1.2 Artificial Intelligence

Artificial intelligence (AI) can perform cognitive tasks with the development of computational solutions. The concepts of AI are usually confused, and in fact, AI is a broad term, and any kind of algorithms or models involved in decision-making with computation fall in the domain of AI. For example, AI can be fuzzy logic and optimization algorithms, which are formulated with human design and involved in the computer decision-making processes. Another subset of AI is called machine learning (ML), but generating an ML model is different from generating a fuzzy logic model. ML uses learning algorithms to generate a model via learning from the historical or large amount of data without being explicitly programmed. ML algorithms can be classified into three categories, which are Supervised, Unsupervised, and Reinforcement learning. In the training process of supervised learning, input variables (x) and output variables (Y) will be provided. The model will learn from the provided datasets to map the x to the Y . A supervised model can generate a prediction based on the new input data (i.e., also called the unseen data). Unsupervised learning is when the dataset is not labeled, the model can learn to infer patterns in the dataset without reference to the known outputs. This type of algorithm can find similarities and differences in the data. In reinforcement learning, models are designed to constantly interact with the environment in a try-and-error way and receive rewards and punishments based on the purpose of the tasks. Generating an optimal action to achieve the lowest penalties is the primary function of a reinforcement learning model. Supervised learning is commonly used for machine learning in water quality control strategies. Regression is a supervised machine learning technique used to predict continuous values. A regression model can estimate the relationship between the input variables in the system and the output target from given datasets and then use the non-linear relationship to map the unseen input data to predicted output data. This type of applications best fits for water quality prediction (Librantz et al., 2018), and sensor fault detection (Cecconi and Rosso, 2021), etc.

2.1.3 Machine learning and deep learning

In machine learning, popular models which researchers frequently use for training predictive models are Supporting Vector Machine (SVM), Random Forest (RF), and Artificial Neural Networks (ANN). RF models are popular due to their superior performance compared to typical machine learning algorithms. Xu et al. (2021) built an RF-based model to predict total nitrogen concentration in water bodies and proved RF models outperformed models such as K nearest neighbor (KNN), Ridge Regression, and Multilayer Perceptron (MLP). The other two widely used models, ANN and SVM, were compared carefully with the reliability and accuracy in predicting 1-day interval T-N concentration in a WWTP (Guo et al., 2015). The results showed that the SVM model has higher accuracy while the ANN model is more reliable for decision-making. Although most of the studies did not focus on the underlying causes of why SVM, RF, and ANN models have more excellent model performance, it would still seem that these models are reliable options for predicting water quality empirically.

As the computing power doubles every 18 months according to Moore's law, implementing Deep Learning (DL)—a subset of machine learning, requires less and less computing time and becomes universal for researchers to solve everyday tasks. One way to explain a DL model is with the definition of having neural networks with more than two hidden layers (i.e., the model complexity increased and required more computing power to calculate). In DL, various architectures are specifically structured based on the problems we attempt to solve. For image processing, Convolutional Neural Network (CNN) is designed to extract essential features from the image vectors. Another famous DL architecture is the Recurrent Neural Network (RNN), which is powerful in solving time series-related applications and Natural Language Processing (NLP) tasks (Li et al., 2018). In particular cases, different DL architectures can be stacked in series to solve specific tasks. A rainfall-runoff prediction model was built using CNN and RNN (Li et al., 2022). The raw data features were extracted by convolution and entered into the RNN models for processing time-series patterns. The results showed a low Kling–Gupta efficiency (KGE) of 0.75 in the high-water period. DL models can be compelling when multiple architectures are stacked into a single model to perform a specific task, which machine learning models cannot realize. That being said, DL models can achieve higher model performance in terms of prediction accuracy compared to ML models.

2.2 Water quality control with machine learning

A drinking water treatment plant (DWTP) produces potable (i.e., drinking water) water for human consumption by removing contaminants from the source water, such as lakes or streams, or from underground aquifers. The raw water enters DWTPs and goes through treatment units of coagulation, flocculation, sedimentation, filtration, and disinfection in sequence as the primary treatment scheme in the conventional DWTPs (Li et al., 2021). During the treatment process, colloids, suspended matter, pathogenic microorganisms, and organic matter are removed to meet the regulated standard. However, raw water quality is not always stable, and corresponding actions must be promptly adopted when events like the surge of pollutants or the large variability of the influent flow. In any event, the treated water from DWTPs should generate drinking water that complies with the World Health Organization's Guidelines (i.e., WHO guideline) for drinking water quality. Otherwise, the treated drinking water would either be discharged, resulting in the short-term outage of water supply to the downstream cities; or the users will receive contaminated drinking water, which can transmit diseases and cause illness.

Turbidity is one of the critical water quality indicators, which can be defined as the "optical quality" of water. The unit describing the turbidity is the Nephelometric Turbidity Unit (NTU). High turbidity levels in raw water can impede the effectiveness of filtration and chlorination processes and potentially cause short-term outages of the water supply. Heavy rainfall and fissures within the aquifer can also lead to turbidity events that are most likely to cause high turbidity (World Health Organization, 2017). The challenge in the event of high turbidity in raw water is that it occurs rapidly, and mitigating activities must be actionable immediately. To address the sudden event of such, Stevenson and Bravo (2019) trained forecasting models based on general linear model (GLM) and RF to predict the time when the turbidity reaches higher than 7 NTU. The results indicate that both models can successfully predict the events (i.e., with accuracy between 0.81 and 0.86), and the RF model is found to have higher precision due to its ability to capture the nonlinear relationship between rainfall (mm) and turbidity (NTU).

To maintain operational costs and water quality in the coagulation process, the amount of coagulant, mainly subject to the turbidity and alkalinity in the raw water, is traditionally determined through manual sampling and analysis. Jar tests are designed to find the optimal chemical dosage for coagulation to remove the turbidity in raw water. The entire

process includes on-site sampling and more than 40 minutes of laboratory work (Gani et al., 2017). To replace the laborious jar tests, Wang et al. (2022) proposed using principal component regression (PCR), support vector regression (SVR), and long short-term memory (LSTM) neural network to build predictive models for estimating daily chemical dosage. Compared with the linear PCR model, nonlinear SVR and LSTM models capture more relationships between the chemical dose (e.g., ferric sulfate) and the raw water quality based on a higher R-squared value of 0.70.

Disinfection is the last step of water treatment processes in drinking water treatment plants to generate safe potable water. In this step, chemical disinfectants such as chlorine, chloramine, or chlorine dioxide are added into the water to inactivate any remaining pathogenic microorganisms. However, the chlorination process requires precise dosing of disinfectant—too high will lead to the formation of disinfection byproducts (DBPs), and too low will result in insufficient levels of the residual disinfectant concentration. In both scenarios, the treated drinking water can pose health threats to the end-users. Although the PID controller can achieve automatic dosing of disinfectants according to the change in water quality, Wang et al. (2020) proposed a model predictive control based on machine learning models to improve the dosing process further. The study indicated that the predicted chlorine dosage from a Support Vector Regression (SVR) model could help the free chlorine residual in the water reach the setpoint concentration in a shorter time compared to the PID controller in both simulations and experimental conditions. Machine learning models can not only reduce the time required to reach setpoint concentration but also decrease the chemical usage required in DWTPs. An Artificial Neural Network-based model has proved to optimize the treatment operation by reducing the chemical usage in a chlorination dosing control system compared to using PID controller (Librantz et al., 2018).

The invariability of the raw water quality is always a big issue for disinfection. For instance, chlorine dose can be excessively dosed when the treated water contains fewer pollutants (e.g., non-organic matters and ammonia nitrogen). Excessive chlorine in water results in the waste of chemicals, which is reflected in the increased operational cost and the generation of undesired disinfection by-products (e.g., trihalomethanes (THMs), which are carcinogenic to humans). Xu et al. (2022) trained an ANN model for predicting the occurrence of THMs in tap water using simple and straightforward water quality parameters (e.g., pH, temperature, UVA_{254} and residual chlorine (Cl_2)). Despite the fact

that the results showed a good model accuracy in predicting for THMs (i.e., T-THMs, TCM, and BDCM), the applications of the model are largely limited in reality due to the lack of dataset regarding quantity and quality. In fact, the lack of high-quality datasets for training ML models is a common issue, which explains, until recently, mathematical or empirical-based AI models like fuzzy logic (Gamiz et al., 2020; Godo-Pla et al., 2021) is still widely used for process control in WTPs.

2.2.1 Wastewater treatment plants

Human activities produce wastewater and discharge it from homes, businesses, factories, and commercial activities to the sewage systems which connect to wastewater treatment plants (WWTPs). The function of WWTPs is to remove contaminants from sewage and water so that the treated water can be returned to the natural water body without endangering any living beings residing in the ecosystem. Undertreated wastewater can lead to harmful algal blooms or cause oxygen deficit in the water (i.e., low oxygen content can kill the fish). The steps for treating municipal wastewater involve three major categories—primary treatment, secondary treatment, and tertiary treatment. Most of the particular matters will be removed in primary treatment via settling or floating; a secondary treatment is mainly responsible for removing BOD_5 in the biological processes; in the final tertiary treatment, membrane filtration, adsorption by activated carbon, and addition of disinfectant are applied optionally to further eliminate the undesired pollutants in the water.

Wastewater is categorized and defined according to its sources of origin. Domestic wastewater is water discharged from residential sources generated by kitchen wastewater, cleaning, and personal hygiene. Industrial/commercial wastewater is generated and discharged from manufacturing and commercial activities, such as the textile industry and food and beverage processing wastewater. Institutional wastewater is generated by large institutions such as hospitals and educational facilities. Regardless of the source of the wastewater, WWTPs have to achieve at least three sustainability targets: environmental protection (i.e., minimum pollutants discharge), social acceptance (i.e., human sanitary protection), and economic development (i.e., feasible operational and management costs) (Mannina et al., 2019). To effectively achieve these goals, process control is required to reduce energy consumption, improve effluent quality, and save costs in plant operation and

management. The focus of this study is on discussing the development of using process control for treatment operation and management.

Under known operational conditions of a WWTP, machine learning models can be trained to assist the plant operators in optimizing treatment processes to improve effluent quality. Wang et al. (2021) proposed a machine learning framework, utilizing a model based on Random Forest to predict the effluent Total Suspended Solid (TSS) and phosphate (PO_4). This study uses data from six on-line sensors (i.e., flow rate, TSS, pH, PO_4 , temperature, and total solids (TS) meters) across the treatment line to train the RF model. The results indicated that the influent temperature is the most influential variable for both TSS and PO_4 in the effluent, and PO_4 depends strongly on the TSS in aeration basins, etc. It has been suggested that the combined use of the RF model and analytical tools allows the author to pinpoint the critical factors influencing the effluent quality, which is regarded as an innovative approach. However, several significant drawbacks hinder such model developments using on-line sensors to collect training data. The term "training data" is a dataset used to feed into the model for the model to learn and pick up hidden patterns in the data. Many of the existing WWTPs and DWTPs are not equipped with on-line sensors, and a lack of automation and instrumentation is universal. The difficulties in installing on-line sensors include the extra costs of purchasing hardware, extra labor works for maintenance, and most importantly, the optimal locations for sensor installation.

In secondary treatment, the relationships between the sludge and wastewater quality are complex due to the complex interactions between the microorganisms and the organic matters in the reactor (Wilén et al., 2018). To fully understand and describe the interactions in such systems requires a substantial amount of data. However, installing on-line sensors everywhere in the system is impossible. Zaghloul et al. (2021) attempted to find out the ideal locations and adequate number for on-line sensor installation. The author used the data collected from the on-line sensors installed in three lab-scaled secondary treatment reactors to train machine learning models to predict effluent quality. In addition, considering the intricacy of operational conditions in the secondary treatment, the author claimed that with the use of feature selection and ensemble model (i.e., average results from multiple model outputs), overfitting could be prevented. The issue of overfitting can be understood as the model memorizing the noises too much in a training dataset, resulting the poor performance when the model is used to predict outputs from

unseen data.

Similar to the secondary treatment units, an electrocoagulation reactor is also a complex system in which the operation and management are based on pH value, current density, flow rate, and the initial concentration of heavy metal ions, etc. Interestingly, instead of using an ensemble model to prevent the overfitting issue claimed by Zaghloul et al. (2021), Zhu et al. (2021) used a deep learning Long and Short-term model (LSTM) and an error compensate Autoregressive Integrated Moving Average model (ARIMA) to predict the removal rate of heavy metal ion concentration in wastewater. An LSTM-ARIMA model has strengthened the model performance compared to the solely used LSTM or ARIMA model in predicting removal rate shown by the Results. A possible rationalization of using an LSTM model without worrying about model overfitting is that deep learning is sophisticated enough for learning the nonlinear patterns in complex systems, while machine learning models like RF might fail to capture the intricate relationships, resulting in overfitting.

Technological advancement allows easy access to real-time water quality data via on-line sensors. The collected real-time data can be used to train predictive models and assist the plant operation and management. Despite the advantages of what on-line sensors are capable of, sensor calibration and maintenance are critical. The malfunctioned sensor can induce wrong decisions for plant operation, ultimately deteriorating treatment efficiency in WWTPs. Haimi et al. (2015) suggested that reliable and moderately-priced on-line sensors are not always available; in addition, sensor malfunctions (i.e., fouling or erroneous measurement) can cause the down-time of the sensors. For the unavailable sensors (i.e., "hard-to-measure" or expensive sensors), many research works have proposed building "soft sensors." Instead of using hardware sensors to measure the water parameters, the soft sensor generates real-time values through a machine learning model, which is trained by other easy-to-measure water quality data. In the works of Wang et al. (2019), easy-to-measure variables such as pH, flow rate, TSS, and ammonium nitrate ($\text{NH}_4\text{-N}$) are input to machine learning models to predict hard-to-measure water quality parameters of COD and total phosphate (TP). Pattnaik et al. (2021) also used DO, pH, conductivity, turbidity, and temperature to train a model to predict BOD. It is believed that both research works can solve the issues of the unavailability of specific water quality sensors.

The automated treatment operation and management heavily rely on the reliability

of the on-line sensors; thus, preventing and the early detection of sensors malfunctioned is the utmost concern to the plant operators. Sensor fault detections can be categorized into three groups which are (1) individual faults— outlier data that can be distinguished concerning other data points; (2) contextual faults—an anomalous instance in a specific context and normal in another; (3) collective faults—a cluster of rare instances with respect to other data trends (Chandola, 2009). Many research papers have proposed using machine learning models to help identify sensor fouling.

Two main algorithms, regression and classification, can be used to find fouling signals. A regression algorithm can identify fouling signals by calculating the difference between model-predicted outputs (e.g., ammonium or COD concentration) to the actual signals. A classification algorithm can distinguish fouling signals through the direct outputs of the model (i.e., the model outputs 2 class labels, one represents normal, and the other is abnormal). Cecconi and Rosso (2021) proposed an ammonium fault detection mechanism, utilizing a regression ANN model, along with principal component analysis (PCA) and Shewhart monitoring charts (i.e., statistical control chart). The remarkable idea of this study is to analyze the residual between the predicted ammonium and the actual ammonium sensor signal and identify the individual and contextual faults with the help of statistical tools. Despite the fact that the accuracy of the fault detection mechanism can reach R^2 value of 0.87, the method comes with significant limitations. The author points out that to maintain the high accuracy of the predictive model, the quality of the input data needs to be carefully attended to by performing manual cleaning procedures on a weekly basis.

Research has focused on solving collective faults in sensor fault detection rather than collective faults. The primary reason is that collectives faults are hidden in regular signals, and the expert can only discover the irregularity by comparing sets of signals in series. Thus classification technique using deep learning is proposed to address collective faults in the works of Mamandipoor et al. (2020). It is believed that this is the first research paper using an LSTM network to achieve a fully automatic fault detection method in WWTPs. In contrast to other works, input variables for model training heavily rely on the experts' manual selection before inputting into models like PCA and fuzzy neural networks. The significance of using a deep learning network is its capability to capture long-term temporal dependencies from a large dataset compared to machine learning models (i.e., PCA-SVM model). The results showed that the accuracy (i.e., F1-score) from the LSTM model is

92%, outperforming the PCA-SVM model of 87%. This finding suggests that using DL models in classification problems is promising for solving collective faults.

2.2.2 Water reclamation system

The increasing demand for water in cities is mainly attributed to the rapid urbanization and the population moving from rural to urban centers. In many major cities, the evergrowing water usage and wastewater discharge drive the development of water reclamation (Lyu et al., 2016). In WWTPs, the technologies applied in water reuse include disinfecting with chlorine addition, ultra-violet (UV) irradiation, biological treatment, membrane filtration, etc (Norton-Brandão et al., 2013). However, even with the most advanced water treatment technology, the treated reclaimed water quality is still subject to the variability and variations of pollutant contents in wastewater effluent (Chen et al., 2003), and can potentially fail to meet the reclaimed water standard. The research studies propose to apply machine learning techniques to assist the disinfection process in water reclamation can be categorized into three groups (1) optimize the treatment management in WWTPs to alleviate the loadings of water reclamation process (Al-Ghazawi and Alawneh, 2021; Viet et al., 2021); (2) actively branch out the desired, and undesired wastewater effluent for subsequent disinfection process of water reuse or direct disposal into water body (Chen et al., 2003); (3) adapt process control methods to stabilize the disinfection performance in the reclaimed water system (Demir and Woo, 2014).

Technology advancement and research studies on water reuse have been discussed for more than two decades. However, there are not too many research publications that aim to improve the reclaimed water system as a whole in recent years. The economic reasons behind constructing water reuse facilities could be a major obstacle for the government sectors. The economic burden of either building new reclaimed water institutions in new locations or retrofitting existing WWTPs is overwhelming (Adewumi et al., 2010). To discover more values and reusable resources from water reuse, Chojnacka et al. (2020) takes the circular economy perspective into accelerating the process of adopting water reuse systems for agriculture production. The author introduces the potential of gradually replacing chemical fertilizers with partially treated wastewater for sustainable crop production despite there are many limitations to be overcome. In Italy, the circular concept is also studied by Colella et al. (2021). Four different resource recovery senarios were

brought up, and two of the scenarios include the nutrient recovery turned into nitrogen and phosphorus fertilizers. Several researchers in recent years have provided the overall potential and challenges of treated wastewater reuse in the world; it is believed the day of using reused water universally will soon come with collaboration across different disciplines.

Reclaimed water for non-potable reuses can serve for irrigation for agriculture, toilet flushing, and irrigation for landscaping, etc. Water Supply Department (WSD) will soon implement a reclaimed water supply system in SWHEPP by disinfecting the tertiary-treated sewage (i.e., MBR permeate). The produced reclaimed water will be served for non-potable reuse and is required to satisfy the water quality standards shown in Table. 2.1.

Table 2.1: Endorsed Reclaimed Water Quality Standards from Water Supply Department.

Parameter	Unit	Requirement ^a
<i>E. coli</i>	cfu/100 mL	Not detectable
Colour	Hazen Unit	≤ 20
Ammoniacal Nitrogen ($\text{NH}_3\text{-N}$)	mg/L as N	≤ 1
Total Residual Chlorine	mg/L	≥ 0.2
Dissolved Oxygen	mg/L	≥ 0.2
Turbidity	NTU	≤ 5
5-day Biochemical Oxygen Demand	mg/L	≤ 1
pH	-	6-9
Threshold Odour Number	-	≤ 100
Synthetic detergents	mg/L	≤ 5

^a The water quality standards for all parameters are applicable at the point-of-use of the system.

2.3 Tools and techniques for enhancing the performance of machine learning modeling

2.3.1 Programming languages

Matrix Laboratory (Matlab) is a proprietary programming and numeric computing platform used across industries and academia for data analysis, algorithm developments, and model buildings. In the wastewater treatment industry, Matlab is known for using an add-on software called Simulink for modeling, simulating, and analyzing the dynamic

system (i.e., chemically enhanced primary clarifier (Bachis et al., 2015). The use of Matlab-Simulink in the wastewater treatment industry is known for the development of control strategies for WWTP automation. In 1987, International Water Association (IWA) developed the first mathematical model for simulation-based evaluation, which is Activated Sludge Model 1 (ASM 1), and the modified activated sludge models and Benchmark Simulation Models (BSM) was further developed in the following years (bin Talib, 2011). The difference between the two is that ASM is designed for developing control strategies exclusively in the activated sludge treatment process, and BSM 1 is to develop the automation in the entire WWTP (Ballhysa et al., 2020).

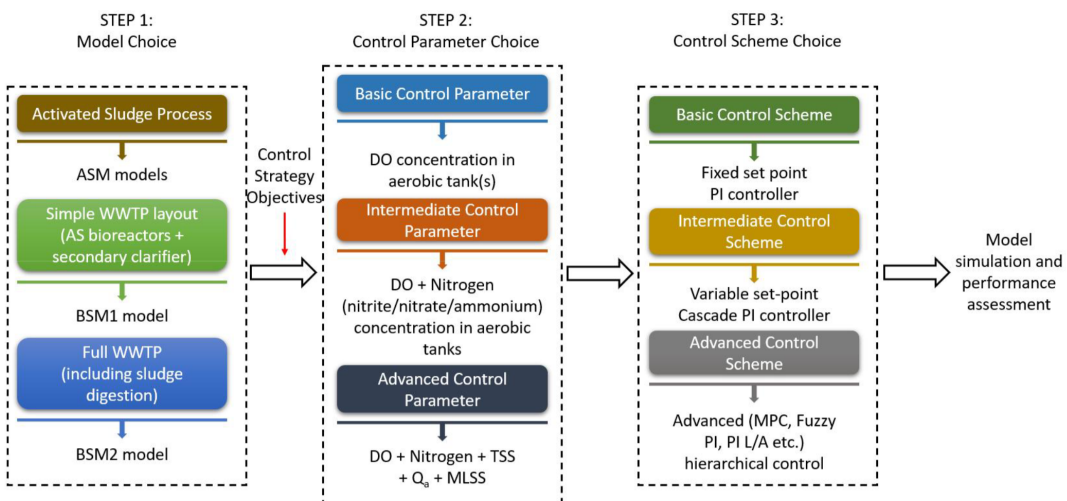


Figure 2.1: Proposed framework for control strategy design by Ballhysa et al. (2020).

In recent years, many publications presented an exciting way to demonstrate how machine learning-based model predictive control (MPC) can outperform the conventional PID controller in WWTPs using BSM. Fig. 2.1 shows how the control scheme choice iterates from fixed set point PI controller to advanced MPC control, etc. The researchers use Matlab-Simulink to simulate the treatment processes in WWTPs. At the same time, the block of PID controllers is replaced with machine learning models, and the effluent quality or treatment system performance can be differentiated via BSM simulated results. Wang et al. (2020) compared the stability of chlorinated water quality in the effluent of a DWTP with two control strategies, which are PID feedback controls and a predictive model-based support vector machine (SVM). The BSM simulated results showed that the SVM model required 21 minutes less to reach the residual chlorine setpoint than PID feedback controls. A proposed neuro-fuzzy PID controller (i.e., a hybrid machine learning model consisting of neural networks and fuzzy logic) also showed superior performance

in optimizing the chlorine dosing rate to minimize the chance of errors (Hong et al., 2012). The significance of using BSM in Matlab-Simulink enables the performance of traditional and machine learning-based control strategies can be compared in objective and fair scenarios, also providing the practicability of machine learning to the experts in the field. Matlab is a powerful and resourceful platform providing various machine learning functions, including point-and-click apps for training and evaluation, available classification and regression algorithms, Automatic machine learning (AutoML), etc (MathWorks, 2022c). The direct access to the abundant features and the integration of Simulink make Matlab an appealing option for many researchers in the wastewater treatment industry, especially in the research domain of machine learning and control strategy simulation. Despite the countless benefits of using Matlab, the Python programming language stands out differently.

