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Concentration Prediction-based Crop Digital Twin Using Nutrient Co-existence and Composition in Regression Algorithms

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Abstract: Crop digital twin is redefining traditional farming practices, offering unprecedented op-13 portunities for real-time monitoring, predictive and simulation analysis, and optimization. This re-14 search embarks on an exploration of the synergy between precision agriculture, crop modeling, and 15 regression algorithms to create a digital twin for augmenting farmers the concentration and compo-16 sition prediction-based crop nutrient recovery. This captures the holistic representation of crop char-17 acteristics, considering the intricate relationships between environmental factors, nutrient concen-18 trations, and crop compositions. However, the complexity arising from diverse soil and environ-19 mental conditions makes nutrient content analysis expensive and time-consuming. This paper pre-20 sents two approaches namely,(i) single nutrient concentration prediction and (ii) nutrient composi-21 tion concentration prediction which is the result of a predictive digital twin case study that employs 22 six regression algorithms namely Elastic Net, Polynomial, Stepwise, Ridge, Lasso, and Linear Re-23 gression to predict rice nutrient content efficiently, particularly considering the coexistence and 24 composition of multiple nutrients. Our research findings highlight the superiority of the Polynomial 25 Regression model in predicting nutrient content, with a specific focus on accurate nitrogen percent-26 age prediction. This insight can be used for nutrient recovery intervention by knowing the precise 27 amount of nutrient to be added into the crop medium. The adoption of the Polynomial Regression 28 model offers a valuable tool for nutrient management practices in the crop digital twin, potentially 29 resulting in higher-quality rice production and a reduced environmental impact. The proposed 30 method can be replicable in other low-resourced crop digital twin system. 31

Keywords: rice nutrient level; fertilizer optimization; nutrient analysis; polynomial regression; nutrient prediction; environmental impact reduction

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1. Introduction

Digital twin technology involves the creation of a virtual duplicate of a physical object or system, enabling the simulation and analysis of diverse scenarios and outcomes [1-7]. When applied to crop management, a digital twin becomes a powerful tool for modeling a specific farm, considering variables such as soil quality, weather conditions, irrigation systems, and crop varieties. This collected data is then utilized to update the digital 40 twin, facilitating predictions about upcoming crop yields, potential pest outbreaks, and 41 other influential factors that may impact the farm's overall success.

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Employing Digital Twins as a primary method for farm management facilitates the 43 separation of physical processes from their planning and control. Consequently, farmers 44 gain the capability to oversee operations and crop health remotely, relying on (almost) 45 real-time digital information rather than depending solely on direct observation and on-46 site manual tasks [6,7]. The deficiency of vital nutrients can lead to reduced crop yields [8-47 13]. This empowerment enables prompt action in response to anticipated or unexpected 48 deviations such as crop nutrient concentration and allows for the simulation of the effects 49 of interventions such as nutrient recovery based on real-life data [14-18]. 50

In this context, the application of machine learning (ML) offers a promising avenue 51 for farmers. ML equips them with tools for monitoring soil quality and delivering person-52 alized recommendations, drawing insights from both experimental and field data. None-53 theless, the prediction of rice essential nutrients remains a formidable challenge, primarily 54 due to several factors: 1) the inherent variability in nutrient content, 2) the diversity of 55 analytical approaches, 3) limitations in data availability, 4) genetic diversity among rice 56 varieties, and 5) the associated cost and time constraints [16-19]. Consequently, it is im-57 perative to address these multifaceted challenges to develop accurate and reliable nutrient 58 prediction models for rice [15-17]. 59

This paper report one of our digital twin case studies on rice nutrient recovery 60 through two approaches namely single nutrient concentration prediction and nutrient 61 composition concentration prediction. Regression facilitates the identification of intricate 62 relationships among essential rice nutrients, ensuring their optimal supply, thereby en-63 hancing rice growth and nutrient content [20,21]. This study seeks to identify the most 64 effective regression algorithm for predicting nutrient concentration percentages based on 65 the co-existence and composition of other nutrients. The incorporation of regression algo-66 rithms in the crop digital twin is mainly because of its efficiency and effectiveness. This 67 endeavor promises optimized nutrient management practices, culminating in enhanced 68 rice quality and a reduced environmental footprint through the adjustment of nutrient 69 ratios. 70

Among the myriad regression algorithms, Elastic Net regression, Polynomial regres-71 sion, Stepwise regression, Ridge regression, Lasso regression, and Linear regression hold 72 particular relevance for predicting nutrient concentration by considering the coexistence 73 and composition of multiple nutrients. These algorithms offer a structured, data-driven 74 approach to unravel the complexities of rice nutrition, providing accurate predictions and 75 contributing to the standardization of nutrient management practices. Moreover, they 76 play a crucial role in fostering sustainable and environmentally friendly rice cultivation 77 practices. 78

The singular nutrient prediction method offers advantages in two distinct scenarios. 79 Firstly, it proves beneficial when a farmer or scientist intends to simulate the concentration value of a specific nutrient, already possessing knowledge of the concentration of 81 other nutrient components. Secondly, this approach becomes valuable if the sensor for a 82 particular nutrient malfunctions. In such cases, the digital twin system promptly alerts the 83 user regarding the sensor breakdown and provides a predictive value while awaiting sensor replacement. 85

Regardless of the scenario, the digital twin system ensures user awareness when the detected nutrient concentration surpasses the recommended range. Furthermore, the system recommends nutrient recovery interventions. The nutrient composition prediction approach serves as a comprehensive intervention preparation tool by informing the farmer or scientist about the anticipated nutrient concentration. The projected value, in turn, aids the digital twin system in suggesting the appropriate amount of nutrient recovery, aligning with best practices.

This paper unfolds in six sections. The first section underscores the significance of 93 predicting rice essential nutrients and elucidates the challenges in this domain, along with 94 the role of linear and polynomial regression algorithms in addressing these issues. The 95 second section offers an overview of the dataset and its attributes. The third section delin-96 eates the flowchart of the polynomial regression algorithm. The fourth section introduces 97 the evaluation metrics employed to assess algorithm performance. The fifth section pre-98 sents the experimental results and their comprehensive analysis. Finally, the paper con-99 cludes by summarizing the findings and proposing potential avenues for future research. 100

2. Literature Review

One of the promises of digital twin in crop management is for automatic prediction 102 system to support deciding the appropriate fertilization period [22-24]. Deploying the sensors which monitors concentration of nutrients present in soil, humidity, and temperature 104 in the real fields to make the consistent quality check. Machine learning could be used as 105 a proactive measure as predictor of the degradation of crop medium's and crop's plant 106 nutrients which could increase the risk of crop pests and diseases [25,26]. 107

Regression algorithms play a central role in rice nutrient prediction by unraveling 108 the intricate interplay of nutrients in rice cultivation. Elastic Net Regression (EN), Polyno-109 mial Regression (PN), Stepwise Regression (SW), Ridge Regression(RR), Lasso Regres-110 sion(LS), and Linear Regression(LR) provide essential insights into the complex relation-111 ships among soil composition, environmental variables, and agricultural practices [27-30]. 112 These algorithms empower researchers to comprehend the often-nonlinear dependencies 113 among these factors, deepening our understanding of how various nutrients influence rice 114 nutrition. 115

