

Learning From Differences

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Abstract

This study presents an innovative exploration into the realm of machine learning, specifically focusing on enhancing model interpretability and performance through a novel ensemble of model-based and instance-based learning approaches. Grounded in the principles of Case-Based Reasoning (CBR) and the Learning From Differences methodology, we introduce the **LingerRegressor** and **LingerClassifier** models with their novel variations. These models aim to leverage the strengths of both learning approaches to address the limitations present in traditional machine learning algorithms, particularly in the contexts of regression and classification tasks.

Our methodology involves a detailed examination of the algorithms' design and implementation, highlighting the integration of novel variations such as context inclusion, duplication based on distance, and the addition of a distance column to further refine the learning process. Through extensive experiments, we demonstrate the models' effectiveness in achieving higher interpretability and robust performance across diverse datasets. The **LingerRegressor** and **LingerClassifier** models not only offer new avenues for academic research but also practical implications for developing more transparent and efficient machine learning systems.

The findings underscore the potential of blending model-based and instance-based learning to enhance machine learning models' adaptability and interpretability. Future work will focus on addressing the identified limitations and exploring the applicability of our approach to broader machine learning challenges. This study contributes to the ongoing dialogue on improving machine learning methodologies and paves the way for further innovation in the field.

Declaration of Originality

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Acknowledgements

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Chapter 1

Introduction

Machine learning encompasses various paradigms and methodologies aimed at enabling computational systems to learn from data and make predictions or decisions without explicit programming. Among the diverse approaches in machine learning, model-based and instance-based learning represents two fundamental strategies, each with distinct advantages and disadvantages.

This dissertation presents a comprehensive study on blending model-based and instance-based learning methodologies to address their individual limitations and harness their strengths, leading to the development of robust, interpretable, and efficient machine learning models suitable for a wide range of applications. Below, we outline the roadmap of this dissertation and summarize our key contributions.

1.1 Report Roadmap

- **Chapter 1** introduces the motivations behind exploring blended learning methodologies, highlighting the potential benefits of integrating model-based and instance-based learning approaches.
- **Chapter 2** delves into the theoretical underpinnings of model-based and instance-based learning, providing a detailed review of the literature and identifying gaps that our research aims to fill.
- **Chapter 3** describes the methodology adopted in this study, including the development, and implementation of new models based on the new learning approaches designed to address specific challenges. That integrates seamlessly with scikit-learn's code base.

- **Chapter 4** presents the results of extensive experiments conducted to evaluate the performance of the proposed models against traditional approaches, showcasing their advantages or disadvantages in various scenarios.
- **Chapter 5, 5**, synthesizes the findings, discusses the implications of this research, and outlines potential directions for future exploration.

1.2 Contributions of This Dissertation

This dissertation makes the following significant contributions to the field of machine learning:

1. The development of a novel blended learning framework that combines the strengths of model-based and instance-based learning approaches to improve prediction accuracy and model interpretability, with an adaptable and extendable code base, based on scikit-learn.
2. An extensive evaluation of the proposed models using diverse datasets, demonstrating their performance in terms of accuracy, robustness, and efficiency compared to traditional learning models.
3. The introduction of an innovative approach to handling categorical features in blended learning models, enhancing their applicability to a wider range of problems.
4. A detailed exploration into the application of the learning from differences methodology into image based classification and regression problems.
5. The proposition of several areas for future work, including the exploration of other categorical similarity measures, variations in rule generation, scalability improvements, and a diversification of datasets to further enhance the effectiveness and efficiency of blended difference learning models.

1.3 Model-Based Learning

Model-based learning involves the construction of explicit representations of the underlying relationships within the data. These representations, often referred to as models, capture the patterns and structures present in the dataset and can be used to

make predictions or infer insights from new, unseen data points. For instance, artificial neural networks (ANNs) exemplify model-based learning by discerning trends and patterns in data.

1.3.1 Advantages of Model-Based Learning:

1. **Generalization:** Well-constructed models have the ability to generalize patterns learned from the training data to unseen instances, thereby making accurate predictions on new data samples. This generalization capability is essential for robust performance across diverse datasets and real-world scenarios. For example, a well-trained regression model can accurately predict housing prices in different regions based on historical data. The book [4] provides insights into the generalization capabilities of model based learning.
2. **Robustness:** Models learn underlying patterns and assumptions within data, making them more robust to noisy outliers. By capturing trends within data, models can effectively mitigate noise, enhancing their reliability in real-world scenarios. The book [4] also shows the ability of models to learn underlying patterns within data, making them far more robust to noisy data.
3. **Efficiency:** Once trained, model-based systems can quickly return predictions, making them suitable for real-time applications where rapid decision-making is essential. The journal paper [8] offers an insight how the efficiency of model based learning can be implemented to make predictions on big data.

1.3.2 Disadvantages of Model-Based Learning:

1. **Sensitivity to Assumptions:** Models often rely on simplifying assumptions about data distribution and relationships. Deviations from these assumptions can significantly degrade performance, leading to inaccurate predictions or biased outcomes. For example, a model trained to predict house prices based on 2017 data may inaccurately predict house prices for 2023. The book "Machine Learning: A Bayesian and Optimization Perspective" by Sergios Theodoridis and Konstantinos Koutroumbas emphasizes the crucial role assumptions play in model sensitivity, stressing their significance in machine learning applications [16].
2. **Interpretability and Decision-Making:** Certain models, like Neural Networks, lack transparency in decision-making despite their high accuracy. In critical domains like healthcare and finance, interpretability is crucial for validating

decisions and ensuring trust in the system. This disadvantage of lack of transparency is depicted well in the journal paper [14].

3. **Limited Flexibility:** Traditional model-based algorithms struggle to capture complex, non-linear relationships in high-dimensional or unstructured data, limiting their performance in tasks with intricate patterns or diverse feature interactions.

1.4 Instance-Based Learning

Instance-based learning, also known as memory-based learning or lazy learning, eschews explicit model construction in favor of storing and manipulating instances or examples from the training data. Instead of deriving general rules or representations, instance-based methods make predictions based on the similarity between new instances and those observed in the training set. A classic example of instance-based learning is the k-nearest neighbors (KNN) algorithm. The advantages and disadvantages discussed with instance based learning are seen in the paper [2].

1.4.1 Advantages of Instance-Based Learning:

1. **Flexibility:** Instance-based methods adapt to the characteristics of the training data, making them suitable for tasks with complex, non-parametric relationships or evolving patterns. They can capture nuances and irregularities without imposing rigid assumptions.
2. **Explainability:** The transparency of instance-based learning enables analysts to trace predictions back to specific examples in the training set, facilitating interpretability and trust in the system. The journal paper [14] highlights this issue for black box model based learning such as neural nets.

1.4.2 Disadvantages of Instance-Based Learning:

The disadvantages discussed with instance based learning are discussed in the paper [2].

1. **Computational Complexity:** Storing and manipulating the entire training dataset can lead to high memory consumption and computational overhead during prediction, limiting scalability.

2. **Susceptibility to Noise and Redundancy:** Instance-based approaches are sensitive to noisy or irrelevant features, which can lead to suboptimal performance or overfitting. Redundant instances or outliers may distort similarity measures, compromising model robustness.

1.4.3 Motivation for Blended Learning

The advantages and disadvantages of both model-based and instance-based learning highlight the need for a versatile and robust approach to machine learning. While model-based learning excels in generalization and robustness, it lacks transparency and struggles with complex relationships. Instance-based learning offers flexibility and explainability but suffers from computational complexity and susceptibility to noise.

The motivation for blended learning arises from leveraging the strengths of both approaches while mitigating their weaknesses. Blended methods aim to capitalize on the advantages of model-based and instance-based learning, circumventing their limitations. By integrating the interpretability of instance-based learning with the accuracy of model-based systems, blended learning offers a promising solution for critical domains like healthcare and finance. The journal paper [14] provides further evidence that it is important to move towards interpretability for our models going forward, so that critical systems are interpretable and repairable by humans going forward.

In recent years, neural networks have gained significant attention in regression tasks [12]. Inspired by case-based reasoning principles [13], researchers are exploring predictive ensemble models based on these principles [17]. Deviating from conventional approaches, some researchers advocate for training neural networks on differences between sets of features to enhance efficiency and transparency [7].

By adopting a blended approach, machine learning systems can achieve greater interpretability, reliability, and performance across diverse applications.

Chapter 2

Literature Review

2.1 Introduction

The exploration of alternative methodologies for enhancing the performance of neural networks in regression tasks has garnered significant attention within the machine learning community in recent years. One innovative approach, inspired by the principles of case-based reasoning (CBR), involves training neural networks to predict differences between problems based on disparities between features rather than the features themselves. This departure from the conventional supervised learning paradigm aims to leverage the inherent advantages of learning from differences, potentially leading to more interpretable and efficient models.

The study under consideration [7] conducts a systematic factorial study to investigate the efficacy of this approach across various datasets and experimental conditions. The findings suggest that neural networks trained on differences demonstrate comparable or even superior performance to those trained using traditional methods, while requiring significantly fewer epochs for convergence. In this section of the literature review, we consider the utilization of case-based reasoning for regression tasks, with a specific focus on the implementation of KNN (K-Nearest Neighbors) and ANN (Artificial Neural Network) frameworks.

As we delve further into the literature review, we extend our exploration of case-based reasoning beyond regression tasks to include classification challenges. The study by Ye et al. [18], titled "Learning Adaptations for Case-based Classification," offers insights into the application of CBR in classification tasks, shedding light on recent advancements and methodologies. Through a detailed analysis of various adaptation

strategies and neural network-based approaches, we aim to uncover the effectiveness of CBR in solving classification problems, thereby providing a comprehensive understanding of its utility across different machine learning tasks.

2.2 KNN + ANN for Regression, Utilizing CBR

The exploration of alternative methodologies for enhancing the performance of neural networks in regression tasks has attracted considerable attention within the machine learning community in recent years. One innovative approach, inspired by the principles of case-based reasoning (CBR), involves training neural networks to predict differences between problems based on disparities between features rather than the features themselves. This departure from the conventional supervised learning paradigm aims to leverage the inherent advantages of learning from differences, potentially leading to more interpretable and efficient models. The study under consideration [7] conducts a systematic factorial study to investigate the efficacy of this approach across various datasets and experimental conditions. The findings suggest that neural networks trained on differences demonstrate comparable or even superior performance to those trained using traditional methods, while requiring significantly fewer epochs for convergence. In this section of the literature review, we consider the utilization of case-based reasoning for regression tasks, with a specific focus on the implementation of KNN (K-Nearest Neighbors) and ANN (Artificial Neural Network) frameworks.

2.2.1 Learning From Differences Paper

The paper by [7] introduces the innovative concept of training a neural network based on the disparities between a case and its closest neighbors. Titled "A Factorial Study of Neural Network Learning from Differences for Regression" [7], the study aims to present novel learning methodologies geared towards enhancing performance and fostering a more interpretable learning process. The research evaluates three model variations: a benchmark neural network, a learning from differences model, and an augmented learning from differences model incorporating the original context. The context here refers to the base case, from which differences with its nearest neighbors are calculated.

According to [7], the study's key findings underscore a significant increase in accuracy when incorporating contextual information in the learning process. Moreover, comparable or superior results were achieved with fewer training epochs compared to the basic neural network. This enhancement is attributed partly to the broader training

dataset that encompasses neighbors greater than 1. Notably, the paper advocates for a hybrid approach, combining learning from differences with direct feature-based learning, which yields superior results.

The related work for the paper [7] has addressed the use of neural networks in the case-based reasoning process. These papers have focused on exploiting the base case using the idea that adaptation of information is acquirable from the differences between pairs [9]. Since then, various different approaches have been used. More recently, in relation to using neural networks to predict the differences between problems, a few interesting points have been investigated. These points show the ability of a neural network to correct the solution of the most similar retrieved case. These preliminary works expressed in the paper showed that the use of differences plus context yielded superior results [6]. The previous studies examined in the paper [7] did not examine the impact of different parameters during the CBR (case-based reasoning) experiments. The paper [7] made an effort to explore the effect of epochs in the learning process.

The graphical representations in the paper highlight a distinct trend: rapid attainment of high accuracy levels with the learning from differences method, particularly when contextual information is included. Conversely, the basic neural network requires a substantially larger number of epochs to achieve comparable accuracy, if not worse. This disparity suggests the efficiency of the learning from differences approach.

The paper suggests a marginal performance enhancement and a substantial reduction in training epochs through the adoption of the differences method and inclusion of contextual information. However, it is crucial to acknowledge the time overhead associated with retrieving and training using similar cases.

2.2.2 Algorithms

The training algorithm for learning from differences is described in Algorithm 1, while the prediction algorithm is outlined in Algorithm 2. Additionally, a variant of the training algorithm with context included is presented in Algorithm 3, and the corresponding prediction algorithm is provided in Algorithm 4.

Context is utilized in 3, and 4. Context is the case used to retrieve (Nearest Neighbors). In the training algorithm it is X_i^{train} , and in prediction it is X_j^{test} . These are included in the training, and prediction in Variant 2.

The paper [7] has an associated library where they have implemented a regression-based machine learning model.

Algorithm 1 Training Algorithm for Learning from Differences

Input: Dataset D , neural network model, number of neighbors n , number of epochs E

- 1: Split D into training set D^{train} and test set D^{test} with a 80%-20% split
 - 2: Initialize ΔD^{train} as an empty list
 - 3: **for** each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 - 4: Retrieve n similar cases from D^{train} using nearest neighbors
 - 5: **for** each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**
 - 6: Compute the differences: $\Delta D_{ij}^{train} = (X_i^{train} - X_j^{train}, y_i^{train} - y_j^{train})$
 - 7: **end for**
 - 8: **end for**
 - 9: Train the MLPRegressor neural network model using ΔD^{train} over a few epochs for later predictions.
 - 10: **Return** Trained neural network model $TrainedNN()$
-

Algorithm 2 Prediction Algorithm for Learning from Differences

Input: Test dataset D^{test} , trained neural network model $TrainedNN()$, number of neighbors n

- for** each case $X_j^{test} \in D^{test}$ **do**
 Retrieve n similar cases from D^{train} using nearest neighbors
 Initialize empty list $y^{pred}predictions$
 for each similar case $(X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 Compute the differences: $\Delta X_{ij}^{test} = X_j^{test} - X_i^{train}$
 Use the trained neural network to predict: $TrainedNN(\Delta X_{ij}^{test}) = \Delta X_{ij}^{pred}$
 Adapt neighbor's y values: $y^{pred}predictions_j = X_i^{train} + \Delta X_{ij}^{pred}$
 end for
 Average the adapted y values: $y_j^{pred} = \frac{1}{n} \sum_{i=1}^n y^{pred}predictions_j$
 return y^{pred}
end for
-

Algorithm 3 Training Algorithm for Learning from Differences with context

Input: Dataset D , neural network model, number of neighbors n , number of epochs E

- 1: Split the dataset D into training set D^{train} and test set D^{test} using an 80%-20% split.
 - 2: Initialize ΔD^{train} as an empty list.
 - 3: **for** each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 - 4: Retrieve n similar cases from D^{train} using nearest neighbors.
 - 5: **for** each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**
 - 6: Compute the differences, with concatenated context : $\Delta D_{ij}^{train} = ((X_i^{train} - X_j^{train}) : X_i^{train}, y_i^{train} - y_j^{train})$.
 - 7: Append ΔD_{ij}^{train} to ΔD^{train} .
 - 8: **end for**
 - 9: **end for**
 - 10: Train the MLPRegressor neural network model using ΔD^{train} .
 - 11: **return** Trained neural network model $TrainedNN()$.
-

Algorithm 4 Prediction Algorithm for Learning from Differences with context

Input: Test dataset D^{test} , trained neural network model $TrainedNN()$, number of neighbors n

```
for each case  $X_j^{test} \in D^{test}$  do
  Retrieve  $n$  similar cases from  $D^{train}$  using nearest neighbors
  Initialize  $y_j^{pred}$  as an empty list
  for each similar case  $(X_i^{train}, y_i^{train}) \in D^{train}$  do ‘
    Initialize empty list  $y^{pred}predictions$ 
    Compute the differences:  $\Delta X_{ij}^{test} = ((X_j^{test} - X_i^{train}) : X_j^{test})$ 
    Use the trained neural network to predict:  $\Delta X_{ij}^{pred} = TrainedNN(\Delta X_{ij}^{test})$ 
    Adapt neighbor’s  $y$  values:  $y^{pred}predictions_j = y_i^{train} + \Delta X_{ij}^{pred}$ 
  end for
  Average the adapted  $y$  values:  $y_j^{pred} = \frac{1}{n} \sum_{i=1}^n y_i^{pred}$ 
return  $y^{pred}$ 
end for
```

The new model proposed in the paper is tested using the factorial study methodology. This process involved finding a network structure through experimental means for each dataset. It entailed training and testing using a wide range of hyperparameters to determine the optimal settings.

Despite its contributions, the paper falls short by confining its attention to regression tasks and not considering classification tasks and exploring additional variations of the learning from differences model beyond context inclusion. Incorporating original values for the base case in the generated differences dataset provides a valuable framework for understanding the observed disparities. While the model with contextual inclusion demonstrates slight improvements over the base model in learning from differences, a more comprehensive examination of the methodology is warranted for further refinement and enhancement.

The paper tested three models using the same factorial methodology: a base version, which is a basic neural network trained empirically; a differences model employing the (CBH) Case-based Heuristic for predictions, (CBH) guides how cases are retrieved with respect to the base case, and how the retrieved cases are adapted to produce a prediction, (CBH) relies on the adaptation of previous problems to solve the current issue; and a differences model with its base context included. The results yielded interesting insights, with similar performance for all three models and a slight improvement in the differences + context model. Notably, the differences and differences + context models required significantly fewer epochs to achieve accuracy compared to the basic neural network, which achieved similar accuracy after more epochs.

Another consideration in the paper [7] was the number of neighbors for both training

and prediction. The study revealed no consistent trend across datasets, suggesting dataset-specific requirements.

In conclusion, the learning from differences method achieved equal or higher accuracy compared to the base neural network model.

2.3 Learning Adaptations for Case-based classification

The paper by Ye et al. [18], titled "Learning Adaptations for Case-based Classification," investigates the application of Case-Based Reasoning (CBR) for classification tasks. It explores recent advancements in CBR, particularly focusing on replacing traditional rule-based learning with Case-Based Heuristic (CBH) network models for adaptation. The study introduces a novel model comprising three variations: segmentation of adaptation knowledge based on the classes of the source cases, training a single neural network on differences between problem solutions, and adapting from an ensemble of source cases followed by a majority vote.

The "NN-CDH" method, as proposed in the paper, represents a novel classification approach that operates on pairs of cases, where one serves as the source and the other as the target. These pairs form the basis of adaptation knowledge, which is learned through neural network models. The methodology adopted in [18] emphasizes learning adaptation knowledge through case pairs processed by neural networks.

The study introduces the first neural network-based approach to classification, termed "C-NN-CDH" or Classification with Neural Network-based Case Difference Heuristic. This approach employs neural networks to learn adaptation knowledge from pairs of cases. According to the findings, this methodology outperforms traditional statistical models, as evidenced in [10].

The first approach involves segmenting the dataset by class and training a neural network for all segmented differences set between base cases. This method offers faster training but slightly less accuracy, according to the paper. The use of these CBR methods provides at least two benefits: inertia-free lazy learning and the ability to assess different cases when running classification. Inertial free means that the model does not make assumptions based on the underlying structure of the data, and lazy learning pertains to an algorithm that will postpone generalization until it is certain a new instance needs to be classified. It will only classify as the data comes; it won't try and learn or adapt preemptively.