Python is a high-level, interpreted, and object-oriented programming language and features simple and easy-to-learn syntax providing good readability (Fundation, 2022). The large developer community (e.g., GitHub and Stackoverflow) and open-source access (i.e., free of charge) have made Python an ideal tool for machine learning starters. The most cutting-edge research in the field of Artificial Intelligence is often led by Tech Giants like Google and Amazon, which conduct research via Python (e.g., machine learning frameworks of TensorFlow (Google)in Python), as well as the big research community using Python. All the latest updates and developments relating to machine learning architectures and techniques are usually accessible in the open-source Python community, including the example codes. Contrary to Python, users of commercial software Matlab need to wait for the software engineers working in Matlab to update the latest machine learning applications onto the Matlab platform, which is a time-consuming process and creates a delay of time and accessibilities to many resources (Castro, 2018). Machine learning developers in the wastewater treatment industry can freely choose between the programming methods based on the research need. Those looking for mature machine learning algorithms can simply use Matlab and be satisfied with the functionalities; on the other hand, those who intend to incorporate more new techniques and architectures in machine learning models can consider using Python as the programming language. Interestingly, MathWorks recently announced using Python functions in Simulink Model (MathWorks, 2022a); despite the update from Matlab, to the best of my knowledge, there are no research papers developing machine learning algorithms on Python and running

on Matlab-Simulink.

2.3.2 Data pre-processing

The ubiquitous sensors installed in WWTPs for treatment automation generate a massive amount of data on a daily basis. Before being used for any purposes, the data must be understandable for explanation and relevant enough for water experts to extract valuable information (Kehrein et al., 2020). Without the help of Artificial Intelligence, data manipulation before training machine learning models can be time-consuming and challenging. The specifically designed algorithms can perform data evaluation and augmentation to improve data quality. Any statistical or machine learning algorithms which can complete these tasks are known as the data pre-processing techniques. The causes of sensors rendering undesired data with low quality are the limitations of the hardware sensors and the dynamics of the sampling locations. In general, the false data generated by sensors can be described in eight distinct states (Rosen et al., 2008; Newhart et al., 2019):

- 1) Operational: Sensor is working properly with normal measurement noise.
- 2) Excessive drift: When a sensor outputs a value progressively further from the true-value.
- 3) Shift: When the output of the sensor is a constant amount away from its true value.
- 4) Fixed value: When the sensor is stuck and keeps repeating the same value.
- 5) Complete failure: Similar to a fixed value fault, but the sensors either give off the maximum or minimum, value, zero or no value at all.
- 6) Wrong gain: When signals away from the calibration point are under- or over-amplified by the sensor.
- 7) Calibration: The sharp change in sensor output directly following a calibration.
- 8) Isolated fault: When a single point in a series shows an incorrect value.

The researchers and experts have been proposing solutions for filling the data gaps created by sensor faults and maintenance operations. However, the number and length of missing values are mainly subject to the dynamics of the system being monitored and other factors. In their open-source wastewater data treatment toolkit, De Mulder et al. (2018) has recommended five data imputation strategies aimed at data generated from

water resource recovery facilities:

- 1) Interpolate.
- 2) Use a correlation with other available measurement signals.
- 3) Replace with a corresponding value in an average daily profile.
- 4) Repeat the values obtained on the preceding day.
- 5) Replace with the output of a model.

The efficient monitoring of sensors and proper use of the data for developing control strategies in the wastewater treatment industry rely on careful data quality control. In recent years, automated data evaluation has drawn the attention of experts and researchers in this field as manual detection of sensor fouling is unrealistic because the tasks are labor-intensive and laborious. Alferes et al. (2013) presented three practical approaches for data quality validation, which are capable of automated calculating single abnormal values and collective faults over a long period. The author claimed that the significance of the research work is performing a data quality validation scheme on the multivariate dataset. The pitfall of the study is that despite the promising approaches proposed, the validity still depends on the thresholds or acceptability limits in the actual WWTPs. Similar to the data imputation strategies, the real situation differs tremendously across different WWTPs. That being said, instead of providing general guidance on how to manipulate data, the focus should be emphasized on how to use algorithms to help users understand, analyze, and process the fouling data.

2.3.3 Feature engineering

Feature engineering aims to enrich the raw dataset by selecting, manipulating, and transforming data, which forms a better dataset relating to the underlying targets to be learned by the machine learning model. Feature engineering and data pre-processing are easily confused with each other. The fundamental difference between the two is that the former creates essential features not included in the raw data; the latter is a data noise removal and cleaning process. In the study of Mamandipoor et al. (2020), feature engineering was performed to generate five extra features, which are the statistical metrics of mean, maximum, minimum, variance, and standard deviation of a specific input feature. However, in comparing the final results, the author only emphasized evaluating model

accuracies across varied machine learning models (i.e, PCA-SVM and LSTM models). Another interesting technique used by Zaghloul et al. (2021) is to create the gradient values of an input variable to assist the model in better learning the trend of the historical removal rate of water parameters in aerobic granular sludge reactors. Similar to the results shown in the work of Mamandipoor et al. (2020), the influence of how engineered features affect the ultimate model accuracy is excluded in the results and discussion part. Thus, creating a lack of knowledge in how significant the feature engineered inputs are to the model accuracy and which techniques can be used in which scenarios.

There is considerable ambiguity concerning the necessity of using feature-engineered inputs in training predictive models in WWTPs. In predicting total nitrogen (TN) in the effluent, Guo et al. (2015) input nine features and performed feature sensitivity analysis, which can capture the change of the output values attributed to the change input. The result showed that the top three most significant inputs, temperature, TN flow, and pH, are critical in predicting TN. The author claimed physical related cause-and-effect relationships between the effluent TN and those top three effective features could be elucidated by the machine learning model. In another work on predicting influent BOD concentration, the study clearly stated that using five inputs instead of three will cause model overfitting. Three inputs for model training were considered sufficient (Alsulaili and Refaie, 2021). Variables created from feature engineering have no physical properties, leading to extra unexplainable essence in addition to the black-box nature of machine learning models. Besides, extra model inputs from feature engineering can also cause overfitting if the data quality is not carefully evaluated. Said by Andrew Ng, "Coming up with features is difficult, time-consuming, requires expert knowledge. Applied machine learning is basically feature engineering". From the quote and the recent studies, we are uncertain how feature engineering techniques can practically help the development of machine learning models in the wastewater treatment industry. More research is required to elucidate further the effectiveness of performing feature engineering.

CHAPTER 3

METHODS AND MATERIALS

3.1 Wastewater treatment plant description

3.1.1 Process and data sources in SWHEPP

Shek Wu Hui Effluent Polish Plant (SWHEPP) is a secondary sewage treatment plant that treats the municipal wastewater from Sheung Shui/Fanling Districts and the treated leachate effluent from North East New Territories (NENT) leachate treatment plant. The plant was designed for 300,000 population equivalents (PE) in 2001, and in 2009, the daily treatment capacity was expanded from 80,000 m³/day to 93,000 m³/day. SHWEPP is operated and maintained by Drainage Services Department (DSD), and the plant will be upgraded to a tertiary treatment level to increase the treatment capacity of 190,000 m³/day by the end of 2025. As shown in Fig. 3.1, the treatment plant consists of primary sedimentation, secondary biological treatment, and final sedimentation, followed by a membrane bioreactor (MBR), which provides an advanced level of organic and suspended solids removal. A low volume of the MBR effluent is pumped to an effluent container n the MBR location to monitor the effluent quality in real-time. An ammoniacal nitrogen on-line sensor and a colour level on-line analyzer are installed in the effluent container, indicated as (a) and (b) in Fig. 3.1.

3.2 Data collection and preparation

3.2.1 On-line data monitoring and collection

To enable us to perform on-line monitoring of ammonium concentration (NH₃-N) in the MBR effluent, an Ammonium and Potassium Probe, AmmoLyt®Plus 700 IQ (Xylem Company) is installed as Fig. 3.3a in the effluent container, as shown in Fig. 3.2. The operation was commenced on 27 April 2021 and completed on 27 March 2022. The ion-selective electrode (ISE) probe provides continuous and reagentless monitoring of ammonium and potassium at the configured interval of one measurement per minute. Due

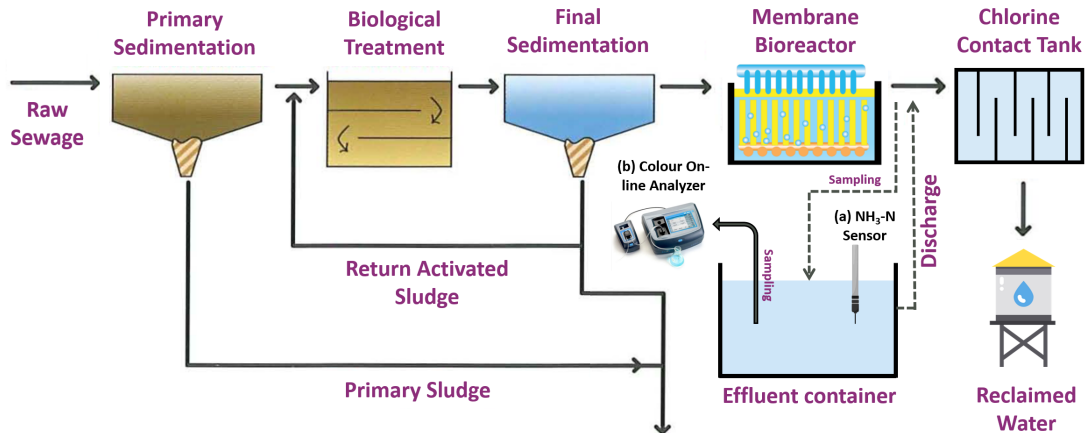


Figure 3.1: Sewage treatment process flowchart at SWHEPP (adapted from Drainage Services Department 2020)



Figure 3.2: Colour levels and ammonia concentration are measured in the effluent container (i.e., on the right of the image.) A water pump transports MBR effluent to the effluent container continuously in real-time. The black vault on the left of the image contains a laptop and a colour spectrophotometer.

to the ISE probe cannot differentiate the potential difference caused by ammonium and potassium ions in the electrodes, the on-line monitoring of ammonium concentration requires continuous calibration using potassium concentration.

The instrument records ammonium concentration as NH₄-N mg/L, a form to express the sum of nitrogen found in reduced nitrogen (III) form. Ammonia has a reported pKa of 9.25 (National Center for Biotechnology Information, 2022), meaning ammonium is a

primary species under the pH of 9.25 in water. In WWTPs, the pH in water typically ranges from pH of 7–8, making the NH₄-N concentration the dominant species. Both ammonia and ammonium contain one nitrogen atom; 1 mg/L NH₃-N is the same as 1 mg/L NH₄-N. Thus, to prevent confusion, in the following paragraph, the unit of NH₄-N will be expressed by NH₃-N, which is the unit used in the water quality standard. The collection of on-line ammonia data is achieved through downloading CSV files from the website connected to the IQ Sensor Controller (Xylem Company), as shown in Fig. 3.3b.



(a) AmmoLyt®Plus 700 IQ,
Xylem.
(b) DIQ/S 284-EF controller,
Xylem.

Figure 3.3: Instrument of on-line ammonium monitoring system.

Hourly monitoring of the colour levels of MBR effluent was conducted from 5 October 2021 to 26 February 2022 by using a custom-made on-line colour analyzer. The default spectrophotometer as Fig. 3.4a and a peristaltic pump as Fig. 3.4b is only capable of initiating a single measurement of colour level by pressing the "READ" button on the DR3900 panel. To achieve continuous sampling and analyzing colour levels without human intervention, an actuator with a programmable time function was mounted on the panel of DR3900, as shown in Fig. 3.4c.

The automatic sampling and analyzing of the colour level begins with the actuator clicking on the "READ" button to initiate the colour analysis at a fixed interval of 30 minutes. 3 mL of sample was collected from the effluent container and delivered to the spectrophotometer cell. After the spectrophotometer analyzed the sample, the data was transmitted to an automatic data acquisition and storage software pre-installed on the laptop. The DR3900 device is connected to a laptop, which receives the real-time data and stores it on data management software from Hach company. To access the real-time data from the laptop, Google Remote Desktop is used to operate the laptop via Internet cloud

services using any devices having access to the Internet. The entire process is illustrated in Fig. 3.5. After the measurement, the sample will be discharged to the effluent container, and the on-line colour monitoring system is left idle until the subsequent measurement.



(a) SIP10 peristaltic pump,
Hach Company



(b) DR3900 spectrophotometer,
Hach Company



(c) Customized clicker/actuator with programmable timer

Figure 3.4: Instruments of on-line colour analysis system.

The maintenance and calibration of the DR3900 spectrophotometer are performed on a weekly basis. During the maintenance, the DR3900 device was shut off, and 100 mg/L chlorine solution was pumped into the sampling tubes and the plastic cuvette for disinfection and cleansing. The cleansing of the tubes and cuvette were manually inspected with eyes to make sure no foreign objects were stuck inside. De-ionized water was brought to the site to perform the spectrophotometer calibration after the reboot of DR3900.

In the proposed model training methods, ammonia and colour data are input into the training forecasting models. Thus, the colour and ammonia data as features should be collected from the same period of time with the same dataset size. In addition, abnormal data caused by sensor downtime should also be excluded. Thus, we chose the ammonia and colour data from 23 December 2021 to 22 January, as shown in Fig. 3.6.

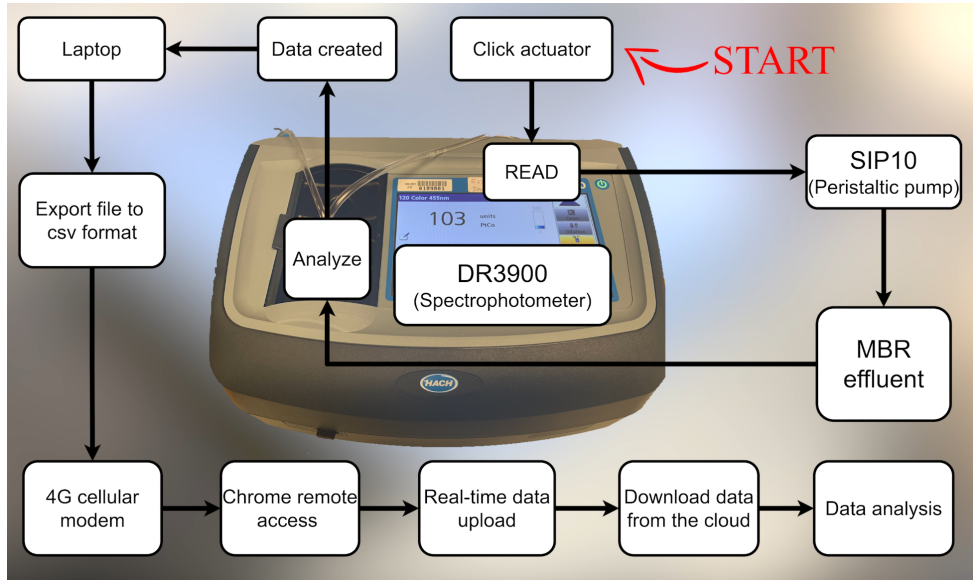


Figure 3.5: Schematic diagram of the custom-made on-line colour analysis system.

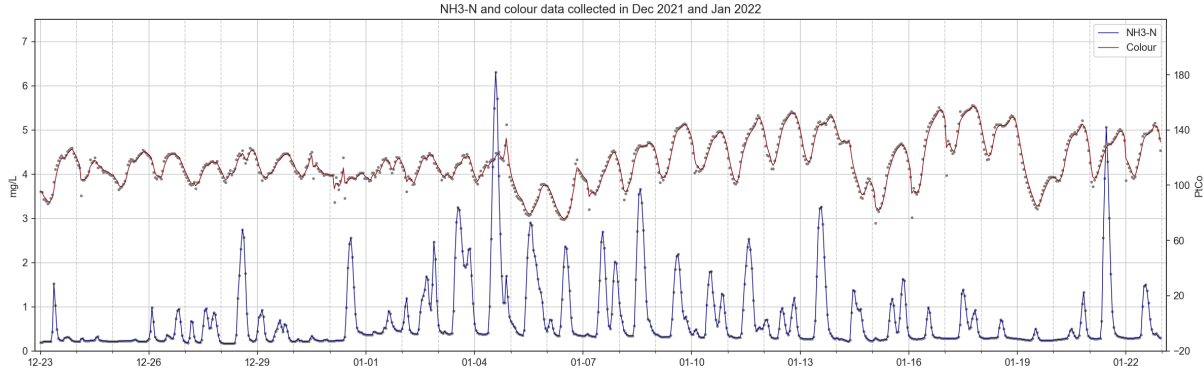


Figure 3.6: Ammonia and colour data collected from 23 December 2021 to 22 January 2022.

3.2.2 Loss function for model evaluation

Loss functions are used to determine the error between the model outputs (i.e., prediction or forecasting values) and the given target value (DeepAI, 2022). The bigger the difference between the ground truth \mathbf{y} and the model outputs $\hat{\mathbf{y}}$, the higher the value of the loss function is, meaning the model performed poorer. A low value for the loss means the model performed well. The selection of the types of loss function is essential for training the model to perform specific tasks. This study uses Mean Squared Error (MSE) to evaluate the regression models. The values of MSE will never be negative and are formally defined by the following equation:

$$MSE = \frac{\sum(y_i - \hat{y}_i)^2}{n} \quad (3.2.1)$$

3.2.3 Data cleaning and pre-processing

In this study, ammonia concentrations and colour levels forecasting models will be trained, and the model training steps are shown in Fig. 3.7. The training processes are split into two sections; one is the baseline model training steps, and the other is the proposed model training steps. The training steps of the first section used cleaned data to train forecasting models and generated baseline model performance, which will be further compared with the model performance generated in the second section. The second section includes using pre-processed datasets (i.e., data smoothing) and feature engineering enhanced datasets to train the forecasting model.

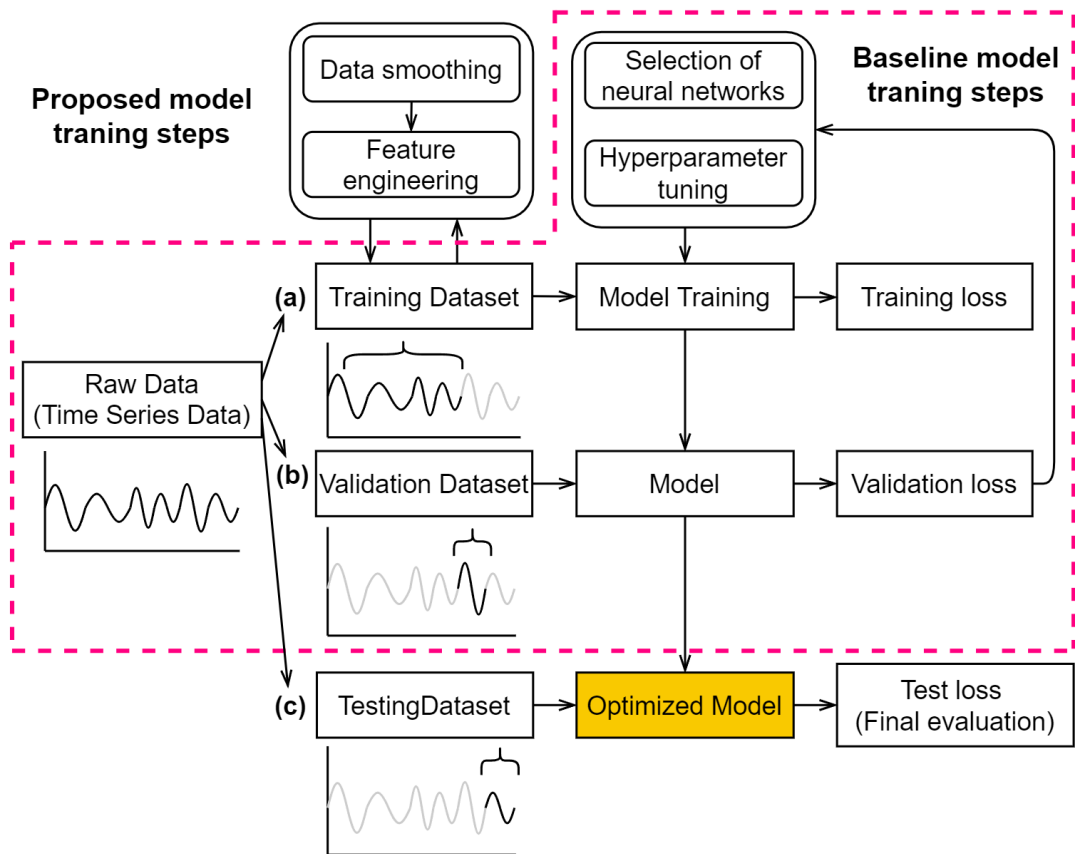


Figure 3.7: Machine learning model training steps.

The raw data embedded in the original CSV files has many problems, such as missing values, extreme low or high values, unreadable texts, etc. Thus, data cleaning and pre-processing are necessary for a more effective model training process. Python programming

language and related libraries such as Numpy and Pandas were used to clean and pre-process the raw dataset for further usage. The raw ammonia dataset collected from the instrument contained 44,640 samples (data points) with eight variables, giving a matrix size of 44,640 x 8, and the samples were collected in time series at 1-minute intervals. The colour level raw dataset collected from the laptop contained 1488 samples with 34 variables, giving a matrix size of 1488 x 34, and the samples were collected in time series at 30-minute intervals.

Extreme values were manually removed before the colour and ammonia datasets were averaged into time-series data at 1-hour intervals. For the ammonia dataset, we replaced the values higher than 7.0 mg/L with NaN (i.e., Not a number), and further interpolation was used to fill up the NaN along with the missing values in the dataset. For colour dataset, we manually took out the relatively low data points on the days when the maintenance and calibration tasks were performed; extremely values higher than 300 Hazen Unit were also replaced by NaN. Same as the data cleaning method used for the ammonia dataset, the missing values and NaN were filled up with interpolation.

3.2.3.1 Data smoothing with Savitzky-Golay and EWMA filter

Data smoothing was performed using the same methods on ammonia concentrations and colour levels datasets. One of the effective ways to remove the noise from the dataset is to apply data smoothing filters. Two filters were applied in this study, Savitzky-Golay (SG) and Exponentially Weighted Moving Average (EWMA) filters.

An SG filter is a digital filter that can be applied to a set of digital data points for the purpose of smoothing the data without distorting the data tendency. This is achieved via convolution by fitting successive subsets of adjacent data points with a low-degree polynomial using linear least-squares (Wikipedia, 2022b). The illustration is shown in Fig. 3.8a, and the procedures of how data points are smoothed are presented in the following steps:

- 1) Extract short-time window (i.e., blue dots in Fig.3.8a)
- 2) Determine polynomial degree (e.g., different polynomial degree is compared in Fig. 3.8a).
- 3) Find the smoothed data point (i.e., at center of the window).
- 4) Repeat for shifted window (e.g., similar to moving average).