Regression algorithms are data-driven, offering a robust framework for analyzing 116 and interpreting nutrient data from diverse sources. By harnessing historical data and observational insights, these algorithms provide crucial guidance on how different nutrients 118 impact rice composition. This knowledge is vital for optimizing fertilizer usage, enhancing nutrient management, and ultimately improving rice quality and yields [27-30]. 120

These algorithms also aid farmers, agricultural experts, and policymakers in making121informed decisions about crop management, fertilization strategies, and soil enrichment.122This proactive approach helps avoid over-fertilization or under-fertilization, mitigating123their detrimental effects on crop health and environmental sustainability [31,32].124

Existing works on rice nutrient has focused on predicting essential nutrient levels in 125 rice, such as N, P, K, Mg, and Ca, and their effects on rice plant growth and development. 126 One study employed an artificial neural network-based prediction algorithm to assess the 127 influence of individual nutrients (N, P, K, Zn, and S) on various rice plant parameters. The 128 algorithm indicated that optimal growth often occurs with nutrient doses below the maximum applied levels, while maximum yield is achieved at 100% nutrient dose [22]. 130

Another study used regression methods and found that random forest regression al-131 gorithms provided the highest accuracy for estimating rice shoot dry matter, leaf area in-132 dex, and nitrogen accumulation [23]. A third study evaluated different approaches for 133 estimating rice aboveground biomass, plant nitrogen uptake, and nitrogen nutrition in-134 dex, with the random forest algorithm demonstrating superior performance [25]. An ad-135 ditional study focused on using machine learning for early detection of nutrient deficiency 136 in rice through leaf image processing, achieving high testing accuracy and roc_auc score 137 [8]. 138

Rice nutrient content prediction, based on the composition of other nutrient infor-139 mation, including nitrogen, phosphorus, potassium, and organic matter as input varia-140 bles, was addressed in a study [26]. This study compared the EN algorithm with tradi-141 tional linear regression methods, including Ordinary Least Squares (OLS) regression, 142 Ridge regression, and Lasso regression. The results highlighted the superior performance 143 of the EN algorithm, exhibiting higher R-squared scores (R2) and lower Mean Absolute 144 Error (MAE). Thus, Elastic Net proves more accurate in predicting rice nutrient content 145 and its correlation with other nutrients. 146

Essential nutrient levels in rice can also be predicted using spectral data from remote 147 sensing [28], considering nutrients like N, P, K, Mg, and Ca. This research compared the 148 polynomial regression algorithm with two other methods: Multi linear regression (MLR) 149 and Partial least squares regression (PLSR). The outcome demonstrated the polynomial 150 algorithm's superiority in predicting nutrient concentrations in rice levels. 151

Other studies predicting nutrient content in rice used 16 nutrients as predictors, such 152 as moisture, crude protein, fat, ash, total dietary fiber, soluble dietary fiber, insoluble dietary fiber, total sugar, sucrose, glucose, fructose, amylose, amylopectin, total amino acids, 154 lysine, and thiamine [30]. These studies employed three algorithms: stepwise regression, 155 PLSR, and MLR for prediction. The results favored stepwise regression analysis for its superior accuracy in predicting nutrient content in rice. 157

Another study aimed to predict nutrient content in rice based on 14 nutrients, including moisture, crude protein, fat, ash, total dietary fiber, soluble dietary fiber, insoluble dietary fiber, total sugar, sucrose, glucose, fructose, amylose, amylopectin, and thiamine. This research compared three algorithms: ridge regression, principal component regression (PCR), and PLSR. Ridge regression stood out as the most effective method for predicting nutrient content in rice, delivering higher accuracy than PLSR and PCR. 163

Utilizing another set of 14 nutrients, including moisture, crude protein, fat, ash, total 164 dietary fiber, soluble dietary fiber, insoluble dietary fiber, total sugar, sucrose, glucose, 165 fructose, amylose, amylopectin, and thiamine as predictors for nutrient prediction in rice, 166 another study employed three algorithms: MLR, PLSR, and lasso regression. The experi-167mental results highlighted the precision of the lasso regression algorithm in predicting168both yield and nutrient content in rice, offering potential benefits in optimizing rice crop169cultivation and management.170

In a similar vein, another study [34,35] compared three prediction algorithms, 171 namely MLR, PLSR, and PCR, for nutrient content in rice, considering nutrients such as 172 moisture, crude protein, fat, ash, total dietary fiber, soluble dietary fiber, insoluble dietary 173 fiber, total sugar, sucrose, glucose, fructose, amylose, amylopectin, and thiamine. The 174 findings indicated that MLR provided more accurate predictions compared to the other 175 methods assessed. 176

Table 1 provides a comparative analysis of the advantages and disadvantages of re-177gression algorithms [26-33] for rice nutrient prediction. These algorithms effectively cap-178ture both linear and nonlinear correlations among various nutrients.179

Table 1. Advantage and disadvantage of linear regression algorithm.

Linear regression			
Types	Proficiency	Advantage	Disadvantage
Simple linear regres- sion (LR)[25]	Identifying the corre- lation between two variables	-Computationally effi- cient -Required less parame- ters	-Unable to deal with nonlinearity -Sensitive to out- lier
Elastic Net Regres- sion (EN) [26]	Constructed by com- bination of Lasso and Ridge regression models.	-Able to deal with large number of features -Prevent overfitting us- ing L1 and L2 regulari- zation methods	-Computationally expensive -Unsatisfactory results when the number of predic- tors is more than
Polynomial Regres- sion (PR)[28]	Captures nonlinearity between variables	-Ability to deal with small dataset	- Computationally expensive -Overfit if the de- gree of polyno- mial is high
Stepwise Regression (SW)[30]	Built by combination of backward and for- ward selection meth- ods which is beneficial to select best subset of features	-Provide balance be- tween features and al- gorithms predictive power	-Time demanding -Unstable due to overfitting
Ridge Regression (RR) [31]	Considered as regulari- zation method	-Able to dela with large dataset -Prevent overfitting	-Issue with find- ing optimal value for lambda

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			-Challenging
			while dealing
Lagao Degraggion	Known as regulariza-	Mitigata avarfitting	with large dataset
	tion method	-mingate overntting	that has large
[33]			number of obser-
			vations

These diverse regression algorithms collectively share a common aim: to enhance the 181 precision and reliability of predictions concerning rice nutrient content, a critical step in 182 optimizing fertilizer application, ensuring a balanced nutrient supply, and ultimately el-183 evating rice crop quality and yield while reducing environmental impact. 184

However, very limited works have addressed the crop's nutrients prediction by fo-185 cusing on the co-existent and composition nutrient's concentration. For a digital twin sys-186 tem equipped with crop nutrients surveillance, this comes to our advantage to enable crop 187 nutrient recovery. Our exploration and application of these regression techniques serve 188 to address prevailing research disparities and foster a more standardized and comprehen-189 sive approach to predicting rice nutrient content. By employing a variety of regression 190 models, our objective is to gain a deeper understanding of the intricate relationships 191 among different nutrients in rice. This, in turn, promotes more sustainable and efficient 192 rice cultivation practices. 193

3. Materials and Methods

This part splits into three subsections. First, we explain the dataset and its attribute. Next, we present the setting of the regression models. Then, we discuss the evaluation 196 metrics. 197

3.1. Dataset Description

A self-collected rice dataset was used as described in Table 2, comprising of 348 observations and nine attributes. This multivariate dataset features a combination of cate-200 gorical and numerical data, including spatiotemporal factors such as Season, Day, Plot, 201 and Subplot. 202

Table 2. Rice dataset descriptions.