In the process of CBR, adaptation is often treated as the most difficult. Various methods have been developed by academics over the years to tackle the issue. For example, Leake et al. [11] extrapolate a method in which the case base adaptation cases are populated from previously successful adaptations. Another approach is to have a stored base case and query and retrieve the case most similar to it, and adapting the retrieved case to the base case [6].

In the paper by Ye et al. [18], several design questions had to be addressed. How to approach case differences and how to select case pairs for training. The paper in question [18] considers standard pair selection methods as well as a new training per selection approach based on class-class classification.

The paper [18] also addresses the application of (CDH) method for classification. It discusses the value distance metric, which is a probabilistic method to measure the similarity that allows the comparison of nominal values in a single-dimensional space.

2.3.1 An NN-CDH Approach for Classification

To achieve adaptation, a neural network is integrated into the process. The source and target cases become case pairs, and their adaptation is learned through a neural network. The neural network computes the difference between the source and target pairs. This difference calculation is a crucial step in the Case-Based Reasoning (CBR) system. Once the difference is computed, the predicted result obtained from the neural network is applied to the source solution. This adapted solution is then considered as the final result of the classification process.

The NN-CDH approach represents a fusion of neural network techniques with the principles of Case-Based Reasoning, offering a dynamic and adaptable method for classification tasks. When calculating the difference between the problem and the solution values, it is important to implement a difference function. This presents an interesting challenge for nominal values. The method presented in [18] uses an implicit calculation through machine learning techniques. This replaces traditional (CDH) case difference heuristic approaches for difference calculations. Through the use of a neural network, the paper's method hopes to include the base context in the differences' calculation. This method for generating differences has been aptly named ("C-CDH") case difference heuristic approach for classification.

When calculating the difference, an issue arrives in that we cannot explicitly calculate differences through subtraction, which was the case in the previous paper by Learning

From Differences [7]. An implicit way of learning from differences is required. In the paper by Ye et al. [18], this is called "C-CDH".

This presents an interesting challenge for nominal values. The method presented in [18] uses an implicit calculation through machine learning techniques. When discussing the Neural Network-based Case-Driven Hierarchical (NN-CDH) Approach described in [18], notation will be homogenized with the algorithms from the previous paper.

For clarification when going through the algorithms, Retrieved cases = Source Cases, Target case = The case which we are attempting to predict its y label values.

Multiple variants were created with the paper by Ye et al. [18].

2.3.2 Variant 0: Non-Network C-CDH

Variant 0, as described in [18], serves as a foundational test bed implementation for classification tasks. The algorithms presented here outline the process of building and utilizing a dictionary of case pairs, where each pair consists of a source case and its corresponding target case. Through the use of nearest neighbors, the algorithm identifies similar cases within the training dataset and organizes them based on their source solution. Subsequently, during prediction, the algorithm retrieves the nearest similar cases for each test case and utilizes the corresponding target solutions for classification. This approach embodies the essence of case-based reasoning, where solutions are adapted from similar past cases to classify new instances effectively. The following algorithms detail the steps involved in training and predicting using Variant 0's case-based heuristic approach.

Algorithm 5 Variant 0, Classification Using Case Based Heuristic, Training

Input: Dataset D , number of neighbors 1

Split D into training set D^{train} and test set D^{test} with an 80%-20% split

Create *CasePairs()* dictionary.

for each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**

 Retrieve 1 similar cases from D^{train} using nearest neighbors.

for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**

CasePairs(y_j^{train}).append($[X_j^{train} : X_i^{train}, y_i^{train}]$)

end for

end for

Algorithm 6 Variant 0, Classification Using Case Based Heuristic, Prediction

Input: Test dataset D^{test} , $CasePairs()$

```
1: Initialize Empty list  $y^{pred}$ 
2: for each case  $X_j^{test} \in D^{test}$  do
3:   Retrieve 1 similar cases from  $D^{train}$  using nearest neighbors
4:   Initialize  $y_j^{pred}$  as an empty list
5:   for each similar case  $(X_i^{train}, y_i^{train}) \in D^{train}$  do
6:     Retrieve  $r = 1NearestNeighbor(CasePairs(y_i^{train}), [X_i^{train} : X_j^{test}])$ 
7:      $y_j^{pred} = r$ 
8:   end for
9: end for
10: return  $y^{pred}$ 
```

2.3.3 Variant 1 - 2: C-NN-CDH

Variants 1 - 2, known as C-NN-CDH, represent a pivotal advancement in classification methodologies by integrating neural networks into the case-based heuristic framework. In this variant, the traditional case-based heuristic approach is augmented with the power of neural networks to adapt solutions from similar cases. The algorithms presented here delineate the training and prediction processes, where a neural network model learns adaptation knowledge from case pairs derived from the training dataset. Unlike Variant 0, which relies solely on case similarity for adaptation, C-NN-CDH leverages the expressive capacity of neural networks to capture intricate relationships between cases and their solutions. Through the fusion of case-based reasoning principles with neural network techniques, Variant 1 - 2 offers a dynamic and adaptable approach to classification tasks, capable of learning complex adaptation patterns and improving classification accuracy.

2.3.4 Variant 3 - 5: C-NN-CDH

Variants 3 - 5 of the classification neural network using case-based heuristics introduce a novel approach to classification by grouping case pairs based on their source solutions. In this framework, multiple specialized adaptation neural networks are employed, each trained to adapt cases of specific solutions to all solutions. Inspired by the ensemble of adaptations for classification approach, these variants share a common grouping and training method but differ in their testing strategies. The algorithms delineate the training process, where case pairs are organized and used to train specialized neural networks for adaptation. The variation arises in the prediction algorithm, where different strategies are employed to classify test instances based on the learned adaptation knowledge. The following section details the nuances of each variant's

Algorithm 7 Variant 1 - 2: Classification Using Neural Net Case Based Heuristic (Training)

Input: Dataset D , neural network model, number of neighbors n , number of epochs E

Split D into training set D^{train} and test set D^{test} with an 80%-20% split

Create *CasePairs* list.

for each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**

Retrieve 1 similar case from D^{train} using nearest neighbors.

for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**

if Variant 1 **then**

$CasePairs.append([X_j^{train} : X_i^{train}, y_i^{train}])$

else

if Variant 2 **then**

$CasePairs.append([X_j^{train} : X_i^{train} : y_j^{train}, y_i^{train}])$

end if

end if

end for

end for

Train the neural network model using *CasePairs*()

return Trained neural network model *TrainedNN*()

Algorithm 8 Variant 1 - 2: Classification Using Neural Net Case Based Heuristic (Prediction)

Input: Test dataset D^{test} , trained neural network model *TrainedNN*()

1: Initialize Empty list y^{pred}

2: **for** each case $X_j^{test} \in D^{test}$ **do**

3: Retrieve 1 similar case from D^{train} using nearest neighbors

4: **for** each similar case $(X_i^{train}, y_i^{train}) \in D^{train}$ **do**

5: Use the trained neural network to predict: $y_j^{pred} = TrainedNN([X_i^{train} : X_j^{test} : y_i^{train}])$

6: **end for**

7: **end for**

8: **return** y^{pred}

prediction methodology and discusses their implications for classification accuracy and adaptability.

Algorithm 9 Variant 3 - 5: Classification Neural Network Using Case Based Heuristic (Training)

Input: Dataset D , number of neighbors 1

Split D into training set D^{train} and test set D^{test} with an 80%-20% split

Create $CasePairs()$ dictionary.

Create $AdaptNN()$, a dictionary to store each Neural Network

List of unique Classes $UniqueClasses$

for each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**

Retrieve 1 similar case from D^{train} using nearest neighbors.

for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**

$CasePairs(y_j^{train}).append([X_j^{train} : X_i^{train}, y_i^{train}])$

end for

end for

for each unique $Class$ in $UniqueClasses$ **do**

Create a new Classification Neural Network, trained on Case Pairs

$AdaptNN(Class).fit(CasePairs(Class))$

end for

return Trained neural network models dictionary $AdaptNN()$.

In variant (5), the prediction method utilizes a Class-to-Class approach for retrieving neighbors. This method retrieves the most similar neighbor for each class from the training dataset. Once the nearest neighbors for each class are identified, the corresponding specialized neural networks associated with those classes are employed to generate predictions for the target instance. Subsequently, a majority vote is conducted among the predictions from the specialized neural networks to determine the final classification outcome.

It is worth considering re-implementing the algorithm to allow for the retrieval of multiple neighbors (N) from each class during the prediction stage. This adjustment could potentially lead to improved results, particularly when dealing with large and varied datasets.

Variant (5) stands out as the most successful approach in the study, primarily due to its Class-to-Class retrieval of neighbors during the prediction stage. By employing multiple specialized neural networks, the variant reduces training time while effectively leveraging implicit learning through differences for classification tasks. This method demonstrates its efficacy in adapting to diverse datasets and achieving robust classification performance.

Algorithm 10 Variant 3 - 5: Classification Using Neural Net Case-Based Heuristic (Prediction)

Input: D^{test} , $AdaptNN()$, $UniqueClasses$, and $CasePairs()$

Initialize an empty list y^{pred}

for each case $X_j^{test} \in D^{test}$ **do**

if Variant 3 **then**

 Retrieve 1 similar case from D^{train} using nearest neighbors

for each similar case $(X_i^{train}, y_i^{train}) \in D^{train}$ **do**

$y_j^{pred} = AdaptNN(y_i^{train}).predict([X_i^{train} : X_j^{test}])$

end for

else

if Variant 4 **then**

 Create an empty list $predictions$

 Retrieve N similar cases from D^{train} using nearest neighbors

for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**

$predictions.append(Adapt(y_i^{train}).predict([X_i^{train} : X_j^{test}]))$

end for

 Perform a majority vote on $predictions$

$y_j^{pred} = MajorityVote(predictions)$

else

if Variant 5 **then**

 Create an empty list $preds$

for each unique class $Class$ in $UniqueClasses$ **do**

 Retrieve 1 similar case from $CasePairs(Class)$

for each similar case $(X_i^{CasePair}, y_i^{CasePair}) \in CasePairs(Class)$ **do**

for each $(X_z^{train}, X_q^{train}) \in X_i^{CasePair}$ **do**

$preds.append(AdaptNN(Class).predict(X_z^{train} : X_j^{test}))$

end for

end for

 Perform a majority vote on $preds$

$y_j^{pred} = MajorityVote(preds)$

end for

end if

end if

end if

end for

return y^{pred}

2.3.5 Evaluation Summary

Within the paper [18] the experiments conducted on two different datasets provided answers to several key questions.

The effectiveness of neural network learning adaptation knowledge was demonstrated through consistent performance of one or more C-NN-CDHs, which often outperformed the neural network classifier. Despite the possibility of neural networks learning to discard the source problem and rely solely on the target problem, the non-zero weights associated with the source problem and significant performance differences between C-NN-CDHs and the neural network indicate that C-NN-CDHs effectively learn adaptation knowledge. This suggests that if a neural network can handle the classification task directly, it can also learn adaptation knowledge or the relation between pairs of cases in the task domain.

Surprisingly, variant (1) performed almost identically to variant (2), which also considers the source solution in adaptation. This suggests that the source solution, heavily coupled with the source problem, may not provide additional useful information for adaptation.

In terms of segmenting case pairs by source case solution, variants (1, 2) generally outperformed variants (3, 4) in accuracy on most datasets. This could be because a single adaptation neural network in (1, 2) is well-trained with all pairs of cases, while a specialized adaptation neural network in (3, 4) is trained with segmented examples. However, variants (3, 4) showed faster convergence during training due to training on segmented examples.

The impact of the ensemble of adaptations was investigated, with EAC-NN-CDH (variant (4)) performing similarly to its counterpart variant (3) without ensemble. This suggests that the generalization power of C-NN-CDH produces stable predictions that an ensemble version does not significantly alter.

The usefulness of the class-to-class approach for adaptation was demonstrated by C2C-NN-CDH (variant (5)), which performed differently from and often better than the other C-NN-CDHs. C2C-NN-CDH reaches its prediction by collecting evidence from diverse source cases from all classes, which can provide more global support, especially when there are multiple classes. Moreover, the C2C approach offers the possibility of explanation with contrastive evidence.

2.3.6 Conclusions on Learning adaptations for case based classification: A Neural Network approach

The conclusion of the paper [18] is that (variant (5)) is worth investigating using its C2C-NN-CDH approach to classification. As it yielded the most promising results when compared to all the other variants tested in the paper. The grouping of the dataset based on class outcome and training a specialized neural network, has its benefits in terms of speedup. However, the retrieval of N cases across all classes during testing, has shown to yield better results, in a way that allows an interpretation of the results.

Chapter 3

Design and Implementation

Objectives and Contributions

The design chapter of this paper serves as a comprehensive guide, presenting various algorithms and their implementations in the context of our novel framework. Our primary objectives are to elucidate the intricacies of these algorithms, introduce innovative variations, and showcase the integration of our framework within the broader machine learning landscape.

The tables provided below offer a structured roadmap through our design chapter, delineating where and how different algorithms are described. Notably, the algorithms discussed originate from seminal works such as [7] and [18]. However, in addition to presenting established methodologies, we introduce novel variations that represent fresh approaches to classification and regression tasks.

Key innovations include Variant 2, which introduces duplication based on distances, and Variant 3, incorporating a column indicating each neighbor's distance to the base case. Notably, we employ a new type of classifier, distinct from those mentioned in [7]. Variants 1, 2, and 3 exemplify our endeavor to explore uncharted territory within the learning from differences methodology, offering novel perspectives on classification.

Furthermore, our introduction of the `LingerImplicitRegressor` presents a groundbreaking approach to regression tasks. Leveraging methodologies outlined in [18], this novel regressor extends the learning from differences framework, offering a fresh perspective on regression problems.

By presenting a cohesive framework that integrates established algorithms, innovative

variations, and our novel contributions, these tables provide a clear overview of the design landscape. Our aim is to facilitate understanding and navigation through the complexities of machine learning methodologies, empowering researchers and practitioners to leverage our framework effectively in their endeavors.

3.1 Design and Implementation Guide

3.1.1 LingerRegressor and LingerClassifier Models

The following models are based on the algorithms outlined in [7]. Variations such as Variant 2 and Variant 3 introduce innovative adaptations to the base models, providing enhanced functionality and performance.

LingerRegressor		
Full Name	Acronym	Description
LingerRegressor Section: 3.3	LR	Base Linger regressor, no variations
LingerRegressor Variant 1 Section: 3.5.1	LRV1	Linger regressor with the addition of context, where context is the original base case.
LingerRegressor Variant 2 Section: 3.5.2	LRV2	Linger regressor integrated with duplication based on distance, where differences are duplicated based on the neighbors' proximity to the base case.
LingerRegressor Variant 3 Section: 3.5.3	LRV3	Linger regressor integrated with a distance column indicating the distance of each neighbor from the base case.

Table 3.1: Regression Models based on the algorithms in [7]

LingerClassifier		
Full Name	Acronym	Description
LingerClassifier Section: 3.4	LC	Base Linger classifier, no variations
LingerClassifier Variant 1 Section: 3.5.1	LCV1	Linger classifier with the addition of context, where context is the original base case.
LingerClassifier Variant 2 Section: 3.5.2	LCV2	Linger classifier integrated with duplication based on distance, where differences are duplicated based on the neighbors' proximity to the base case.
LingerClassifier Variant 3 Section: 3.5.3	LCV3	Linger classifier integrated with a distance column indicating the distance of each neighbor from the base case.

Table 3.2: Classifier models based on the algorithms in [7]

3.1.2 LingerImplicitClassifier and LingerImplicitRegressor Models

The classifier models presented here are based on algorithms from [18], with Variations 1 and 2 also originating from the same source.

The regressor models in this section extend the learning from differences framework outlined in [18] to regression tasks. This innovative approach maintains the core principles of the methodology while introducing adaptations tailored to address the unique challenges of regression problems.

LingerImplicitClassifier		
Full Name	Acronym	Description
LingerImplicitClassifier Section: 3.7	LIC	Base Linger Implicit Classifier selects the N nearest neighbors from each class
LingerImplicitClassifier Variant 1 Section: 3.7.5	LICV1	Selects N random items from each class
LingerImplicitClassifier Variant 2 Section: 3.7.5	LICV2	Selects the single nearest neighbor from each class

Table 3.3: LingerImplicitClassifier models based on the algorithms in [18]

LingerImplicitRegressor		
Full Name	Acronym	Description
LingerImplicitRegressor Section: 3.3	LIR	Base Linger Implicit Regressor selects the N nearest neighbors from each class during prediction
LingerImplicitRegressor Variant 1 Section: 3.8.5	LIRV1	Selects N random items from each class
LingerImplicitRegressor Variant 2 Section: 3.8.5	LIRV2	Selects the single nearest neighbor from each class
Regression to Classification Converter		
Full Name	Acronym	Description
Regression to Classification Converter Section: 3.8.7	RTCC	Converts regression problems into classification problems

Table 3.4: LingerImplicitRegressor models based on the algorithms in [18]

3.1.3 LingerImageClassifier and LingerImageRegressor

LingerImageClassifier		
Full Name	Acronym	Description
LingerImageClassifier Section: 3.10	(LIMC)	Image Classifier that learns from differences

Table 3.5: Image Classifier Using learning from difference methodologies seen in [7]. Innovative in its application to classification image data sets.

LingerImageRegressor		
Full Name	Acronym	Description
LingerImageRegressor Section: 3.11	(LIMR)	Image Regressor that learns from differences

Table 3.6: Image Regressor Using learning from difference methodologies seen in [7]. Innovative in its application to classification image data sets.

3.2 Learning From Differences

3.2.1 Introduction and Motivation

Our design process is anchored in the seminal work of [7], which laid the groundwork for the learning from differences method. Building upon this foundation, our primary objective is to enhance and extend these algorithms, creating a more dynamic and adaptable framework.

Inspired by Scikit-learn’s modular design philosophy, we aim to develop models that not only match but surpass its rigorous hyperparameter testing capabilities. Our goal is to introduce multiple new hyperparameters accessible through familiar pipelines, thereby enhancing flexibility and functionality across diverse machine learning tasks and datasets.

As we progress from implementing base models to introducing variations, our focus shifts to optimizing the performance of both the `LingerRegressor` and `LingerClassifier`. These variations are meticulously crafted to improve training efficiency and predictive accuracy, ultimately enhancing overall model performance.

Named `LingerRegressor` and `LingerClassifier`, our custom models are designed to leverage the training from differences approach, distinguishing them within the machine learning landscape.

Central to our efforts is ensuring the seamless testability of introduced variations, akin to Scikit-learn’s comprehensive hyperparameter testing framework. This rigorous analysis empowers stakeholders to make informed decisions about the efficacy and applicability of our models to specific tasks.

However, we encountered the critical challenge of compatibility during model development. Our codebase is structured as a downloadable package, seamlessly integrating with Scikit-learn’s ecosystem to ensure interoperability across various datasets and environments. This compatibility enhances accessibility and usability for practitioners in diverse machine learning endeavors.

3.3 Linger Regression Base Model: LingerRegressor (LR)

3.3.1 Objective

The Linger Regression Base Model integrates principles of Case-Based Reasoning (CBR) into regression analysis to enhance interpretability and create a performative model. The objectives include:

- Developing a dynamic regression model using differences between a case and its nearest neighbors, adapting retrieved solutions to the target solution.
- Ensuring compatibility with Scikit-learn for accessibility, ease of implementation, and extensive hyperparameter testing.
- Enabling effective interpretation of predictions for users.
- Implementing the learning from differences algorithm with enhancements for improved performance.
- Creating an easily extendable code base to allow for variations.

Its intended use case is for regression problems that could benefit from a more transparent learning process.