The equation to describe the smoothed value of \mathbf{Y}_j can be expressed in Eq. 3.2.2:

$$Y_j = (C \otimes y)_j = \sum_{i=\frac{1-m}{2}}^{\frac{m-1}{2}} C_i y_{j+i}, \frac{m+1}{2} \leq j \leq n - \frac{m-1}{2} \quad (3.2.2)$$

where Y_j corresponds to the j^{th} smoothed data point, m to the window size (i.e., numer of data points intended to smooth out) and C_i to the convolution coefficients (i.e., determined by Savitzky and Golay (1964)).

Exponentially weighted moving average (EWMA), also known as autoregressive (AR) filtering, is a technique that filters measurements. An EWMA filter smoothes a measured data point by exponentially averaging that particular point with all previous measurements. The EWMA equation can be expressed in Eq. 3.2.3:

$$\begin{aligned} \alpha &= \frac{2}{span + 1} \\ y_0 &= x_0 \\ y_t &= (1 - \alpha)y_{t-1} + \alpha x_t \end{aligned} \quad (3.2.3)$$

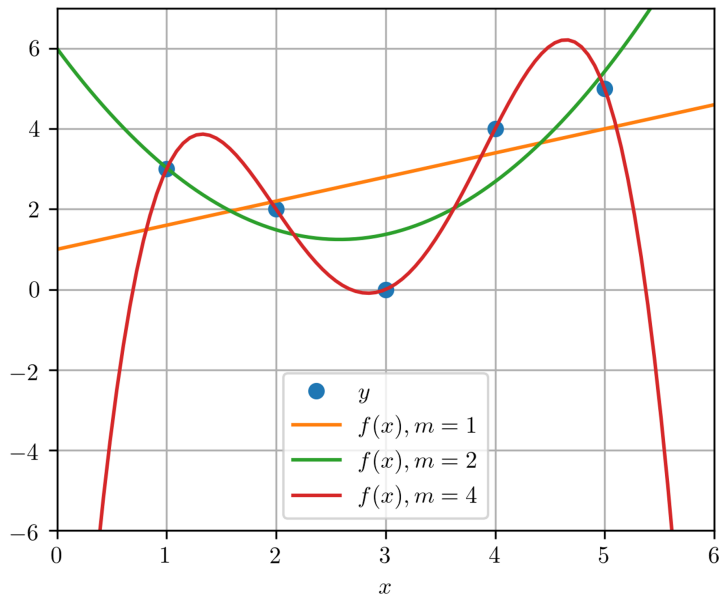
where α corresponds to the decay paratmeters, x_t to the value at a time period, y_t to the value of the EWMA at any time period t, span to the window size.

Both SG and EWMA filters are required to select the hyperparameters, the selected values are presented in Table. 3.1.

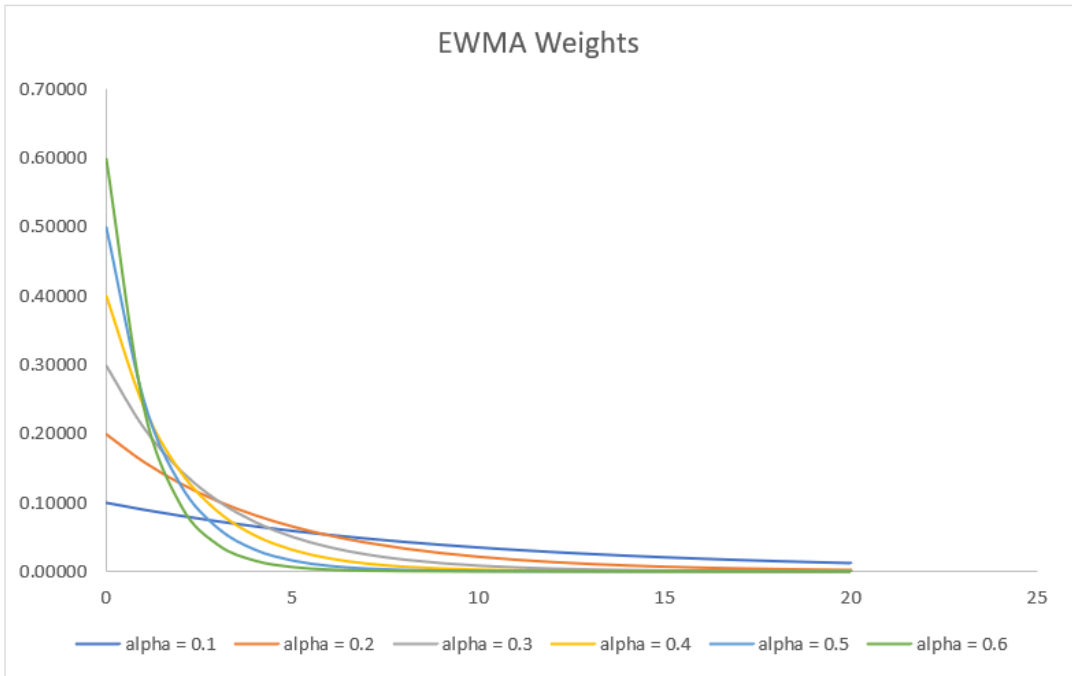
Table 3.1: The selected hyperparameters for SG and EWMA filters.

Group Name	Window size	Polynomial degree
SG-5	5	2
SG-7	7	2
SG-9	9	2
EWMA-2	2	-
EWMA-3	3	-
EWMA-4	4	-

Fig. 3.9 shows the influences of different windows sizes on SG filters as in Fig. 3.9a and on EWMA filters as in Fig. 3.9b.



(a) SG filter with different polynomial degree (Taal, 2017).

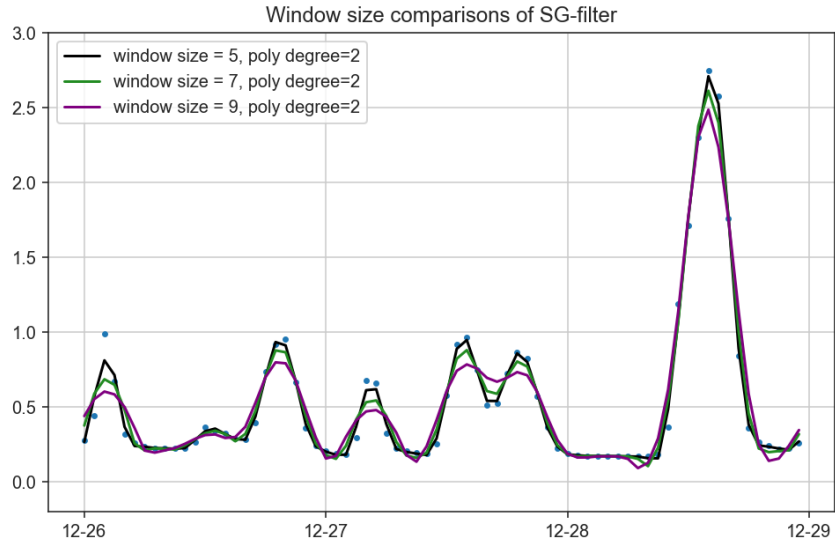


(b) Examples of weights with exponential decay at varied alpha values (CFI, 2022).

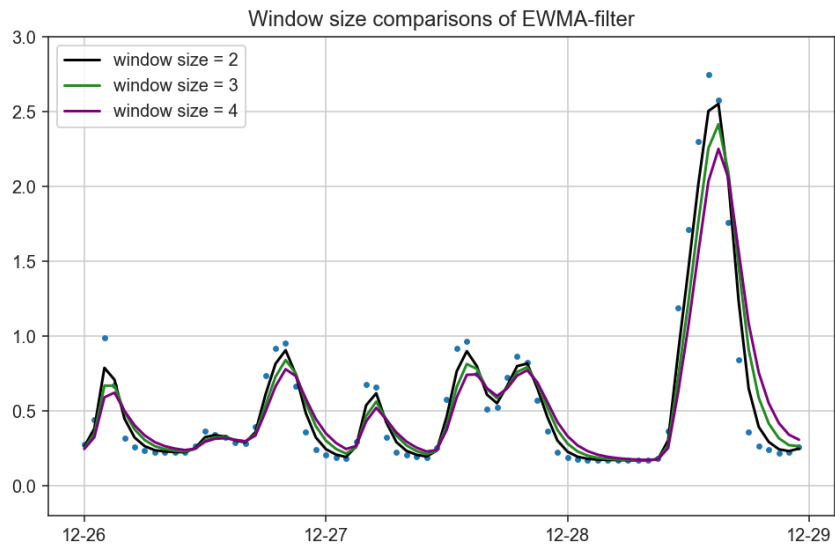
Figure 3.8: Illustration of the influence of different polynomial degrees in the fitting of SG filter and the weight decay with varied alpha values in EWMA filter.

3.2.3.2 Outlier Removal

Although the extreme values in the raw ammonia dataset were removed based on basic rules (i.e., concentration higher than 7.0 mg/L), the ammonia sensor can still collectively



(a) Ammonia data filtered by SG filters with different window sizes.



(b) Ammonia data filtered by EWMA filters with different window sizes.

Figure 3.9: Illustration of the influence of different polynomial degrees in the fitting of SG filter and the weight decay with varied alpha values in EWMA filter.

capture unideal data points. In the outlier removal process, we intended to identify the collective faults of ammonia data in the unit of an entire day. Two abnormal conditions are defined to determine whether the ammonia data on a specific day shows collective fault:

- 1) $\text{NH}_3\text{-N}$ fluctuation ≤ 0.1 (i.e., lower than the sensor resolution).
- 2) No diurnal fluctuation (i.e., Fluctuation = peak value – bottom line value).

Peak analysis was performed on the daily ammonia data to automatically identify normal or abnormal signals. The analysis takes a one-dimension array (i.e., the data form of ammonia in a day) and finds all local maximum values by comparing neighbouring values. This function will also provide information such as width and prominence, as in Fig. 3.10 to help us identify whether the diurnal fluctuation exists.

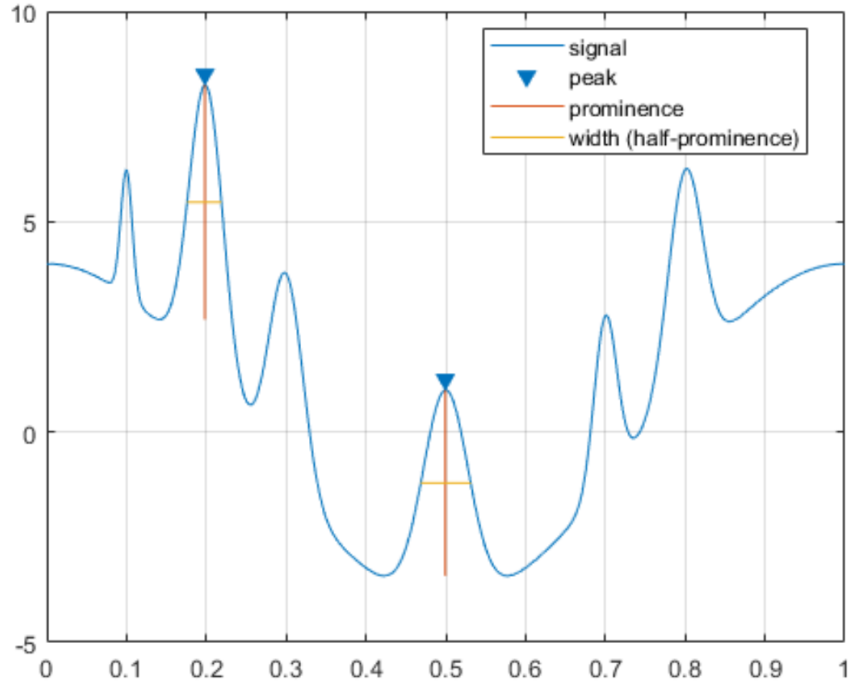


Figure 3.10: Illustration of peak analysis. Four important elements are automatic calculated by the function (MathWorks, 2022b).

3.2.3.3 Feature Engineering

We have carefully observed and analyzed the SWHEPP influent to create new features from the raw datasets based on our domain knowledge. We discovered that the SWHEPP influent consists of treated landfill effluent from NENT landfill leachate site and municipal wastewater, as shown in Fig. 3.11. We observed that with a higher blending ratio, which is calculated from the daily volume of treated leachate effluent divided by the daily inflow volume of SHWEPP, the colours level are also higher, as shown in Fig 3.13a. With the Pearson coefficient of 0.68, the increased volume of treated leachate effluent in the public sewage system is proportional to the increase of the colour levels in the SHWEPP influent, while the ammonia concentration is mainly from the municipal wastewater. During the mixing of both types of wastewater, as in Fig. 3.12a, substances contributing to colour levels were diluted by the municipal wastewater; at the same time, the ammonia

concentration was also diluted by the treated leachate effluent. In Fig. 3.13b, we can observe that the time when the lowest colour level of the day occurred is close to when the highest ammonia concentration was observed. The changes in colour levels and ammonia concentration are interactive. Thus, in feature engineering, colour level data was selected for training the ammonia forecasting model; ammonia data was selected for the training colour forecasting model, as shown in Fig. 3.18.

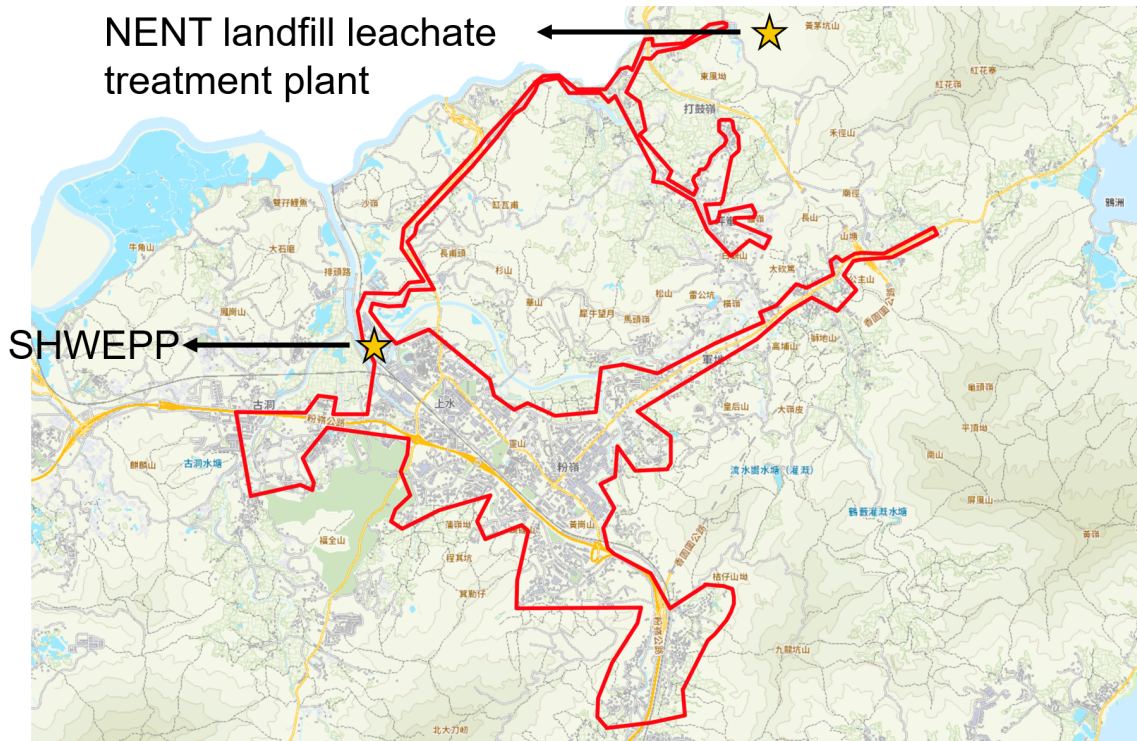
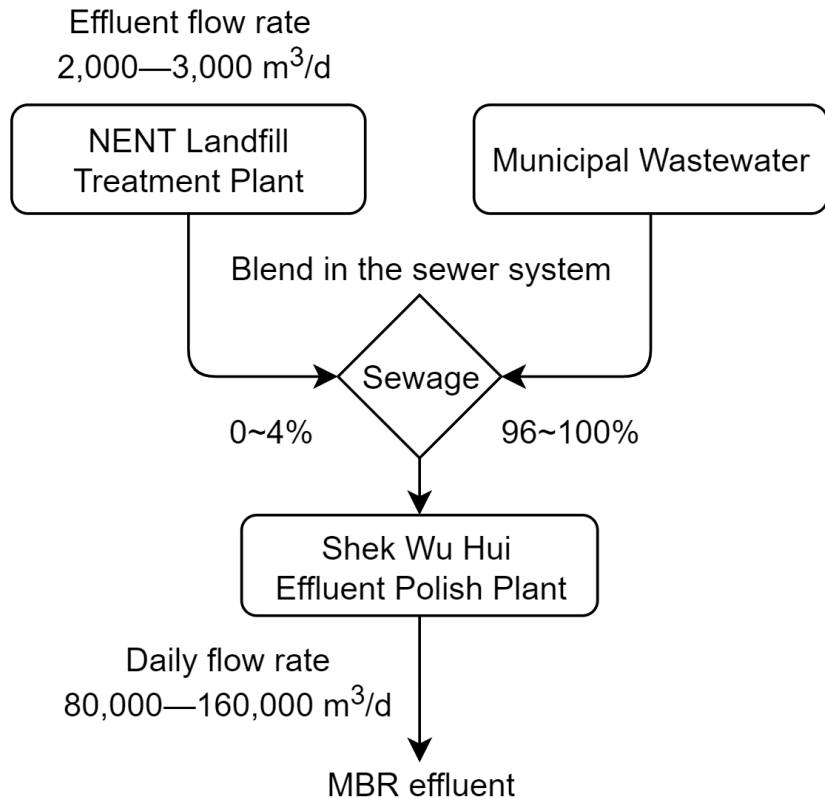
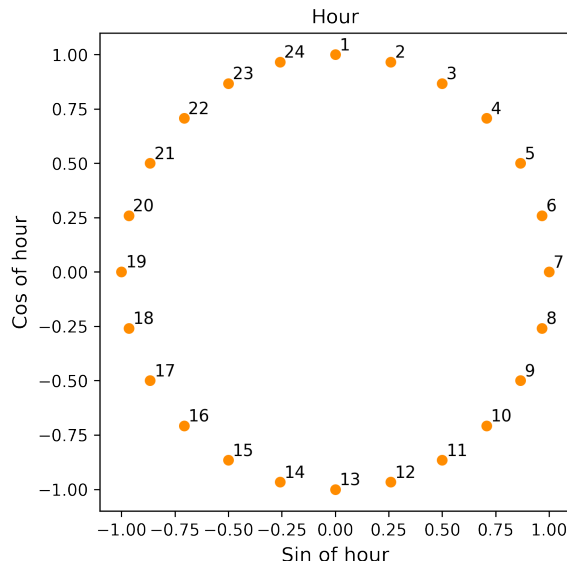


Figure 3.11: Sewer system coverage of SHWEPP. The covered areas (i.e., area circled in red boundary) include Fanling/Sheung-Shui new towns and NENT landfill leachate treatment plant.

The new features are inspired by the research work of Abu-Bakar et al. (2021). The author summarized the four types of hourly household water consumption patterns as in Fig. 3.14, which correlates the specific time of the day to the volume of water consumed in households. In other words, as fresh water is consumed, wastewater is generated simultaneously; the wastewater then enters the public sewage system and increases ammonia concentration. As shown in Fig. 3.15, the peak analysis tool helped us to identify the ammonia concentration's peak hour, which occurred around 13:00 to 14:00, and 20:00 to 21:00. Thus, it is convinced that time features will help the machine learning models correlate better and predict the change of ammonia concentration in the wastewater. Time feature is created through a technique called positional encoding (POS). The positional



(a) Flowchart showing the blending of treated leachate effluent with municipal wastewater.

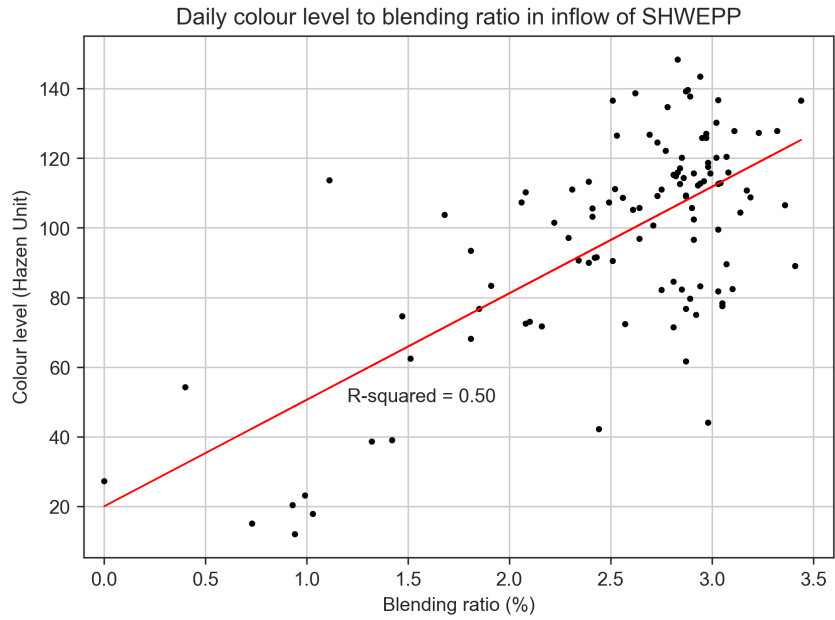


(b) Positional encoding of hour components.

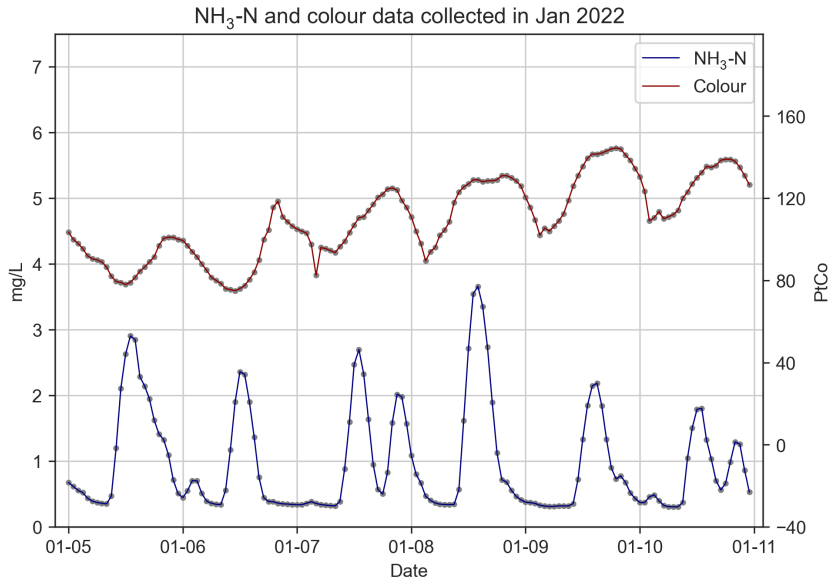
Figure 3.12: Analysis of influent quality composition and the illustration of the positional encoding.

encoded feature was achieved in the following steps:

- 1) The timestamp is represented as three elements—hour, day and month.



(a) Coefficient between blending ratio and colour levels.



(b) Trend comparison of ammonia concentration and colour levels.

Figure 3.13: Observed ammonia concentration and colour levels in SHWEPP influent.