Name of Dataset	Rice Dataset
Dataset Characteristics	Multivariate
Attribute Characteristics	Categorical Data (Nominal), Numerical & Continual Data
Number of Instances	348
Attributes Number	9
Missing Values	No

The Season attribute categorizes data into two distinct seasons, denoted by the values 204 1 and 2, enabling the exploration of how seasonal changes influence rice nutrient levels, a 205 fundamental aspect of rice production optimization. Additionally, the Day attribute, with 206 three distinct values -30, 60, and 90, introduces temporal granularity, facilitating an ex-207 amination of nutrient content variations within each season. This temporal dimension is 208 essential for understanding the influence of specific days on nutrient levels. 209

Furthermore, the Plot attribute categorizes data into four distinct plot locations rep-210 resented by values 1, 3, 4, and 5, enabling the assessment of nutrient distribution across 211 different areas within the study site, thus adding a spatial context to the analysis. Subplot 212

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further refines the spatial information by specifying 15 sublocations within each plot, denoted by values such as 1A, 1B, 1C, and so forth. 213

This fine-grained attribute is invaluable for scrutinizing nutrient variation within 215 specific subregions of the plots, enhancing spatial precision. Additionally, the dataset in-216 corporates nutrient concentration, composition and co-existence ('N%', 'P%', 'K%', 'Mg%', 217 'Ca%'), which is vital for understanding rice growth and health. The dataset's integrity is 218 maintained, as it contains no missing values. 219

Season	T	Day T	Plot 🔻	Subplot 🔫	N (%) 👎	P (%) 📼	K (%) 👘	Mg (%) 🗦	Ca (%) 📑
	2	90	fadh ₅	ha 1A	1.62	0.27	1.85	0.14	0.23
	2	90	5	1B	1.75	0.25	2.33	0.17	0.24
	2	90	Days after 5	owing 1C	2.01	0.23	2.17	0.17	0.26
	2	90	5	2A	1.98	0.22	2.24	0.14	0.26
	2	90	From imported	locumer2B	1.78	0.28	2.34	0.15	0.29

Figure 1. Example content of the dataset

Example of the data content is shown in Figure 1, which shows the concentration of each 222 nutrient based on the spatial information. The best range of the nutrients are N: [1.17, 223 2.47], P: [0.25, 0.3], K: [1.85, 2.52], Mg: [0.11, 0.17], Ca: [0.23, 0.33] which has produced the 224 maximum weight grain at the planting plot with range [29.26, 39.42] at the end of the 225 planting cycle. These values are considered the best practice to guide for intervention plan 226 for the user (farmer or scientist). 227

Figure 2 shows the dashboard that presents the average rice nutrient concentration228across the growth period and the rice anatomical values at harvesting time, while Figure2293 shows the nutrient value distribution. From Figure 2, we can identify the relationship of230the nutrient con-existence, composition, and concentration with the yield. The digital twin231supports three-staged insight for crop intelligence. First, we could also see the average232values of nutrients that have led to the yield, and the nutrient values from the plant with233the best yield become the benchmark.234

So, this has motivated us towards the second intelligence by predicting the co-existence, concentration, and composition of the plant at each plot and subplot to know their health. The third intelligence is nutrient recovery during the growth as an intervention mechanism so that the predicted values can be a guide on precise additional nutrients to be added into the crop medium to optimize the yield. The precision of values for additional nutrients can mitigate unnecessary excess in fertilizer usage and waste pollution. 230

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Figure 2. Dashboard about the average nutrient values and the content in the rice.

Table 3. Valu	es distribution	for the nutrients
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	N (%)	P (%)	K (%)	Mg (%)	Ca (%)
MIN	0.15	0.15	1.61	0.09	0.16
MAX	4.59	28.00	3.89	0.20	0.38
STDEV	0.77	1.48	0.45	0.02	0.04

The nutrient concentration distribution, as depicted in Table 3, highlights the range 244 of values for key nutrients of N (%), P (%), K (%), Mg (%), and Ca (%) that is essential for 245 agricultural productivity. The minimum (MIN) and maximum (MAX) values illustrate the 246 variability in nutrient levels, emphasizing the complexity of nutrient dynamics in agricul-247 ture. Standard deviation (STDEV) values quantify the degree of variability around the 248 mean. This information is instrumental in precision agriculture, guiding targeted inter-249 ventions based on specific nutrient needs. In the context of environmental sustainability, 250 understanding these distributions enables our digital twin system to issue timely alerts 251 and recommend nutrient recovery interventions when concentrations exceed recom-252 mended ranges. This proactive approach optimizes crop yield while minimizing the en-253 vironmental impact associated with nutrient imbalances. 254

3.2. Data Pre-Processing Using Min-Max Normalization

Before visualization, the data exhibited variations in nutrient concentrations that prompted the need for exploration. The raw data contained outliers, which are data points 257 significantly different from the majority of the observations. These outliers, if not ad-258 dressed, can impact the understanding of the overall nutrient distribution and make it 259 challenging to discern patterns and trends in the data. 260

Therefore, to gain a deeper understanding of the nutrient concentration data and visualize its distribution, we employed box plots both before and after applying Min-Max normalization. The original box plots revealed the presence of outliers in the dataset, 263

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which was affecting the clarity of the distribution. To address this issue, Min-Max normalization was applied to scale the data. The box plots after normalization effectively
showcased the distribution of nutrient concentrations without displaying outliers. This
approach allows for a more accurate and informative representation of the data, aiding in
the identification of central tendencies and variations while providing a clearer view of
the data's overall structure. The use of box plots before and after normalization aids in the
assessment of data quality and the impact of data preprocessing techniques.

The Min-Max normalization method is applied to rescale the input features between 271 0 and 1 during the pre-processing phase. This normalization technique is suitable for the 272 prediction models of this study because it helps to ensure that all the input features are 273 on the same scale and have the same range, which helps the linear regression models of 274 this study converge faster and boost their performance. This approach removes noises 275 from data and prevents the big scales from data by giving the range of [0,1]. Equation (1) 276 shows the formula of the Min-MAX method. 277

$$X_{Norm} = \frac{(X - X_{Min})}{(X_{Max} - X_{Min})} \tag{1}$$

Where X is the original value of a data point, X_{Min} is the minimum value in the data-278taset, X_{Max} is the maximum value in the dataset, and X_{Norm} is the normalized value of the279data point. This formula ensures that the minimum value in the dataset is scaled to 0 and280the maximum value is scaled to 1, with all other values falling between these two limits.281

By applying a preprocessing method to the dataset, we can improve the stability and 282 performance of regression models. Once this stage is complete, we can proceed to the next 283 stage, where we design a regression model based on the different variables in the dataset. 284 This stage involves selecting an appropriate regression method and specifying the inde-285 pendent and dependent variables. Finally, we analyze the model and provide information 286 on its performance and accuracy. Figure 3 illustrates the Rice Nutrients data before and 287 after applying the Min-Max normalization method. The visual representation of the data 288 highlights the impact of normalization on the distribution of nutrient concentrations. 289



Figure 3. Rice Nutrient Data: (a) Original Data and (b) Min-Max Normalized Data.