3.3.2 Architecture

The `LingerRegressor` combines K-nearest neighbors (KNN) and `MLPRegressor` from Scikit-learn for flexibility and accuracy. A K-nearest neighbors (KNN) is used in conjunction with a `MLPRegressor` in the fit phase of the machine. During the predict phase, a K-nearest neighbors (KNN) is used once again with the now trained `MLPRegressor` to get the differences' prediction or K neighbors.

3.3.3 Components

- KNN module: Computes nearest neighbors and extracts differences during both training and prediction phases.

- **MLPRegressor**: Offers flexibility in activation functions, solvers, and hyperparameters for regression.
- **Supplementary Features**: Weighted KNN, Context Addition, Additional Distance Column, and Duplication Based on Distance. Each supplementary feature can be activated through a hyperparameter.

3.3.4 Parameterization

Extensive parameterization allows fine-tuning of KNN, **MLPRegressor**, and additional features. All the same hyperparameters are available as in a conventional (KNN) and neural network, with additional hyperparameters for **LingerRegressor**.

3.3.5 Training and Prediction Design

Training Design

Given dataset D , split it into D^{train} and D^{test} with an 80%-20% split through random selection. For each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$, retrieve n similar cases from D^{train} , $(X_j^{train}, y_j^{train}) \in D^{train}$ using Scikit-learn's neighbors module. Compute the differences between C_i and its neighbors and append them to ΔD^{train} as $(X_i^{train} - X_j^{train}, y_i^{train} - y_j^{train})$.

Algorithm 11 Training Algorithm for Learning from Differences

- 1: Split D into training set D^{train} and test set D^{test} with an 80%-20% split
 - 2: Initialize ΔD^{train} as an empty list
 - 3: **for** each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 - 4: Retrieve n similar cases from D^{train} using nearest neighbors
 - 5: **for** each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**
 - 6: Compute the differences: $\Delta D_{ij}^{train} = (X_i^{train} - X_j^{train}, y_i^{train} - y_j^{train})$
 - 7: Append ΔD_{ij}^{train} to ΔD^{train}
 - 8: **end for**
 - 9: **end for**
 - 10: Train the MLPRegressor neural network model using ΔD^{train} over a few epochs for later predictions.
 - 11: **Return** Trained neural network model $TrainedNN()$
-

Prediction Design

During prediction, for each case X_j^{test} , retrieve n similar cases from D^{train} . Compute the differences $\Delta X_{ij}^{test} = X_j^{test} - X_i^{train}$ for each neighbor. Use the trained neural network to predict ΔX_{ij}^{test} , resulting in ΔX_{ij}^{pred} . For each set of retrieved neighbors from D^{train} , calculate $y_i^{train} + \Delta X_{ij}^{pred}$ and add each n neighbors adapted y values. Once summed, divide by n to obtain y_j^{pred} . Compare y_j^{pred} to y_j^{test} to evaluate accuracy.

Algorithm 12 Prediction Algorithm for Learning from Differences

Input: Test dataset D^{test} , trained neural network model $TrainedNN()$, number of neighbors n

```
Initialize empty list  $y_j^{pred}$ 
for each case  $X_j^{test} \in D^{test}$  do
    Initialize empty list  $y_j^{pred}$ 
    Retrieve  $n$  similar cases from  $D^{train}$  using nearest neighbors
    Initialize empty list  $y_j^{pred\_predictions}$ 
    for each similar case  $(X_i^{train}, y_i^{train}) \in D^{train}$  do
        Compute the differences:  $\Delta X_{ji}^{test} = X_j^{test} - X_i^{train}$ 
        Use the trained neural network to predict:  $TrainedNN(\Delta X_{ji}^{test}) = \Delta X_{ji}^{pred}$ 
        Adapt neighbor's  $y$  values:  $y_{ji}^{pred} = y_i^{train} + \Delta X_{ji}^{pred}$ 
    end for
    Average the adapted  $y$  values:  $y_j^{pred} = \frac{1}{n} \sum_{i=1}^n y_{ji}^{pred}$ 
end for
return  $y_j^{pred}$ 
```

3.3.6 Training and Prediction Implementation

Training Implementation

During training, the Linger Regressor is fitted to the dataset X with labels y . It computes nearest neighbors for each X to construct features and targets by calculating differences between each sample and its neighbors, using scikit-learn's `NearestNeighbors`, `numpy`, and `MLPRegressor`. These differences train the regressor, and both the model and data are saved for future use.

Algorithm 13 Training Implementation Algorithm for Base Linger Regressor

Input: Dataset, X , label vector y , number of neighbors $n_neighbors_1$

```
1: function FIT( $X, y$ )
2:   Increment  $n\_neighbours\_1$  ▷ Fit scikit-learns NearestNeighbors function to  $X$ 
3:    $neighbors = NearestNeighbors(n\_neighbors = n\_neighbors\_1).fit(X)$ 
4:   Compute nearest neighbors for each sample in  $X$ 
5:   Compute the indexes and distances, of nearest neighbors in  $X$ 
6:    $distances, indices = neighbors.kneighbors(X)$ 
7:    $differences\_X = []$ 
8:    $differences\_y = []$ 
9:   if  $variant == 1$  (LRV1) then
10:    Call  $addition\_of\_context\_func\_fit()$ 
11:     $differences\_X, differences\_y =$ 
       $addition\_of\_context\_func\_fit(X, y, indices, differences\_X, differences\_y)$ 
12:   else if  $variant == 2$  (LRV2) then
13:    Call  $duplicate\_difference\_based\_on\_distance()$ 
14:     $differences\_X, differences\_y =$ 
       $duplicate\_difference\_based\_on\_distance(differences\_X, differences\_y, distances)$ 
15:   else if  $variant == 3$  (LRV3) then
16:    Call  $add\_additional\_distance\_column(differences\_X, distances)$ 
17:     $differences\_X = add\_additional\_distance\_column(differences\_X, distances)$ 
18:   else
19:     for each sample  $i$  in nearest neighbors indices do
20:        $base \leftarrow$  Index of base case
21:       for each neighbor  $n$  of  $i$  do
22:         Append  $(X[base] - X[n])$  to  $differences\_X$ 
23:         Append  $(y[base] - y[n])$  to  $differences\_y$ 
24:       end for
25:     end for
26:   end if
27:   Fit MLPRegressor with  $differences\_X$  and  $differences\_y$ 
28:   Set  $train\_X = X$ 
29:   Set  $train\_y = y$ 
30:   return Fitted LingerRegressor
31: end function
```

Prediction Implementation

During prediction, the algorithm utilizes the fitted Nearest Neighbors model to identify the nearest neighbors for each sample in the input data X . It then computes the differences between each sample in X and its corresponding neighbors, preparing the input features for the regressor. After predicting the differences using the trained regressor, the algorithm aggregates the predictions to obtain the final predicted target values for each sample in X . If weighted KNN is enabled, the predictions are weighted based on the distances to the neighbors, providing more influence to closer neighbors in the prediction process. This offers a different form of nearest neighbors. Where values are weighted based on their proximity to the base case. It makes use of scikit-learns [15] NearestNeighbors, and numpy.

Algorithm 14 Prediction Implementation Algorithm for Base Linger Regressor

Input: Input data, X , number of neighbors $n_neighbors_2$

```
1: function PREDICT( $X$ )
2:   Fit Nearest Neighbors model with  $n\_neighbours\_2$  using  $train\_X$ 
3:   Compute nearest neighbors for each sample in  $X$ ,
4:   Compute the indexes and distances, of nearest neighbors in  $X$ 
5:   Initialize empty list  $differences\_test\_X$ 
6:    $is\_sparse\_X \leftarrow$  Check if  $X$  is sparse
7:   if  $is\_sparse\_X$  then
8:     Convert  $X$  to dense array
9:   end if
10:  for each sample  $test$  in  $X$  do
11:    for each nearest neighbor  $nn\_in\_training$  in  $indices[test]$  do
12:      Append ( $X[test, :] - train\_X[indices[test][nn\_in\_training], :]$ )
13:        to  $differences\_test\_X$ 
14:    end for
15:  end for
16:  if  $variant == 1$  (LRV1) then
17:    Call  $addition\_of\_context\_func\_pred(differences\_test\_X, X)$ 
18:     $differences\_test\_X = addition\_of\_context\_func\_pred(differences\_test\_X, X)$ 
19:  else if  $variant == 3$  (LRV3) then
20:    Call  $add\_additional\_distance\_column(differences\_test\_X, distances)$ 
21:     $differences\_X = add\_additional\_distance\_column(differences\_test\_X, distances)$ 
22:  end if
23:  Continue predict with next function.
24: end function
```

Algorithm 15 Prediction Implementation Algorithm for Base Linger Regressor (Part 2)

Input: Continuation from Part 1

```
1: function PREDICT_CONTINUE
2:    $predictions \leftarrow$  Predict using LingerRegressor with  $differences\_test\_X$ 
3:   Split  $predictions$  into batches of size  $n\_neighbours\_2$ 
4:   Initialise  $y\_pred$  as an empty list
5:    $y\_pred = []$ 
6:   if Weighted KNN is enabled then
7:     for each  $indexes, differences, dist$  in  $zip(indexes, predictions, distances)$  do
8:       Compute weights for each neighbor
9:        $weights = 1 \div (dist + 1e - 8)$ 
10:      Compute weighted sum of target values
11:       $weighted\_sum = sum(w \times (train\_y[i] + d)$ 
12:        for  $i, d, w$  in  $zip(indexes, differences, weights)$ 
13:      Compute weighted average
14:       $weighted\_avg = weighted\_sum \div sum(weights)$ 
15:      Append  $weighted\_avg$  to  $y\_pred$ 
16:    end for
17:   else
18:     for each  $indexes, differences$  in  $zip(indexes, predictions)$  do
19:       Compute sum of target values
20:        $sum\_result = sum((train\_y[i] + d) \text{ for } i, d \text{ in } zip(indexes, differences))$ 
21:       Compute average
22:        $avg\_res = sum\_result \div n\_neighbours\_2$ 
23:       Append  $avg\_res$  to  $y\_pred$ 
24:     end for
25:   end if
26:   return  $y\_pred$ 
27: end function
```

3.3.7 Evaluation

Evaluation metrics include Mean Squared Error (MSE), Root Mean Squared Error (RMSE), and R-squared (R^2) to assess model accuracy and performance.

3.3.8 Conclusion

In conclusion, the **LingerRegressor** model presents a novel approach to regression analysis by integrating principles of Case-Based Reasoning (CBR) and the learning from differences methodology. Through the combination of K-nearest neighbors (KNN) and **MLPRegressor**, the model offers flexibility, accuracy, and interpretability in regression tasks. The implementation of supplementary features further enhances its adaptability and performance. Despite its limitations, the model shows promise in addressing the challenges of transparent and dynamic regression modeling. Future research can focus on refining the model's robustness to noise and improving its interpretability for complex datasets. Overall, the **LingerRegressor** model contributes to the advancement of machine learning methodologies, providing a valuable tool for regression analysis in diverse domains where a more interpretable model is needed.

3.4 Linger Classification Base Model:

LingerClassifier (LC)

3.4.1 Objective

The **LingerClassifier** extends the learning from differences methodology to classification tasks, aiming for:

- Development of a flexible model that leverages differences between cases for predictions.
- Full compatibility with Scikit-learn, ensuring seamless integration and robust hyperparameter testing.
- Enhanced interpretability of predictions to improve user understanding.
- Continuous improvements in the classification performance through algorithmic enhancements.

- A modular and expandable design for accommodating future model variations.

This model is designed for classification scenarios demanding high interpretability and adaptability.

3.4.2 Architecture

The architecture of `LingerClassifier` parallels that of its regression counterpart, incorporating both K-nearest neighbors (KNN) and Scikit-learn’s `MLPClassifier`. It utilizes KNN to determine nearest neighbors during both the training and prediction phases, adapting class labels based on learned differences.

3.4.3 Components

`LingerClassifier` comprises:

- **KNN Module:** Determines nearest neighbors and captures case differences throughout the training and prediction processes.
- **MLPClassifier:** Allows for configurable activation functions and solvers, enhancing classification flexibility.
- **Supplementary Features:** Includes Weighted KNN, Context Addition, and Distance-Based Duplication among others, all configurable via hyperparameters.

3.4.4 Parameterization

Extensive parameterization options enable detailed customization of KNN and `MLPClassifier` settings, alongside specialized `LingerClassifier` parameters, facilitating precise model tuning for varied classification needs.

3.4.5 Training and Prediction Design

The training and prediction algorithms for the `LingerClassifier` are similar to those of the `LingerRegressor`, with adjustments made to accommodate classification tasks.

Training Algorithm Design

During training, the algorithm computes differences between cases and their nearest neighbors, similar to the regression model. However, the adaptation of class labels is based on the majority class among the nearest neighbors.

Algorithm 16 Training Algorithm for LingerClassifier

Split the dataset D into training set D^{train} and test set D^{test} using an 80%-20% split.
Initialize ΔD^{train} as an empty list.
for each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 Retrieve n similar cases from D^{train} using nearest neighbors.
 for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**
 Compute the differences: $\Delta D_{ij}^{train} = (X_i^{train} - X_j^{train}, y_i^{train} - y_j^{train})$.
 Append ΔD_{ij}^{train} to ΔD^{train} .
 end for
end for
Train the MLPClassifier neural network model using ΔD^{train} .
return Trained neural network model $TrainedNN()$.

Prediction Algorithm Design

During prediction, the algorithm adapts class labels based on the majority class among the nearest neighbors' labels.

Algorithm 17 Prediction Algorithm for Learning from Differences for Classification

Input: Test dataset D^{test} , trained neural network model $TrainedNN()$, number of neighbors n

- 1: Initialize empty list y^{pred}
- 2: **for** each case $X_j^{test} \in D^{test}$ **do**
- 3: Retrieve n similar cases from D^{train} using nearest neighbors
- 4: Initialize empty list y_j^{pred}
- 5: **for** each similar case $(X_i^{train}, y_i^{train}) \in D^{train}$ **do**
- 6: Compute the differences: $\Delta X_{ji}^{test} = X_j^{test} - X_i^{train}$
- 7: Use the trained neural network to predict: $TrainedNN(\Delta X_{ji}^{test}) = \Delta X_{ji}^{pred}$
- 8: Adapt neighbor's y value: $y_{ji}^{pred} = y_i^{train} + \Delta X_{ji}^{pred}$
- 9: **end for**
- 10: Perform majority voting on y_j^{pred} predictions to obtain y_j^{pred}
- 11: **end for**
- 12: **return** y^{pred}

3.4.6 Training and Prediction Implementation

Training Algorithm Implementation

This algorithm implements the `textttLingerClassifier` by fitting a Nearest Neighbors model, computing nearest neighbors for the training data, and calculating differences for feature preparation. Differences are used to train the `MLPClassifier`, accounting for sparse data and employing variants like context addition or distance-based duplication as needed.

Algorithm 18 Training Implementation Algorithm for Base Linger Classifier

Input: Dataset, X , label vector y , number of neighbors $n_neighbors_1$

```
1: function FIT( $X, y$ )
2:   Increment  $n\_neighbors\_1$ 
3:    $neighbors \leftarrow$  Fit scikit-learn's NearestNeighbors function to  $X$ 
4:   Compute nearest neighbors for each sample in  $X$ 
5:    $distances, indices \leftarrow neighbors.kneighbors(X)$ 
6:    $differences\_X \leftarrow []$ 
7:    $differences\_y \leftarrow []$ 
8:   if  $variant == 1$  (LCV1) then
9:     Call  $addition\_of\_context\_func\_fit()$ 
10:     $differences\_X, differences\_y$ 
11:     $= addition\_of\_context\_func\_fit(X, y, indices, differences\_X, differences\_y)$ 
12:  else if  $variant == 2$  (LCV2) then
13:    Call  $duplicate\_difference\_based\_on\_distance() =$ 
14:     $differences\_X, differences\_y =$ 
15:     $duplicate\_difference\_based\_on\_distance(differences\_X, differences\_y,$ 
16:     $distances)$ 
17:  else if  $variant == 3$  (LCV3) then
18:    Call  $add\_additional\_distance\_column(differences\_X, distances)$ 
19:     $differences\_X = add\_additional\_distance\_column(differences\_X, distances)$ 
20:  else
21:    for each sample  $i$  in nearest neighbors indices do
22:       $base \leftarrow$  Index of base case
23:      for each neighbor  $n$  of  $i$  do
24:        Append  $(X[base] - X[n])$  to  $differences\_X$ 
25:        Append  $(y[base] - y[n])$  to  $differences\_y$ 
26:      end for
27:    end for
28:  end if
29:  Fit MLPClassifier with  $differences\_X$  and  $differences\_y$ 
30:  Set  $train\_X = X$ 
31:  Set  $train\_y = y$ 
32:  return Fitted LingerClassifier
33: end function
```

Prediction Algorithm Implementation

The prediction process utilizes the trained model and Nearest Neighbors to predict outcomes for new data. It includes adjustments for sparse data and leverages weighted KNN if enabled, ultimately aggregating predicted values based on computed differences.

Algorithm 19 Prediction Algorithm for Linger Classifier (Part 1)

Input: Input data, X , number of neighbors $n_neighbours_2$

```
1: function PREDICT( $X$ )
2:   Fit Nearest Neighbors model with  $n\_neighbours\_2$  using  $train\_X$ 
3:   Compute nearest neighbors for each sample in  $X$ 
4:   Initialize empty list  $differences\_test\_X$ 
5:    $is\_sparse\_X \leftarrow$  Check if  $X$  is sparse
6:   if  $is\_sparse\_X$  then
7:     Convert  $X$  to dense array
8:   end if
9:   for each sample  $test$  in  $X$  do
10:    for each nearest neighbor  $nn\_in\_training$  in  $indices[test]$  do
11:      Append ( $X[test, :] - train\_X[indices[test]][nn\_in\_training, :]$ )
12:        to  $differences\_test\_X$ 
13:    end for
14:  end for
15:  if  $variant == 1$  (LCV1) then
16:    Call  $addition\_of\_context\_func\_pred(differences\_test\_X, X)$ 
17:     $differences\_test\_X = addition\_of\_context\_func\_pred(differences\_test\_X, X)$ 
18:  else if  $variant == 3$  (LCV3) then
19:    Call  $add\_additional\_distance\_column(differences\_test\_X, distances)$ 
20:     $differences\_X = add\_additional\_distance\_column(differences\_test\_X, distances)$ 
21:  end if
22: end function
```

Algorithm 20 Prediction Algorithm for Linger Classifier (Part 2)

Input: Continuation from Part 1

```
1: function PREDICT_CONTINUE
2:    $predictions \leftarrow$  Predict using LingerClassifier with  $differences\_test\_X$ 
3:   Split  $predictions$  into batches of size  $n\_neighbours\_2$ 
4:   Initialise  $y\_pred$  as an empty list
5:    $y\_pred = []$ 
6:   if Weighted KNN is enabled then
7:     for each  $indexes, differences, distance$  in  $zip(indexes, differences\_test\_X, distances)$ 
8:       do
9:         Compute weights for each neighbor
10:         $weights = 1 \div (dist + 1e - 8)$ 
11:        Compute weighted sum of target values,
12:         $weighted\_sum = []$ 
13:         $weighted\_sum = [w \times (train\_y[i] + d)]$  for  $i, d, w$  in  $zip(indexes, differences, weights)$ 
14:        Find the item with the highest weighted sum
15:         $max\_sum = max(weighted\_sum)$ 
16:        Append  $max\_sum$  to  $y\_pred$ 
17:      end for
18:    else
19:      for each  $indexes, differences$  in  $zip(indexes, predictions)$  do
20:         $results = [train\_y[i] + d \text{ for } i, d \text{ in } zip(indexes, differences)]$ 
21:        Find the most common item in  $results$ 
22:         $result = most\_common(results)$ 
23:        Append the  $result$  to  $y\_pred$ 
24:      end for
25:    end if
26:  return  $y\_pred$ 
27: end function
```

3.4.7 Evaluation

Evaluation metrics for the `LingerClassifier` include accuracy, precision, recall, F1-score, and confusion matrix analysis. These metrics provide insights into the model's performance and its ability to classify instances accurately.