- 2) Each element will bed decomposed into sine and cosine components.
- 3) Last step is applied to hours and days to make all elements represented cyclically.

Due to the size of the datasets used in this study for training ammonia and colour forecasting model being 31 days, only the hour element was transformed into sine and cosine components as in Fig. 3.12b.

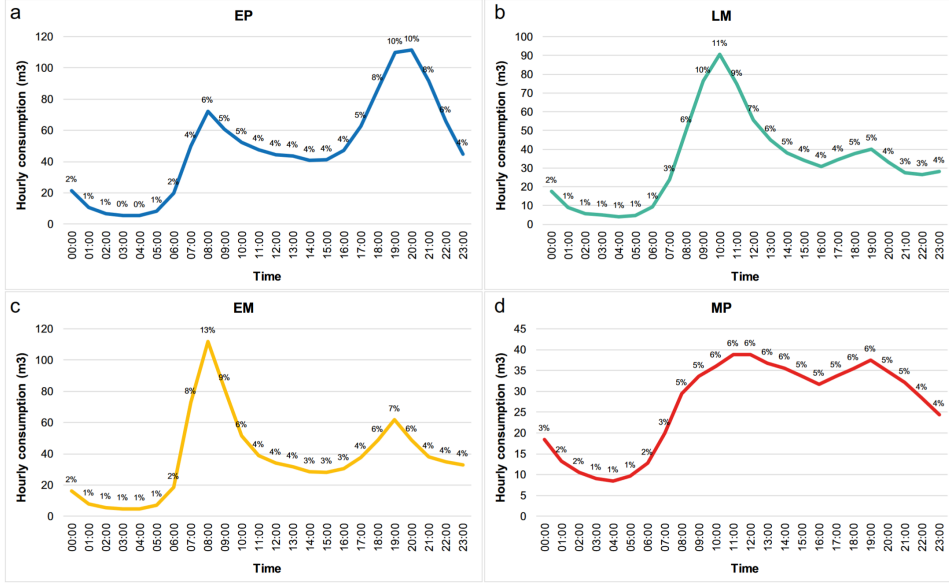


Figure 3.14: Hourly water consumption patterns in households (Abu-Bakar et al., 2021). (a) Cumulative pattern and percentage of hourly consumption for households in the “Evening Peak (EP)” cluster (b) Cumulative pattern and percentage of hourly consumption for households in the “Late Morning Peak Peak (LM)” cluster. (c) Cumulative pattern and percentage of hourly consumption for households in the “Early Morning Peak (EM)” cluster. (d) Cumulative pattern and percentage of hourly consumption for households in the “Multiple Peak (MP)” cluster. Consumption is in (m^3).

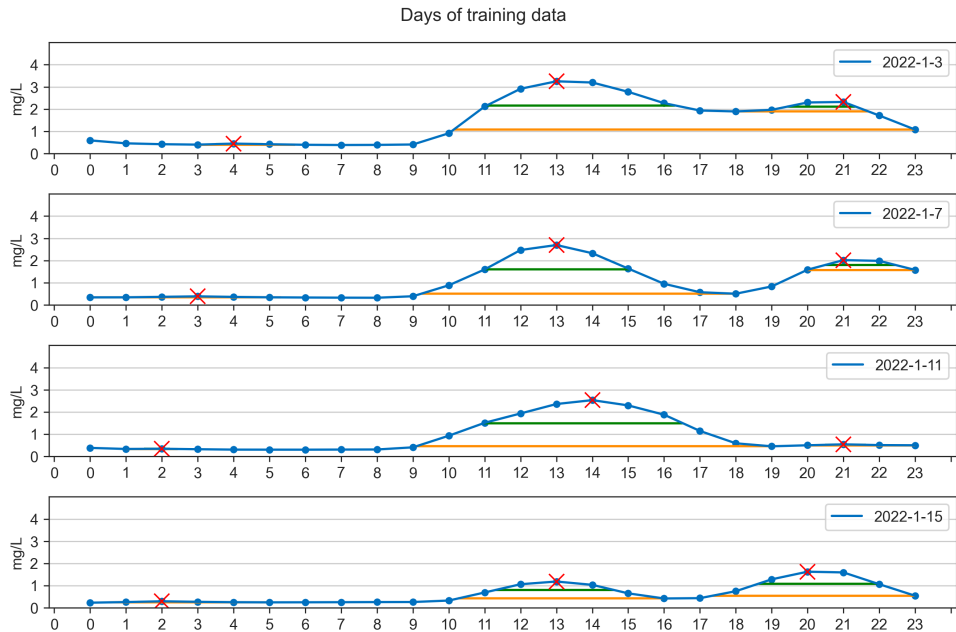


Figure 3.15: The daily patterns of ammonia concentration on 3, 7, 11, 15 January 2022.

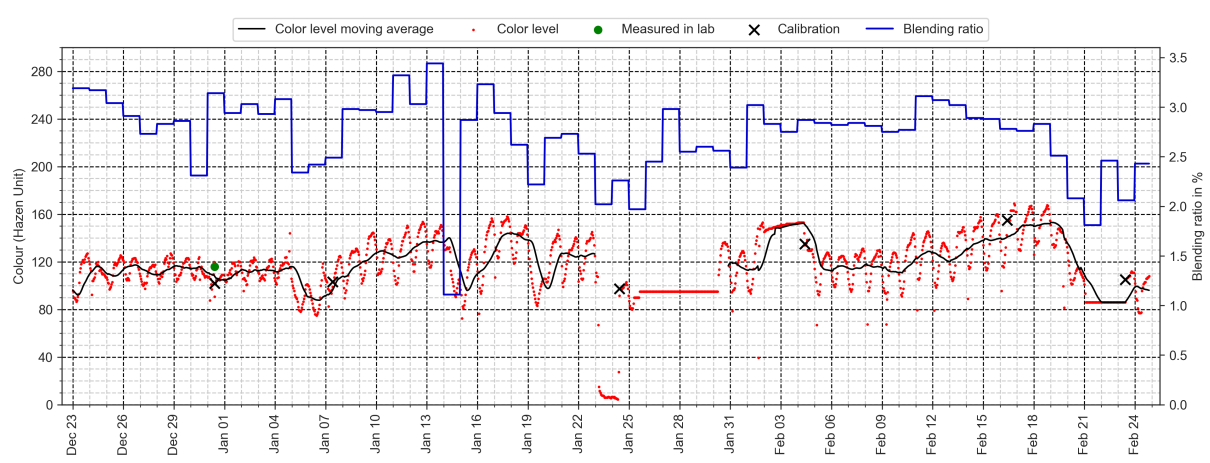
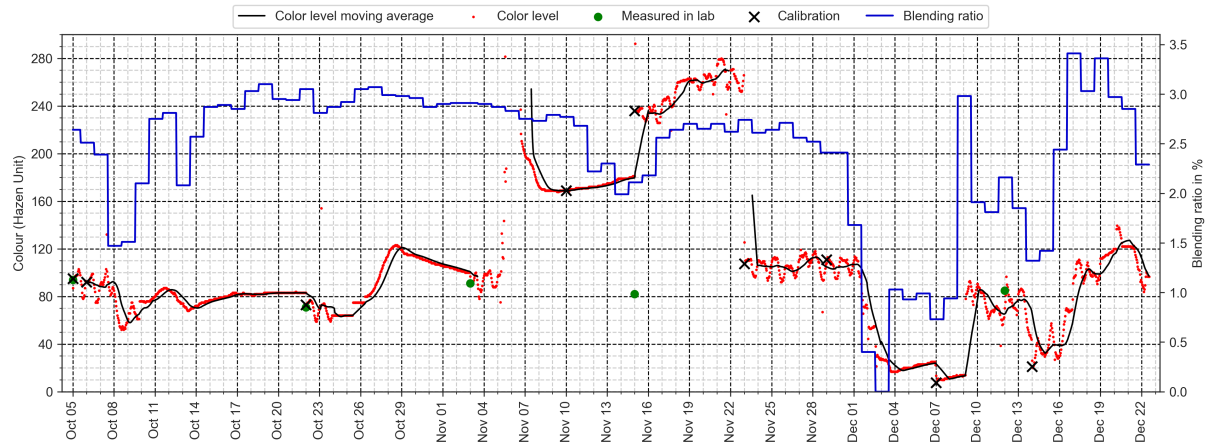


Figure 3.16: Monitored colour levels in MBR effluent and the changes in blending ratio (v/v) of treated leachate effluent to municipal wastewater in the inflow of SWHEPP from December 2021 to January 2022. The date of manually calibration and colour level measured in the laboratory are also provided as a black cross and green dot. The moving average of colour level is calculated by averaging the colour level in the past 24 hours. Note: The colour levels analyzed by the on-line colour monitoring system were compared to the manually measured data obtained from the laboratory, which showed errors of 2.08%, 4.05%, 1.11%, 65.25%, 4.94% and 11.0% in the effluent samples collected 5 Oct, 22 Oct, 3 Nov, 15 Nov, 12 Dec, and 31 Dec 2021, respectively.

3.2.4 Data transformation

Before the pre-processed data is fed into the models for training, we need to split the data into three clusters, which are training (60%), validation (20%), and testing dataset (20%). Among each cluster, the data will be further split into input variables \mathbf{X} and output variables \mathbf{Y} (i.e., training X/training Y, testing X/testing Y). During the training

process, machine learning algorithms will learn a target function \mathbf{f} to best map \mathbf{X} to \mathbf{Y} . A training dataset is a set of examples (e.g., historical data) for models to learn the hidden trends and information in the data, shown in (a) in Fig. 3.7. Training loss is calculated by taking the sum of loss for each pair of input and output in the training dataset after every training cycle (i.e., epoch).

In this study, the model is designed to forecast values three hours into the future using the values from the past 24 hours. Fig. 3.17 illustrates a forecasting model's training and forecasting process. The length of the sliding time window in this study is set to be 25 (hours). In training set 1 (i.e., the first 24 hours from the training dataset), the blue block represents the observed values of 24 hours, while the yellow block is the first data point from the testing dataset (i.e., equivalent to the 25th hour of the training dataset). The model is required to learn how to map the blue block to the yellow block; the times of model learning is equivalent to the length of the training dataset deducted by the length of the sliding time window (i.e., the second to the last 25th hour will be mapped to the last hour of the training dataset). Once the training process is complete, the model will be able to generate a value, known as the prediction or forecast, given an input of 24 hours of data.

For forecasting one hour into the future, the model will be input with 24 hours of observed values from the testing dataset, and the model will generate a value known as the forecasted values of the 25th hour. For predicting two hours into the future, the model will be input with 23 hours of observed values and the first forecasted values (i.e., the 25th hour), as shown in Forecast Set 2 in Fig. 3.17. For forecasting three hours into the future, the model will be input with 22 hours of observed values and two forecasted values from the last two forecasting processes to generate the value, known as the 26th hour. As the sliding time window moves toward the future forecast horizons, the model forecasted results would rely more on the forecasted values instead of the observed values, making the forecasted values less reliable. In this study, a forecast horizon of three is selected for testing the reliability of the model forecasting performance.

The function of a validation dataset, as in (b) in Fig. 3.7, is used to assess the model performance until we obtain the optimized hyperparameter settings, including the number of neurons in machine learning models, epoch, etc. The hyperparameter settings for each model will be discussed in the next section. The validation loss plays a vital role during

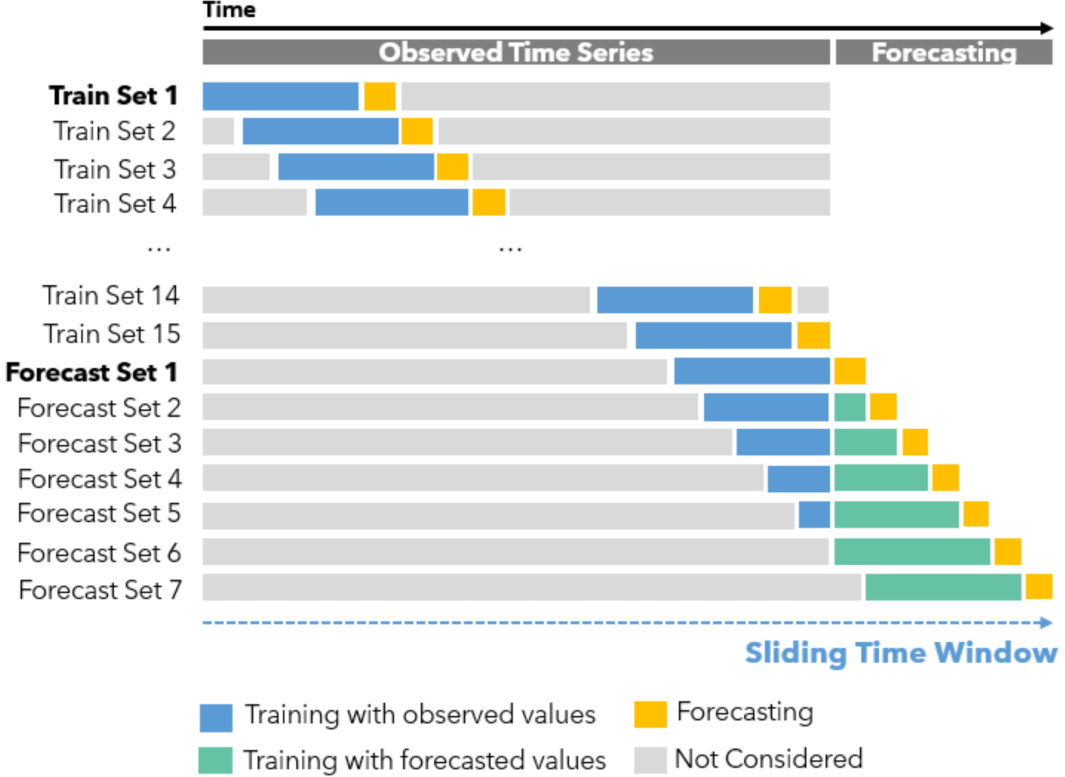


Figure 3.17: Concept of forecasting models (Liu, 2020).

the model training. The adjustments of the hyperparameters will directly reflect on the change of the validation loss; the lower the values, the better the model performance is. As the optimized model is obtained, a testing dataset is used to evaluate the performance of the forecasting model, as shown in (c) in Fig. 3.7. The testing datasets will only be input into the models when the models are tuned to the optimized settings and ready for the final evaluation. The testing datasets are also known as the unseen datasets, which can fairly evaluate the model performance. If the model tuning process was performed on the testing dataset, the model performance would be biased since the hyperparameters are adjusted in favour of the evaluation of the testing dataset.

In Fig. 3.7, the hyperparameters will remain the same once the optimized values are found, thus generating a baseline model performance from different machine learning algorithms. The baseline results will be further compared with the results from the model trained by the proposed model training steps, which include datasets that have been performed data smoothing and feature engineering techniques.

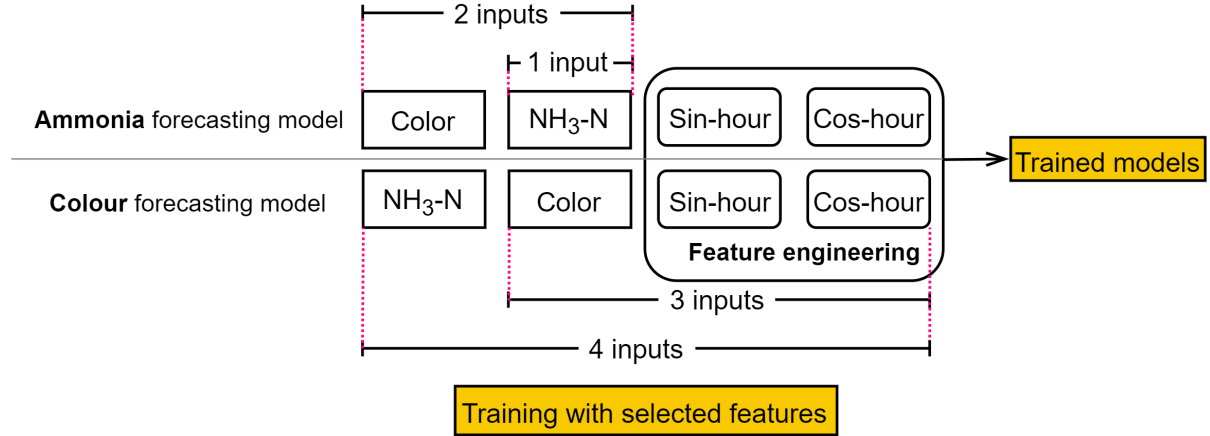


Figure 3.18: Illustration of feature selections for model training.

3.2.5 Feature selection

Fig. 3.18 illustrates which features are selected during the model training processes. In baseline model training steps, for both ammonia and colour forecasting models, only one feature is used for training for each model, which is ammonia and colour data, respectively. Following the baseline model training steps, the model trained by a single feature will generate baseline models. The results from the final evaluation will be defined as the baseline model performance, which will be compared with the model evaluated results from the proposed model training steps. Once the baseline model performance is obtained, more features will be input to the model training processes in the order of two features, three features, and four features.

3.3 Machine learning models

3.3.1 Random Forest

The machine learning model used in this study (i.e., not deep learning models) is random forest (RF). It is an ensemble method in which the final output is obtained by averaging the results from multiple tree learners (Wang et al., 2021), as shown in Fig. 3.19a. The training algorithm applies the general technique of bootstrap aggregating, also known as bagging, to tree learners. Given a training set $X = x_1, \dots, x_n$ with targets $Y = y_1, \dots, y_n$, bagging repeatedly (B times) selects a random sample with replacement (i.e., not putting the samples back to the population) of the training set and fits trees to these samples (Wikipedia, 2022a), RF generate outputs through the following steps:

For $b = 1, \dots, B$:

- 1) Sample (with replacement) n training examples from X, Y , call these X_b, Y_b .
- 2) Train a regression tree f_b on X_b, Y_b .
- 3) Predict unseen samples x' by averaging the predictions from all the regression tree learners on x' as in Eq. 3.3.1:

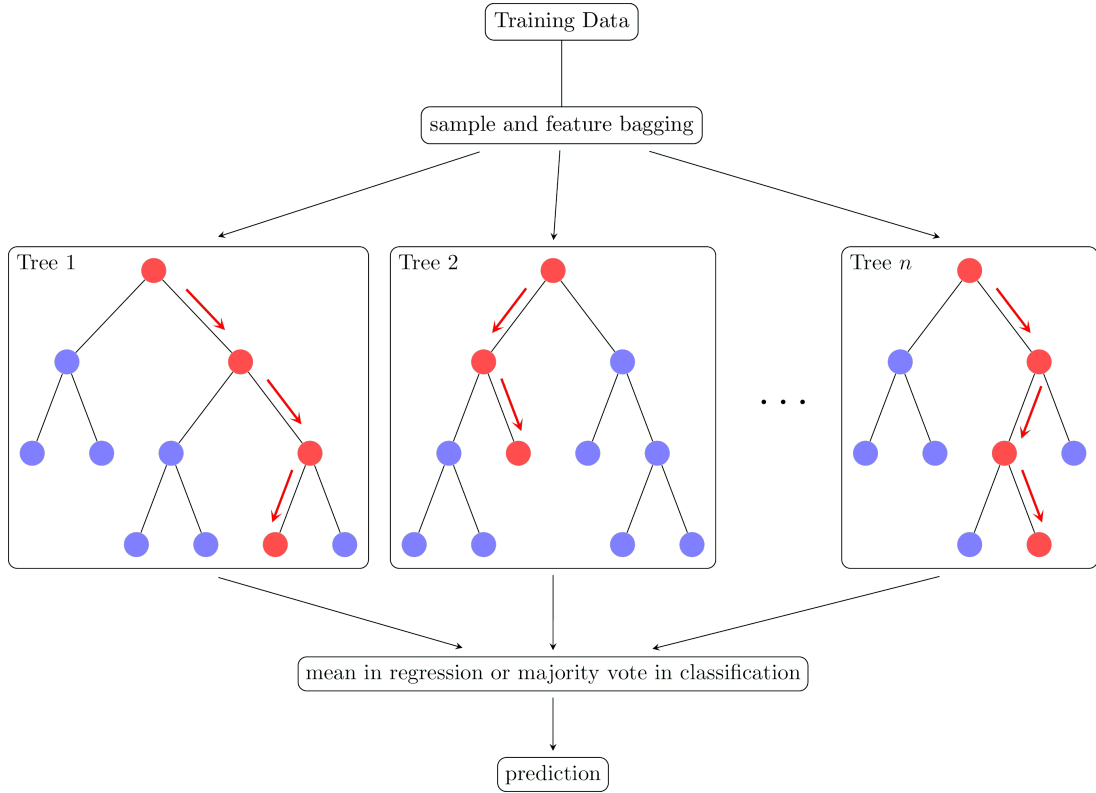
$$\hat{f} = \frac{1}{B} \sum_{b=1}^B f_b(x') \quad (3.3.1)$$

3.3.2 Deep Neural Networks

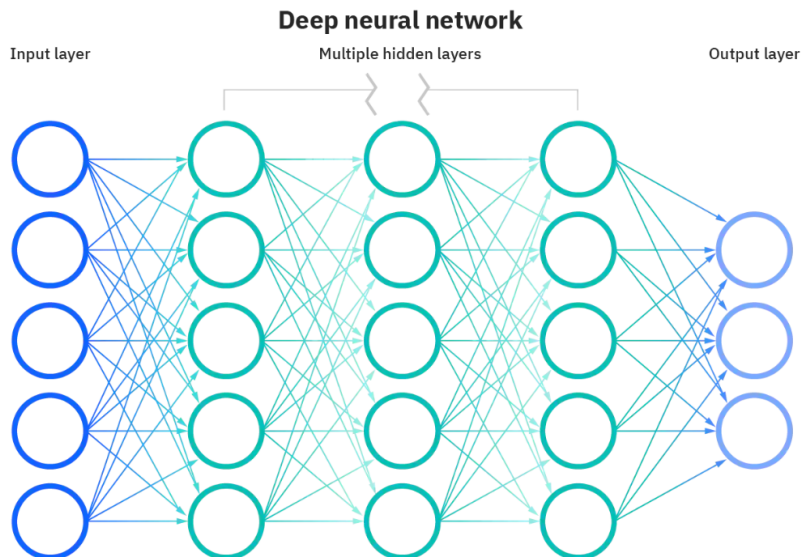
Artificial Neural Network (ANN) is a broad term that encompasses any form of Deep Learning model. A typical ANN consists of input, hidden, and output layers, and each layer comprises multiple neurons (i.e., nodes). The connected neurons simulate the human brain by processing and transmitting input signals to the next nodes (Mohseni-Dargah et al., 2022). What sets it apart from an ANN model and a DNN model is that the former contains only one hidden layer while the latter has more than one, as shown in Fig. 3.19b. The DNN models are nonlinear, which finds the correct mathematical manipulation to turn the input into the output (Bangaloreai, 2018).

3.3.3 Recurrent Neural Network

A recurrent neural network (RNN) is a type of Artificial Neural Network designed to work with sequence data. For instance, sequence data are time series, DNA, language, speech, sequences of user actions data, etc. The ammonia concentration and colour level data are time-series data, a series of data points listed in minute orders (Donges, 2021). A distinguishing characteristic of RNN is that they share parameters across each layer of the network by allowing information to be passed from the last step of the network to the next. Unlike RNN, feedforward networks like DNN have different weights across each node. The reuse of previous information for making the decision on RNN makes it capable of "learning" from the previous inputs. The realization of the memorizing function is through a memory unit called hidden state (i.e., a vector contains weights) in RNN architecture,



(a) Random Forest (RF) (Riebesell, 2022).