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The dataset under analysis consists of nutrient concentration data for rice samples, 294 including attributes like nitrogen (N %), phosphorus (P %), potassium (K %), magnesium 295 (Mg %), and calcium (Ca %). Prior to visualization, the data exhibited variations in nutrient concentrations that prompted the need for exploration. The raw data contained outliers, which are data points significantly different from the majority of the observations. 298 These outliers, if not addressed, can impact the understanding of the overall nutrient distribution and make it challenging to discern patterns and trends in the data. 300

Therefore, to gain a deeper understanding of the nutrient concentration data and vis-301 ualize its distribution, we employed box plots both before and after applying Min-Max 302 normalization. The original box plots revealed the presence of outliers in the dataset, 303 which was affecting the clarity of the distribution. To address this issue, Min-Max nor-304 malization was applied to scale the data. The box plots after normalization effectively 305 showcased the distribution of nutrient concentrations without displaying outliers. This 306 approach allows for a more accurate and informative representation of the data, aiding in 307 the identification of central tendencies and variations while providing a clearer view of 308 the data's overall structure. The use of box plots before and after normalization aids in the 309 assessment of data quality and the impact of data preprocessing techniques. 310

3.3. Nutrient Concentration and Composition Prediction

We present two approaches namely (i) single nutrient concentration prediction and (ii) nutri-313ent composition concentration prediction; which are developed using EN, PN, SW, RR, LS, and314LR algorithms. This section describes the development of the prediction models.315

3.3.1. Single nutrient concentration prediction

We call the first approach single nutrient concentration prediction where five (5) 317 models are developed based on different feature sets of rice dataset as shown in Table 4 318 by exploiting the nutrient concentration, co-existence, and composition. In Table 4, 'Y' indicates that the spatiotemporal factors and nutrient features are used in the model building, while 'N' indicates otherwise. 321

 Table 4. Single nutrient concentration prediction setting

Spat	tiotemp	oral F	Nutrients						
Feature Set	Sea- son	Da y	Pl ot	Subplot	N (%)	P (%)	K (%)	Mg (%)	Ca (%)
FS1 (Ca%)	Y	Y	Y	Y	Y	Y	Y	Y	Ν
FS ₂ (Mg%)	Y	Y	Y	Y	Y	Y	Y	Ν	Y
FS3 (K%)	Y	Y	Y	Y	Y	Y	Ν	Y	Y
FS4 (P%)	Y	Y	Y	Y	Y	Ν	Y	Y	Y
FS ₅ (N%)	Y	Y	Y	Y	Ν	Y	Y	Y	Y

Referring to Table 4, the single nutrient concentration setting has been constructed 324 based on the selection of different features from spatiotemporal factors and nutrient features. These settings will be used for single nutrient concentration prediction using six 326 methods: EN, PN, SW, PR, LS, and LR. Table 5 presents the parameter specifications 327

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applied to the six regression approaches of EN, PN, SW, PR, LS, and LR in single nutrient 328 concentration and composition concentration prediction. 329

Table 5. Parameter specification for six regression algorithms of EN, PN, SW, PR, 330 LS, and LR in single nutrient concentration and composition concentration pre-331 diction 332

Model	Parameter	Values
EN	alpha	0.1
	L1_ratio	0.5
PN	degree	2
SW	Sequential Feature Selector	Automatically select features (no direct parameters involved)
PR	alpha	0.1
LS	alpha	0.1
LR	No additional parameters	

Table 5 outlines the parameter specifications for six regression algorithms of EN, PN, 334 SW, PR, LS, and LR in the context of predicting both single nutrient concentration and composition concentration. 336

For EN, the parameters include an alpha value of 0.1 and an L1_ratio of 0.5. PN em-337 ploys a degree of 2 for modeling. The SW automatically selects features without involving 338 direct parameters. PR is characterized by an alpha value of 0.1, and LS also utilizes an 339 alpha value of 0.1. LR, on the other hand, involves no additional parameters, as indicated 340 by the dash line in the "Values" column. 341

The steps for the single nutrient concentration prediction are described in Algorithm 342 1, based on the parameters setting for the machine learning algorithms described in Table 343 5. 344

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Algorithm 1: Single nutrient concentration prediction

Input: Nutrient concentration dataset

Process:

- 1. Apply the Min-Max normalization method (Eq. 1)
- Set training ratio= 80% 2.
- 3. For each feature set, fs in Table 4: $FS_1,...,FS_5$
 - Load FS_x to be the predictors a.

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- b. ModelEN_x=Develop Elastic Net regression using FS_x with parameters in Table 5
- ModelSW_x=Develop Polynomial regression using FS_x with parameters in Tac. ble 5
- d. ModelSW_x=Develop Stepwise regression using FS_x with parameters in Table 5
- ModelRR_x=Develop Ridge regression using FS_x with parameters in Table 5 e.
- f. ModelLS_x=Develop Lasso regression using FS_x with parameters in Table 5
- ModelLR_x=Develop Linear regression using FS_x with parameters in Table 5 g.

End For 4.

Output: ModelEN_{Ca}, ModelEN_{Mg}, ModelEN_K, ModelEN_P, ModelEN_N, ModelPN_{Ca}, ModelPN_{Mg}, ModelPN_K, ModelPN_P, ModelPN_N, ModelSW_{Ca}, ModelSW_{Mg}, ModelSW_K, ModelSW_P, ModelSW_N, ModelSW_{Ca}, ModelSW_{Mg}, ModelSW_K, ModelSW_P, ModelSW_N, ModelRR_{Ca}, ModelRR_{Mg}, ModelRR_K, ModelRR_P, ModelRR_N, ModelLS_{Ca}, ModelLS_{Mg}, ModelLS_K, ModelLS_P, ModelLS_N, ModelLR_{Ca}, ModelLR_{Mg}, ModelLR_K, ModelLR_P, ModelLR_N.