3.4.8 Conclusion

The `LingerClassifier` offers a novel approach to classification tasks by integrating the learning from differences methodology. Its modular design and extensible architecture make it well-suited for a wide range of classification problems, particularly those where interpretability and adaptability are paramount.

For more details on the `LingerClassifier`, and `LingerRegressor` variations, refer to Table ??.

3.5 Variations For `LingerRegressor` (LR) and `LingerClassifier` (LC)

3.5.1 Variation 1: Addition of Context Design and Implementation (LRV1, LCV1)

Objective

Enhance model learning by including both the original attribute values and the differences with nearest neighbors in the dataset. This is applied similarly in both `LingerRegressor` and `LingerClassifier` models.

During both training and prediction, the algorithm appends the source case attributes to the differences array, enriching the feature set with contextual data. This is intended to improve prediction accuracy by preserving information about the original attributes.

Training Design

Algorithm 21 Training Algorithm for Learning from Differences, with context included in `LingerClassifier` and `LingerRegressor`

- 1: Split the dataset D into training set D^{train} and test set D^{test} using an 80%-20% split.
 - 2: Initialize ΔD^{train} as an empty list.
 - 3: **for** each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 - 4: Retrieve n similar cases from D^{train} using nearest neighbors.
 - 5: **for** each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**
 - 6: Compute the differences, with concatenated context:
 - 7: $\Delta D_{ij}^{train} = ((X_i^{train} - X_j^{train}) : X_i^{train}, y_i^{train} - y_j^{train})$.
 - 8: Append ΔD_{ij}^{train} to ΔD^{train} .
 - 9: **end for**
 - 10: **end for**
 - 11: Train the `MLPClassifier` or `MLPRegressor` neural network model using ΔD^{train} .
 - 12: **return** Trained neural network model $TrainedNN()$.
-

Prediction Design

Algorithm 22 Prediction Algorithm for Variant 1, context included.

Input: Test dataset D^{test} , trained neural network model $TrainedNN()$, number of neighbors n

```
1: Initialize  $y^{pred}$  as an empty list.
2: for each case  $X_j^{test}$  in  $D^{test}$  do
3:   Retrieve  $n$  similar cases from  $D^{train}$  using nearest neighbors.
4:   Initialize  $y_j^{pred}$  as an empty list.
5:   for each similar case  $(X_i^{train}, y_i^{train})$  in  $D^{train}$  do
6:     Compute the differences:  $\Delta X_{ji}^{test} = ((X_j^{test} - X_i^{train}) : X_j^{test})$ .
7:     Use the trained neural network to predict:  $Trained\ NN(\Delta X_{ji}^{test}) = \Delta X_{ji}^{pred}$ .
8:     Adapt neighbor's  $y$  values:  $y_{ji}^{pred} = y_i^{train} + \Delta X_{ji}^{pred}$ .
9:   end for
10:  if Regression then
11:    Average the adapted  $y$  values:  $y_j^{pred} = \frac{1}{n} \sum_{i=1}^n y_{ji}^{pred}$ .
12:  else if Classification then
13:    Perform majority voting on  $y_{ji}^{pred}$  to obtain  $y_j^{pred}$ .
14:  end if
15: end for
16: return  $y^{pred}$ 
```

Training Implementation

This algorithm is called during the Fit stage of both the `LingerClassifier`, and `LingerRegressor`.

Algorithm 23 Context Addition Function for Base Linger Classifier

Input: Dataset X , label vector y , nearest neighbor indices $indices$, differences in X $differences_X$, differences in y $differences_y$ **Output:** Updated differences in X $differences_X$, updated differences in y $differences_y$

```
1: function addition_of_context_func_fit( $X, y, indices, differences\_X, differences\_y$ )
2:   for  $i$  in  $indices$  do
3:      $base \leftarrow i[0]$ 
4:      $neighbors \leftarrow i[1:]$ 
5:     for  $n$  in  $neighbors$  do
6:        $base\_case \leftarrow X[base]$ 
7:        $diff \leftarrow X[base] - X[n]$ 
8:        $combined\_list \leftarrow []$ 
9:       for  $pair$  in  $zip(diff, base\_case)$  do
10:        for  $item$  in  $pair$  do
11:          Append  $item$  to  $combined\_list$ 
12:        end for
13:      end for
14:      Append  $np.array(combined\_list)$  to  $differences\_X$ 
15:      Append  $y[base] - y[n]$  to  $differences\_y$ 
16:    end for
17:  end for
18:  return  $differences\_X, differences\_y$ 
19: end function
```

Prediction Implementation

This algorithm is called during the predict algorithm for both the `LingerClassifier`, and `LingerRegressor`.

Algorithm 24 Context Addition Function for Base Linger Classifier during Prediction

Input: Differences in test data $differences_test_X$, input data X **Output:** Combined differences in test data $combined_differences_test_X$

```
1: function addition_of_context_func_pred(differences_test_X, X)
2:   combined_differences_test_X  $\leftarrow$  []
3:   duplicated_test_data  $\leftarrow$  [item for item in X for _ in range(n_neighbours_2)]
4:   num_items  $\leftarrow$  length of duplicated_test_data
5:   for item_pos in range(num_items) do
6:     combined_list  $\leftarrow$  []
7:     for pair in zip(differences_test_X[item_pos], duplicated_test_data[item_pos])
8:       do
9:       for item in pair do
10:        Append item to combined_list
11:      end for
12:    end for
13:    Append np.array(combined_list) to combined_differences_test_X
14:  end for
15: return combined_differences_test_X
16: end function
```

3.5.2 Variation 2: Duplication Based on Distance Design and Implementation (LRV2, LCV2)

Objective

This study introduces a variation that duplicates differences based on distance, enhancing the influence of neighbors closely resembling the base case. This novel approach diverges from traditional methods by prioritizing differences that strongly correlate with the base case, thus reducing the impact of outliers and enhancing model robustness. Integrated into the fitting process of both the **LingerClassifier** and **LingerRegressor**, this method emphasizes the importance of similar neighbors, improving the model's ability to identify meaningful patterns and boosting predictive accuracy and generalization.

1. **Normalize Distances:** Normalize the distances from the base case to its nearest neighbors represented by $Distances_{ij}$. We apply a logarithmic transformation to the distances: $Distances_{ij} = \log_e(Distances_{ij} + 1)$, with the addition of 1 to

prevent a logarithm of 0.

2. **Determine Maximum and Minimum Distances:** Determine the maximum distance $MaxDistance$ and minimum distance $MinDistance$ in $Distances_{ij}$.
3. **Compress Distances:** Compress the distances within $Distances_{ij}$ to a range between 0 and 10 using the formula:

$$Distances_{ij} = \text{round} \left(10 - \frac{(Distances_{ij} - MinDistance)}{(MaxDistance - MinDistance) \times 10} \right)$$

4. **Duplicate Items:** Duplicate each item ΔD_{ij}^{train} by the corresponding distance in $Distances_{ij}$. It's important to note that this duplication process can significantly increase the size of the dataset and, consequently, the training time overhead. The process of duplication is only used in the training process of the algorithm.

Training Design

Algorithm 25 Duplication Based on Distance in LingerClassifier and LingerRegressor

Input: Distances $Distances_{ij}$, Base dataset ΔD^{train}

- 1: Normalize Distances:
 - 2: **for** each distance $Distances_{ij}$ **do**
 - 3: $Distances_{ij} \leftarrow \log_e(Distances_{ij} + 1)$ \triangleright Apply logarithmic transformation
 - 4: **end for**
 - 5: Determine Maximum and Minimum Distances:
 - 6: $MaxDistance \leftarrow$ maximum value of $Distances_{ij}$
 - 7: $MinDistance \leftarrow$ minimum value of $Distances_{ij}$
 - 8: Compress Distances:
 - 9: **for** each distance $Distances_{ij}$ **do**
 - 10: $Distances_{ij} \leftarrow \text{round} \left(10 - \frac{(Distances_{ij} - MinDistance)}{(MaxDistance - MinDistance) \times 10} \right)$
 - 11: **end for**
 - 12: Duplicate Items:
 - 13: **for** each item ΔD_{ij}^{train} in ΔD^{train} **do**
 - 14: Duplicate ΔD_{ij}^{train} by the corresponding distance in $Distances_{ij}$
 - 15: **end for**
-

Training Implementation

Algorithm 26 Difference Duplication Based on Distance (Part 1)

```
1: function duplicate_difference_based_on_distance(differences_X, differences_y, distances
2:   distances_X  $\leftarrow$  []
3:   for d in distances do
4:     neighbors  $\leftarrow$  d[1 :]
5:     for n in neighbors do
6:       append n to distances_X
7:     end for
8:   end for
9:   distances_array  $\leftarrow$  np.array(distances_X)
10:  log_values  $\leftarrow$  np.log(distances_array + 1)
11:  min_val  $\leftarrow$  np.min(log_values)
12:  max_val  $\leftarrow$  np.max(log_values)
13:  if min_val == max_val then
14:    scaled_distances_rounded  $\leftarrow$  np.ones_like(log_values).astype(int)
15:  else
16:    scaled_distances  $\leftarrow$   $10 - ((\log\_values - \min\_val) /$ 
17:       $(\max\_val - \min\_val) \times 10)$ 
18:    scaled_distances_rounded  $\leftarrow$  np.round(scaled_distances).astype(int)
19:  end if
20:  Algorithm continues
21: end function
```

Algorithm 27 Difference Duplication Based on Distance (Part 2)

```
1: function DUPLICATE_DIFF_CONTD(filtered_data_X, filtered_data_y, scaled_dist_rounded)
2:   duplicated_list_X  $\leftarrow$  []
3:   duplicated_list_y  $\leftarrow$  []
4:   for each item, count in zip(filtered_data_X, scaled_dist_rounded) do
5:     for  $\_ \leftarrow 1$  to count do
6:       append item to duplicated_list_X
7:     end for
8:   end for
9:   for each item, count in zip(filtered_data_y, scaled_dist_rounded) do
10:    for  $\_ \leftarrow 1$  to count do
11:      append item to duplicated_list_y
12:    end for
13:  end for
14:  return duplicated_list_X, duplicated_list_y
15: end function
```

3.5.3 Variation 3: Addition of Distance Column Design and Implementation (LRV3, LCV3)

Objective

This novel variation incorporates the distance between each base case and its nearest neighbor into both the training and prediction phases of the neural network. By leveraging this proximity information, the approach enhances the learning process and predictive accuracy of the model. Integrating distance information helps the model capture nuanced patterns and relationships, enhancing its robustness and reliability. It also improves robustness by identifying outliers via distance.

Design for Training

In the training phase, distances between each base case D_i^{train} and its nearest neighbor are calculated and stored in $Distances_{ij}$. These distances are then concatenated with the corresponding differences ΔD^{train} , forming a new dataset where each difference is paired with its respective distance.

Algorithm 28 Training Algorithm for Learning from Differences, with Distance Column

Input: Dataset D , Neural network model, Number of neighbors n , Number of epochs E

- 1: Split D into training set D^{train} and test set D^{test} with an 80%-20% split
 - 2: Initialize ΔD^{train} and $Distances$ as empty lists
 - 3: **for** each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**
 - 4: Retrieve n similar cases from D^{train} using nearest neighbors
 - 5: **for** each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**
 - 6: Compute the differences: $\Delta D_{ij}^{train} = (X_i^{train} - X_j^{train}, y_i^{train} - y_j^{train})$
 - 7: Compute the distance: $Distances_{ij} = \|X_i^{train}, X_j^{train}\|$ (Euclidean distance)
 - 8: Concatenate $Distances_{ij}$ as a new attribute to ΔD_{ij}^{train}
 - 9: **end for**
 - 10: **end for**
 - 11: Train the MLPRegressor neural network model using ΔD^{train} over E epochs
 - 12: **return** Trained neural network model $TrainedNN()$.
-

Design for Prediction

During the prediction phase, distances between each test case X_j^{test} and its nearest neighbors are calculated and stored in $Distances_{ji}$. These distances are then added as attributes to the differences ΔX^{test} , creating a dataset that maintains a one-to-one ratio between distances and differences.

Algorithm 29 Prediction Algorithm for Learning from Differences with Additional Distance Column

Input: Test dataset D^{test} , Trained neural network model, Number of neighbors n , $TrainedNN()$

```
1: Initialize  $\Delta D^{train}$  and  $Distances$  as empty lists
2: for each case  $X_j^{test} \in D^{test}$  do
3:   Retrieve  $n$  similar cases from  $D^{train}$  using nearest neighbors
4:   for each similar case  $(X_i^{train}, y_i^{train}) \in D^{train}$  do
5:     Compute the differences:  $\Delta X_{ij}^{test} = (X_j^{test} - X_i^{train})$ 
6:     Compute the distance:  $Distances_{ji} = ||X_j^{test}, X_i^{train}||$  (Euclidean distance)
7:     Add  $Distances_{ji}$  as a new attribute to  $\Delta X_{ji}^{test}$ 
8:     Use the trained neural network to predict:  $TrainedNN(\Delta X_{ji}^{test}) = \Delta X_{ji}^{pred}$ 
9:     Adapt neighbor's  $y$  values:  $y_j^{pred} = y_i^{train} + \Delta X_{ji}^{pred}$ 
10:  end for
11:  if Regression then
12:    Average the adapted  $y$  values:  $y_j^{pred} = \frac{1}{n} \sum_{i=1}^n y_j^{pred}$ 
13:  else if Classification then
14:    Perform majority voting on  $y_j^{pred}$  to obtain  $y_j^{pred}$ 
15:  end if
16:  return  $y^{pred}$ 
17: end for
```

Implementation for Training, and Prediction

The same function is called during both predict and training.

Algorithm 30 Add Additional Distance Column

Input: Differences dataset $differences_X$, Distances of the nearest neighbors $distances$

```
1: function add_additional_distance_column(differences_X, distances)
2:   Adds a column with distance values associated with each differences_X during
   fitting.
3:   Output: differences_X with an additional column.
4:   for  $i$  in  $\text{range}(\text{len}(\text{distances}))$  do
5:      $diff\_res \leftarrow$  Convert  $distances[i]$  to numpy array
6:     Append  $diff\_res$  to  $differences\_X[i]$ 
7:   end for
8:   return differences_X
9: end function
```

3.6 Learning From differences Implicitly

3.6.1 Introduction and Motivation

This design segment aims to implement and extend the algorithms proposed by Ye et al. [18] for both classification and regression tasks. The primary objective is to develop a unified framework where a single set of algorithms can be employed for both classification and regression tasks, leveraging implicit learning from differences within case pairs.

As outlined in the literature review (see Section 2), the algorithms (Algorithm 9, Algorithm 10) proposed by Ye et al. were chosen as the basis for our model implementation. Variant 5, as demonstrated in the paper, exhibited superior performance among the variants explored, prompting its adoption in our implementation.

Moreover, while the paper focuses on classification tasks, it overlooks the potential application of implicit difference learning within case pairs for regression tasks. Thus, our secondary motivation is to explore the adaptability of Variant 5 to regression models. We seek to investigate whether the framework outlined in Algorithm 9 and Algorithm 10 can be extended and optimized for regression tasks effectively.

By extending the application of Variant 5 to regression tasks, we aim to provide a comprehensive and versatile framework for both classification and regression tasks, harnessing the power of implicit learning from differences within case pairs.

3.7 Linger Implicit Classification Model:

`LingerImplicitClassifier` (LIC)

3.7.1 Objective

The objective of the Linger Implicit Classification Base Model (`LingerImplicitClassifier`) is to implement the learning from differences methodology seen in [18] for classification tasks. The primary goals include:

- Developing a versatile classification model that learns through implicit differences in case pairs.
- Ensuring compatibility with Scikit-learn for seamless integration and extensive hyperparameter testing.
- Providing interpretable predictions to facilitate user understanding, for a variety of tasks.
- Implementing enhancements to the learning from differences algorithm for improved classification performance through hyperparameter testing.
- Designing a modular and extensible codebase to support future variations and enhancements.

The `LingerImplicitClassifier` is tailored for classification problems where interpretability and adaptability are critical, Where explicit differences may not be easily calculable. It adds another layer of interpretability, as both the target case and retrieved case are visible within the case pair which the Neural Nets are trained on.

3.7.2 Architecture

The `LingerImplicitClassifier` combines K-nearest neighbors (KNN) and `MLPClassifier`, utilizing KNN for generating informative case pairs during both

training and prediction phases. This hybrid approach leverages the strengths of both models to enhance classification accuracy.

3.7.3 Components

- **KNN Module:** Identifies nearest neighbors to form case pairs.
- **MLPClassifier:** Provides a robust framework for learning from the identified case pairs.
- **Additional Features:** Includes class-to-class pair formation and selection mechanisms for neighbors.

3.7.4 Parameterization

The model provides extensive parameterization options for fine-tuning KNN, MLPClassifier, and additional features. Hyperparameters for the classification model mirror those available in standard KNN and neural networks, with additional parameters specific to LingerClassifier.

3.7.5 Training and Prediction Design

The training and prediction algorithms for the LingerImplicitClassifier.

Training Algorithm Design

Algorithm 31 Training Algorithm for LingerImplicitClassifier

Input: Dataset D , number of neighbors 1

Split D into training set D^{train} and test set D^{test} with an 80%-20% split

Create *CasePairs()* dictionary.

Create *AdaptNN()*, a dictionary to store each Neural Network

List of unique Classes *UniqueClasses*

for each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**

 Retrieve 1 similar case from D^{train} using nearest neighbors.

for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**

CasePairs(y_j^{train}).append($[X_j^{train} : X_i^{train}, y_i^{train}]$)

end for

end for

for each unique *Class* in *UniqueClasses* **do**

 Create a new Classification Neural Network, trained on Case Pairs

AdaptNN(*Class*).fit(*CasePairs*(*Class*))

end for

return Trained neural network models dictionary *AdaptNN*().

Prediction Algorithm Design

During the prediction phase, various methods exist for retrieving neighbors within each class. These methods can be categorized into three variants:

1. **Variant 1:** (LICV1) Selecting N random items from each class.
2. **Variant 2:** (LICV2) Selecting the single nearest neighbor from each class.

Each variant may be suitable depending on the specific characteristics of the dataset and the objectives of the prediction task.

Algorithm 32 Prediction Algorithm for LingerImplicitClassifier

Input: Dataset D^{test} , Dictionary $AdaptNN()$, Set $UniqueClasses$, and Dictionary $CasePairs()$

Initialize an empty list y^{pred}

for each case $X_j^{test} \in D^{test}$ **do**

 Create an empty list $preds$

for each unique class $Class$ in $UniqueClasses$ **do**

if Base version: (LIC) **then**

 Retrieve N nearest neighbors from $CasePairs(Class)$

else if variant 1: (LICV1) **then**

 Select N random items from $CasePairs(Class)$

else if variant 2: (LICV2) **then**

 Select the single nearest neighbor from $CasePairs(Class)$

end if

for each selected case $(X_i^{CasePair}, y_i^{CasePair})$ **do**

for each $(X_z^{train}, X_q^{train}) \in X_i^{CasePair}$ **do**

$preds.append(AdaptNN(Class).predict(X_z^{train} : X_j^{test}))$

end for

end for

 Perform a majority vote on $preds$

$y_j^{pred} = MajorityVote(preds)$

end for

end for

return y^{pred}

3.7.6 Training and Prediction Implementation

Training Algorithm Implementation

Both training and prediction phases share similar structural implementations, promoting code reuse and maintainability. The training phase involves preparing the neural networks by forming and utilizing case pairs, while the prediction phase focuses on applying these trained networks to new data.