(b) Deep Neural Network (DNN) (IBM, 2022).

Figure 3.19: Illustration of RF and DNN model structure.

which enables RNN to persist data, thus capturing short-term dependencies. The RNN architecture is presented in Fig. 3.20a. The general formulation of a RNN is expressed in Eq. 3.3.2 (Mamandipoor et al., 2020):

$$h_t = \sigma(W^h h_{t-1} + W^x x_t + b) \quad (3.3.2)$$

where x_t is the current input, h_t is the current hidden state (output), h_{t-1} is the previous output, W^x is the weights of the hidden state, W^h is the weight of the input, b is the bias, σ is the sigmoid activation function.

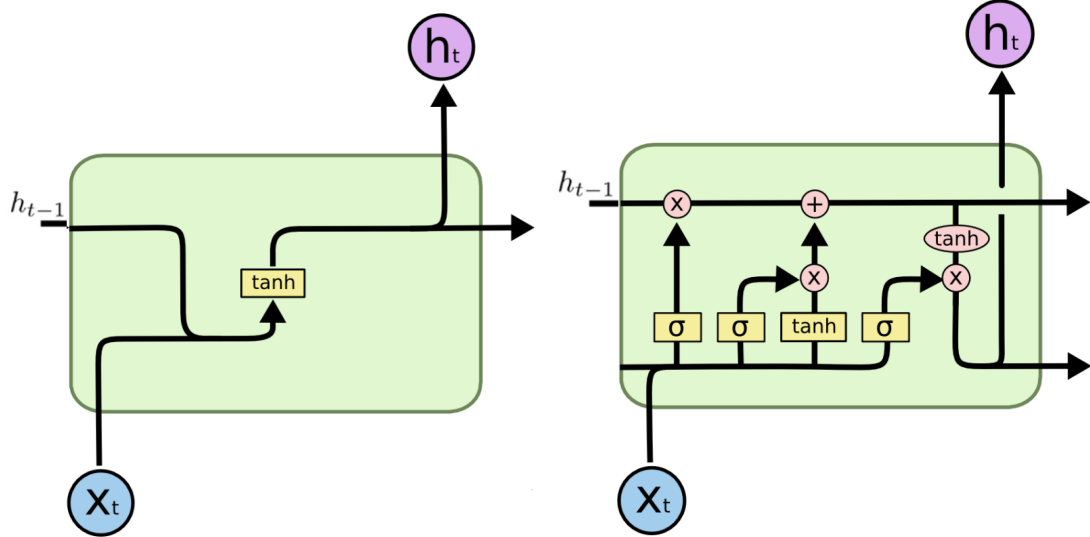


Figure 3.20: Variant architectures of Recurrent Neural Networks (adapted from Olah (2015)). x_t corresponds to the current input, h_{t-1} to the last hidden state (output), h_t to the current output, \tanh is the tangent activation function, σ is the sigmoid activation function, \times is the vector pointwise multiplication, $+$ is the vector pointwise addition.

3.3.4 Long Short-term Memory

Long Short-term Memory (LSTM) is a deep recurrent neural network (RNN), an advanced and improved version of RNN. The advent of LSTM solves problems requiring long-term temporal dependencies that RNN cannot learn due to the simple model architecture. The fundamental LSTM network is built on memory blocks called "cells", which are responsible for transferring and receiving the states (i.e., vectors) recording the information from the previous cells. In a cell block, there is an input gate, a forget gate, and an output gate. The function of these three gates is to control the movement of the information into and out of the cell via the sigmoid function. The inputs of the cell will first go through a forget gate (f_t) as Eq. 3.3.3a, where the function will multiply each element in the input states by values ranging from 0 to 1 to realize the effect of "forget." Next, an input gate (i_t) as in Eq. 3.3.3b will decide whether the new information should be updated or ignored by the sigmoid function (i.e., 0 or 1), followed by a tangent function giving the weight of importance (i.e., -1 to 1) to the values which passed by as in Eq. 3.3.3c. New memory then is appended to the previous memory C_{t-1} resulting a new C_t . Lastly, output values (h_t) is obtained based on output cell state (O_t) as in Eq. 3.3.3e and Eq. 3.3.3f (Le et al., 2019). The equations for LSTM structure are shown in Eq. 3.3.3:

$$f_t = \sigma(W_f[h_{t-1}, X_t] + b_f) \quad (3.3.3a)$$

$$i_t = \sigma(W_i[h_{t-1}, X_t] + b_i) \quad (3.3.3b)$$

$$\tilde{C}_t = \tanh(W_n[h_{t-1}, X_t] + b_n) \quad (3.3.3c)$$

$$C_t = C_{t-1}f_t + \tilde{C}_ti_t \quad (3.3.3d)$$

$$O_t = \sigma(W_o[h_{t-1}, X_t] + b_o) \quad (3.3.3e)$$

$$h_t = O_t \tanh(C_t) \quad (3.3.3f)$$

where f_t corresponds to the forget gate, i_t to the input gate, \tilde{C}_t to the candidate cell state, C_t to the current cell state, O_t to the output cell state, h_t to the output values, σ to the sigmoid function, X_t to the current input, \tanh to the tangent function, W and b are the weight matrices and bias of the corresponding output gate, respectively.

3.3.5 Gate Recurrent Unit

Gated Recurrent Unit (GRU) model is a variant of the LSTM model; by combining the forget gate and input gate into an update gate as in Fig. 3.20c, GRU has fewer parameters compared to LSTM. The advantage of GRU over LSTM is less computing power required while maintaining a similar model performance compared to LSTM. The inputs of the GRU model first enter the update gate (z_t) as in Eq. 3.3.4a, where the function will help the model determine how much of the past information needs to be passed along to the future via sigmoid functions, and then followed by the reset gate (r_t) as in Eq. 3.3.4b, which is used to decide how much of the past information to forget. Although Eq. 3.3.4a and Eq. 3.3.4b have the same inputs of X_t and h_{t-1} , the usages of the gates are different. The outputs of the reset gate will be used to determine the candidate hidden state (\tilde{h}_t) as in Eq. 3.3.4c, where the tangent function will determine the importance of the current input (X_t), reset gate output, and previous hidden state (h_t). At the last step, the output values (h_t) is calculated from the candidate hidden state (\tilde{h}_t), previous hidden state (h_{t-1}), and the outputs of update gate as in Eq. 3.3.4d. The equations of GRU structures are presented in Eq. 3.3.4 (Cheng et al., 2020):

$$z_t = \sigma(X_t W_{xz} + h_{t-1} W_{hz} + b_z) \quad (3.3.4a)$$

$$r_t = \sigma(X_t W_{xr} + h_{t-1} W_{hr} + b_r) \quad (3.3.4b)$$

$$\tilde{h}_t = \tanh(X_t W_{xh} + (r_t \circ h_{t-1}) W_{hh} + b_h) \quad (3.3.4c)$$

$$h_t = z_t \circ h_{t-1} + (1 - z_t) \circ \tilde{h}_t \quad (3.3.4d)$$

where z_t corresponds to the update gate, r_t to the reset gate, \tilde{h}_t to the candidate hidden state, h_t to the output values, σ to the sigmoid function, \tanh to the tangent function, X_t to the current input, W and the b are the weight matrices and bias of the corresponding output gate, respectively.

3.3.6 Configuration of models

Hyperparameters are variables that we need to set before applying a learning algorithm to a dataset (Agrawal, 2019). For different tasks and datasets, the optimized hyperparameters vary, which makes the seeking of hyperparameters challenging. For RF models,

only one hyperparameter needs to be selected—the number of estimators. As shown in Fig. 3.19a, each estimator, known as the tree in the forest, makes a decision. Therefore, we need to set the number of estimators for making a forecast. In this study, we tried different numbers of estimators and selected 500 estimators ultimately.

For training neural networks (NNs), the selection of hyperparameters is much more. The hyperparameters in NNs can be split into two categories, as shown in the followings:

Optimized hyperparameters

- 1) Learning rate
- 2) Number of epochs
- 3) Mini batch size

Model-specific hyperparameters

- 1) Number of hidden units (neurons)
- 2) Number of layers

The learning rate is a tuning parameter in an optimization algorithm that determines the step size at each iteration while moving toward a minimum of a loss function. An iteration describes the number of times a batch of data passed through the algorithm. In our study, the training data has a length of 432, with a batch size of one; the model will iterate 432 times to complete one epoch. There is a trade-off between the rate of convergence and overshooting when determining an optimal learning rate. A too high learning rate leads to a learning step jump over minima as in Fig. 3.21c, yet a too low learning rate will either be too slow to converge or get stuck in a local minimum loss as in Fig 3.21a. A good size of learning rate should reach the minimum loss at a reasonable time, as in Fig. 3.21b. However, searching for the most optimal learning rate can be time-consuming and a waste of computing power. In this study, we used a learning rate scheduler to achieve the same effect of using a decent learning rate. The scheduler can be set to reduce the learning rate as the epoch increases. When the algorithm detects the test loss is not reducing during the training within a designated epoch time, the learning rate will be multiplied by a customized factor. A factor of 0.5 and a patience of 10 were used in this study. The effect of using a learning rate scheduler is shown in Fig. 3.21d.

In model-specific hyperparameter tuning, the number of neurons and the number of

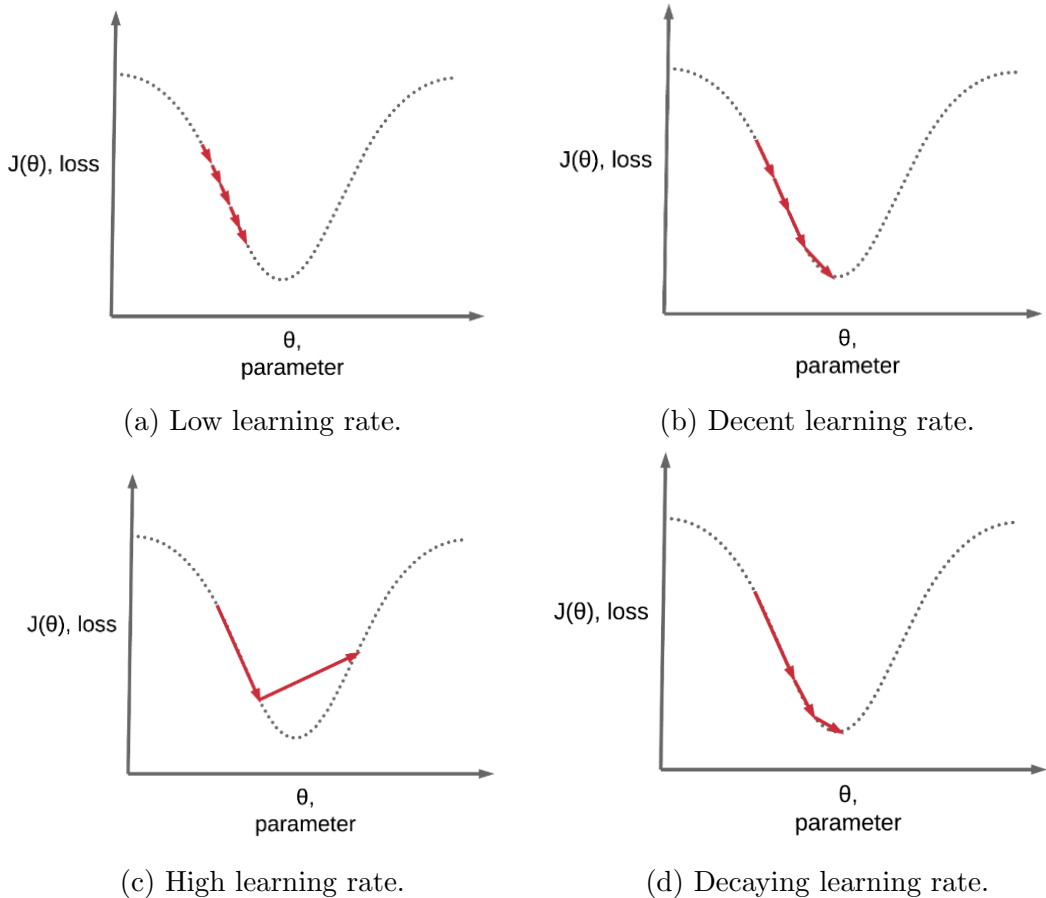


Figure 3.21: Illustration of how different step sizes of learning rate reach the minimum loss (Ritchie Ng, 2019).

layers need to be determined based on the complexity of our training dataset. The ammonia and colour datasets are considered simple and small datasets. In the hyperparameter tunings of the deep learning models, we simplified the model structure by lowering the number of layers to 1 except for the DNN model. If the number of hidden layers decreased to one, the DNN models would be called the ANN models according to the definition. The number of neurons was set to 10 to maintain simple deep learning models to prevent overfitting.

The settings of the optimized hyperparameters are listed in the followings in the final iteration of model hyperparameter tuning:

Optimized hyperparameters

- 1) Learning rate: 5e-05
- 2) Number of epochs: 100
- 3) Batch size: 1

Table 3.2: Final model configuration.

Model	Input	h.d ^a	Output	Num. of Exp ^b	Comments
RF	24 ^c	-	3	3	Estimators = 500
DNN	24	2	1	3	h.d = 10 neurons
RNN	24	1	1	3	h.d = 10 neurons
GRU	24	1	1	3	h.d = 10 neurons
LSTM	24	1	1	3	h.d = 10 neurons

^a Hidden layer.

^b The times the experiments were repeated.

^c 24 hourly data points were input into the models for training.

CHAPTER 4

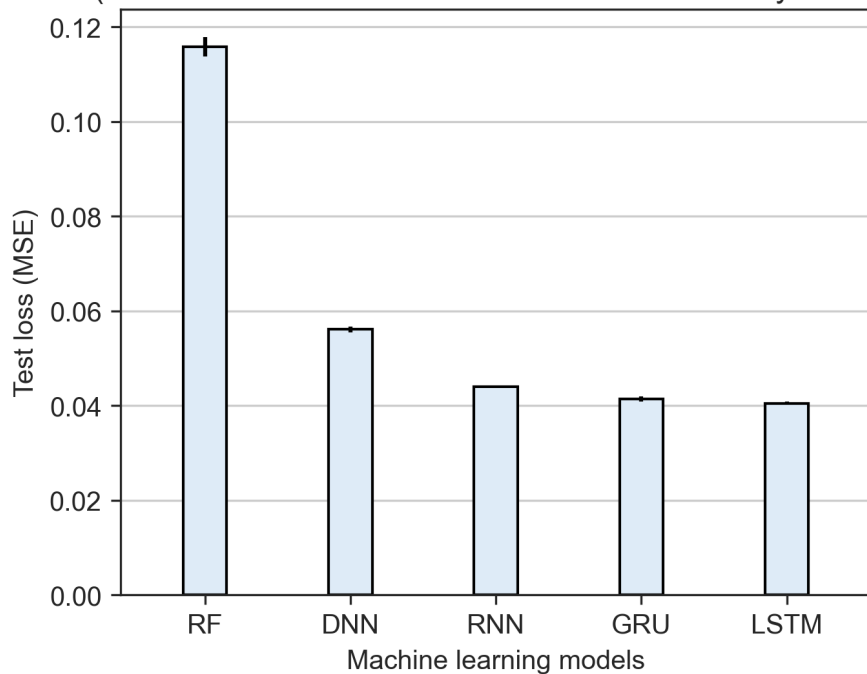
RESULTS AND DISCUSSION

4.1 Baseline performance of the forecasting models

In this study, five machine learning algorithms were trained with univariate datasets to predict the ammonia concentrations and colour levels in the reclaimed water system. All baseline models are trained by training datasets which were not applied with data pre-processing and feature engineering techniques. The forecasting model performance is presented in Fig. 4.1. As shown in Fig. 4.1a, the test loss values of RF, DNN, RNN, GRU, and LSTM models are 0.1158, 0.0561, 0.0440, 0.0414, and 0.0405, respectively. RF model is the least capable model in forecasting ammonia concentrations, given that its test loss is significantly higher than all the other four deep learning models. The cause of poor RF model performance can be attributed to its simple model structure. RF model generates results based on the averaging results from each decision tree (i.e., each decision tree will generate a prediction based on entropy and information gain). There is only one available hyperparameter for tuning RF models: the estimators (i.e., the number of the decision tree). Therefore, throughout the entire model tuning process. We observed the RF model had the lowest test loss at the beginning among all the models, and the increased estimators did not help lower the test loss values. Meanwhile, several iterations of hyperparameter tunings help the deep learning models to reduce the test loss values to critical values, which were lower than the test loss of the RF model. The gradual reductions of test loss values for deep learning models can be attributed to the nature of their complex model architectures (i.e., a good quantity of neurons, neurons are designed to perform unique functions) and the available hyperparameters for tuning. For instance, the number of hidden layers, number of neurons, learning rate, and epoch are adjustable. The customizable hyperparameters in the deep learning models allow the researchers to fully explore the possibilities of training better models, and the superior performance is reflected in the values of test loss obtained from the optimized hyperparameter settings.

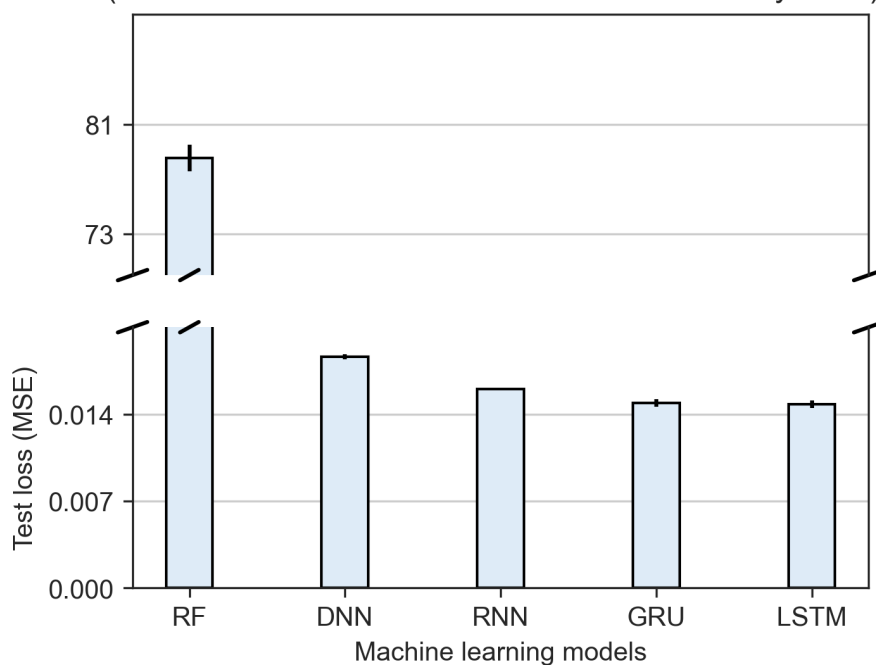
GRU and LSTM models learn the data in similar ways by utilizing memorizing cells to pass and receive critical information from the previous memorizing cells, known as the

Baseline model performance in forecasting NH₃-N.
(Evaluated on test dataset from 16 to 22 January 2022)



(a) Test loss values from five ammonia forecasting models.

Baseline model performance in forecasting colour.
(Evaluated on test dataset from 16 to 22 January 2022)



(b) Test loss values from five colour forecasting models.

Figure 4.1: Baseline performance of ammonia and colour forecasting models.

architecture of recurrent neural network. Compared to RNN models, both models contain more "gates" in the architectures to help control the flow of information, enabling the models to capture more details. The number of gates in RNN, GRU, and LSTM is one, three, and four; theoretically, GRU and LSTM can learn more information from the data based on a greater number of gates. The results in Fig. 4.1a showed good agreement with our understanding that LSTM performed better than GRU, followed by RNN models based on the values of test loss. For DNN models, the lack of memorizing cells in the model architecture relates to the poorer capability of learning information hidden in time-series datasets. In other words, DNN models cannot comprehend the information hidden in each datapoint in sequence, making the time-series dataset merely a common set of data. The DNN model with a test loss of 0.0440, higher than the 0.0414 of the RNN models, fully justifies the need to use the architecture of recurrent neural networks for training ammonia forecasting models.

In Fig. 4.1b, the test loss from colour forecasting models are 78.5296, 0.0186, 0.0160, 0.0149, and 0.0148 for RF, DNN, RNN, GRU and LSTM models, respectively. We first noticed the highest test loss value of 78.5296 in the RF model compared to the other four, making RF model the worst model in forecasting colour levels. The extremely high MSE values were caused by the colour levels fluctuating in a wider range of 80 to 160 Hazen Units. The large discrepancy between the actual and predicted colour levels increases the error values, which are further amplified as the MSE values are calculated by the average of the squares of the errors. As shown in Fig. 4.3a, on 20 January 2022, the errors between the ground truth and forecasted values are up to around 30 Hazen Units, which contribute to a large increase of MSE values in the test loss. RF model is regarded as an inferior model for forecasting colour levels using the data collected in SWHEPP.

The performance of DNN, RNN, GRU, and LSTM models, from the best to the least, are identical to what we observed in the results of ammonia forecasting models. LSTM model has the lowest test loss of 0.0148, followed by the GRU, RNN, and DNN models. In colour forecasting models, the model performance of LSTM is very close to GRU, with a difference of less than 0.0001 (i.e., less than 1%). However, the lowest test loss generated from the LSTM model in all the experiment runs (i.e., three runs) is 0.0143, which is lower than 0.0146 from the GRU model. Indicating LSTM model has more potential in forecasting time-series data.

The significantly higher test loss of RF models compared to other models can be visualized by plotting the forecasted values with the ground truths (i.e., observed values). In Fig. 4.2 and Fig. 4.3, one-step-ahead forecast horizon of ammonia concentrations and colour levels are plotted by RF as in Fig. 4.2a and Fig. 4.3a and LSTM models as in Fig. 4.2b and Fig. 4.3b. It is easier to observe that the RF models are less capable of predicting the water quality parameters.

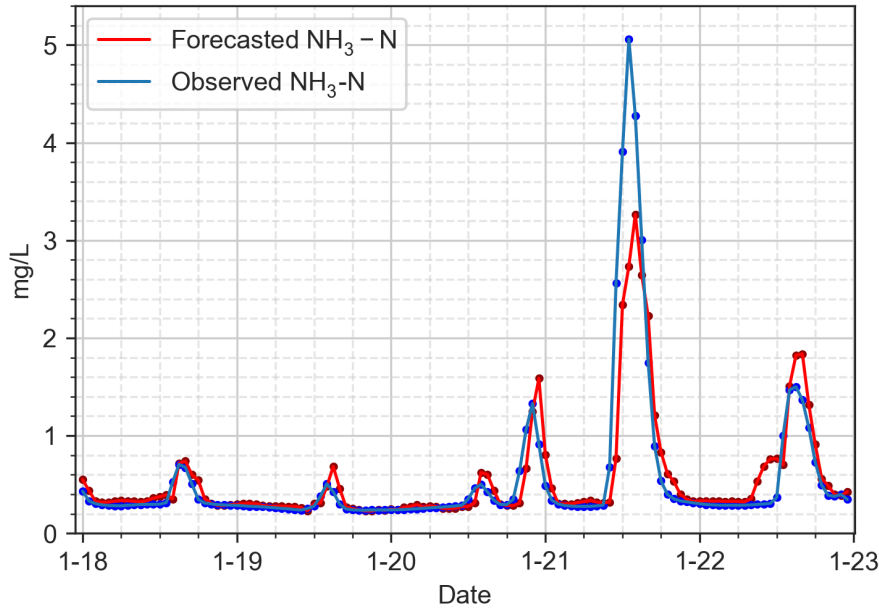
4.2 Improved performance on forecasting models using data pre-processing techniques

4.2.1 Models trained by pre-processed datasets

In this study, we investigate whether the datasets treated by the proposed data pre-processing techniques can improve the baseline model performance using the same hyperparameter settings. As shown in Table. 4.1 and Table. 4.3, we listed all the test loss values of five machine learning algorithms trained with each proposed pre-processed technique for ammonia concentrations and colour levels forecasting. The machine learning algorithm trained by datasets that were applied with SG filters at different window sizes is denoted as model-sg5, model-sg7, and model-sg9. The naming rule applies the same to EWMA filtered dataset; the method of outlier removal for ammonia data is denoted as model-or; models trained with the raw datasets are denoted as model-obs (i.e., observed dataset).