Regards to Algorithm 1, the process for single nutrient concentration prediction, out-347 lined in Algorithm 1, involves applying Min-Max normalization to the nutrient concen-348 tration dataset and setting an 80% training ratio. For each of the five feature sets (FS1 to 349 FS5) detailed in Table 3, the algorithm loads the respective features and employs six re-350 gression models (Elastic Net, Polynomial, Stepwise, Ridge, Lasso, Linear), each with its 351 parameters specified in Table 4. The result is a set of trained models for predicting nutrient concentrations (Ca, Mg, K, P, N) denoted by prefixes such as ModelENCa, ModelENMg, 353 and so on. The models are developed using various regression techniques tailored to each 354 feature set, creating a comprehensive framework for nutrient concentration prediction. 355

3.3.2. Nutrient composition concentration prediction

In the second approach, model is developed based on different feature sets of rice 357 dataset as shown in Table 6 based on solely the spatiotemporal factors. 358

Table 6. Nutrient composition concentration prediction setting

Spatiotemporal Factors						Nutrients			
Feature Set	Sea- son	Day	Plo t	Subplot	N (%)	P (%)	K (%)	Mg (%)	Ca (%)
FS ₆ (All)	Y	Y	Y	Y	N	Ν	Ν	Ν	Ν

Referring to Table 6, the nutrient composition concentration prediction setting has 361 been constructed by incorporating features from both spatiotemporal factors and nutrient 362 features. 363

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These settings will be utilized for nutrient composition concentration prediction using364six methods: EN, PN, SW, PR, LS, and LR. The parameter specifications for these models365in nutrient composition concentration prediction are consistent with those applied for sin-366gle nutrient concentration prediction (refer to Table 5).367

The steps outlined in Algorithm 2 illustrate the processes for nutrient composition con-368centration prediction, developed based on the similar parameter specifications listed in369Table 4 for single nutrient concentration prediction.370

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Algorithm 2: Nutrient composition concentration prediction

Input: Nutrient concentration dataset Process:

- 1. Apply the Min-Max normalization method (Eq. 1)
- 2. Set training ratio= 80%
- 3. Load FS₆ from Table 6
- ModelENx=Develop Elastic Net regression using FSx with parameters in Table
 5
- 5. ModelSW_x=Develop Polynomial regression using FS_x with parameters in Table 5
- 6. ModelSW_x=Develop Stepwise regression using FS_x with parameters in Table 5
- 7. ModelRRx=Develop Ridge regression using FSx with parameters in Table 5
- 8. ModelLS_x=Develop Lasso regression using FS_x with parameters in Table 5
- 9. ModelLRx=Develop Linear regression using FSx with parameters in Table 5

Output: ModelENAII, ModelPNAII, ModelSWAII, ModelRRAII, ModelLSAII, ModelLRAII

Algorithm 2, designed for nutrient composition concentration prediction, starts by nor-373 malizing the input nutrient concentration dataset using the Min-Max method and setting 374 an 80% training ratio. It then exclusively utilizes features from FS6 in Table 6 to develop 375 six regression models – Elastic Net, Polynomial, Stepwise, Ridge, Lasso, and Linear – each 376 configured with parameters specified in Table 5. The resulting output comprises compre-377 hensive models denoted as ModelENAll, ModelPNAll, ModelSWAll, ModelRRAll, Mod-378 elLSAll, and ModelLRAll. This algorithm provides an efficient means of predicting nutri-379 ent composition concentrations based on the designated features and regression tech-380 niques. 381

4. Experimental Setting

This section presents the experimental results for Elastic Net Regression, Polynomial 383 Regression, Stepwise Regression, Ridge Regression, Lasso Regression, and Linear Regression to predict rice nutrient levels using FS 1 until 6. Table 4 and Figure 4 display the 385 RMSE score of all six models where polynomial regression has the best performance in 386

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four models to predict Ca%, K%, P% and N% with an average of 0.1502 RMSE, except387Model 2 (prediction of Mg%), with very little standard deviation (0.1980).388

5.1 Performance of single nutrient concentration approach

We present Table 7 until Table 11 to explain the performance of the single nutrient 392 concentration approach by using R², MAE and RMSE. A larger R2 value is generally con-393 sidered better. An R2 value closer to 1 suggests that a larger proportion of the variation in 394 the dependent variable is accounted for by the independent variables in the model, indi-395 cating a better fit. However, it's important to note that a high R² does not necessarily imply 396 causation or the absence of model errors, and other factors should be considered in eval-397 uating the overall validity of the regression model. MAE represents the average absolute 398 difference between the predicted values and the actual values. The smaller the MAE, the 399 better the model performance. MAE is less sensitive to outliers compared to RMSE. Lower 400 values of MAE and RMSE indicate better model performance. 401

Table 7. Performance of Ca prediction using approach 1

Algorithm	R ² score	MAE	RMSE
ModelENca	0.0	0.0297	0.0362
ModelPN Ca	0.5017	0.0204	0.0255
ModelESW Ca	0.0257	0.0292	0.0357
ModelRR Ca	0.0869	0.0281	0.0345
ModelLS _{Ca}	0.0	0.0361	0.0297
ModelLR _{Ca}	0.0931	0.0279	0.0345
AVG	0.1179	0.0286	0.0327
STDEV	0.1942	0.0050	0.0042

Based on Table 7, the best model for Ca prediction is ModelPN_{Ca} with consistent performance in all three evaluation metrics, and that the PN algorithm has far better performance compared to the other algorithms which indicate the superior performance of this algorithm to capture the nutrients values variability. Two algorithms, EN and LS could not capture the variability in the dataset for predicting Ca, based on the zero R² value. 408

Table 8. Performance of Mg prediction using approach 1

Algorithm	R ² score	MAE	RMSE	
ModelEN _{Mg}	0.0	0.0154	0.0193	
ModelPN _{Mg}	-3.1900	0.0301	0.0395	
ModelESW _{Mg}	0.0879	0.0151	0.0184	
ModelRR _{Mg}	0.1734	0.0142	0.0176	
ModelLS _{Mg}	0.0	0.0154	0.01934	
ModelLR _{Mg}	0.1742	0.0141	0.0175	
AVG	-0.451	0.0174	0.0219	

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STDEV	1.3401	0.0063	0.0086

Contrary to its performance in Table 7, the PN algorithm shows a bad performance 411 for Magnesium. The best for Magnesium prediction is the LR algorithm. The negative R² 412 value of PN implies that the model is so inadequate that it is worse than a naive model 413 that merely predicts the mean of the dependent variable for all observations. This indicates that PN could have been overfit and too complex for the given data, and it fits noise 415 rather than the underlying patterns. 416

The performance of LR and RR are very similar which reflect their high similarity. 417 Both algorithms assume a linear relationship between the independent variables and the 418 dependent variable. The models are expressed as linear combinations of the input features. Both methods aim to minimize a certain objective function to find the optimal set of 420 coefficients that best fits the data. In LR, this is typically done by minimizing the sum of 421 squared differences between the predicted and actual values. In RR, the objective function 422 includes an additional regularization term. 423

The primary difference between RR and LR lies in how they handle multicollinearity 424 and overfitting. RR uses regularization term penalizes large coefficients, helping to mitigate the effects of multicollinearity and prevent overfitting. The regularization term is 426 controlled by a hyperparameter (usually denoted as "alpha" or "lambda"). LR does not 427 include a regularization term in the objective function. It is more prone to overfitting when 428 dealing with highly correlated features (multicollinearity) or when the number of features 429 is close to or exceeds the number of observations. 430

Algorithm	R ² score	MAE	RMSE	
ModelENĸ	0.1967	0.3101	0.3991	
ModelPNκ	0.8496	0.1275	0.1726	
ModelESWĸ	0.0926	0.3464	0.4241	
ModelRRκ	0.5873	0.2266	0.2860	
ModelLSκ	0.1391	0.3235	0.4131	
ModelLRĸ	0.5895	0.2261	0.2852	
AVG	0.4091	0.2600	0.3300	
STDEV	0.3087	0.0823	0.0993	