Algorithm 33 Fit Function for LingerImplicitClassifier

Input: Training data X , target labels y

```
1: function FIT( $X, y$ )
2:    $is\_sparse\_X \leftarrow$  Check if  $X$  is sparse
3:    $is\_sparse\_y \leftarrow$  Check if  $y$  is sparse
4:   if  $is\_sparse\_X$  then
5:     Convert  $X$  to dense array
6:   end if
7:   if  $is\_sparse\_y$  then
8:     Convert  $y$  to dense array
9:   end if
10:   $classes\_ \leftarrow$  Get unique classes from  $y$ 
11:   $adaptation\_networks\_ \leftarrow \{\}$ 
12:   $neighbours \leftarrow$  Fit Nearest Neighbors with  $X$ 
13:   $distances, indices \leftarrow$  Compute nearest neighbors for each sample in  $X$ 
14:  Initialize  $case\_pairs\_X$  and  $case\_pairs\_y$ 
15:  for each  $i$  in  $indices$  do
16:     $base \leftarrow i[0]$ 
17:     $neighbors \leftarrow i[1:]$ 
18:    for each  $n$  in  $neighbors$  do
19:       $case\_x \leftarrow [X[base], X[n]]$ 
20:       $case\_y \leftarrow [y[base], y[n]]$ 
21:      Append  $case\_x$  to  $case\_pairs\_X$ 
22:      Append  $case\_y$  to  $case\_pairs\_y$ 
23:    end for
24:  end for
25:  Initialize dictionaries:  $case\_pairs\_by\_class, Cases\_without\_concatenation, case\_pairs\_y$ 
26:  for each ( $source, target$ ) in  $zip(case\_pairs\_X, case\_pairs\_y)$  do
27:    Append  $source[0]$  to  $cases\_without\_concatenation[target[1]]$ 
28:     $problem\_query \leftarrow$  Concatenate  $source[0]$  and  $source[1]$ 
29:    Append  $problem\_query$  to  $case\_pairs\_by\_class[target[1]]$ 
30:    Append  $target[0]$  to  $case\_pairs\_y[target[1]]$ 
31:  end for
32:  for each  $class\_label, cases$  in  $case\_pairs\_by\_class$  do
33:    Initialize MLPClassifier with parameters
34:    Fit MLPClassifier with  $cases$  and  $case\_pairs\_y[class\_label]$ 
35:    Store the fitted MLPClassifier in  $adaptation\_networks\_ [class\_label]$ 
36:  end for
37: end function
```

Prediction Algorithm Implementation

The `predict` function of the `LingerImplicitClassifier` model predicts class labels for the input samples. It creates nearest neighbor finders for each class, then iterates through the test samples to predict class labels. Depending on the chosen variant, it either selects nearest neighbors using (LICV1) or a single nearest neighbor using (LICV2). Finally, it performs majority voting to decide the final classification.

Algorithm 34 Predict Function for `LingerImplicitClassifier`

Input: Test data X

```
1: function PREDICT( $X$ )
2:    $y^{pred} \leftarrow []vector$ 
3:    $is\_sparse\_X \leftarrow$  Check if  $X$  is sparse
4:   if  $is\_sparse\_X$  then
5:     Convert  $X$  to dense array
6:   end if
7:    $nbrs\_by\_class \leftarrow$  Create nearest neighbor finders for each class
8:   for each  $sample$  in  $X$  do
9:     if  $variant == 2$  (LICV2) then
10:       $class\_predictions \leftarrow$  Predict class labels for  $sample$  using 1 NearestNeighbor
11:    else
12:       $class\_predictions \leftarrow$  Predict class labels for  $sample$  using  $N$  NearestNeighbors
13:    end if
14:     $majority\_prediction \leftarrow$  Perform majority voting on  $class\_predictions$ 
15:    Append  $majority\_prediction$  to  $y^{pred}$ 
16:  end for
17:  return  $y^{pred}$ 
18: end function
```

Algorithm 35 Helper: Create Nearest Neighbor Finders

```
1: function _CREATE_NBR_BY_CLASS
2:    $nbrs\_by\_class \leftarrow \{\}$ 
3:   for each  $class\_label$  in  $classes\_$  do
4:      $n\_neighbors \leftarrow 1$  if  $variant == 2$  (LICV2) else  $n\_neighbours\_2$ 
5:      $nbrs \leftarrow$  NearestNeighbors( $n\_neighbors$ ).fit( $cases\_without\_concatenation[class\_label]$ )
6:      $nbrs\_by\_class[class\_label] \leftarrow nbrs$ 
7:   end for
8:   return  $nbrs\_by\_class$ 
9: end function
```

Algorithm 36 Helper: Predict Class Labels

```
1: function _predict_class(sample, nbrs_by_class)
2:   class_predictions  $\leftarrow$  {class_label : [] for class_label in classes_}
3:   for each class_label, nbrs in nbrs_by_class.items() do
4:     if variant == 1 (LICV1) then
5:       indices  $\leftarrow$  Randomly select n_neighbours_2 indices from range(len(cases_without_c
6:     else
7:       distances, indices  $\leftarrow$  nbrs.kneighbors([sample])
8:     end if
9:     nearest_neighbors  $\leftarrow$  indices[0]
10:    for each neighbor_index in nearest_neighbors do
11:      neighbor_sample  $\leftarrow$  cases_without_concatenation[class_label][neighbor_index]
12:      merged_case_pair  $\leftarrow$  Concatenate neighbor_sample and sample
13:      adapted_prediction  $\leftarrow$  Predict with adaptation_networks_[class_label]
14:        using merged_case_pair
15:      Append adapted_prediction[0] to class_predictions[class_label]
16:    end for
17:  end for
18:  return class_predictions
19: end function
```

Algorithm 37 Helper: Perform Majority Voting

```
1: function _majority_vote(class_predictions)
2:   all_predictions  $\leftarrow$  [prediction for predictions in
3:     class_predictions.values() for prediction in predictions]
4:   prediction_counter  $\leftarrow$  Counter(all_predictions)
5:   most_common_predictions  $\leftarrow$  prediction_counter.most_common(1)
6:   If the case occurs where there is no distinct most common.
7:   if len(most_common_predictions) > 1 then
8:     majority_prediction  $\leftarrow$  Randomly select from most_common_predictions
9:   else
10:    majority_prediction  $\leftarrow$  most_common_predictions
11:  end if
12:  return majority_prediction
13: end function
```

3.7.7 Conclusion

The `LingerImplicitClassifier` offers a novel approach to classification tasks by integrating the learning from differences implicitly through case pairs. It performs well for a variety of classification tasks. That could benefit from a more interpretable learning methodology

3.8 Linger Implicit Regression Model:

`LingerImplicitRegressor`

3.8.1 Objective

The objective of the `LingerImplicitRegressor` is to pioneer a learning methodology seen in [18] tailored specifically for regression tasks, known as Learning from Differences. The primary objectives encompass:

- Creation of a flexible regression model capable of learning implicitly from differences observed in pairs of cases.
- Adaptation of regression tasks into the Learning from Differences framework to unlock its potential.
- Seamless integration with Scikit-learn to ensure compatibility and enable extensive hyperparameter optimization.
- Provision of interpretable predictions crucial for user comprehension across a wide spectrum of regression scenarios.
- Continuous enhancement of the Learning from Differences algorithm through rigorous hyperparameter testing to achieve superior regression performance.
- Designing a modular and extensible codebase to accommodate future variations and improvements.

The `LingerImplicitRegressor` is designed to address regression challenges where explicit differences may be elusive, prioritizing interpretability and adaptability. It implements the classification methodology described in [18] but facilitates its use for regression tasks. This is a novel approach. It provides added transparency by exposing

both the target case and the retrieved case within the case pair, facilitating a deeper understanding during model training.

3.8.2 Architecture

The `LingerImplicitRegressor` utilizes a hybrid architecture integrating K-nearest neighbors (KNN) with Scikit-learn's `MLPClassifier`. In the training phase, KNN plays a pivotal role in computing nearest neighbors, thereby facilitating the creation of case pairs. During the prediction phase, KNN is again utilized to retrieve the most similar cases from each class.

The process of generating case pairs for predictions involves KNN's functionality to identify the nearest neighbors. These neighbors serve as crucial elements in forming case pairs, enabling the regressor to make informed predictions based on similarities observed within the dataset.

The integration of KNN with the `MLPClassifier` offers an interesting approach to regression tasks, combining the strengths of both algorithms. It places regression values within specified ranges, offering a new type of solution to classic regression issues. While KNN excels in identifying similarities and forming case pairs, the `MLPClassifier` contributes by providing a powerful neural network-based classification model capable of learning intricate patterns and relationships within the data. By placing continuous values, into classes, we essentially construct a classification problem out of a regression issue.

By harnessing the complementary capabilities of KNN and the `MLPClassifier`, the `LingerImplicitRegressor` aims to achieve robust and accurate regression results across diverse datasets while using a classifier, making it a versatile tool for various machine learning applications. This new approach to regression tasks that allows for categorization of regression results. It processes a regression task and converts it to a classification task, this could be useful for problems where ranges of values are highly important. This would make certain regression tasks more interpretable.

3.8.3 Components

The components are analogous to those of the `LingerImplicitClassifier`, focusing on the effective generation of case pairs and leveraging neural networks for predictive modeling.

3.8.4 Parameterization

The model provides extensive parameterization options for fine-tuning KNN, `MLPClassifier`, and additional features. Hyperparameters for the classifier model mirror those available in standard KNN and neural networks, with additional parameters specific to `LingerRegressor`.

3.8.5 Training and Prediction Design

The training and prediction algorithms for the `LingerImplicitRegressor`.

Training Algorithm Design

Algorithm 38 Training Algorithm for `LingerImplicitRegressor`

Input: Dataset D , number of neighbors 1, N Classes

Split D into training set D^{train} and test set D^{test} with an 80%-20% split

Create *CasePairs()* dictionary.

Create *AdaptNN()*, a dictionary to store each Neural Network

Convert training data y values into classes, using the algorithm 39

$D^{train}y, unique_ranges, y_{train}^{max}, y_{train}^{min} = RegToClassConverterFunction(D^{train}, N)$

Convert test y values based on the train y value range. So that a new range is the same for both.

$D^{test}y, unique_ranges, y_{test}^{max}, y_{test}^{min} = RegToClassConverterFunction(D^{train}, N, y_{train}^{max}, y_{train}^{min})$

List of unique Classes *UniqueClasses*

for each case $C_i = (X_i^{train}, y_i^{train}) \in D^{train}$ **do**

 Retrieve 1 similar case from D^{train} using nearest neighbors.

for each similar case $(X_j^{train}, y_j^{train}) \in D^{train}$ **do**

CasePairs(y_j^{train}).append($[X_j^{train} : X_i^{train}, y_i^{train}]$)

end for

end for

for each unique *Class* in *UniqueClasses* **do**

 Create a new Classification Neural Network, trained on Case Pairs

AdaptNN(*Class*).fit(*CasePairs*(*Class*))

end for

return Trained neural network models dictionary *AdaptNN*()

Prediction Algorithm Design

In the prediction phase for regression, several methods exist for retrieving neighbors within each class. These methods can be categorized into three variants:

1. **Variant 1:** (LIRV1) Selecting N random items from each class.
2. **Variant 2:** (LIRV2) Selecting the single nearest neighbor from each class.

The prediction algorithm mirrors Algorithm 32. The key distinction lies in the preprocessing step, where continuous y values are converted into classes to enable accuracy testing.

3.8.6 Training and Prediction Implementation

Both the `LingerImplicitClassifier` and `LingerImplicitRegressor` share the same training and prediction implementations. This shared functionality is encapsulated in a base class, allowing for code reuse and maintenance. The algorithm 39 found in the 3.8.7 section is used during the data preprocessing stage in the same way as `sklearn.LabelEncoder`. To convert the regression problem into a classification problem.

Training Implementation

The training algorithm for `LingerImplicitRegressor` is outlined in 3.7.6. The function 3.8.7 is called before training to convert y values into class labels for both training and test datasets

Prediction Implementation

The prediction algorithm for `LingerImplicitRegressor` is outlined in 3.7.6.

3.8.7 Regression to Classification Converter Design and Implementation

(RTCC) This function serves to categorize continuous y target values for training, validation, and testing. It returns a human-readable range with their corresponding

numeric prediction. This ensures that the results are interpretable to humans at the end of the process. Central to its operation is the parameter equal division, which indicates whether samples should be equally distributed into N classes. This feature enables uniform representation across different segments of the target variable, promoting fairness and consistency in the categorization process.

Regression to Classification Converter Design

The "Regression to Classification Converter" algorithm (Algorithm 39) is designed to transform continuous regression target values into categorical classes. It takes as input a dataset D , the desired number of segments N , and optional parameters such as y_{\max} , y_{\min} , and Equal division.

If Equal division is set, the algorithm divides the data into N segments based on nearly equal counts. Otherwise, it calculates the range for each segment based on the provided or inferred maximum and minimum values of y . It then iterates through each value in y , determining the corresponding category based on the segment ranges. The algorithm returns the transformed y values, along with the unique ranges, minimum and maximum values of y , providing a categorical representation suitable for classification tasks.

Algorithm 39 Converter Algorithm: Regression to Classification (Part 1)

Input: Dataset D , number of segments N , optional: y_{\max} , y_{\min} , Equal division

```

1:  $(X, y) \in D$ 
2: Get the minimum and maximum values of the labels
3: Ensure that the max and min are used from the test set so that  $y$  values are classified
   the same in both fit and predict.
4: if Equal division then
5:   Divide the data into  $N$  segments based on nearly equal counts
6:    $quantiles \leftarrow \text{np.linspace}(0, 1, N + 1)$ 
7:   if  $unique\_ranges$  is None then
8:      $unique\_ranges \leftarrow \text{np.quantile}(y, quantiles)$ 
9:   end if
10:   $categories \leftarrow \text{np.digitize}(y, unique\_ranges[:-1])$ 
11:   $category\_counts \leftarrow \text{Counter}(categories)$ 
12:  Return  $categories, unique\_ranges, y_{\min}, y_{\max}$ 
13: end if

```

Algorithm 40 Converter Algorithm: Regression to Classification (Part 2)

Input: Continuation from Part 1

```
1: if  $y_{\max}$  and  $y_{\min}$  are provided then
2:   Leave  $y_{\max}$  and  $y_{\min}$  as is
3: else
4:    $y_{\max} \leftarrow \text{MAX}(y)$ 
5:    $y_{\min} \leftarrow \text{MIN}(y)$ 
6: end if
7: Calculate the range for each segment for  $y$  values
8:  $\text{segmentRange} \leftarrow \frac{y_{\max} - y_{\min}}{N}$ 
9: Initialize an empty list: categories
10: Initialize an empty dictionary: unique_ranges
11:  $\text{current\_index} \leftarrow 0$ 
12: for each value val in  $y$  do
13:   for  $\text{doi}$  in  $\text{range}(N)$ 
14:     if  $i < N - 1$  then
15:       if  $y_{\min} + i \times \text{segmentRange} \leq \text{val} < y_{\min} + (i + 1) \times \text{segmentRange}$  then
16:          $\text{category} \leftarrow (y_{\min} + i \times \text{segmentRange}) - (y_{\min} + (i + 1) \times \text{segmentRange})$ 
17:         if category not in unique_ranges then
18:            $\text{unique\_ranges}[\text{category}] \leftarrow \text{current\_index}$ 
19:            $\text{current\_index} \leftarrow \text{current\_index} + 1$ 
20:         end if
21:         Append  $\text{unique\_ranges}[\text{category}]$  to categories
22:         break
23:       end if
24:     else
25:       if  $y_{\min} + i \times \text{segmentRange} \leq \text{val} \leq y_{\max}$  then
26:          $\text{category} \leftarrow (y_{\min} + i \times \text{segmentRange}) - (y_{\max})$ 
27:         if category not in unique_ranges then
28:            $\text{unique\_ranges}[\text{category}] \leftarrow \text{current\_index}$ 
29:            $\text{current\_index} \leftarrow \text{current\_index} + 1$ 
30:         end if
31:         Append  $\text{unique\_ranges}[\text{category}]$  to categories
32:         break
33:       end if
34:     end if
35:   end for
36: end for
37: Converted  $y$  values:  $y \leftarrow \text{categories}$ 
38: Return  $y, \text{unique\_ranges}, y_{\min}, y_{\max}$ 
```

Regression to Classification Converter Implementation

This function is encapsulated within a class called `RegressionToClassificationConverter`. The `transform` function carries out the conversion of regression values to classification values.

Algorithm 41 Transform function Initialised

```
1: function TRANSFORMINIT(y, min_val, max_val, unique_ranges)
2:   Input:
3:   y (list or array-like): The continuous target variable.
4:   min_val (float): Minimum value of the target variable.
5:   max_val (float): Maximum value of the target variable.
6:   unique_ranges (array-like): Predefined unique ranges for segments.
7:   Output:
8:   categories (list): A list of categories (segment indices) corresponding
9:     to each value in y.
10:  unique_ranges (dict or array-like): Unique ranges for segments.
11:  min_val (float): Minimum value of the target variable.
12:  max_val (float): Maximum value of the target variable.
13:  if min_val is None and max_val is None then
14:    min_val  $\leftarrow$  min(y)
15:    max_val  $\leftarrow$  max(y)
16:  end if
17: end function
```

Algorithm 42 Transform function (Part 1)

```
1: All variables from transformInit() are available.
2: function    TRANSFORMPARTONE( $y, min\_val$     =     $None, max\_val$     =
    $None, unique\_ranges = None$ )
3:   if equal_division is None then
4:      $segment\_range \leftarrow (max\_val - min\_val) / n\_segments$ 
5:      $categories \leftarrow []$ 
6:      $unique\_ranges \leftarrow \{\}$ 
7:      $current\_index \leftarrow 0$ 
8:     for  $val$  in  $y$  do
9:       for  $i$  in range( $n\_segments$ ) do
10:        if  $i < n\_segments - 1$  then
11:          if  $min\_val + i \times segment\_range \leq val < min\_val + (i + 1) \times$ 
    $segment\_range$  then
12:             $category \leftarrow \text{int}(min\_val + i \times segment\_range)$ 
13:             $\text{int}((min\_val + (i + 1)) \times segment\_range)$ 
14:            if category not in  $unique\_ranges$  then
15:               $unique\_ranges[category] \leftarrow current\_index$ 
16:               $current\_index \leftarrow current\_index + 1$ 
17:            end if
18:             $categories.append(unique\_ranges[category])$ 
19:            break
20:          end if
21:        else
22:          if  $min\_val + i \times segment\_range \leq val \leq max\_val$  then
23:             $category \leftarrow \text{int}(min\_val + i \times segment\_range)$ 
24:             $\text{int}(max\_val)$ 
25:            if category not in  $unique\_ranges$  then
26:               $unique\_ranges[category] \leftarrow current\_index$ 
27:               $current\_index \leftarrow current\_index + 1$ 
28:            end if
29:             $categories.append(unique\_ranges[category])$ 
30:            break
31:          end if
32:        end if
33:      end for
34:    end for
35:    return  $categories, unique\_ranges, min\_val, max\_val$ 
36:  else
37:    Call transformPartTwo() function
38:  end if
39: end function
```

Algorithm 43 Transform Function (Part 2): Equal Division

```
1: function TRANSFORMPARTTWO( $y, min\_val, max\_val, unique\_ranges, n\_segments$ )
2:    $quantiles \leftarrow \text{np.linspace}(0, 1, n\_segments + 1)$ 
3:   if  $unique\_ranges$  is None then
4:      $unique\_ranges \leftarrow \text{np.quantile}(y, quantiles)$ 
5:   end if
6:    $categories \leftarrow \text{np.digitize}(y, unique\_ranges[: -1])$ 
7:   return  $categories, unique\_ranges, min\_val, max\_val$ 
8: end function
```

3.8.8 Conclusion

The `LingerImplicitClassifier` offers a novel approach to regression tasks by integrating the learning from differences implicitly through case pairs. It performs well for a variety of regression tasks, while utilizing a learning from differences classification model. The number of segments greatly affects the outcome of this model. One of the limitations observed the `LingerImplicitClassifier` is its sensitivity to noise, particularly when labeling values based on the maximum and minimum. However, opting to evenly distribute samples into classes can mitigate this sensitivity.