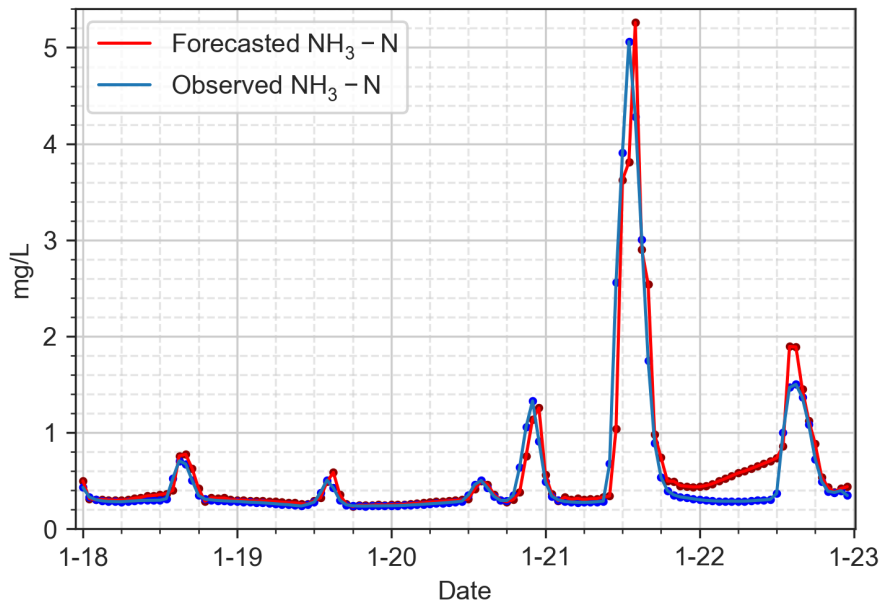
The improvements in the performance of ammonia forecasting models are most significant with models trained by SG filtered datasets. Training GRU models with an sg7 filtered dataset reduced the test loss of GRU-obs from 0.0414 to 0.0383 (-7.5%). LSTM-sg7 also successfully decreased the test loss value of LSTM-obs from 0.0405 to 0.0388 (-4.2%), while RNN-sg5 reduced the test loss value of RNN-obs from 0.0440 to 0.0415 (-5.7%). Using SG filters on the training datasets improves the performance of LSTM, GRU, and RNN models. However, the DNN and RF models trained by sg filtered datasets did not show a superior model performance compared to the test loss values of 0.0561 and 0.1158 of DNN-obs and RF-obs, respectively. Given that DNN and RF models perceive the data points as clusters of individuals, data smoothing using SG filters is not expected to help improve their model performance. SG filter smoothes the data points by convoluting both

The ammonia forecasting results.
(R-squared=0.7743)



(a) Baseline RF model forecasting ammonia concentration.

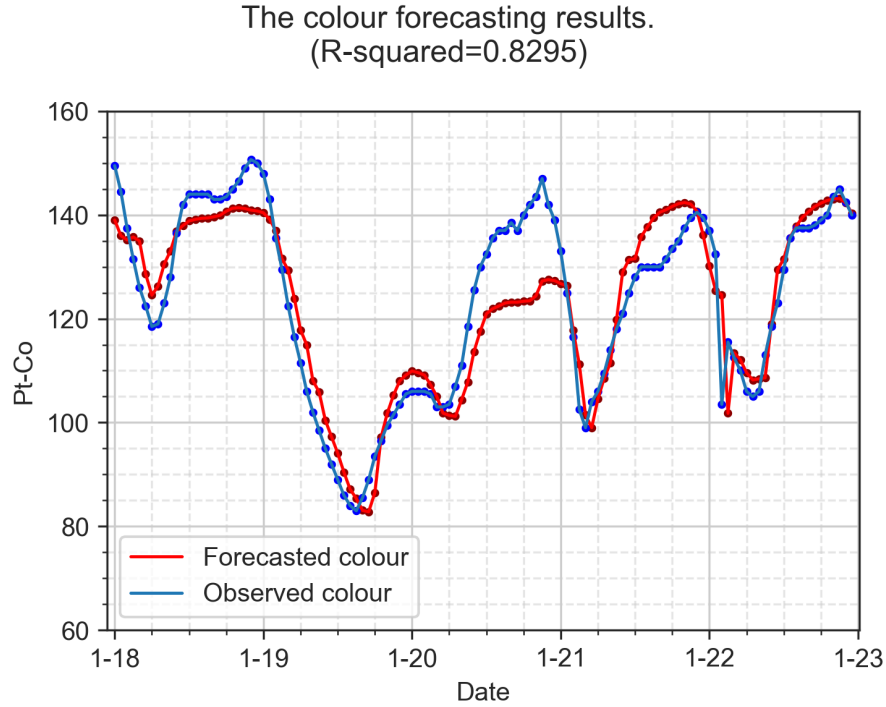
The ammonia forecasting results.
(R-squared=0.8847)



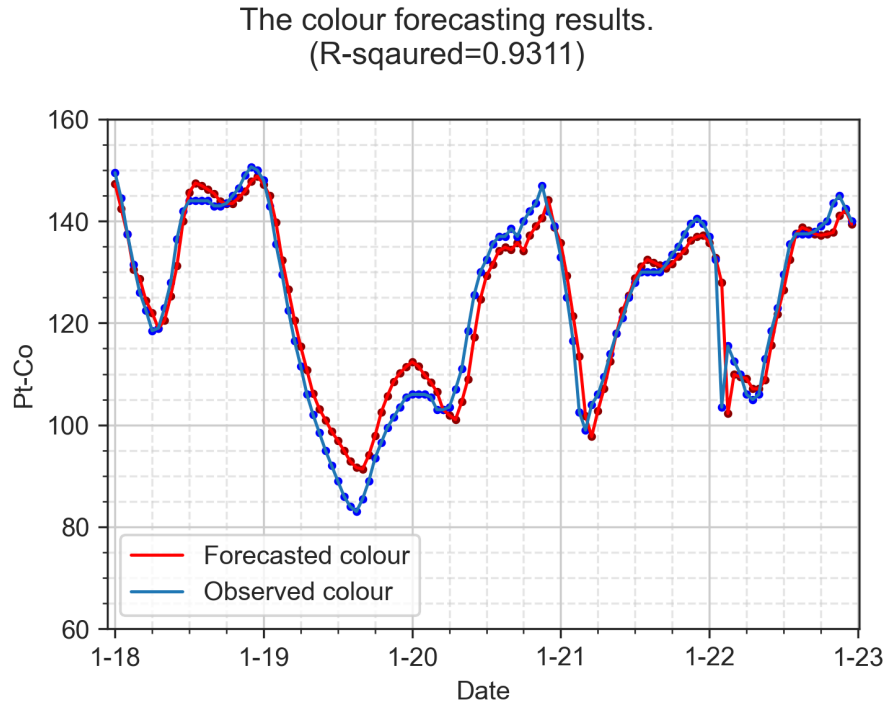
(b) Baseline LSTM model forecasting ammonia concentration.

Figure 4.2: Visualization of the model forecasting results.

previous and subsequent data points, making a series of data points correlated or linked with each other. Such data property is believed to be captured by the memorizing cells in



(a) Baseline RF model forecasting colour levels.



(b) Baseline LSTM model forecasting colour levels.

Figure 4.3: Visualization of the model forecasting results.

recurrent neural networks, such as RNN, GRU, and LSTM models. From the results in Table. 4.1, all the recurrent neural networks-based models outperformed all the DNN and

Table 4.1: Baseline performance of ammonia forecasting model, evaluated on test dataset from **16 to 22 January 2022**. Loss values are calculated by MSE.

Model-Dataset	Test loss	Valid loss	Model-Dataset	Test loss	Valid loss
GRU-sg7	0.0383	1.2508	RNN-or	0.0432	1.6345
GRU-sg5	0.0385	1.2644	RNN-ew3	0.0434	1.6041
LSTM-ew3	0.0388	1.0796	RNN-obs	0.0440	1.6734
LSTM-sg5	0.0388	1.2346	RNN-sg9	0.0442	1.7046
LSTM-sg7	0.0388	1.1804	DNN-obs	0.0561	3.2383
GRU-ew2	0.0389	1.1891	DNN-sg5	0.0562	3.2170
GRU-ew4	0.0391	1.2390	DNN-ew2	0.0563	3.1677
GRU-ew3	0.0392	1.2199	DNN-ew3	0.0569	3.2317
LSTM-ew2	0.0392	1.0969	DNN-sg7	0.0570	3.2014
LSTM-ew4	0.0395	1.1219	DNN-ew4	0.0571	3.2188
GRU-sg9	0.0396	1.3097	DNN-or	0.0572	3.1972
LSTM-or	0.0398	1.2612	DNN-sg9	0.0574	3.2484
LSTM-obs	0.0405	1.3993	RF-obs	0.1158	-
GRU-or	0.0405	1.2366	RF-sg9	0.1196	-
LSTM-sg9	0.0410	1.3076	RF-ew2	0.1286	-
GRU-obs	0.0414	1.3638	RF-or	0.1294	-
RNN-sg5	0.0415	1.5088	RF-sg5	0.1298	-
RNN-ew2	0.0421	1.5425	RF-ew3	0.1313	-
RNN-sg7	0.0423	1.6267	RF-sg7	0.1409	-
RNN-ew4	0.0432	1.5992	RF-ew4	0.1441	-

RF models. It can be concluded that DNN and RF models are poor options for training time-series models, even with the use of the SG filter technique.

The RNN-or, GRU-or, and LSTM-or models, which were trained with datasets applied with outlier removal methods, showed lower test loss values of 0.0432 (-1.8%), 0.0405 (-2.2%), and 0.0398 (1.7%) compared to test loss values of 0.0440, 0.0414, and 0.0405 from RNN-obs, GRU-obs, and LSTM-obs, respectively. We also noticed that the improvements of RNN-or, GRU-or, and LSTM-or are minor compared with the models trained by SG and EWMA filtered datasets. In this method, three days of abnormal data were removed from an 18-day dataset, which accounts for around 15% of the data. Despite the fact that 15% of the data was removed, the improvement in lowering the test loss values was slight. It is suggested that the deep learning models are smart enough to neglect the noise in the training datasets while performing forecasts from the test dataset.

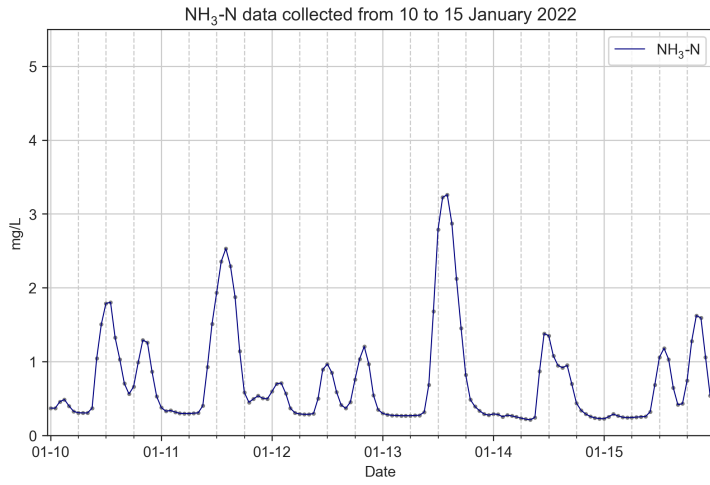
RNN, GRU, and LSTM models trained by EWMA filtered datasets also showed good improvements in the model performance. RNN-ew2, GRU-ew2, and LSTM-ew3 showed

lower test loss of 0.0421 (4.3%), 0.0389 (6.0%), and 0.0388 (4.2%) compared to RNN-obs, GRU-obs, and LSTM-obs of 0.0440, 0.0414, and 0.0405, respectively. EWMA filters modified the data points by averaging the value of the current data points with previous ones, making the data property almost identical to the SG filtered data. Both SG and EWMA filters similarly influenced the baseline models, in which LSTM obtained the lowest test loss values, followed by GRU and RNN models. By far, the results only suggest that both filters are robust techniques in terms of lowering the test loss, yet we cannot draw conclusions about which filter is more effective in improving the model performance. In addition, we discovered our test loss values to be abnormal when inspecting models' validation loss and the test loss values.

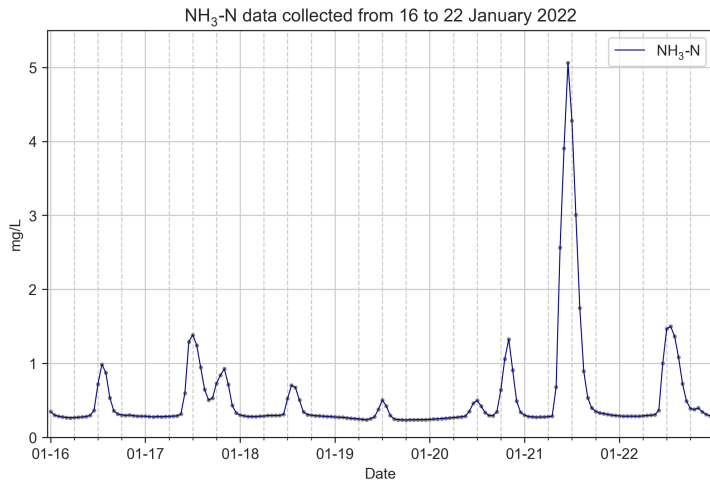
Empirically, the best-performed Model-Dataset combination should match the lowest test with the lowest validation loss values when using the same testing dataset to evaluate a group of models. For instance, the GRU-sg7 model in forecasting ammonia has the lowest test loss of 0.0383, yet the validation loss of 1.2508 only ranks tenth among the validation loss values. The top three lowest validation loss models are LSTM-ew3, LSTM-ew2 and LSTM-ew4, yet the top three lowest test loss models are from GUR-sg7, GRU-sg5, and LSTM-ew3 models. This finding points to the potential heterogeneity between the validation and testing datasets. The limitation of this study's validation and testing datasets is the small dataset size, resulting in specific daily fluctuation patterns of ammonia may only occur in the testing dataset. In all the available ammonia data, we selected the data from October 2021 as the second testing dataset for its high similarity to the validation dataset in January 2022.

As shown in Fig. 4.4, the fluctuation patterns of NH₃-N in validation dataset as in Fig. 4.4a is much resemble to the testing dataset from Fig. 4.4c compared to testing dataset from Fig. 4.4b. Further tests were carried out using a testing dataset from October to re-evaluate the model performance from Table. 4.1. It is expected that the Model-Dataset ranks of test and validation loss values from the lowest to the highest will change. To the best of my understanding, the comparisons between testing and validation loss are not discussed in the currently available research papers in the modelling of the wastewater treatment industry.

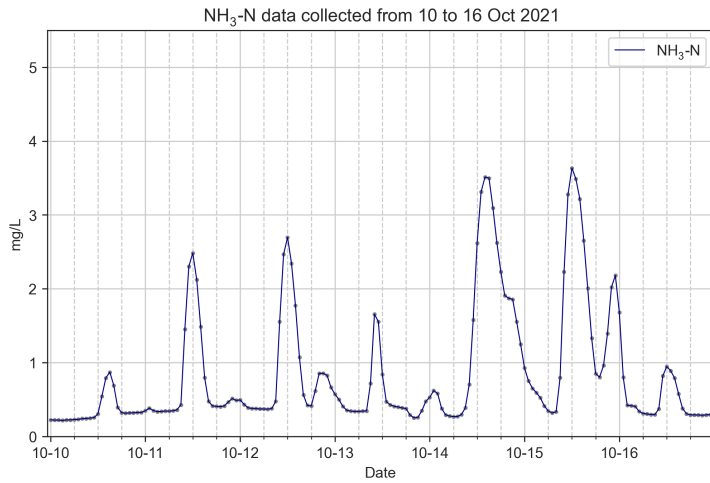
As shown in Table. 4.2, the models with the top lowest test loss values are 0.0158, 0.0161, 0.0163 for LSTM-ew3, LSTM-ew2, and LSTM-ew4, which match the top three



(a) Validation dataset from January 2022.



(b) Testing dataset from January 2022.



(c) Testing dataset from October 2021.

Figure 4.4: Illustration of the heterogeneity and homogeneity between validation and different testing datasets.

lowest validation loss values of 1.0796, 1.0969, and 0.1219. This is in good agreement with how the heterogeneity of the datasets can impact the model performance. The evaluations of the ammonia forecasting models in October 2021 showed completely different outcomes compared to those in January 2022. Instead of GRU, LSTM becomes the best model for training the ammonia forecasting model. For LSTM models, the top three Model-Dataset combinations are LSTM-ew3, LSTM-ew2, and LSTM-ew4; for GRU models, they are GRU-ew3, GRU-ew4, and GRU-ew2; for RNN models are RNN-ew4, RNN-ew2, and RNN-ew3. It is evident that EWMA filters have a more significant influence on the model performance for all the recurrent neural network models than SG filters. However, given the small dataset size, caution must be taken if the EWMA filter is applied in future works.

Table 4.2: Baseline performance of ammonia forecasting model, evaluated on test dataset from **10 to 16 October 2021**. Loss values are calculated by MSE.

Model-Dataset	Test loss	Valid loss	Model-Dataset	Test loss	Valid loss
LSTM-ew3	0.0158	1.0796	RNN-or	0.0197	1.6345
LSTM-ew2	0.0161	1.0969	RNN-sg7	0.0201	1.6267
LSTM-ew4	0.0163	1.1219	RNN-sg9	0.0205	1.7046
LSTM-sg5	0.0166	1.2346	RNN-obs	0.0206	1.6734
GRU-ew3	0.0167	1.2199	DNN-ew3	0.0316	3.2317
GRU-ew4	0.0169	1.2390	DNN-or	0.0316	3.1972
GRU-ew2	0.0170	1.1891	DNN-sg7	0.0316	3.2014
GRU-sg9	0.0174	1.3097	DNN-ew2	0.0318	3.1677
LSTM-obs	0.0175	1.2366	DNN-ew4	0.0319	3.2188
LSTM-or	0.0177	1.2612	DNN-obs	0.0319	3.2383
GRU-sg5	0.0178	1.2644	DNN-sg5	0.0319	3.2170
GRU-sg7	0.0180	1.2508	DNN-sg9	0.0319	3.2484
LSTM-sg7	0.0180	1.1804	RF-sg9	0.1307	-
GRU-or	0.0187	1.3993	RF-sg7	0.1311	-
LSTM-sg9	0.0188	1.3076	RF-sg5	0.1343	-
GRU-obs	0.0189	1.3638	RF-ew2	0.1346	-
RNN-ew4	0.0190	1.5992	RF-ew3	0.1368	-
RNN-ew2	0.0191	1.5425	RF-obs	0.1443	-
RNN-ew3	0.0193	1.6041	RF-ew4	0.1451	-
RNN-sg5	0.0195	1.5088	RF-or	0.1477	-

The test loss values of the colour forecasting models are presented in Table. 4.3. The top six lowest test loss models are LSTM-ew4, LSTM-ew2, LSTM-ew3, GRU-ew3, GRU-ew2, and GRU ew4 with the values of 0.0136, 0.0138, 0.0138, 0.0140, 0.0142, and 0.0143, respectively. LSTM models are shown to be the best-performed model in forecasting

colour levels. The results also suggest that all the top lowest test loss models are trained by EWMA filtered datasets. We found that LSTM, GRU, and RNN models trained by EWMA filtered datasets generated the top lowest test loss values compared to the same models trained by SG filtered datasets. Interestingly, in both colour and ammonia forecasting models, LSTM models trained by EWMA filtered dataset showed the most superior performance, as shown in Table. 4.2 and Table. 4.3. LSTM models trained with EWMA filtered datasets are proved to be the best model and pre-processing techniques for training colour forecasting models in this study.

In the investigation of how a small dataset can influence the model results, we found that the top three lowest validation loss values are LSTM-sg9, LSTM-sg7, and LSTM-ew4, which rank the 7th, 20th, and 1st as the lowest test loss values. In this study, there is no extra colour testing dataset we can retrieve from the historical dataset, despite the fact that we were keen to investigate the homogeneity and heterogeneity of the colour validation and testing dataset. Compromises have to be made during the analysis of colour forecasting models.

Table 4.3: Baseline performance of colour forecasting model, evaluated on test dataset from **16 to 22 Janurary 2022**. Loss values are calculated by MSE.

Model-Dataset	Test loss	Valid loss	Model-Dataset	Test loss	Valid loss
LSTM-ew4	0.0136	0.7515	RNN-obs	0.0160	1.0623
LSTM-ew2	0.0138	0.8011	LSTM-sg7	0.0161	0.7439
LSTM-ew3	0.0138	0.7547	LSTM-sg5	0.0168	0.8355
GRU-ew3	0.0140	0.8068	DNN-sg5	0.0180	1.4702
GRU-ew2	0.0142	0.8330	DNN-sg7	0.0180	1.4823
GRU-ew4	0.0143	0.7694	DNN-sg9	0.0180	1.4574
LSTM-sg9	0.0143	0.7137	DNN-ew4	0.0181	1.4632
RNN-ew3	0.0144	0.8492	DNN-ew3	0.0182	1.4716
RNN-ew4	0.0147	0.8476	DNN-ew2	0.0183	1.4946
RNN-sg9	0.0147	0.8363	DNN-obs	0.0186	1.5397
LSTM-obs	0.0148	0.9744	RF-sg9	63.6847	-
GRU-obs	0.0149	0.9927	RF-sg7	73.8263	-
RNN-ew2	0.0150	0.9083	RF-ew3	75.1974	-
GRU-sg9	0.0151	0.7575	RF-ew4	77.8829	-
RNN-sg5	0.0158	0.8846	RF-obs	78.5296	-
RNN-sg7	0.0158	0.8755	RF-ew2	78.8753	-
GRU-sg7	0.0159	0.7791	RF-sg5	81.0696	-
GRU-sg5	0.0160	0.8080	-	-	-

By comparing the baseline performance and the influences of data pre-processing tech-

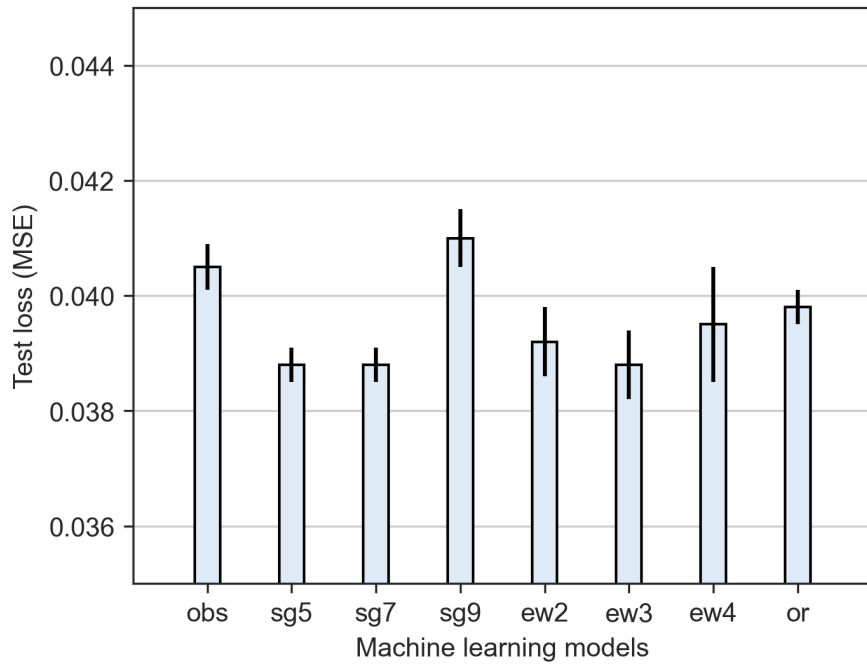
niques on machine learning models, our findings appear to be well substantiated by using LSTM models for training ammonia and colour forecasting models due to their outstanding model performance evaluated by test loss values. Although EWMA filters showed surprising effects on improving the performance of most models, the conclusions of determining which pre-processing techniques are the optimum option should be treated with caution. Thus, the testings of the proposed model training processes will include all the pre-processing techniques for model training, and LSTM will be used as the only machine learning model.