Table 9. Performance of K prediction using approach 1

PN maintains the best algorithm for K prediction, and again, the performance of RR 433 and LR are very similar for predicting K. As explained, RR is a modified version of LR 434 that adds a regularization term to address certain issues, particularly multicollinearity. If 435 the correlation between independent variables is high, RR can provide more stable and 436 reliable coefficient estimates compared to LR. Since the performance of RR is better in 437 predicting K, this indicates that the dataset for the training possesses multicollinearity. 438

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Algorithm	R ² score	MAE	RMSE
ModelEN _P	0.0	0.0529	0.0651
ModelPN _P	0.8308	0.0212	0.0267
ModelESW _P	0.4180	0.0377	0.0497
ModelRR _P	0.6193	0.0311	0.0402
ModelLS _P	0.0	0.0529	0.0651
ModelLR _P	0.6202	0.0312	0.040
AVG	0.4147	0.0378	0.0478
STDEV	0.3468	0.0128	0.0153

Table 10. Performance of P prediction using approach 1

Likewise, the best technique for P prediction is PN, and it is observed that the performance of PN in this nutrient prediction is the best compared to other nutrients. All the other algorithms also had better scores, which indicates that the values in the features used for training the P prediction are more homogeneous compared to the earlier models. 445

Table 11. Performance of N prediction using approach 1

Algorithm	R ² score	MAE	RMSE
ModelEN N	0.3006	0.4524	0.6326
ModelPN _N	0.5862	0.3808	0.4866
ModelESW _N	0.4240	0.4388	0.5741
ModelRR _N	0.5508	0.3657	0.5070
ModelLS _N	0.1994	0.4948	0.6768
ModelLR N	0.5532	0.3661	0.5056
AVG	0.4357	0.4164	0.5638
STDEV	0.1574	0.0535	0.0777

Similarly, PN achieved the best performance in comparison to the other models. All 448 models had lower performance in prediction N compared to predicting P. It is also observed that the performance of SW in predicting N are similar in predicting P, when compared against RR and LR. Although LR and RR show stability and generalizability across 451 different datasets, SW has better performance in this nutrient compared to Ca and Mg 452 because of its simplicity drawback and tendency of assumption that the relationship between variables is best represented by a combination of selected features. 454

Figures 4 to 8 depict the Streamlit outputs for the single-nutrient prediction of Ca, 455 Mg, K, P, and N, respectively, based on the best-performing model, PN. The predicted 456 values for each nutrient are computed utilizing the PN model, taking into account spatialtemporal parameters and other relevant nutrient inputs. The diagrams illustrate that the 458 predicted nutrient concentrations are used to recommend the amount of nutrient recovery, by comparing against the benchmark nutrient values. 460

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Rice Ca(%) Nutrient Prediction

Season Selection	
\bigcirc 1	
• 2	
Day Selection:	
30	~
Plot:	
4	~
Subplot (A=1,B=2,C=3):	
1A	~
Nutrient N:	
1.60	
0.93	4.59
Nutrient P:	
0.20	
0.15	0.46
Nutrient K:	
1.79	
1.61	3.89
Nutrient Mg:	
0.09	0.20
Predict	

On season 2, Day=30, Plot=4, Subplot=11, P=0.2, N=1.6, K=1.79, Mg=0.1 the values of Ca(%) nutrient is as follows:

Nutrient	Predicted	Best practice (Range)	Best practice (Average)	Intervention
Ca(%)	0.748022	[0.23, 0.33]	0.280000	-0.468022

Figure 4. Rice Ca Nutrient prediction based on other nutrients of N, P, K, and Mg.



Rice Mg(%) Nutrient Prediction

Season Selection	
O 1	
• 2	
Day Selection:	
30	~
Plot:	
4	*
Subplot (A=1,B=2,C=3):	
2B	~
Nutrient N: 1.00	
0.93	4.59
Nutrient P: 0.16	
0.15	0.46
Nutrient K: 1,82	
1.61	3.89
Nutrient Ca:	
0.16	0.38
Predict	

On season 2, Day=30, Plot=4, Subplot=22, P=0.16, N=1.0, K=1.82, Ca=0.21 the values of Mg(%) nutrient is as follows:

Nutrient	Predicted	Best practice (Range)	Best practice (Average)	Intervention
Mg(%)	0.170428	[0.11, 0.17]	0.150000	-0.020428

Figure 5. Rice Mg Nutrient prediction based on other nutrients of N, P, K, and Ca.

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Rice K(%) Nutrient Prediction

Season Selection	
• 1	
○ 2	
Day Selection:	
30	~
Plot:	
1	~
Subplot (A=1,B=2,C=3):	
1A	~
Nutrient N:	
1.08	
0.93	4.59
Nutrient P:	
0.17	
0.15	0.46
Nutrient Mg:	
0.10	
0.09	0.20
Nutrient Ca:	
0.18	
0.16	0.38
Predict	

On season 1, Day=30, Plot=1, Subplot=11, N=1.08,Mg=0.1,P=0.17, Ca=0.18 the values of K (%) nutrient is as follows:

Nutrient	Predicted	Best practice (Range)	Best practice (Average)	Intervention
K(%)	2.257959	[1.85, 2.52]	2.480000	0.222041

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Rice P(%) Nutrient Prediction

Season Selection	
• 1	
○ 2	
Day Selection:	
30	~
Plot:	
5	~
Subplot (A=1,B=2,C=3):	
2A	~
Nutrient N:	
1.07	
0.93	4.59
Nutrient K:	
1.61	
1.61	3.89
Nutrient Mg:	
0.10	
0.09	0.20
Nutrient Ca:	
0.17	
0.16	0.38
Predict	

On season 1, Day=30, Plot=5, Subplot=21, N=1.07, K=1.61, Mg=0.1, Ca=0.17 the values of P (%) nutrient is as follows:

Nutrient	Predicted	Best practice (Range)	Best practice (Average)	Intervention
P(%)	6.200514	[0.25, 0.3]	0.380000	-5.820514

Figure 7. Rice P Nutrient prediction based on other nutrients of N, K, Mg, and Ca.

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Rice N (%) Nutrient Prediction

Season Selec	tion				
01					
0 2					
Day Selection	n:				
30					~
Plot:					
1					~
Subplot (A=1	,B=2,C=3):				
1A					~
Nutrient P: 0.16					
0.15					0.46
Nutrient K: 1.73					
1.61					3.89
Nutrient Mg: 0.10					
0.09					0.20
Nutrient Ca: 0.17					
0.16					0.38
Predict					
On season 1, as follows:	Day=30, Plot=	1, Subplot=11, P=0.16, K=1	73, Mg=0.1, Ca=0.17 the val	ues of N (%) nutri	ent is
Nutrient	Predicted	Best practice (Range)	Best practice (Average)	Intervention	
N(%)	0.222117	[1.17, 2.47]	2.340000	2.117883	

N(%) 0.222117 [1.17, 2.47] 2.340000 2.117883

Figure 8. Rice N Nutrient prediction based on other nutrients of P, K, Mg, and Ca.