3.9 Learning from Differences For Images

3.9.1 Introduction and Motivation

The introduction of the `LingerImageClassifier` and `LingerImageRegressor` represents a novel approach to image classification and regression tasks. Motivated by the innovative methodology outlined in "Learning from Differences 2022," these tools are specifically designed to enhance interpretability and explainability in processing image data. The aim was to ensure these models not only integrate smoothly into the existing software framework but also support the customization of Keras convolutional neural networks. This strategy diverges from traditional methods, such as those employed by Scikit-learn's `MLPClassifier` and `MLPRegressor`, offering a fresh perspective on addressing image-centric challenges.

3.10 Linger Image Classification Model

`LingerImageClassifier` (LIMC)

3.10.1 Objective

The motivation behind the development of (LIMC) stems from the growing demand for more sophisticated image classification tools capable of handling the complexities and nuances of real-world data. Traditional models often fall short in terms of explainability, and transparency. (LIMC) aims to address these gaps by:

- Incorporating state-of-the-art convolutional neural network (CNN) designs that

improve upon the limitations of existing models in capturing intricate patterns in image data.

- Utilizing a novel training methodology seen in [7] that enhances model learning from image differences, inspired by recent research findings.
- Aiming to demystify the black-box nature of deep neural networks by providing insights into the decision-making process, making the model more trustworthy and applicable in sensitive areas such as medical diagnosis and autonomous driving.

3.10.2 Architecture

The `LingerImageClassifier` employs a sophisticated, modular architecture designed to leverage the nuances of image data, enhancing both predictive accuracy and interpretability. This architecture consists of three integral components:

- **Custom Fit Method:** Developed to integrate with the scikit-learn ecosystem, this Fit method is central to the classifier's functionality. It generates a unique dataset of image differences, effectively capturing variations between image pairs. This method not only emphasizes the importance of contrastive learning but also enriches the model's understanding of nuanced features within the data, such as changes in texture, lighting, and spatial relationships. The process involves computing the differential aspects of images, focusing on attributes that significantly influence the y label values, thereby enabling a more focused and effective learning strategy. It also guarantees that we know what it is precisely learning from, its visual difference to its neighbor.
- **Convolutional Neural Network (CNN):** At the heart of the classifier lies a customizable Convolutional Neural Network, tailored from Keras's comprehensive library. This CNN is specifically designed to process and predict the differences highlighted by the custom Fit method. The architecture features a combination of convolutional layers, activation functions, and pooling layers, strategically organized to maximize feature extraction and minimize information loss. Special attention is given to the network's depth, filter sizes, and the incorporation of advanced techniques such as batch normalization and dropout for regularization. This careful construction ensures optimal performance in discerning subtle differences in image data, thus significantly contributing to the precision of subsequent predictions.

- **Custom Predict Method:** Complementing the custom Fit method, the Predict function stands as a testament to the model's innovative approach to classification. This component takes the CNN's predictions on image differences and synthesizes them with the original data to formulate the final classification outcomes. The method employs an algorithm that accurately applies predicted differences to their corresponding instance, accounting for the model's understanding of contrastive features within the dataset. This not only enhances the model's accuracy but also its adaptability to various image classification scenarios, ensuring that the final predictions are both precise and contextually grounded.

Integration and Workflow: The synergy between these components forms the cornerstone of the `LingerImageClassifier`'s architecture. Initially, the custom Fit method preprocesses the input images to create a differential dataset, which is then fed into the CNN for feature extraction and difference prediction. The custom Predict method subsequently interprets these predictions, applying them to derive the final classification results. This integrated workflow ensures a coherent and efficient processing pipeline, from initial data preparation to final outcome generation.

Innovations and Contributions: The architecture of the `LingerImageClassifier` introduces several key innovations to the field of image classification. By focusing on image differences, the model uncovers deeper insights into the data, facilitating a more nuanced understanding and interpretation of the factors driving classification decisions. This approach not only improves model performance across diverse datasets but also enhances the explainability of its predictions, addressing a critical need within the domain of deep learning. Visually we could Identify exactly what the image is learning from based on the difference image generated.

3.10.3 Components

The key components of the `LingerImageClassifier` include:

- KNN Module: Computes nearest neighbors used to extract differences during training phase and prediction phase.
-
- **Convolutional Neural Network (CNN):** The heart of the classification process is a Convolutional Neural Network (CNN), meticulously designed to analyze and interpret the differences identified by the KNN module. This CNN is

notable for its flexibility, allowing for the customization of activation functions, solvers, and an extensive range of hyperparameters to optimize classification performance. The adaptability of the CNN component ensures that the model can be fine-tuned to suit various datasets and classification challenges. It excels in processing the contrastive features extracted by the KNN module, effectively learning from these differences to make accurate predictions. The CNN's architecture, which can include layers specifically tailored to enhance feature extraction and reduce overfitting, such as dropout and batch normalization layers, is instrumental in capturing the complex patterns inherent in image data.

3.10.4 Parameterization

The `LingerImageClassifier` is designed with flexibility and adaptability at its core, offering a comprehensive suite of parameters that can be fine-tuned to optimize performance across a wide range of image classification tasks. The user can vary the design of the (CNN) convolutional Neural Network, as well as vary the number of neighbors used in the fit and prediction methods.

3.10.5 Training and Prediction Design

Training Design

The Image Classification Training Algorithm outlines the steps involved in training a convolutional neural network (CNN) for image classification tasks using the differences between nearest neighbor images.

Algorithm 44 Image Classification Training Algorithm

```

function TRAINIMAGECLASSIFIER( $X, y$ )
  Input: Dataset of images  $X$ , Target labels  $y$ 
  Output: Trained CNN model
  Generate the image difference datasets.
   $differences\_X, differences\_y \leftarrow \text{GenerateDifferences}(X, y)$ 
  Initialize (CNN) model.
  Train CNN model using  $differences\_X$  and  $differences\_y$ 
  return Trained CNN model
end function

```

The ‘GenerateDifferences’ 45 algorithm encapsulates the process of computing differences between nearest neighbor images and their base image, preparing the data for training the k-nearest neighbors classifier. It iterates through each image in the training dataset, identifies its nearest neighbors, and calculates the absolute differences between their pixel values. These differences, along with the target value differences, are stored in lists for subsequent use in the training process.

Algorithm 45 Generate Differences Algorithm

```

function GENERATEDIFFERENCES( $X, y$ )
  Input: Training images  $X$ , Target values  $y$ 
  Output: List of arrays representing differences  $differences_X$ ,
            List of target differences  $differences_y$ 
  if Shape of  $X$  is More than 2-dimensional then
    Reshape  $X$  to 2-dimensional array
  else
    Keep  $X$  as it is
  end if
  Increment  $n\_neighbours\_1$  by 1
  Initialize empty list  $differences\_X$ 
  Initialize empty list  $differences\_y$ 
  Fit nearest neighbors to find indices of nearest neighbors
  Compute nearest neighbor indices for each image in  $X$ 
  for each image  $I$  in  $X$  do
    Get nearest neighbors indices for  $I$ 
    for each neighbor index  $N$  do
      Get neighbor image and target value
      Compute absolute difference between  $I$  and neighbor image
      Append difference to  $differences\_X$ 
      Append target value difference to  $differences\_y$ 
    end for
  end for
  return  $differences\_X, differences\_y$ 
end function

```

Prediction Design

This algorithm, named "Image Classification Prediction Algorithm," takes a dataset of images X and a trained convolutional neural network (CNN) model $trainedCNN$ as input. It utilizes the trained CNN model to predict the differences between images in the dataset. The predicted differences are then processed further using the FinalPrediction algorithm (referenced as Algorithm 47) to compute the final predictions for the images. Finally, the algorithm returns the predicted target values $y_p predictions$.

Algorithm 46 Image Classification Prediction Algorithm

```
1: function PREDICTIMAGECLASSIFIER( $X$ ,  $trainedCNN$ )
2:   Input: Dataset of images  $X$ , Trained CNN model  $trainedCNN$ 
3:   Output: Predictions
4:   Use trained CNN on  $X$  to predict differences
5:    $difference\_pred \leftarrow trainedCNN(X)$ 
6:   Compute final predictions using Algorithm:
7:    $y^{pred} \leftarrow \text{FinalPrediction}(difference\_pred)$ 
8:   return  $y^{pred}$ 
9: end function
```

This algorithm, titled "FinalPrediction," takes input data X , a trained CNN model $model$, and the input shape $input_shape$. It computes the predicted target values y^{pred} based on the differences between the input data and the training images using the trained CNN model.

First, it fits nearest neighbors to the training data and then computes the absolute differences between each test image and its nearest neighbors from the training set. These differences are reshaped to match the input shape and used to make predictions using the trained CNN model.

Finally, it computes the predicted results for each nearest neighbor and determines the most common item among them to form the final prediction y^{pred} . If no items are found, it handles this case by appending "None" to y^{pred} .

Algorithm 47 Image Classification Prediction Algorithm

```
1: function FINALPREDICTIONCLASSIFICATION( $X$ ,  $model$ ,  $input\_shape$ )
2:   Input: Input data  $X$ , Trained CNN model  $model$ , Input shape  $input\_shape$ 
3:   Output: Predicted target values  $y\_pred$ 
4:   Fit nearest neighbors to the training data
5:   Initialize empty list  $differences\_test\_X$ 
6:   Compute differences between test and training images
7:   for  $test$  in range(length of  $X$ ) do
8:     for  $nn\_in\_training$  in range(length of nearest neighbors) do
9:       Compute difference between test and nearest neighbor:
10:       $diff \leftarrow$  absolute difference between  $X[test]$ 
11:        and  $self.train\_X[indices[test][nn\_in\_training]]$ 
12:      Append  $diff$  to  $differences\_test\_X$ 
13:    end for
14:  end for
15:  Reshape  $differences\_test\_X$  to match input shape
16:  Make predictions using the trained CNN model:
17:   $predictions \leftarrow model.predict(differences\_test\_X)$ 
18:  Initialize empty list  $y\_pred$ 
19:  for  $indexes, differences$  in zip( $indices, predictions$ ) do
20:    Compute predicted results for each nearest neighbor:
21:     $results \leftarrow [$ 
22:       $self.train\_y[i][0] + d$ 
23:      for  $i, d$  in zip( $indexes, differences$ )
24:    Compute counts of each result:
25:     $counts \leftarrow$  count of each value in results
26:    Retrieve most common item:
27:     $most\_common\_item \leftarrow$  most common value in  $results$ 
28:    if  $most\_common\_item$  exists then
29:      Append  $most\_common\_item$  to  $y^{pred}$ 
30:    else
31:      Append None to  $y^{pred}$  ▷ Handle cases with no items
32:    end if
33:  end for
34:  return  $y^{pred}$ 
35: end function
```

3.10.6 Training and Prediction Implementation

Training Implementation

Algorithm 48 Initialize LingerImageClassifier

```
1: Number of nearest neighbors for difference computation
2: Input:  $n\_neighbours\_1, n\_neighbours\_2$ 
3: procedure INITIALIZECLASSIFIER( $n\_neighbours\_1, n\_neighbours\_2$ )
4:    $self.n\_neighbours\_1 \leftarrow n\_neighbours\_1$ 
5:    $self.n\_neighbours\_2 \leftarrow n\_neighbours\_2$ 
6:   return self
7: end procedure
```

Algorithm 49 Generate Differences

```
1: procedure GENERATEDIFFERENCES( $X, y, n\_neighbours$ )
2:   Input:  $X$ : dataset of images,  $y$ : target labels,  $n\_neighbours$ : number of neighbors
3:   Differences in images and labels
4:   Output:  $differences\_X, differences\_y$ 
5:   Initialize  $differences\_X$  and  $differences\_y$  as empty lists
6:    $neigh \leftarrow$  Fit NearestNeighbors on  $X$  with  $n\_neighbours$ 
7:   for each image  $I$  in  $X$  do
8:     Find  $n\_neighbours$  nearest neighbors for  $I$ 
9:     for each neighbor  $N$  do
10:      Compute absolute difference between  $I$  and  $N$ 
11:      Append difference to  $differences\_X$ 
12:      Compute difference in target values and append to  $differences\_y$ 
13:     end for
14:   end for
15:   return  $differences\_X, differences\_y$ 
16: end procedure
```

Algorithm 50 Train with LingerImageClassifier

```
1: procedure TRAIN LINGERIMAGECLASSIFIER( $X_{train}, y_{train}, X_{test}$ )  
2:   Load dataset into  $X_{train}, y_{train}, X_{test}, y_{test}$   
3:   Initialize LingerImageClassifier with  $n\_neighbours\_1$  and  $n\_neighbours\_2$   
4:    $differences\_X, differences\_y \leftarrow \text{GenerateDifferences}(X_{train}, y_{train})$   
5:   Train CNN model on  $differences\_X, differences\_y$   
6: end procedure
```

Prediction Implementation

Algorithm 51 Prediction Implementation with LingerImageClassifier

```
1:  $X$  = test data set,  $model$  = Trained CNN model,  $input\_shape$  = image shape
2: procedure PREDICT( $X, model, input\_shape$ )  $\triangleright$  Pythonic steps for predicting
   target values using image differences.
3:   Input:  $X$  – Input data,  $model$  – Trained CNN,  $input\_shape$  –
   Shape for CNN input.
4:   Output:  $y\_pred$  – Predicted targets.
5:    $nbrs \leftarrow$  NearestNeighbors( $n\_neighbors = self.n\_neighbours\_2$ )
6:    $.fit(self.train\_X.reshape(len(self.train\_X), -1))$ 
7:    $indices \leftarrow nbrs.kneighbors(X.reshape(len(X), -1), return\_distance = False)$ 
8:    $differences\_test\_X \leftarrow [np.abs(X[test].flatten() -$ 
9:      $self.train\_X[indices[test][nn]].flatten())$ 
10:    for test in range(len(X)) for nn in range(len(indices[0]))]
11:    $differences\_test\_X \leftarrow np.array(differences\_test\_X).reshape(-1, *input\_shape)$ 
12:    $predictions \leftarrow model.predict(differences\_test\_X)$ 
13:    $y^{pred} \leftarrow []$  vector
14:   for  $indexes, differences$  in zip( $indices, predictions$ ) do
15:      $results \leftarrow [self.train\_y[i][0] + d$  for  $i, d$  in zip( $indexes, differences$ )]
16:      $counts \leftarrow$  Counter( $results$ )
17:      $most\_common \leftarrow counts.most\_common(1)$ 
18:     if  $most\_common$  then
19:        $y^{pred}.append(most\_common[0][0])$ 
20:     else
21:        $y^{pred}.append(None)$ 
22:     end if
23:   end for
24:   return  $y^{pred}$ 
25: end procedure
```

3.11 Linger Image Regressor Model

LingerImageRegressor (LIMR)

3.11.1 Objective

The motivation behind the development of (LIMR) stems from the growing demand for more sophisticated image regression tools capable of image data and solving regression issues. Traditional models often fall short in terms of explainability, and transparency. (LIMR) aims to address these gaps by:

- Incorporating state-of-the-art convolutional neural network (CNN) designs that

improve upon the limitations of existing models in capturing intricate patterns in image data.

- Utilizing a novel training methodology seen in [7] that enhances model learning from image differences, inspired by recent research findings.
- Aiming to demystify the black-box nature of deep neural networks by providing insights into the decision-making process, making the model more trustworthy and applicable in sensitive areas such as medical diagnosis and autonomous driving.

3.11.2 Architecture

Its architecture is similar to the classifier, only that it uses a convolutional neural net to predict continuous values rather than classes. The method of applying the generated differences to the data during prediction varies slightly. The structure of the `LingerImageClassifier`, and `LingerImageRegressor` is identical, the functionality of their predict methods vary. See section 3.10.2 for a more indepth analysis.

3.11.3 Components

The key component within the `LingerImageRegressor` are identical to those in the `LingerImageClassifier`. For more details see section 3.10.3.

3.11.4 Parameterization

The `LingerImageRegressor` is designed with flexibility and adaptability at its core, offering a comprehensive suite of parameters that can be fine-tuned to optimize performance across a wide range of image classification tasks. The user can vary the design of the (CNN) convolutional Neural Network, as well as vary the number of neighbors used in the fit and prediction methods.

3.11.5 Training and Prediction Design

Training Design

The Image Regression Training Algorithm outlines the steps involved in training a convolutional neural network (CNN) for image regression tasks using the differences between nearest neighbor images. For regression, it uses the same algorithm ?? to generate the image differences datasets as `LingerImageClassifier`. . To finalize its

Algorithm 52 Image Regression Training Algorithm

```
function TRAINIMAGEREGRESSOR( $X, y$ )  
  Input: Dataset of images  $X$ , Target labels  $y$   
  Output: Trained CNN model  
  Generate the image difference datasets.  
   $differences\_X, differences\_y \leftarrow \text{GenerateDifferences}(X, y)$   
  Initialize (CNN) model.  
  Train CNN model using  $differences\_X$  and  $differences\_y$   
  return Trained CNN model  
end function
```

prediction it takes an average of nearest neighbors with their predicted differences added to them.

This algorithm outlines a method for image regression prediction using a Convolutional Neural Network (CNN). It involves fitting nearest neighbors to training data, calculating differences between test images and their nearest training images using, and using these differences as inputs to a pre-trained CNN model to predict target values. The algorithm computes predictions by averaging adjustments made to the nearest neighbors' target values based on the CNN's output 53, thereby producing a final prediction for each input image. This approach combines instance-based learning with deep learning to enhance prediction accuracy for image-based regression tasks. The method of learning from differences is based on the algorithms in the paper [7],

Algorithm 53 Image Regression Prediction Algorithm

```
1: function FINALPREDICTION( $X$ ,  $model$ ,  $input\_shape$ )
2:   Input: Input data  $X$ , Trained CNN model  $model$ , Input shape  $input\_shape$ 
3:   Output: Predicted target values  $y\_pred$ 
4:   Fit nearest neighbors to the training data
5:   Initialize empty list  $differences\_test\_X$ 
6:   Compute differences between test and training images
7:   for  $test$  in range(length of  $X$ ) do
8:     for  $nn\_in\_training$  in range(length of nearest neighbors) do
9:       Compute difference between test and nearest neighbor:
10:       $diff \leftarrow$  absolute difference between  $X[test]$ 
11:        and  $train\_X[indices[test][nn\_in\_training]]$ 
12:      Append  $diff$  to  $differences\_test\_X$ 
13:    end for
14:  end for
15:  Reshape  $differences\_test\_X$  to match input shape
16:  Make predictions using the trained CNN model:
17:   $predictions \leftarrow model.predict(differences\_test\_X)$ 
18:  Initialize empty list  $y^{pred}$ 
19:  for  $indexes, differences$  in zip( $indices, predictions$ ) do
20:    Compute predicted results for each nearest neighbor:
21:     $results \leftarrow [self.train\_y[i][0] + d \text{ for } i, d \text{ in zip}(indexes, differences)]$ 
22:    Compute average of each result:  $average \leftarrow$  average of values in  $results$ 
23:    Append  $average$  to  $y^{pred}$ 
24:  end for
25:  return  $y^{pred}$ 
26: end function
```

3.11.6 Training and Prediction Implementation

Training Implementation

The same algorithm 49 is used to generate differences for the training process in the `LingerImageRegressor` as the `LingerImplicitClassifier`.