4.2.2 The effect of window size of data smoothing filters

The influences of window sizes in the data smoothing process are investigated using LSTM models and illustrated in Fig. 4.5. Larger and smaller SG window sizes have different impacts on ammonia and colour forecasting models. In ammonia forecasting models, as shown in Fig. 4.5a, LSTM models trained with SG filtered datasets with window sizes of 5, 7, and 9 have the test loss values of 0.0388, 0.0388, and 0.0410. The results suggested that modifying data points at higher degrees may negatively affect the model training process. The results from models trained by EWMA filtered datasets showed good agreement with this finding. The model trained with EWMA filtered datasets with the windows size of 2, 3, and 4 have the test loss values of 0.0392, 0.0388, and 0.0395. A higher test loss value is observed in LSTM-ew4 compared to LSTM-ew3.

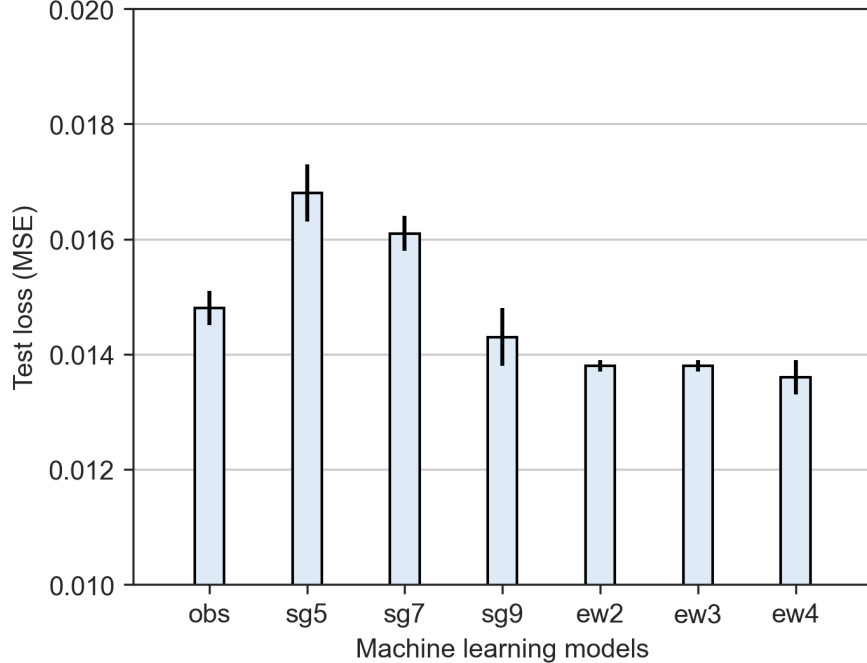
For colour forecasting models, as shown in Fig. 4.5b, LSTM models trained by SG filtered datasets with window sizes of 5, 7, and 9 have test loss values of 0.0168, 0.0161, and 0.0143. LSTM models trained by EWMA filtered datasets with window sizes of 2, 3, and 4 showed test loss values of 0.0138, 0.0138, and 0.0136. From these results, we observed that larger window sizes helped the models achieve lower test loss for colour forecasting models, which does not support what we have concluded for the ammonia forecasting models. One possible explanation for the contradictory results is that ammonia and colour data have different sensitivity toward the data smoothing filters. For instance, ammonia concentrations change between the values of 1.0 to 7.0 mg/L, while colour levels vary from 80 to 160 Hazen Units, making the values of filtered data points less significant in colour data. In other words, if ammonia data points are shifted from the original values after applying data smoothing techniques, the values might be biased considering

LSTM model trained by pre-processed datasets
 (Evaluated on test dataset from 16 to 22 January 2022)



(a) Baseline performance of ammonia forecasting models trained by LSTM.

LSTM model trained by pre-processed datasets
 (Evaluated on test dataset from 16 to 22 January 2022)



(b) Baseline performance of colour forecasting models trained by LSTM.

Figure 4.5: Baseline performance of ammonia and colour forecasting models.

the fluctuated range of ammonia is small, while the shifted colour level data can be less biased among the sample regarding the fluctuation range of colour level is much larger. By far, we can not conclude how to select the window sizes of the data smoothing filters. The unpredictable influences of applying data smoothing filters on forecasting models impede the determination of the optimum data smoothing techniques in the subsequent experiments.

4.3 Exploit hidden patterns in MBR effluent water quality to enhance model performance

4.3.1 Ammonia forecasting models

In the section of feature engineering, we have introduced the selection and creation of the extra input features for training forecasting models, as shown in Fig. 3.18. In this study, a forecasting model trained by one feature is called an univariate model and denoted as LSTM-1; a forecasting model trained by two features is called a multivariate model and denoted as LSTM-2. For models trained by three and four features are denoted as LSTM-3 and LSTM-4. In Fig. 4.6, the performance of ammonia forecasting models trained by two to four inputs (i.e., LSTM-2, LSTM-3, LSTM-4) is compared with the baseline performance (i.e., LSTM-1-obs) to demonstrate how the feature engineered features influenced on the model outputs.

As shown in Fig. 4.6, LSTM-4-obs, LSTM-3-obs, LSTM-2-obs, and LSTM-1-obs have the test loss values of 0.0432, 0.0426, 0.0411, and 0.0405, respectively. This result indicates that LSTM models trained with more features resulted in poorer model performance. Based on our understanding to the extra features such as color levels and sine/cosine features, models trained with more features are expected lower test values. The model performance from LSTM-sg7 and LSTM-sg9 fits well with what we hypothesized. The test loss values of LSTM-4-sg7, LSTM-3-sg7, LSTM-2-sg7, LSTM-1-sg7 are 0.0369, 0.0373, 0.0379, 0.0388, respectively. For LSTM-4-sg9, LSTM-3-sg9, LSTM-2-sg9, and LSTM-1-sg9, the test loss values are 0.0384, 0.0391, 0.0409, 0.0410, respectively. These findings showed that the test loss values of the LSTM models trained by sg7 and sg9 filtered datasets followed the trends of $LSTM-4 < LSTM-3 < LSTM-2 < LSTM-1$. The most remarkable results are from LSTM models trained by SG filtered dataset at a window

size of 7. Comparing to the baseline model performance (i.e., LSTM-1-obs), the test loss values of LSTM-1-sg7, LSTM-2-sg7, LSTM-3-sg7 and LSTM-4-sg7 reduced by 4.2%, 6.4%, 7.9%, and 8.9%, respectively.

Our findings in the ammonia forecasting models suggest that colour level is an indispensable feature for improving the model performance. LSTM-2 models trained by datasets applied with any pre-processing techniques showed lower test loss compared to LSTM-1, except LSTM-2 trained by dataset without applying any methods. Strong evidence leads us to believe that the fluctuation of ammonia concentration is highly correlated with the colour levels in SHWEPP influent even without direct evidence.

The methods of training LSTM models on pre-processed datasets have proved their benefits in improving baseline model performance. Yet, the test loss values were only reduced slightly for those models trained with EWMA filtered datasets. As shown in Fig. 4.6, LSTM-3-ew2, LSTM-4-ew2, LSTM-3-ew4, and LSTM-4-ew4 shared very similar test loss values to LSTM-1-obs, indicating the advantages of enhanced training datasets were not fully reflected on the model performance when LSTM models were trained with EWMA filtered datasets.

4.3.2 Colour forecasting models

As shown in Fig. 4.7, the baseline performance is LSTM-1-obs with test loss value of 0.0148, and many models trained by both SG and EWMA filtered datasets show lower test loss values. The performance of models trained by SG filtered datasets was rather disappointing. In the results of models trained by sg-5 and sg7 filtered datasets, only LSTM-3-sg5, LSTM-3-sg7, and LSTM-4 sg-7 showed lower test loss values of 0.0144, 0.0143, and 0.0136, respectively, compared to LSTM-1-obs. Models trained by sg9 and all the EWMA filtered datasets showed improvement over LSTM-1-obs. In LSTM-3-sg9, we observed the lowest test loss value of 0.0129, which is 28.6% lower than the test loss values of 0.0148 from LSTM-1-obs.

The test loss values of LSTM-4-sg9, LSTM-4-ew2, LSTM-4-ew3, and LSTM-4-ew4 are higher than LSTM-3-sg9, LSTM-3-ew2, LSTM-3-ew3, and LSTM-3-ew4, by 0.0009, 0.0009, 0.0002, and 0.0002, respectively. This finding indicates that training with ammonia and the sine/cosine features deteriorate the model performance for color forecasting models. From what we found in the results of ammonia forecasting models, we concluded

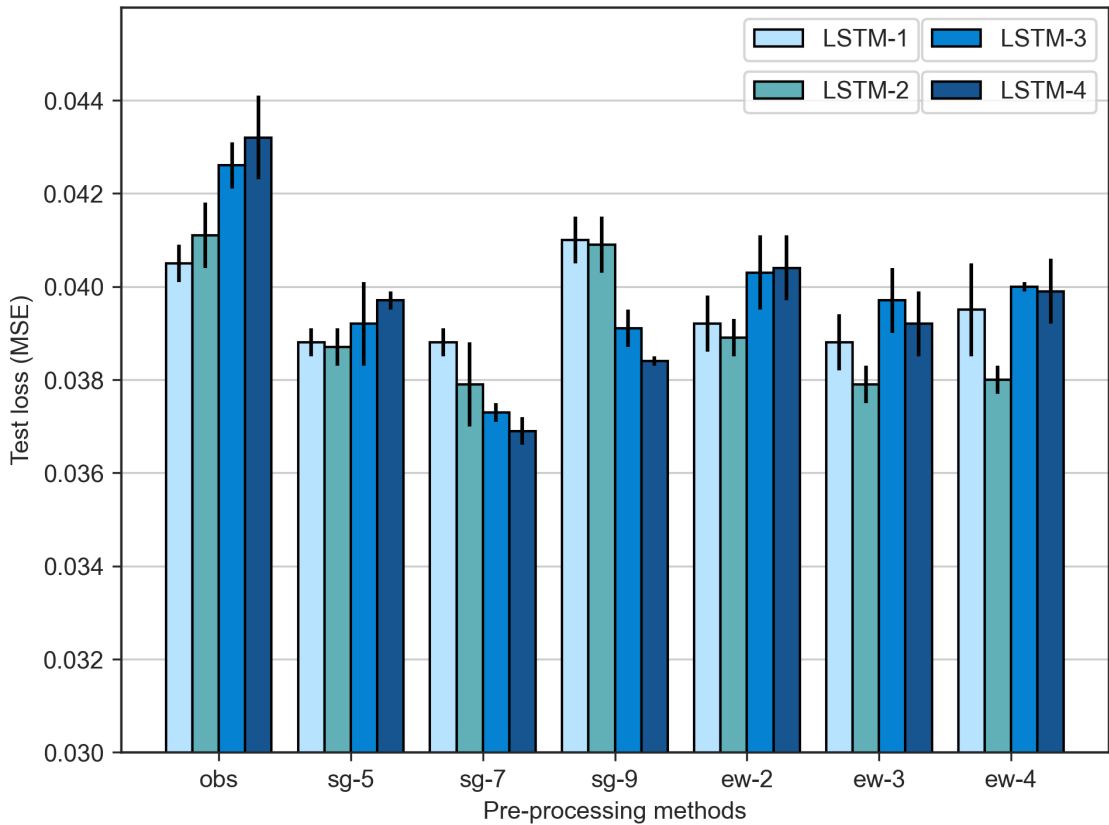


Figure 4.6: Comparisons of the model performance in forecasting ammonia concentrations.

that the test loss values increase more when more features were input to the training datasets. In the colour forecasting results, the finding contrasts what we have found previously.

The interpretation for the higher test loss in LSTM-4 models in sg9, ew2, ew3, and ew4 filtered datasets compared to LSTM-3 and LSTM-2 models is that ammonia and sine/cosine features are irrelevant to the development of colour forecasting models. In the process of generating feature engineering, we observed that colour substances are mixed with municipal wastewater at the volume to volume ratio of 1 to 50. Hence, we can infer that the model outputs of forecasted colour levels are highly subject to the input of ammonia concentration. In the training process of the machine learning model, the model treats each input feature with equivalent importance; however, when the model is trained and input with unseen data, the model cannot differentiate which input feature actually influences more on the model outputs. The results suggest that it is best to train features of colour data and sin/cosine features for training color forecasting models.

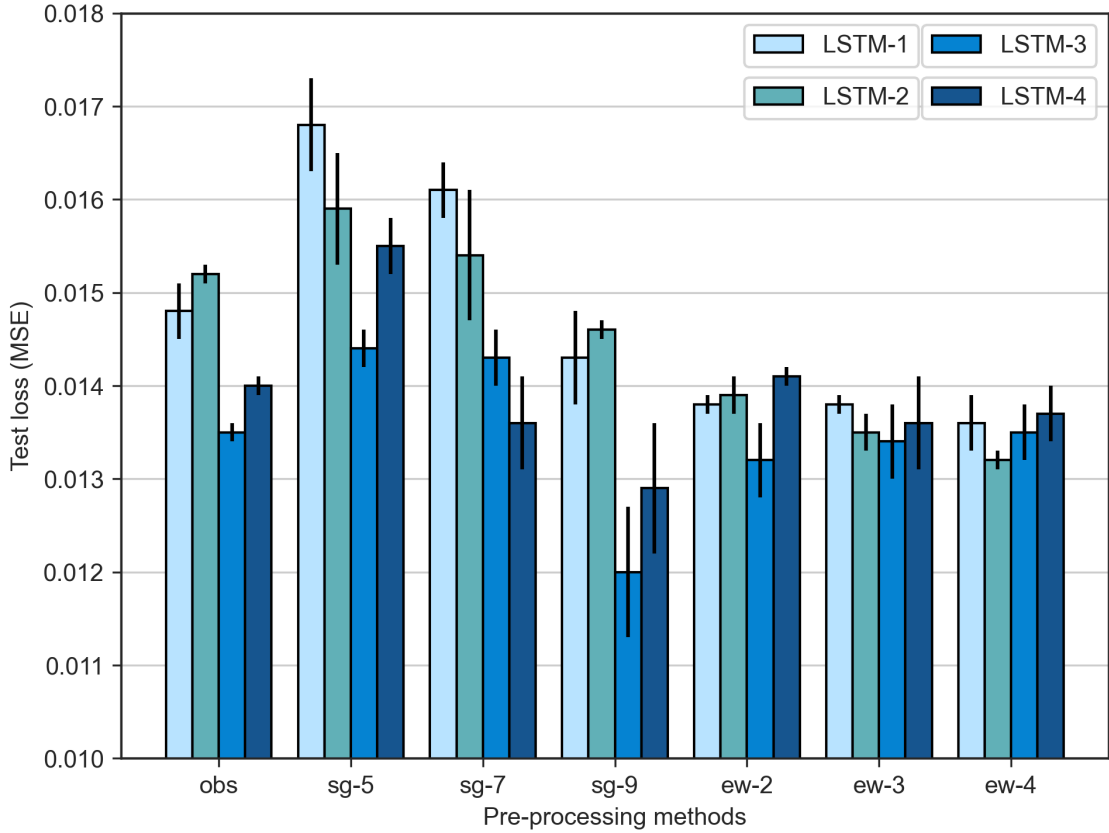
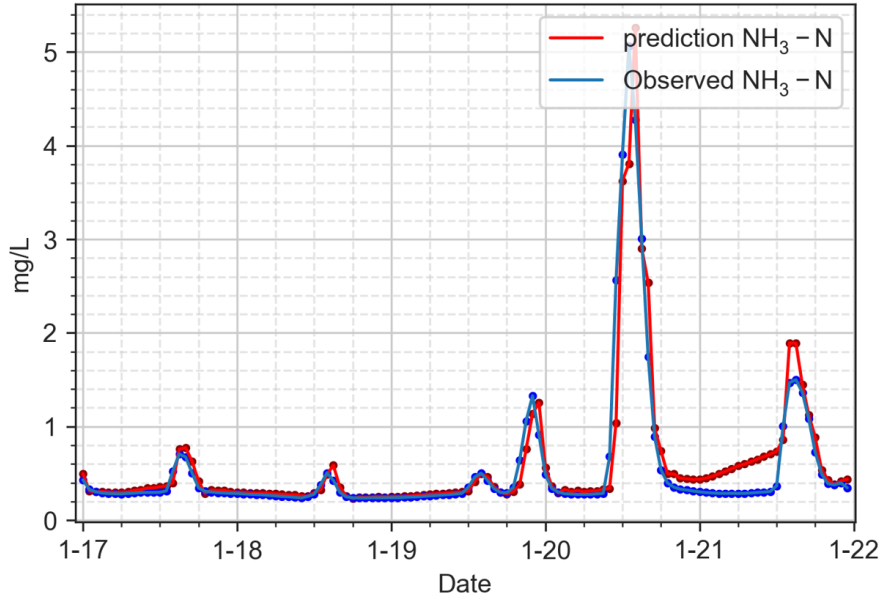


Figure 4.7: Comparisons of model performance in forecasting colour levels.

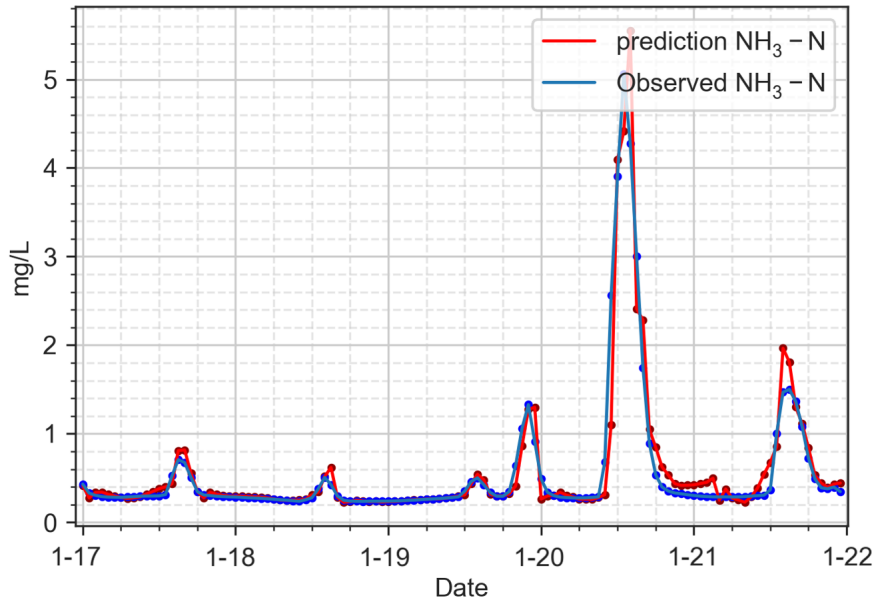
4.3.3 Model forecasting results on different forecast horizon

In this study, ammonia and colour forecasting models were input with data from the past 24 hours to forecast the values three hours into the future. To demonstrate how the proposed model training methods improved the baseline model performance, the forecasted results were visualized for easier comparisons. As shown in Fig. 4.8, the proposed model training methods helped the model to forecast better on 21 January as in Fig. 4.8b during the low ammonia concentration period. On other days, both LSTM-1-obs and LSTM-4-sg7 shared similar accuracy in forecasting ammonia concentration.

In forecasting ammonia concentration in the second hour into the future as in Fig. 4.9, both model showed much higher MSE values of 0.2916 and 0.2351 compared to the MSE values of 0.0647 and 0.0529 from Fig. 4.8. Both models forecasted the ammonia concentration fairly on 17, 18, 19, and 20 January but forecasted poorly on 21 January. During the last two days of forecasting, the patterns of ammonia concentration were quite different compared to the previous four days. For instance, on the 20 January, the peak concentration of ammonia during the day reached to 5.0 mg/L. Both models seemed unable to



(a) LSTM-1-obs, MSE = 0.0647

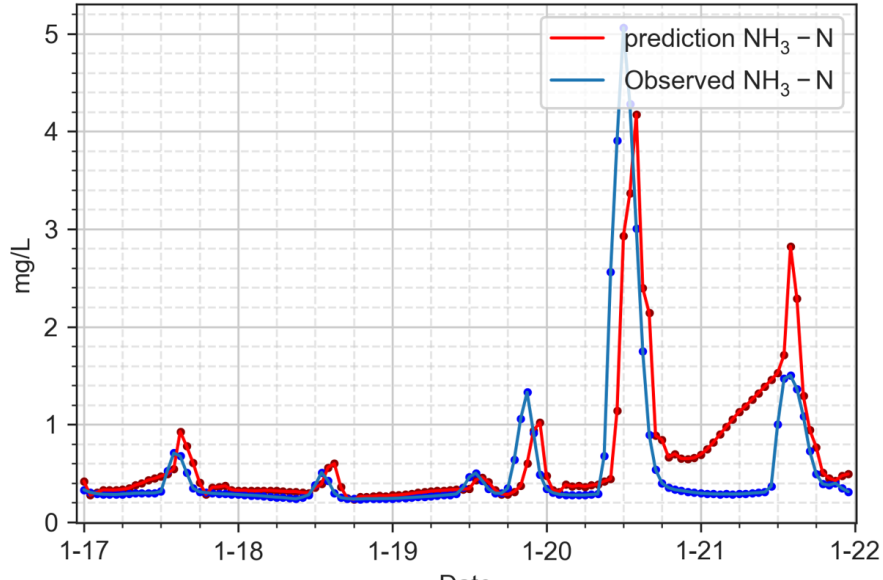


(b) LSTM-4-sg7, MSE = 0.0529

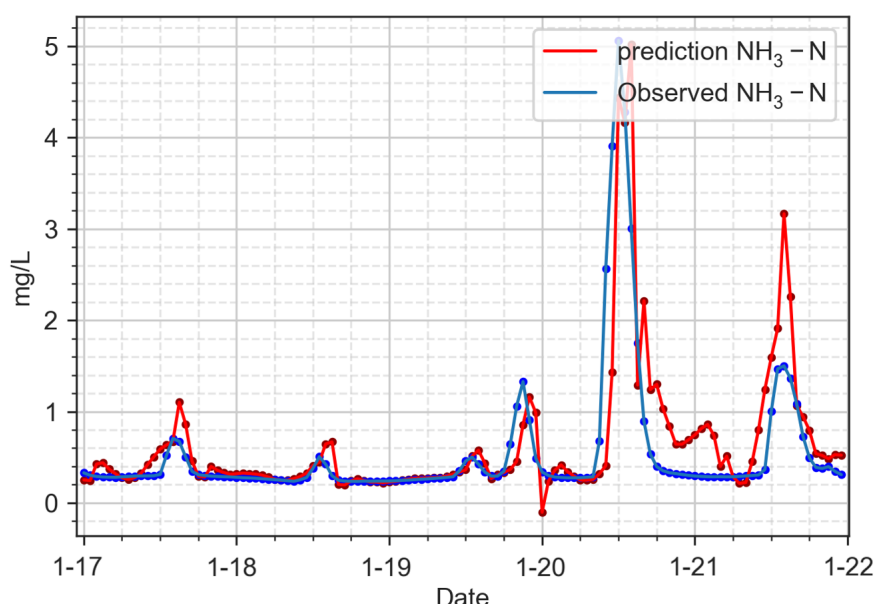
Figure 4.8: Visualization of ammonia forecasting models at forecast horizon of one.

precisely forecast the trend of the ammonia concentrations, resulting in overestimated ammonia concentration around noon on 21 January. The proposed model training methods did not seem to forecast better than the baseline model. Forecasting longer time horizons requires an adequate training dataset size in terms of the number of training features and the length of the dataset. The ammonia forecasting model as in Fig. 4.9b was trained with four features with a dataset length of 18 days. Yet, the results suggested that the

quantity of training dataset is not sufficient enough for forecasting two hours into the future.