Referring to the aforementioned Streamlit interface for individual nutrients, including Ca, Mg, K, P, and N, the application provides essential values for "predicted," "Best practice (Range)," "Best practice (Average)," and "Intervention." The predicted values for each nutrient are computed utilizing the PN model, taking into account spatial-temporal parameters and other relevant nutrient inputs. 487

The "Best practice Range" and "Best practice Average" values specify the optimal 488 range and average of nutrient concentrations, offering valuable benchmarks for nutrient 489 levels. To further enhance precision in nutrient management, the intervention value is 490 calculated by estimating the difference between the best practice average and the pre-491 dicted value derived from the PN model. This intervention value serves as a critical metric 492

for nutrient recovery interventions, providing insights into the precise amount of nutrients required for optimal crop growth. 494

Therefore, in the context of precision agriculture and environmental sustainability 495 the crafted Streamlit tool for predicting individual nutrients, utilizing prior knowledge of 496 other nutrient concentrations, offers advantages to farmers and scientists seeking specific 497 insights into individual nutrient levels. This method proves especially advantageous 498 when a sensor dedicated to a specific nutrient experiences a malfunction. As a result, our 499 digital twin system promptly alerts users about sensor malfunctions and supplies predic-500 tive values while waiting for sensor replacement. This immediate functionality guarantees 501 continuous monitoring and safeguards data accuracy, essential for the effectiveness of 502 precision agriculture practices. 503

5.2 Performance of nutrient composition concentration approach

Algorithm R² score MAE RMSE **ModelEN**AII 0.0771 0.1814 0.2376 **ModelPN**AII 0.5237 0.1211 0.1502 **ModelESW**AII 0.0450 0.2054 0.2572 **ModelRR**AII 0.3066 0.1477 0.1949 ModelLSAII 0.0377 0.1918 0.2494 ModelLRAII 0.3066 0.1949 0.1477 AVG 0.2161 0.1659 0.2140 **STDEV** 0.1957 0.0321 0.0412

Table 12. Performance of approach 2 to predict all nutrients

ModelPNAll appears to be the best-performing model based on R², MAE, and RMSE. 508 It explains a significant proportion of variability and provides accurate predictions. Mod-509 elRRAll and ModelLRAll have the same R², MAE, and RMSE values, indicating similar 510 performance. They both exhibit a moderate level of explained variability and reasonable 511 predictive accuracy. ModelENAll, ModelESWAll, and ModelLSAll have lower R² values, 512 suggesting limited ability to explain variability. They also have higher MAE and RMSE 513 values, indicating higher prediction errors compared to the better-performing models. 514 The choice of features included in the models can significantly impact performance. Mod-515 els that incorporate irrelevant or highly correlated features may exhibit lower accuracy. 516 The results also indicate that the features incorporated have a complex relationship with 517 each other and the target variable. 518

The experiment results led us to the conclusion that regression models have good 519 performance in informing nutrient co-existence, concentration, and composition. This insight allows intervention to increase nutrient recovery to optimize the crop's yield. PN 521 generally outperformed the other tested algorithms in terms of producing higher R² values, and lower MAE and RMSE values for almost all models. This is due to the ability of 523 the polynomial function to capture nonlinear relationships among variables. However, it 524

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should be noted that for Mg, the polynomial regression algorithm produced a negative R² 525 value, indicating that it explained less variance in the dependent variable than a horizon-526 tal line. Therefore, the polynomial function was not well-suited for predicting nutrient 527 content in Mg. In contrast, LR produced better performance compared to the other meth-528 ods for Mg, signifying that this model was better approximated by a straight-line relation-529 ship. This finding highlights the significance of considering the specific nature of the data 530 and the relationships between variables when selecting the most appropriate regression 531 model for nutrient prediction. 532

Figure 9 illustrates the Streamlit outputs for the prediction of nutrient composition533concentrations, based on the best-performing model, PN.534



Rice Nutrients Prediction

Season Selection	
○ 1	
• 2	
Day Selection:	
60	~
Plot:	
5	~
Subplot (A=1,B=2,C=3):	
2B	~
Predict	
On season 2. Dav=60. Plot=5. Subplot=22. the values of nutrient are as follows:	

Nutrient	Predicted	Best practice (Range)	Best practice (Average)	Interventio
N (%)	2.322551	[1.17, 2.47]	2.340000	0.017449
P (%)	0.473045	[0.25, 0.3]	0.380000	-1.942551
K (%)	0.282621	[1.85, 2.52]	2.480000	0.157449
Mg (%)	0.147359	[0.11, 0.17]	0.150000	-2.172551

On season 2, Day=60, Plot=5, Subplot=22, the values of nutrient are as follows:

[0.23, 0.33]

Ca (%)

2.578441

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Figure 9. Rice nutrients composition concentrations prediction based on spatial-temporal parameters.

0.280000

-2.042551

Referring to the Figure 9 interface for nutrient composition concentrations, similar to the single-nutrient prediction (see Figures 4-8), the application furnishes crucial values for "predicted," "Best practice (Range)," "Best practice (Average)," and "Intervention." The predicted values for each nutrient are calculated employing the PN model, considering spatial-temporal parameters and other pertinent nutrient inputs. 537 The "Best practice Range" and "Best practice Average" values delineate the optimum 542 range and average of nutrient concentrations, providing valuable benchmarks for nutrient 543 levels. Furthermore, this information serves as a comprehensive intervention preparation 544 tool by informing farmers or scientists about the anticipated nutrient concentration. The 545 projected value, in turn, facilitates the digital twin system in suggesting the appropriate 546 amount of nutrient recovery, aligning with established best practices. 547

So, the provided streamlit for rice nutrients composition concentrations prediction 548 serves as a powerful intervention preparation tool. By informing farmers and scientists 549 about the anticipated nutrient concentrations, this approach enables the digital twin system to suggest the precise amount of nutrient recovery aligned with best practices. This 551 proactive and informed approach not only optimizes crop yields but also minimizes the 552 environmental footprint associated with excessive fertilizer application. 553

5.3 RMSE analysis and approach performance highlights

To identify the best model, we provide an analysis of RMSE across both approaches. 557

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Table 13. RMSE with Average and STDEV.
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Method	Ca	Mg	К	Р	Ν	All	AVG	STDEV
EN	0.03 62	0.0193	0.3991	0.0651	0.6326	0.2376	0.2305	0.2738
PN	0.02 55	0.0395	0.1726	0.0267	0.4866	0.1502	<u>0.1502</u>	0.1979
SW	0.03 57	0.0184	0.4241	0.0497	0.5741	0.2572	0.2204	0.2601
RR	0.03 45	0.0176	0.2860	0.0402	0.5070	0.1949	0.1771	0.2152
LS	0.02 97	0.0193	0.4131	0.0651	0.6768	0.2494	0.2408	0.2934
LR	0.03 45	0.0175	0.2852	0.0400	0.5056	0.1949	0.1766	0.2146
AVG	<u>0.03</u> <u>27</u>	0.0219	0.3300	0.0478	0.5638	0.2140		
STDEV	0.00 42	0.0086	0.0993	0.0153	0.0777	0.0412		

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Figure 10. RMSE performance for each nutrient prediction models



Figure 11. Stdev performance for each nutrient prediction models

The best performance of algorithm for FS 2 is Linear Regression. In terms of the performance to predict each nutrient, FS 2 is the easiest to be predicted, based on the average (AVG) of RMSE for this model, at 0.0219 (Figure 10). On the contrary, according to Figure 11, the percentage of N is the most difficult and inconsistent performance across the regression models, with an average of RMSE at 0.5638. 567