Algorithm 54 Initialize LingerImageRegressor

```
1: Number of nearest neighbors for difference computation
2: Input:  $n\_neighbours\_1, n\_neighbours\_2$ 
3: procedure INITIALIZEREGRESSOR( $n\_neighbours\_1, n\_neighbours\_2$ )
4:    $self.n\_neighbours\_1 \leftarrow n\_neighbours\_1$ 
5:    $self.n\_neighbours\_2 \leftarrow n\_neighbours\_2$ 
6:   return self
7: end procedure
```

Algorithm 55 Train with LingerImplicitRegressor

```
1: procedure TRAIN LINGERIMPLICITREGRESSOR( $X_{train}, y_{train}, X_{test}$ )
2:   Load dataset into  $X_{train}, y_{train}, X_{test}, y_{test}$ 
3:   Initialize LingerImageRegressor with  $n\_neighbours\_1$  and  $n\_neighbours\_2$ 
4:    $differences\_X, differences\_y \leftarrow \text{GenerateDifferences}(X_{train}, y_{train})$ 
5:   Train CNN model on  $differences\_X, differences\_y$ 
6: end procedure
```

Prediction Implementation

Algorithm 56 Prediction with LingerImageRegressor

```
1: procedure PREDICT( $X, model, dataset, input\_shape$ )
2:   Predicts target values using a regression approach based on image differences.
3:   Input:  $X$ - Input data for prediction,  $model$ - Trained regression model,
4:            $dataset$ - Dataset object containing training data,
5:            $input\_shape$ - Shape of input data expected by the model.
6:   Output:  $y^{pred}$  - Predicted target values for the input data.
7:   if  $len(X.shape) > 2$  then
8:      $X \leftarrow X.reshape(len(X), -1)$ 
9:     Flatten the input if it's multi-dimensional.
10:  end if
11:  Train the NearestNeighbors function on our data.
12:   $nbrs \leftarrow$  NearestNeighbors( $n\_neighbors = self.n\_neighbours\_2$ )
13:   $.fit(dataset.train\_X.reshape(len(dataset.train\_X), -1))$ 
14:  Identify nearest neighbors in the training set.
15:   $indices \leftarrow nbrs.kneighbors(X, return\_distance = False)$ 
16:  Prepare  $differences\_test\_X$  by computing absolute differences between  $X$ 
17:  and its nearest neighbors in the training set.
18:  Reshape  $differences\_test\_X$  to match  $input\_shape$  for
19:  compatibility with the model.
20:   $predictions \leftarrow model.predict(differences\_test\_X)$ 
21:  Predict target values based on computed differences.
22:  Initialize an empty list  $y^{pred}$  to store final predictions.
23:  for each  $indexes, differences$  pair in zipped  $indices, predictions$  do
24:    Calculate the sum of the training target values and the predicted differences.
25:    Compute the average result to obtain the regression prediction.
26:    Append the average result to  $y^{pred}$ .
27:  end for
28:  return  $y^{pred}$ 
29: end procedure
```

3.12 Optimization Strategies

Across all Linger models, a consistent focus on hyperparameter tuning and architectural refinements enhances performance and interpretability. For instance:

- **Hyperparameter Tuning:** Each model undergoes extensive hyperparameter optimization to identify settings that maximize accuracy and efficiency. This includes adjustments to learning rates, neighbor counts, and decision thresholds.
- **Architectural Adjustments:** Modifications in model architectures, especially for variant models, aim to better capture the nuances of learning from differences.

This involves experimenting with different layers, activation functions, and aggregation methods to improve model responsiveness to dataset specifics.

- **CNN Configurations for Image-Based Models:** Image models, such as the LingerImageClassifier and LingerImageRegressor, benefit from tailored Convolutional Neural Network (CNN) configurations. By optimizing filter sizes, stride patterns, and pooling layers, these models achieve significant gains in learning efficiency and prediction accuracy.
- **Computational Efficiency:** Efforts are underway to enhance the computational efficiency of the models, focusing on reducing training times without compromising performance. Techniques include model pruning, efficient data loading, and leveraging hardware accelerations.
- **Dataset Applicability:** A key optimization goal is to ensure models can adapt to a wide range of datasets, encompassing varying sizes, dimensions, and complexity. This involves refining data preprocessing and feature extraction techniques to maintain model robustness across diverse data scenarios.

3.13 Limitations and Future Work

While the Linger models introduce innovative approaches to machine learning, they encounter certain limitations that direct future research:

- **Computational Intensity:** The models, especially when dealing with large datasets or complex image data, can be computationally demanding. Ongoing research seeks to address this through algorithmic improvements that reduce computational overhead without sacrificing accuracy.
- **Data Quality Dependency:** The effectiveness of the models is closely tied to the quality and representativeness of the training data. Efforts to make the models more robust to noisy or incomplete data include developing advanced noise filtering techniques and data augmentation strategies.
- **Scalability to Larger Datasets:** As datasets grow in size and complexity, scaling model training and inference efficiently becomes a challenge. Future developments aim to enhance the scalability of Linger models, potentially through distributed computing techniques and more efficient data handling mechanisms.

- **Extension to New Domains:** Exploring applications of Linger models in new domains, such as time series analysis, natural language processing, and more complex forms of image and video data, represents a promising avenue for future work. This includes adapting the learning from differences approach to tackle domain-specific challenges.
- **Algorithmic Efficiency:** Developing more efficient algorithms to handle the iterative processes inherent in learning from differences is a primary focus. This may involve exploring new mathematical frameworks or leveraging advancements in optimization theory.
- **Broadening Data Types and Applications:** The adaptability of Linger models to various data types and application contexts remains an area for exploration. Future research will investigate the integration of these models into multi-modal learning frameworks and their applicability in interdisciplinary fields.

Chapter 4

Experiments

4.1 Research Objective

The overarching objective of this study is to investigate the efficacy of blended learning approaches in machine learning tasks, particularly focusing on regression and classification problems. Blended learning, which integrates the strengths of model-based and instance-based learning while mitigating their respective weaknesses, offers a promising avenue for enhancing the interpretability, reliability, and performance of machine learning systems.

Specifically, this research aims to achieve the following objectives:

1. **Evaluate Blended Learning Methods:** The first objective is to evaluate the performance of blended learning methodologies in contrast to traditional model-based and instance-based approaches across various regression and classification tasks. This assessment will encompass key metrics including accuracy, interpretability, computational efficiency, and robustness. By comparing the effectiveness of blended learning methods, we aim to determine their potential to enhance the overall performance of machine learning systems.
2. **Investigate Model Interpretability:** The second objective is to explore the interpretability of blended learning models compared to traditional black-box models, such as neural networks. Understanding the interpretability of these models is crucial for enhancing trust and usability in critical domains. By investigating how blended learning methods provide insights into model decisions, we aim to demonstrate their potential to address the interpretability challenges associated with traditional machine learning approaches.

3. **Examine Efficiency and Scalability:** The third objective is to investigate the computational efficiency and scalability of blended learning methods, particularly in scenarios with large datasets. This examination will involve analyzing the memory and computational overhead of blended learning models compared to traditional approaches. By assessing the efficiency and scalability of blended learning methods, we aim to identify their suitability for practical deployment in real-world applications.
4. **Explore Novel Architectures:** The fourth objective is to explore novel architectures and techniques for implementing blended learning, including ensemble models, case-based reasoning principles, and approaches that leverage differences between sets of features. By exploring innovative approaches, we aim to further enhance the performance and interpretability of blended learning methods, thereby advancing the state-of-the-art in machine learning.
5. **Provide Insights for Practical Applications:** The fifth objective is to provide insights and recommendations for the practical application of blended learning methods in real-world scenarios, with a particular focus on critical domains such as healthcare, finance, and decision support systems. By discussing the implications of the findings for stakeholders and suggesting best practices for deploying blended learning models effectively, we aim to facilitate the adoption of these methods in critical domains.
6. **Develop Python Package for Seamless Integration:** The final objective is to develop a Python package that seamlessly integrates with scikit-learn, allowing users to incorporate blended learning methods into their machine learning pipelines and workflows effortlessly. This package should provide compatibility with scikit-learn's tools, enabling users to import it like any other scikit-learn package and use it seamlessly within their existing workflows. By developing such a package, we aim to enhance the accessibility and usability of blended learning methods for the broader machine learning community.

By addressing these objectives, this research seeks to advance our understanding of blended learning approaches and their potential to address the challenges faced by traditional model-based and instance-based methods. Ultimately, the goal is to contribute to the development of more interpretable, reliable, and efficient machine learning systems with broader applicability across various domains.

4.1.1 Data Collection and Preprocessing

All datasets used in this study were collected from the UCI Machine Learning Repository [1], a widely recognized and reliable source of machine learning datasets. These datasets cover various domains and have been extensively studied in the machine learning community.

Preprocessing For Labelled Datasets

The preprocessing procedure involves several steps to ensure the quality and suitability of the data for machine learning tasks. These steps are applied uniformly across all labeled datasets.

1. **Column Naming:** The columns of the dataset are assigned appropriate names for clarity and consistency throughout the analysis.
2. **Handling Missing Values:** Rows containing missing values in specified features are removed from the dataset. This ensures that the dataset used for training and testing is complete and does not contain any incomplete or unreliable data points.
3. **Index Resetting:** After removing rows with missing values, the index of the dataset is reset to maintain a continuous and consistent index structure.
4. **Data Splitting:** The dataset is split into two subsets: a development set and a test set. This splitting is performed using a train-test split ratio of 80 : 20 to ensure a sufficient amount of data for training while also reserving a portion for evaluating the model's performance.
5. **Development Set Copying:** A copy of the development set is created to preserve the original dataset for future reference or comparison.
6. **Feature Transformation:** Numeric features undergo feature scaling using the `StandardScaler` to ensure that they are on a similar scale, which aids in model convergence and performance. Nominal features are transformed using one-hot encoding to convert categorical variables into a numerical format suitable for machine learning algorithms.

These preprocessing steps collectively prepare the data for subsequent modeling and analysis, ensuring that it is clean, complete, and appropriately formatted for use in machine learning tasks.

4.2 Results and Analysis

4.2.1 Experiment 1: Glass Dataset Classification

Methodology

In Experiment 1, we conducted classification tasks on the Glass dataset using a two-step methodology. Initially, we trained a neural network (NN) classifier through grid search and cross-validation to determine the optimal hyperparameters. These hyperparameters were then applied to train a `LingerClassifier` (LC) and its variants for improved robustness, tested across five runs. For comparative analysis, we also evaluated the performance of K-Nearest Neighbors (KNN), weighted (KNN), and (NN) on the same dataset. The results section details the hyperparameters selected and the test accuracies achieved.

The `LingerImplicitClassifier` was only run over 3 iterations due to a lack of resources.

Hyperparameter Optimization

For both the Neural Network and Linger Classifier, a comprehensive exploration of hyperparameters was conducted, as summarized in Table 4.1. This table delineates the varied hyperparameters scrutinized during the optimization phase, spanning hidden layer configurations, activation functions, regularization terms, and more, underscoring our meticulous approach to model tuning.

Table 4.1: Tested Hyperparameters for Neural Network and Linger Classifier

Hyperparameter	Values
Neural Network	
Hidden Layer Sizes	(256, 128), (128, 64), (100,), (200, 100), (300, 200, 100)
Activation	identity, logistic, tanh, relu
Alpha	0.0001, 0.001, 0.01, 0.1
Max Iterations	1000, 1500
Early Stopping	True
Validation Fraction	0.1, 0.2, 0.3
Learning Rate Init	0.001, 0.01, 0.1
Solver	adam, sgd
Beta 1	0.9, 0.95, 0.99
Beta 2	0.999, 0.995, 0.9
Linger Classifier	
Neighbors 1	2, 5, 7, 10, 13, 15, 17, 21
Neighbors 2	2, 5, 7, 10, 13, 15, 17, 21
Weighted KNN	False
Additional Distance Column	True, False
Duplicated on Distance	True, False
Addition of Context	True, False

Results

Table 4.2: Best Parameters for KNN Comparison Models

Classifier	Hyperparameter	Value
KNN	n_neighbors	2
Weighted KNN	n_neighbors	7

Table 4.3: Summary of Best Parameters for `LingerClassifier` and its Variations

Parameter	LC	LCV1	LCV2	LCV3
Activation	relu	relu	relu	relu
Alpha	0.0001	0.1	0.1	0.001
Beta 1	0.9	0.9	0.9	0.99
Beta 2	0.995	0.999	0.995	0.999
Early Stopping	True	True	True	True
Hidden Layer Sizes	(300, 200, 100)	(300, 200, 100)	(300, 200, 100)	(300, 200, 100)
Learning Rate Init	0.01	0.01	0.01	0.01
Max Iter	1000	1500	1000	1000
Solver	adam	adam	adam	adam
Validation Fraction	0.3	0.3	0.3	0.3
Addition of Context	False	True	False	False
Additional Distance Column	False	False	False	True
Duplicated on Distance	False	False	True	False
N Neighbors 1	2	17	21	21
N Neighbors 2	5	17	13	13
Weighted KNN	False	False	False	False

Table 4.4: Summary of Best Parameters for `LingerImplicitClassifier` and Its Variations

Parameter	LIC	LICV1 (Random)	LICV2 (Single)
Activation	tanh	tanh	tanh
Alpha	0.01	0.001	0.001
Beta 1	0.9	0.9	0.9
Beta 2	0.9	0.9	0.9
Hidden Layer Sizes	(128, 64)	(128, 64)	(128, 64)
Learning Rate Init	0.01	0.01	0.01
Max Iter	2000	2000	2000
Solver	sgd	sgd	sgd
Validation Fraction	0.1	0.1	0.1
N Neighbors 1	21	7	10
N Neighbors 2	13	2	21
Random Pairs	False	True	False
Single Pair	False	False	True

Table 4.5: Summary of Training and Test Accuracy Across All Models

Model / Variation	Training Accuracy (%)	Test Accuracy (%)
KNN	68.2	60.46
Weighted KNN	70.0	62.8
Neural Network	73.1	61.4
LC (Base)	70.04	60.46
LCV1 (Context)	69.41	65.70
LCV2 (Duplicated on Distance)	67.29	69.30
LCV3 (Additional Distance Column)	67.65	60.93
LIC (Implicit)	41.4	51.2
LICV1 (Random)	31.17	41.08
LICV2 (Single)	50.39	47.28

Analysis

- **LCV2 (Duplicated on Distance)**

- Highest test accuracy.
- Effective relationship modeling.

- **Neural Network**

- High training accuracy.
- Reduced test performance, indicating potential overfitting on training data.

- **Weighted KNN**

- Better performance than standard KNN.
- Benefits from using neighbor distances as weights.

4.2.2 Experiment 2: Abalone Dataset Regression

Methodology

In Experiment 2, we explored regression tasks on the Abalone dataset with a two-step approach. Initially, a neural network (NN) regressor was trained using grid search and cross-validation to identify optimal hyperparameter combinations. These selected hyperparameters were then applied to train a **LingerRegressor** (LR) and its variants,

targeting improved robustness, with their performance evaluated across four runs. Additionally, to benchmark the `LingerRegressor`'s performance, K-Nearest Neighbors (KNN), weighted KNN, and NN models were similarly assessed on the dataset. The results section will detail the hyperparameters and performance metrics obtained from Experiment 2. The variation in the number of run times is due to resource constraints. For (LIR) and its variant, the continuous y values were broken up into 3 classes with a roughly equal number of samples in each.

- `LingerRegressor` (LR) was run 4 times, its (LRV1), (LRV2), and (LRV3) variants were each run twice
- `LingerImplicitRegressor` (LIR), along with its variants LICV1 and LICV2, were each run twice.

Negative mean squared error is used for LR and all variations. Accuracy % is used for LIR and all variations. As it uses a classifier on regression tasks.

Tested Hyperparameters

Table 4.6: Tested Hyperparameters for Neural Network and LingerRegressor

Hyperparameter	Values
Neural Network	
Hidden Layer Sizes	(256, 128), (128, 64), (200, 100), (300, 200, 100)
Activation	relu
Alpha	0.0001, 0.001, 0.01, 0.1
Max Iterations	2000
Early Stopping	True
Validation Fraction	0.1
Learning Rate Init	0.001, 0.01, 0.1
Solver	adam, sgd
Beta 1	0.9, 0.95, 0.99
Beta 2	0.999, 0.995, 0.9
Linger Regressor	
N Neighbours 1	2, 5, 7, 10, 13, 15, 17, 21
N Neighbours 2	2, 5, 7, 10, 13, 15, 17, 21
Weighted KNN	False
Additional Distance Column	True, False
Duplicated on Distance	True, False
Addition of Context	True, False

Results

Table 4.7: Best Parameters for KNN Comparison Models

Classifier	Hyperparameter	Value
KNN	n_neighbors	17
Weighted KNN	n_neighbors	21

Table 4.8: Summary of Best Parameters for `LingerRegressor` and its Variations

Parameter	LR	LRV1	LRV2	LRV3
Activation	relu	relu	relu	relu
Alpha	0.01	0.001	0.1	0.01
Beta 1	0.95	0.95	0.95	0.99
Beta 2	0.9	0.9	0.9	0.9
Early Stopping	True	True	True	True
Hidden Layer Sizes	(200, 100)	(200, 100)	(256, 128)	(200, 100)
Learning Rate Init	0.01	0.01	0.01	0.01
Max Iter	2000	2000	2000	2000
Solver	sgd	sgd	sgd	sgd
Validation Fraction	0.1	0.1	0.1	0.1
Addition of Context	False	True	False	False
Additional Distance Column	False	False	False	True
Duplicated on Distance	False	False	True	False
N Neighbors 1	5	2	21	5
N Neighbors 2	21	17	21	17
Weighted KNN	False	False	False	False

Table 4.9: Summary of Best Parameters for `LingerImplicitRegressor` and Its Variations

Parameter	LIR	LIRV1	LIRV2
Activation	relu	relu	relu
Alpha	0.01	0.01	0.01
Beta 1	0.99	0.9	0.9
Beta 2	0.9	0.9	0.9
Early Stopping	True	True	True
Hidden Layer Sizes	(128, 64)	(256, 128)	(200, 100)
Learning Rate Init	0.01	0.01	0.01
Max Iter	2000	2000	2000
Solver	sgd	sgd	sgd
Validation Fraction	0.1	0.1	0.1
N Neighbors 1	13	17	15
N Neighbors 2	21	13	7
Random Pairs	False	True	False
Single Pair	False	False	True

Table 4.10: Summary of Training and Test NMSE Across Models

Model / Variation	Training NMSE	Test NMSE
KNN	−1.959	−1.930
Weighted KNN	−1.941	−1.912
Neural Network	−1.930	−1.895
LR (Base)	−1.898	−1.835
LRV1 (Context)	−1.949	−1.897
LRV2 (Duplicated on Distance)	−2.047	−1.949
LRV3 (Additional Distance Column)	−1.9006	−1.8250

Table 4.11: Summary of Training and Test Accuracy Across Models

Model / Variation	Training Accuracy (%)	Test Accuracy (%)
Neural Network with Labels Classified	67.3	64.8
LIR (Implicit)	63.2	57.7
LIRV1 (Random)	52.6	48
LIRV2 (Single)	59.7	52

Analysis

• LRV2 (Duplicated on Distance)

- Best test NMSE.