(a) LSTM-1-obs, MSE = 0.2916

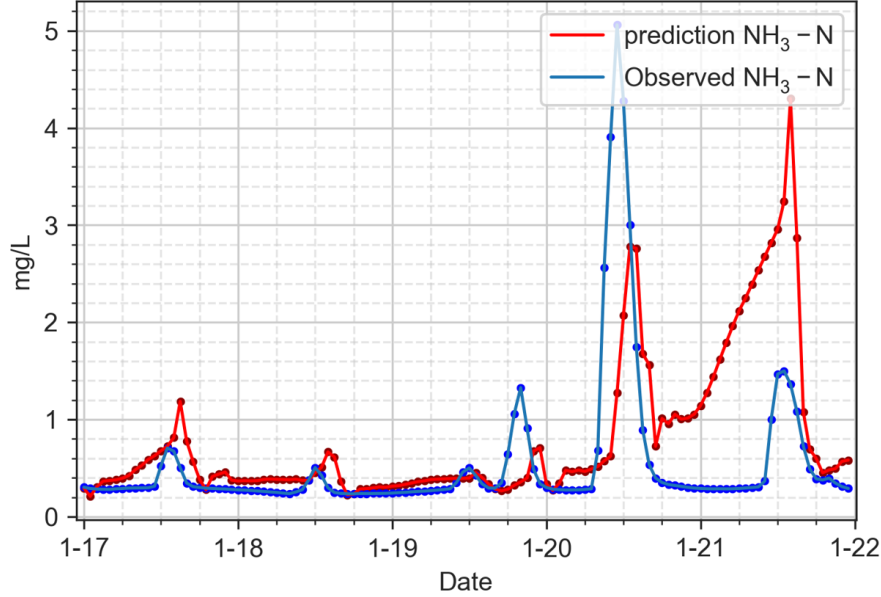


(b) LSTM-4-sg7, MSE = 0.2351

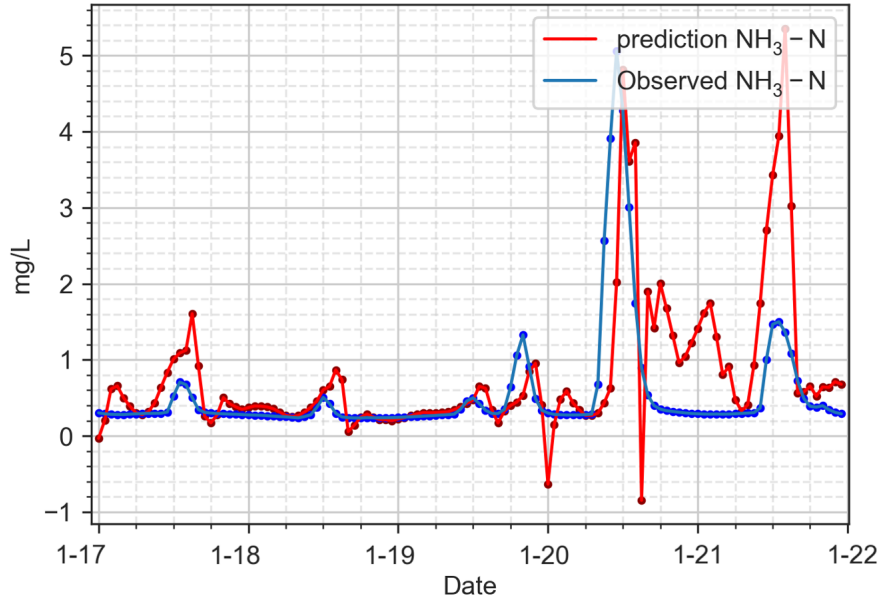
Figure 4.9: Visualization of ammonia forecasting models at forecast horizon of two.

In forecasting ammonia concentration at a forecast horizon of three, although the MSE values of 0.7637 from LSTM-4-sg7 are lower than 0.8025 from LSTM-1-obs, the difference between the two model performances is negligible. For the LSTM-4-sg7 model, we observed ammonia concentrations lower than 0 mg/L were forecasted on 20 January. Both

LSTM-4-sg7 and LSTM-1-obs models poorly forecasted the peak ammonia concentration of over 5.0 mg/L on 21 January, which is 3.0 mg/L higher than the actual ammonia concentration on the same day. The results suggest that even with the use of proposed model training methods, the capability of the model performance is still limited due to the limited size of the training dataset.



(a) LSTM-1-obs, MSE = 0.8025

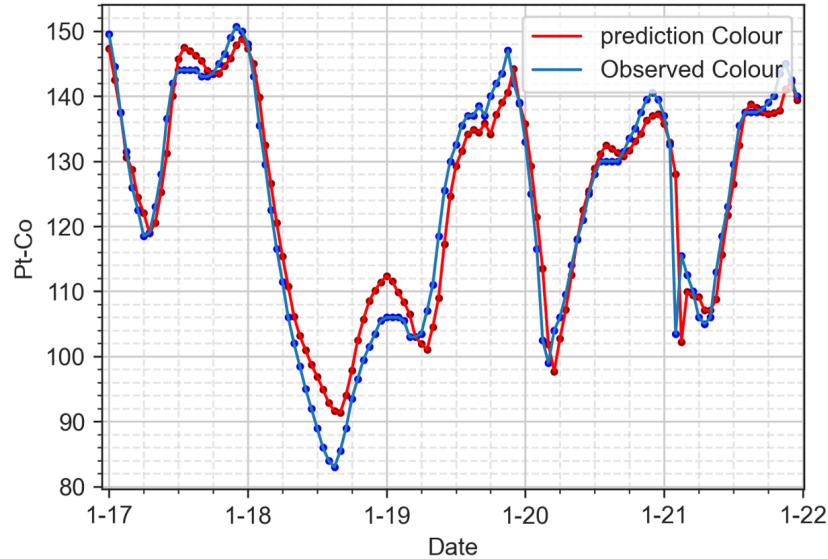


(b) LSTM-4-sg7, MSE = 0.7637

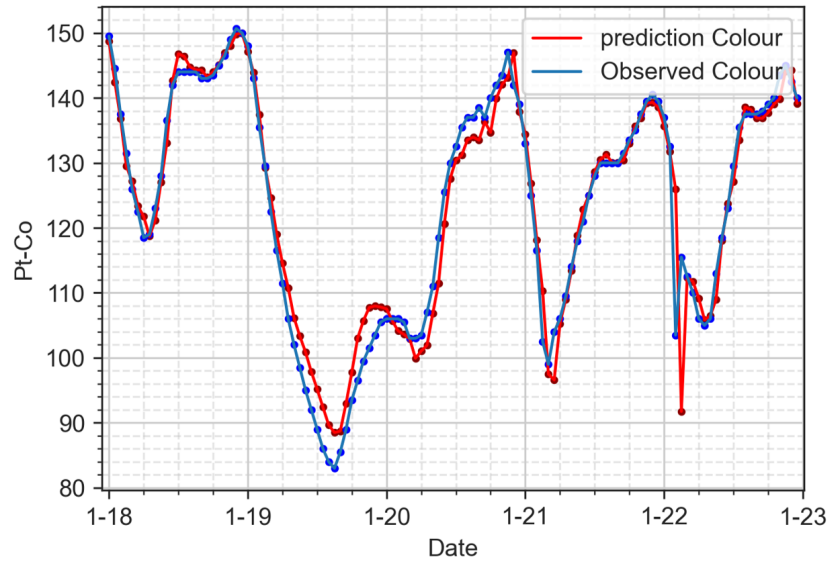
Figure 4.10: Visualization of ammonia forecasting models at forecast horizon of three.

LSTM-1-obs and LSTM-3-sg9 models forecasted colour levels at a forecast horizon of

one with good MSE values of 22.4922 and 17.5955. The errors between the actual and forecasted values are mostly less than 5 Hazen Units. On 18 January, the colour levels dropped to 80 Hazen Units, and both models forecasted colour levels with errors values of up to 5 Hazen Units and higher. Although on 22 January, the LSTM-3-sg9 model forecasted the colour level of 92 Hazen Units, which is 10 Hazen Units off from the actual values, the general model performance is satisfactory.



(a) LSTM-1-obs, MSE = 22.4922

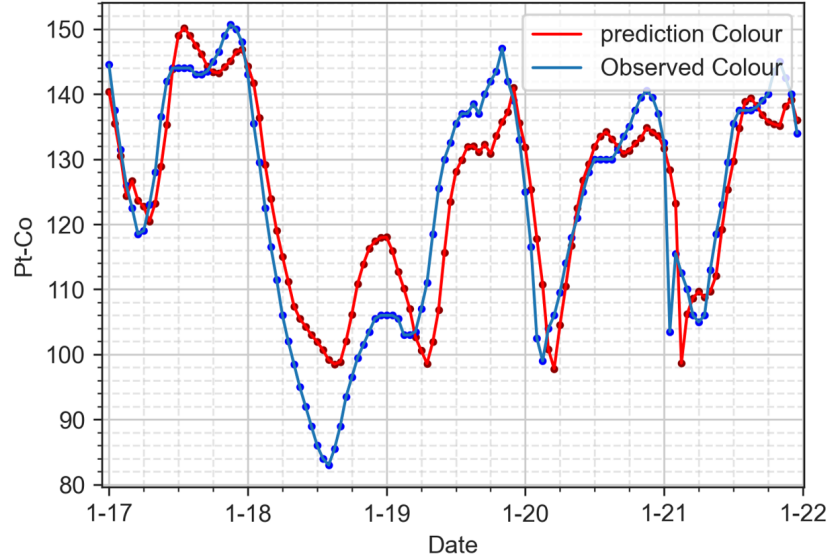


(b) LSTM-3-sg9, MSE = 17.5955

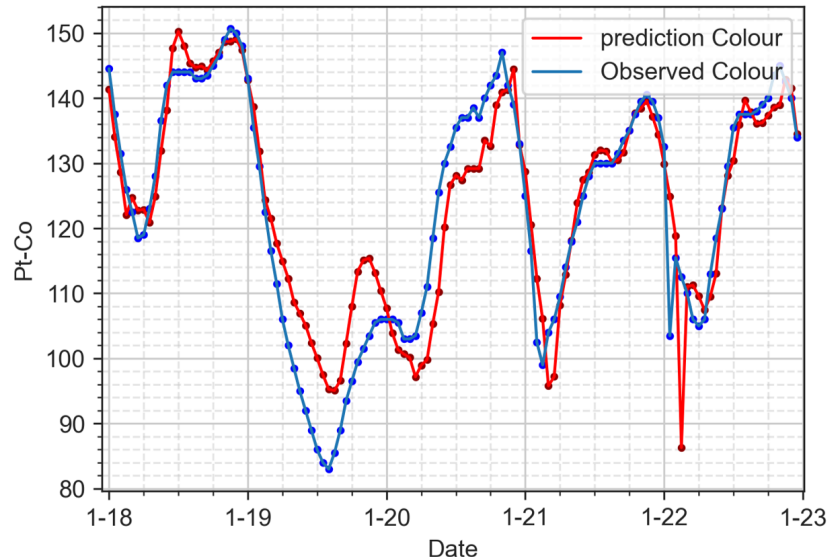
Figure 4.11: Visualization of colour forecasting models at a forecast horizon of one.

In forecasting colour levels at a forecast horizon of two, the MSE values of LSTM-1-obs and LSTM-3-sg9 increased from 22.4922 and 17.5955 to 62.6678 and 47.4252. The

forecasting errors expanded from less than 5 Hazen Units on average to 10 Hazen Units. In Fig. 4.12, LSTM-3-sg9 showed more reliable forecasting results compared to LSTM-1-obs by generating minor errors between the forecasted and actual values. However, the lowest forecasted colour level on 22 January has increased from 10 to 24 Hazen Unis, and we can see clearly that the models were getting less reliable in forecasting two hours into the future in forecasting colour levels. The cause of it can also be attributed to insufficient quantity of training dataset.



(a) LSTM-1-obs, MSE = 62.6678

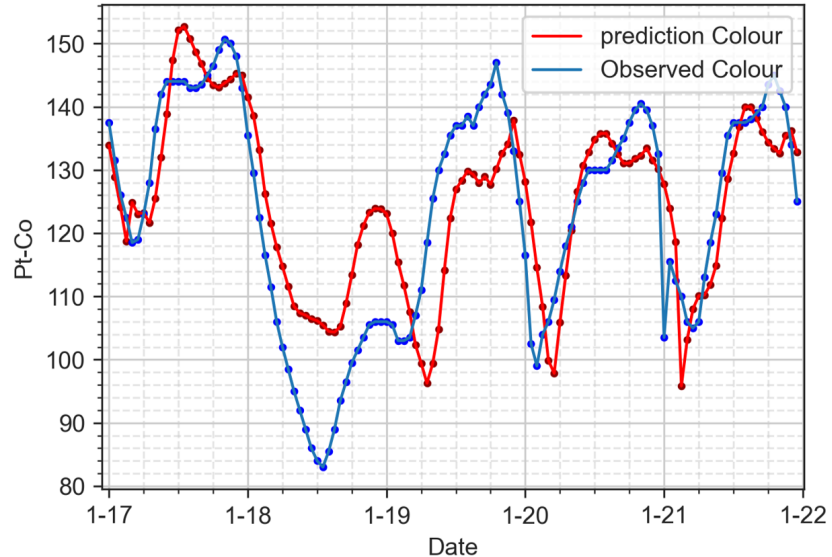


(b) LSTM-3-sg9, MSE = 47.4252

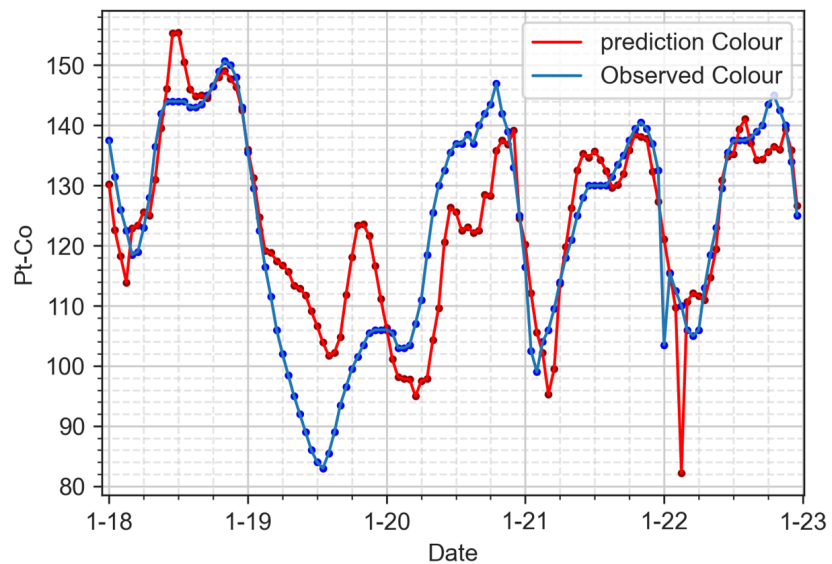
Figure 4.12: Visualization of colour forecasting models at forecast horizon of two.

In Fig. 4.13, the MSE values of LSTM-1-obs and LSTM-3-sg9 have increased to

116.8928 and 103.4329 in forecasting colour levels at forecast horizons of three. We first noticed that both the models failed to forecast the lowest colour levels on 19 January. The significant drop in colour level can be a rare event in which the model did not learn how to react to such a change of colour levels from historical data. On the following days of 20 January, both the models underestimated the colour levels by forecasting up to 20 Hazen Units lower. The model performance deteriorated even faster than using ammonia forecasting models to forecast ammonia concentration at a forecast horizon of three. The results suggest that with much strong fluctuation of colour levels during the day, it is not reasonable to use colour forecasting models trained with only three input features to forecast three hours into the future.



(a) LSTM-1-obs, MSE = 116.8928



(b) LSTM-3-sg9, MSE = 103.4329

Figure 4.13: Visualization of colour forecasting models at forecast horizon of three.

CHAPTER 5

CONCLUSIONS AND RECOMMENDATION

5.1 Conclusions

5.1.1 Machine learning models vs deep learning models

The selection of using which machine learning and deep learning models was not widely discussed to the best of our knowledge in modelling forecasting models in the wastewater treatment industry. This study has investigated the model performance of the machine learning model of RF and four other deep learning models of DNN, RNN, GRU, and LSTM on forecasting ammonia concentrations and colour levels in the reclaimed water system for assisting treatment operation and management. The evidence from this study suggested deep learning models are much capable of learning from historical data and generating more accurate forecasting results. In both ammonia and colour forecasting models, the test loss values of RF are much higher than those of the least-performance deep learning model of DNN. Among all the deep learning models, the results indicate that LSTM and GRU models have the lowest test loss of 0.0405 and 0.0414, respectively. However, further research works suggest that LSTM models trained with pre-processing methods generate the lowest test loss compared to GRU, making the LSTM model the most promising recurrent neural network model for training forecasting models in WWTPs.

5.1.2 Data pre-processing techniques

Our research also highlighted how the model performance could be improved by applying data pre-processing and feature engineering techniques. Generally speaking, all the proposed data smoothing and outlier removal methods reduced the test loss values compared to the baseline model performance (i.e., the window sizes of the data smoothing filters need to be carefully selected), as shown in Fig. 4.5. Ammonia and colour forecasting models trained by EWMA filtered datasets showed the lowest test loss values compared with models trained by SG filtered datasets and datasets applied with outlier removal

methods. Applying an EWMA filter on training datasets can reduce the noise and allow the important patterns to stand out more clearly. The information hidden in the convoluted data points then can be further captured by the memorizing cells in the recurrent neural networks such as GRU and LSTM.

5.1.3 Feature engineering

This study is the first step towards enhancing our understanding of the potential benefits of using created features for model training. The thorough examinations of the Geomap near the SHWEPP and the investigation of water composition in the public sewage system helped us hypothesize that the change of ammonia concentrations and colour levels depend on each other. With the help of an additional colour/ammonia feature for the ammonia/colour forecasting models, the test loss was reduced by 6.4% (i.e., LSTM-2-sg7 compared to LSTM-1-obs) and 10.8% (i.e., LSTM-2-ew4 compared to LSTM-1-obs), respectively.

Moreover, the similarity between the household consumption patterns and the daily fluctuation of ammonia concentrations have unexpectedly helped us formulate the time features via positional encodings. The influence of the sine and cosine hour features on the model performance showed tremendous improvements in both ammonia and colour forecasting models. In the former, test loss dropped by 8.9% (i.e., LSTM-1-obs compared with LSTM-4-sg7) while the latter reduced by 28.6% (i.e., LSTM-1-obs compared with LSTM-3-sg9). The remarkable use of positional encoding features is that they are not limited to ammonia and colour forecasting models. Any time-series data characterized by daily fluctuation patterns can adopt the use of the features of sine and cosine hour as long as the patterns are based on actual events. In addition, the positional encoding features are not limited to the hour component, we can encode time component features from seconds to weeks, and even years, the application of it is unlimited. However, the feature engineering method comes with limitations. In the results of ammonia forecasting models, LSTM-2-obs, LSTM-3-obs, and LSTM-4-obs showed higher test loss compared to LSTM-1-obs, indicating that when the models were not trained with ammonia feature only, the model performance worsened. Our results suggested that feature engineering needs to be carefully evaluated and experimented with before its real application. Despite the limitations, the combination use of feature engineering in building ammonia and colour

forecasting models in this study has fully proved its advantages.

5.2 Recommendations for future research

Due to the insufficient amount of ammonia and colour data, we cannot differentiate whether the undesired model performance was caused by the heterogeneity of the validation and testing datasets or caused by the pre-processing and feature engineering techniques we applied to the datasets. It is recommended a larger dataset (e.g., a larger dataset in length and better data quality with more input features for training) should be used in the future study when evaluating the proposed methods in this study. The insufficient data could also lead to the unstable performance of different models trained by the same data smoothing techniques. For instance, models trained by sg7 filtered dataset (LSTM-4-sg7 and LSTM-3-sg7) have the lowest test loss values; however, LSTM-2-ew4 has a lower test loss than LSTM-4-sg7. We failed to explain why models trained by the sg7 filtered dataset influenced ammonia forecasting models in different ways among LSTM-2, LSTM-3, and LSTM-4. It is necessary to elucidate the influence of each data pre-processing technique to establish robust strategies for smoothing the training datasets.

All the forecasting models in this study only focus on predicting ammonia concentration and colour levels in the reclaimed water system. In future research, more water quality parameters should be included. In reclaimed water systems, the concentration of water quality parameters such as turbidity and E. coli are also regulated by Water Supply Department. Violating any water quality parameter will directly lead to the disqualification of being used as reclaimed water. Using more water quality parameters as features has extra benefits for building forecasting models. The hidden correlations between each water quality parameter will most likely help build more accurate water quality forecasting models.

Previous research studies have demonstrated using Matlab-Simulink to simulate the improved process control strategies using machine learning model controls compared to PID or other traditional mathematical models. In future works, the study will explore writing the physical and operational characteristics of the water reclaimed system into the Matlab-Simulink. By implementing the models developed in this study on Matlab, we can investigate how the improvements in model forecasting accuracy can help the process control strategy in stabilizing the reclaimed water quality. Several metrics can be

used to evaluate the machine learning model control, such as the required time to reach set-point conditions and how much reclaimed water in volume we can generate from the same amount of wastewater effluent recycled.

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