5.1. Statistical Analysis

For this investigation, this study chose to use parametric statistical analysis because 571 the assumptions of normality and equal variance are likely to be met given the data and 572 the fact that we are comparing means within each regression model. Additionally, parametric tests are generally more powerful than non-parametric tests, meaning they have a 574 greater ability to detect differences between groups when they exist. 575

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The normality assumption was evaluated through the Shapiro-Wilk test, which is a 576 commonly used test for normality. This test checks whether the data follows a normal 577 distribution. The equal variance assumption was examined using Levene's test. The 578 Shapiro-Wilk test for normality was applied to the residuals of the regression models, and 579 the results indicated that the residuals were normally distributed (p-value > 0.05). Addi-580 tionally, Levene's test was employed to assess the equality of variances among the groups, 581 and the results did not suggest any significant deviation from homogeneity of variances 582 (p-value > 0.05). 583

The application of these tests supports the validity of the ANOVA results presented 584 in Table 14. These tests, along with the reported F-statistic and p-value, confirm that the 585 assumptions necessary for ANOVA were satisfied. Therefore, we can observe differences 586 among the six designed regression models are statistically significant and not a result of 587 violations of normality or equal variance assumptions. Table 14 presents the ANOVA test 588 for six designed regression models using different regression methods of "Elastic Net Re-589 gression," "Polynomial regression," "Stepwise regression," "Ridge regression," "Lasso re-590 gression," and "Linear Regression." Table 14. ANOVA test for performance analysis. 591

Anova: Si	ingle Fac-				
to	or				
SUMN	MARY				
G	roups	Count	Sum	Average	Variance
	FS1	6	0.1961	0.0327	1.77137E-05
FS2		6	0.13164	0.0219	7.46328E-05
FS3		6	1.9801	0.3300	0.0098
FS4		6	0.2868	0.0478	0.0002
FS 5		6	3.3827	0.5638	0.0060
FS 6		6	1.2842	0.2140	0.0017
ANOV	A				
Source of					
Variation	SS	df	MS	F	P-value
Between					
Groups	1.394	5	0.2787	93.3932	2.3253E-17
Within					
Groups	0.0895	30	0.0030		
Total	1.4833	35			

Based on the ANOVA test with a p-value of 2.3253E-17 and an alpha level of 0.05, we can conclude that there is a statistically significant difference among the six designed regression models. Therefore, we reject the null hypothesis that there is no significant difference and accept the alternative hypothesis that at least one of the regression models has a different performance value than the others. 598

Post-hoc analysis was conducted using the Tukey Honestly Significant Difference 599 (Tukey HSD) test to determine specific pairwise differences between the regression 600

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models. This test accounts for multiple comparisons and provides valuable insights into601which models significantly differ in performance.602

Based on the results of the ANOVA test, Model 5 demonstrated better performance 603 compared to other designed feature set models (Refer to Table 4). As a result, to gain insight into the impact of each nutrient on N% nutrient concentration, we utilized SHAP 605 visualization. Figure 12 illustrates the effect of each nutrient on N% nutrient concentration. 607



Figure 12. Features importance for N% nutrient concentration prediction.

Referring to Figure 12, the attributes K (Potassium), Mg (Magnesium), Day, Season, 610 Ca (Calcium), Plot, SubPlot, and P (Phosphorus) appear to have varying levels of impact 611 on N% nutrient concentration. Potassium (K) has the highest impact, followed by Mag-612 nesium (Mg), indicating that their concentrations in the soil or nutrient supply signifi-613 cantly influence N%. The day and season when measurements are taken also play essen-614 tial roles, while attributes like Calcium (Ca), Plot, SubPlot, and Phosphorus (P) have var-615 ying degrees of influence, with P showing the lowest impact. Therefore, this visualiza-616 tion can be valuable for optimizing agricultural and environmental practices to manage 617 nutrient levels effectively, considering specific local conditions and domain knowledge. 618

6. Conclusion and Future Work

Crop digital twin offers a revolution to monitor and intervene crop health management. The physical twin surveil the condition of the crop and this information can be analysed by the digital twin to provide suggestions for countermeasures such as adding nutrient concentration level. 621

Predicting nutrient levels is crucial for optimizing fertilizer usage and ensuring a balanced nutrient supply, leading to higher-quality and increased yields, and reduced environmental impact. The importance of accurately anticipating essential nutrients, such as Nitrogen (N), Phosphorus (P), Potassium (K), Calcium (Ca), and Magnesium (Mg), in rice cannot be overstated, as it directly impacts crop yield, quality, and environmental sustainability. The challenges in this field stem from the complexities introduced by the for the state of the stem from the complexities introduced for the state of the stem from the complexities introduced by the for the state of the stem from the complexities introduced by the for the state of the state of the state of the stem from the complexities introduced by the for the state of the state of the state of the stem from the complexities introduced by the for the state of the s

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variability in nutrient content, the diversity of analytical approaches, data availability con-630 straints, genetic diversity, and the associated costs and time investments. 631

To address these challenges, this research has present two approaches of namely,(i) 632 single nutrient concentration prediction and (ii) nutrient composition concentration pre-633 diction to explored a range of regression algorithms, including Elastic Net Regression, 634 Polynomial Regression, Stepwise Regression, Ridge Regression, Lasso Regression, and 635 Linear Regression, to predict rice nutrient content. These algorithms have proven to be 636 invaluable tools for capturing both linear and nonlinear correlations among various nu-637 trients, offering a structured, data-driven approach to understanding and managing the 638 complexities of rice nutrition. 639

The findings reveal that the Polynomial Regression algorithm consistently outper-640 forms the other models for predicting several nutrients, particularly Calcium (Ca%), Po-641 tassium (K%), Phosphorus (P%), and Nitrogen (N%). This algorithm's ability to handle 642 both small and large datasets, along with its proficiency in capturing nonlinear relation-643 ships, makes it a favorable choice for optimizing nutrient management practices. It is im-644 portant to note, however, that Model 2, focused on predicting Magnesium (Mg%), demon-645 strated a unique characteristic, as Linear Regression outperformed Polynomial Regres-646 sion. 647

The dashboard in the digital twin visualizes the current nutrient content of the crop 648 as a surveillance mechanism while the predicted nutrient concentration is a valuable in-649 sight for precise fertilisation to be added as nutrient recovery. This may mitigate fertilisation overload and waste pollution. Albeit manual intervention is currently addressed in this research, the regression method's implementation supports low-resourced crop digi-652 tal twin so that fast computation could be performed. 653

In summary, these regression models provide essential insights into rice nutrient pre-654 diction, offering a pathway to optimize fertilizer use, ensure balanced nutrient supply, 655 enhance rice quality, and reduce environmental impact. They contribute to the develop-656 ment of standardized methodologies for nutrient prediction and promote more sustaina-657 ble and environmentally friendly rice cultivation practices. The choice of the most suitable 658 regression model depends on the specific characteristics of the dataset and the nature of 659 the nutrient interactions. Therefore, the selection of the appropriate algorithm is pivotal 660 to achieving the highest predictive accuracy for rice nutrient content. 661

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