- Effective in handling patterns and relationships within the data.
- **Neural Network with Labels Classified**
 - Highest test accuracy.
 - Potential overfitting issues, based on the reduction in accuracy from training to test.
- **LRV3 (Additional Distance Column)**
 - Competitive NMSE to the leading accuracy.
 - Efficient in capturing relationships within the data.
- **Weighted KNN**
 - Consistent performance.
 - Benefits from weighting distances.

4.2.3 Experiment 3: Raisin Dataset Classification

Methodology

In Experiment 3, we conducted a comprehensive analysis on the Raisin dataset sourced from the UCI Machine Learning Repository [3], specifically detailed in [5]. Our methodology was two-fold: initially, we determined the optimal hyperparameters for a neural network (NN) classifier through an exhaustive grid search coupled with cross-validation. This foundational step ensured the NN classifier was finely tuned for the dataset. Subsequently, these hyperparameters were employed to train the **LingerClassifier** (LC) and its variants, focusing on enhancing model robustness. The performance of these classifiers was benchmarked against K-Nearest Neighbors (KNN), Weighted KNN, and the baseline NN classifier through five independent runs to ensure statistical reliability. Our comparative analysis leverages this diverse classifier ensemble to draw insights into the efficacy and robustness of each model relative to the Raisin dataset, showcasing the nuanced trade-offs between model complexity and classification accuracy. Due to time and resource constraints the certain classifiers were run for a different number of iterations.

- **LingerClassifier** (LC) was run 4 times.
- **LingerClassifier** variant 1 (LCV1) was run twice.

- `LingerClassifier` variant 2 (LCV2) was run twice.
- item `LingerClassifier` variant 3 (LCV3) was run twice.
- `LingerImplicitClassifier` (LIC), along with its variants LICV1 and LICV2, were each run twice.

Tested Hyperparameters

Table 4.12: Tested Hyperparameters for Neural Network and `LingerClassifier`

Hyperparameter	Values
Neural Network	
Hidden Layer Sizes	(256, 128), (128, 64), (200, 100), (300, 200, 100), (400, 300, 200, 100)
Activation	identity, logistic, tanh, relu
Alpha	0.0001, 0.001, 0.01, 0.1
Max Iterations	1500, 2000
Early Stopping	True
Validation Fraction	0.1, 0.2, 0.3
Learning Rate Init	0.001, 0.01, 0.1
Solver	adam, sgd
Beta 1	0.9, 0.95, 0.99
Beta 2	0.999, 0.995, 0.9
Linger Regressor	
N Neighbours 1	2, 5, 7, 10, 13, 15, 17, 21
N Neighbours 2	2, 5, 7, 10, 13, 15, 17, 21
Weighted KNN	False
Additional Distance Column	True, False
Duplicated on Distance	True, False
Addition of Context	True, False

Results

Table 4.13: Best Parameters for KNN Comparison Models

Classifier	Hyperparameter	Value
KNN	n_neighbors	13
Weighted KNN	n_neighbors	13

Table 4.14: Summary of Best Parameters for `LingerClassifier` and its Variations

Parameter	LC	LCV1	LCV2	LCV3
Activation	identity	tanh	tanh	identity
Alpha	0.01	0.01	0.01	0.01
Beta 1	0.9	0.95	0.9	0.9
Beta 2	0.999	0.9	0.9	0.9
Early Stopping	True	True	True	True
Hidden Layer Sizes	(128, 64)	(256, 128)	(128, 64)	(300, 200, 100)
Learning Rate Init	0.001	0.001	0.001	0.001
Max Iter	1500	1500	1500	2000
Solver	adam	adam	adam	adam
Validation Fraction	0.1	0.3	0.3	0.3
Addition of Context	False	True	False	False
Additional Distance Column	False	False	False	True
Duplicated on Distance	False	False	True	False
N Neighbors 1	5	15	10	2
N Neighbors 2	13	21	21	13
Weighted KNN	False	False	False	False

Table 4.15: Summary of Best Parameters for `LingerImplicitClassifier` and Its Variations

Parameter	LIC	LICV1	LICV2
Activation	tanh	identity	identity
Alpha	0.1	0.01	0.01
Beta 1	0.95	0.9	0.9
Beta 2	0.999	0.999	0.999
Early Stopping	True	True	True
Hidden Layer Sizes	(128, 64)	(400, 300, 200, 100)	(400, 300, 200, 100)
Learning Rate Init	0.1	0.001	0.001
Max Iter	1500	2000	2000
Solver	sgd	adam	adam
Validation Fraction	0.2	0.3	0.3
N Neighbors 1	2	2	7
N Neighbors 2	13	13	13
Random Pairs	False	True	False
Single Pair	False	False	True

Table 4.16: Summary of Training and Test Accuracy Across All Models

Model / Variation	Accuracy (%)	Accuracy (%)
KNN	91.7	91
Weighted KNN	91.2	91
Neural Network	94.1	91
LC (Base)	91.7	90.5
LCV1 (Context)	92.2	89.5
LCV2 (Duplicated on Distance)	92.3	90.5
LCV3 (Additional Distance Column)	91.7	91
LIC (Implicit)	91.6	90.5
LICV1 (Random)	91.6	89.5
LICV2 (Single)	91.5	91

Analysis

- **Neural Network**

- Highest training accuracy.
- Consistent test performance indicates effective generalization.

- **LCV1 (Context)**

- Slightly lower test accuracy.
- Suggests potential overfitting or inefficacy in utilizing contextual data.

- **LCV2 (Duplicated on Distance)**

- Maintains high training accuracy.
- Solid test performance, indicating good performance.

- **Weighted KNN**

- Consistent test accuracy.
- Shows robustness in weighting approach.

- **LICV1 (Random)**

- Lower test accuracy.
- Randomness may introduce variability reducing robustness; over more runs, results may vary.

4.2.4 Experiment 4: House Prices Dataset Regression

Methodology

The methodology is identical to the Abalone experiment 4.2.2. The difference between the experiments is in the number of run times per regressor and variation.

- **LingerRegressor** (LR) was run 3 times, its (LRV1), (LRV2), were run once and (LRV3) was run 5 times.
- **LingerImplicitRegressor** (LIR) was run 3 times, its variants (LICV1) and (LICV2), were each run twice.

Negative mean squared error is used for LR and all variations. Accuracy % is used for LIR and all variations. As it uses a classifier on regression tasks.

Tested Hyperparameters

Table 4.17: Tested Hyperparameters for Neural Network and **LingerRegressor**

Hyperparameter	Values
Neural Network	
Hidden Layer Sizes	(128, 64), (200, 100), (300, 200, 100) (400, 300, 200, 100)
Activation	relu
Alpha	0.0001, 0.001, 0.01, 0.1
Max Iterations	2000
Early Stopping	True
Validation Fraction	0.1
Learning Rate Init	0.001, 0.01, 0.1
Solver	adam, sgd
Beta 1	0.9, 0.95, 0.99
Beta 2	0.999, 0.995, 0.9
Linger Regressor	
N Neighbours 1	2, 5, 7, 10, 13, 15, 17, 21
N Neighbours 2	2, 5, 7, 10, 13, 15, 17, 21
Weighted KNN	False
Additional Distance Column	True, False
Duplicated on Distance	True, False
Addition of Context	True, False

Results

Table 4.18: Best Parameters for KNN Comparison Models

Classifier	Hyperparameter	Value
KNN	n_neighbors	2
Weighted KNN	n_neighbors	5

Table 4.19: Summary of Best Parameters for `LingerRegressor` and its Variations

Parameter	LR	LRV1	LRV2	LRV3
Activation	relu	relu	relu	relu
Alpha	0.1	0.1	0.1	0.01
Beta 1	0.9	0.9	0.9	0.9
Beta 2	0.999	0.999	0.9	0.9
Early Stopping	True	True	True	True
Hidden Layer Sizes	(400, 300, 200, 100)	(300, 200, 100)	(400, 300, 200, 100)	(400, 300, 200, 100)
Learning Rate Init	0.01	0.001	0.01	0.01
Max Iter	2000	2000	2000	2000
Solver	adam	adam	adam	adam
Validation Fraction	0.1	0.1	0.1	0.1
Addition of Context	False	True	False	False
Additional Distance Column	False	False	False	True
Duplicated on Distance	False	False	True	False
N Neighbors 1	17	2	17	15
N Neighbors 2	13	15	13	7
Weighted KNN	False	False	False	False

Table 4.20: Summary of Best Parameters for `LingerImplicitRegressor` and Its Variations

Parameter	LIR	LIRV1	LIRV2
Activation	logistic	logistic	logistic
Alpha	0.001	0.001	0.001
Beta 1	0.9	0.9	0.9
Beta 2	0.9	0.9	0.9
Early Stopping	True	True	True
Hidden Layer Sizes	(128, 64)	(128, 64)	(128, 64)
Learning Rate Init	0.1	0.1	0.1
Max Iter	2000	2000	2000
Solver	sgd	sgd	sgd
Validation Fraction	0.1	0.1	0.1
N Neighbors 1	7	21	15
N Neighbors 2	5	7	15
Random Pairs	False	True	False
Single Pair	False	False	True

Table 4.21: Summary of Training and Test NMSE Across Models

Model / Variation	Training NMSE	Test NMSE
KNN	−61.60	−48.30
Weighted KNN	−60.37	46.94
Neural Network	−56.06	−48.21
LR (Base)	−51.36	−40.45
LRV1 (Context)	−58.29	−49.87
LRV2 (Duplicated on Distance)	−59.69	− 40.04
LRV3 (Additional Distance Column)	−51.37	−42.22

Table 4.22: Summary of Training and Test Accuracy Across Models

Model / Variation	Training Accuracy (%)	Test Accuracy (%)
Neural Network with Labels Classified	63	61
LIR (Implicit)	47.7	48.42
LIRV1 (Random)	31.61	22.10
LIRV2 (Single)	42.73	47.36

Analysis

- **LRV1 (Context)**
 - Highest NMSE improvement from training to test.
 - Effective in context utilization.
- **Neural Network**
 - Consistently strong in both NMSE and accuracy, showcasing overall robustness.
- **Weighted KNN**
 - Competitive test NMSE.
 - Benefiting from adaptive neighbor weighting.
- **LIRV1 (Random)**
 - Struggles with lowest accuracy.
 - Highlights the challenges with random feature selection.

4.2.5 Experiment 5: Dry Bean Dataset Classification

Methodology

The experimental methodology is identical to that of the other classification experiments. The vary factor is the number of runs, due to time constraints.

- **LingerRegressor** (LR), (LRV1), (LRV2), were run once and (LRV3) was run 3 times.
- **LingerImplicitRegressor** (LIR) was run twice, its variant (LICV1) was run twic and (LICV2) was run once.

Tested Hyperparameters

Table 4.23: Tested Hyperparameters for Neural Network and LingerClassifier

Hyperparameter	Values
Neural Network	
Hidden Layer Sizes	(256, 128), (128, 64), (200, 100), (300, 200, 100), (400, 300, 200, 100)
Activation	identity, logistic, tanh, relu
Alpha	0.0001, 0.001, 0.01, 0.1
Max Iterations	1000, 1500
Early Stopping	True
hline Validation Fraction	0.1, 0.2, 0.3
Learning Rate Init	0.001, 0.01, 0.1
Solver	adam, sgd
Beta 1	0.9, 0.95, 0.99
Beta 2	0.999, 0.995, 0.9
LingerClassifier	
N Neighbours 1	2, 5, 7, 10, 13, 15, 17, 21
N Neighbours 2	2, 5, 7, 10, 13, 15, 17, 21
Weighted KNN	False
Additional Distance Column	True, False
Duplicated on Distance	True, False
Addition of Context	True, False

Results

Table 4.24: Best Parameters for KNN Comparison Models

Classifier	Hyperparameter	Value
KNN	n_neighbors	5
Weighted KNN	n_neighbors	10

Table 4.25: Summary of Best Parameters for `LingerClassifier` and its Variations

Parameter	LC	LCV1	LCV2	LCV3
Activation	tanh	tanh	tanh	tanh
Alpha	0.001	0.01	0.01	0.01
Beta 1	0.95	0.95	0.95	0.95
Beta 2	0.999	0.999	0.999	0.999
Early Stopping	True	True	True	True
Hidden Layer Sizes	(128, 64)	(128, 64)	(128, 64)	(128, 64)
Learning Rate Init	0.01	0.01	0.01	0.01
Max Iter	1000	1000	1000	1000
Solver	adam	adam	adam	adam
Validation Fraction	0.3	0.3	0.3	0.3
Addition of Context	False	True	False	False
Additional Distance Column	False	False	False	True
Duplicated on Distance	False	False	True	False
N Neighbors 1	21	21	15	17
N Neighbors 2	15	7	21	13
Weighted KNN	False	False	False	False

Table 4.26: Summary of Best Parameters for `LingerImplicitClassifier` and Its Variations

Parameter	LIC	LICV1	LICV2
Activation	tanh	tanh	tanh
Alpha	0.1	0.01	0.1
Beta 1	0.9	0.9	0.99
Beta 2	0.9	0.9	0.995
Early Stopping	True	True	True
Hidden Layer Sizes	(300, 200, 100)	(300, 200, 100)	(300, 200, 100)
Learning Rate Init	0.01	0.01	0.01
Max Iter	1500	2000	2000
Solver	adam	sgd	adam
Validation Fraction	0.2	0.3	0.3
N Neighbors 1	13	2	7
N Neighbors 2	2	2	15
Random Pairs	False	True	False
Single Pair	False	False	True

Table 4.27: Summary of Training and Test Accuracy Across All Models

Model / Variation	Accuracy (%)	Accuracy (%)
KNN	96.33	96.36
Weighted KNN	96.55	97.27
Neural Network	97.93	98.18
LC (Base)	98.39	100
LCV1 (Context)	98.86	100
LCV2 (Duplicated on Distance)	98.39	100
LCV3 (Additional Distance Column)	98.4	97.3
LIC (Implicit)	5.9	5
LICV1 (Random)	12.7	1.81
LICV2 (Single)	6.45	0.9

Analysis

- **LC Variants**

- All three variations (Context, Duplicated on Distance, Additional Distance Column) achieved perfect test accuracy.
- Indicates robustness in diverse settings.

- **Neural Network**

- High accuracy suggests effective learning.
- Slightly less test resilience compared to LC variants.

- **Weighted KNN**

- Improved performance over standard KNN.
- Benefiting from distance weighting in classification.

- **LIC Variants**

- Significantly lower accuracy.
- Suggesting poor model fit or insufficient training for implicit methods in this context.

4.2.6 Experiment 6: White Wine Database Regression

Methodology

The methodology is identical to the Abalone experiment 4.2.2. The difference between the experiments is in the number of run times per regressor variation.

- `LingerRegressor` (LR),(LRV1), (LRV2), and (LRV3) were run 5 times.
- `LingerImplicitRegressor` (LIR), (LICV1), and (LICV2) were run 5 times.

Negative mean squared error is used for LR and all variations. Accuracy % is used for LIR and all variations. As it uses a classifier on regression tasks.

Tested Hyperparameters

Table 4.28: Tested Hyperparameters for Neural Network and `LingerRegressor`

Hyperparameter	Values
Neural Network	
Hidden Layer Sizes	(128, 64), (200, 100), (300, 200, 100) (400, 300, 200, 100)
Activation	relu
Alpha	0.0001, 0.001, 0.01, 0.1
Max Iterations	2000
Early Stopping	True
Validation Fraction	0.1
Learning Rate Init	0.001, 0.01, 0.1
Solver	adam, sgd
Beta 1	0.9, 0.95, 0.99
Beta 2	0.999, 0.995, 0.9
Linger Regressor	
N Neighbours 1	2, 5, 7, 10, 13, 15, 17, 21
N Neighbours 2	2, 5, 7, 10, 13, 15, 17, 21
Weighted KNN	False
Additional Distance Column	True, False
Duplicated on Distance	True, False
Addition of Context	True, False

Results

Table 4.29: Best Parameters for KNN Comparison Models

Classifier	Hyperparameter	Value
KNN	n_neighbors	2
Weighted KNN	n_neighbors	13

Table 4.30: Summary of Best Parameters for `LingerRegressor` and its Variations

Parameter	LR	LRV1	LRV2	LRV3
Activation	relu	relu	relu	relu
Alpha	0.1	0.01	0.01	0.01
Beta 1	0.95	0.9	0.95	0.95
Beta 2	0.9	0.999	0.999	0.999
Early Stopping	True	True	True	True
Hidden Layer Sizes	(400, 300, 200, 100)	(300, 200, 100)	(400, 300, 200, 100)	(400, 300, 200, 100)
Learning Rate Init	0.01	0.01	0.01	0.01
Max Iter	2000	2000	2000	2000
Solver	sgd	sgd	sgd	sgd
Validation Fraction	0.2	0.2	0.3	0.3
Addition of Context	False	True	True	False
Additional Distance Column	False	False	False	True
Duplicated on Distance	False	False	False	False
N Neighbors 1	21	2	2	21
N Neighbors 2	21	13	5	21
Weighted KNN	False	False	False	False

Table 4.31: Summary of Best Parameters for `LingerImplicitRegressor` and Its Variations

Parameter	LIR	LIRV1	LIRV2
Activation	relu	relu	relu
Alpha	0.01	0.01	0.01
Beta 1	0.9	0.9	0.9
Beta 2	0.9	0.9	0.9
Early Stopping	True	True	True
Hidden Layer Sizes	(128, 64)	(200, 100)	(256, 128)
Learning Rate Init	0.01	0.01	0.01
Max Iter	2000	2000	2000
Solver	sgd	sgd	sgd
Validation Fraction	0.1	0.1	0.1
N Neighbors 1	7	7	2
N Neighbors 2	5	7	15
Random Pairs	False	True	False
Single Pair	False	False	True

Table 4.32: Summary of Training and Test NMSE Across Models

Model / Variation	Training NMSE	Test NMSE
KNN	-0.5947	-0.53
Weighted KNN	-0.5946	-0.4786
Neural Network	-0.5770	-0.6055
LR (Base)	-0.5549	-0.5819
LRV1 (Context)	-0.6178	-0.6193
LRV2 (Duplicated on Distance)	-0.4926	-0.5287
LRV3 (Additional Distance Column)	-0.5600	-0.5799

Table 4.33: Summary of Training and Test Accuracy Across Models

Model / Variation	Training Accuracy (%)	Test Accuracy (%)
Neural Network with Labels Classified	63	61
LIR (Implicit)	55.35	47.55
LIRV1 (Random)	54.79	52
LIRV2 (Single)	53.37	51

Analysis

- **Weighted KNN**
 - Best NMSE test performance among KNN variants.
 - Shows efficiency in handling data weights.
- **Neural Network**
 - Balanced NMSE and highest accuracy among all models.
 - Indicates strong generalization.
- **LRV1 (Context)**
 - Notably lower NMSE than the leading model.
 - Shows improvements in Linger Regression predictions with context.
- **LIR Variants**
 - Consistently lower test accuracies.
 - Suggest challenges in model complexity or training sufficiency, possibly due to inadequate amounts of data.
- **LRV3 (Additional Distance Column)**
 - Slightly worse results than LRV2.
 - Suggesting an increase in the amount of samples would benefit the learning process.

4.2.7 Experiment 7: Image Classification using CIFAR-100

Methodology

In this study, we utilized the CIFAR-100 dataset, ensuring data security through modified SSL settings. Our methodology focused on the development and comparative analysis of two convolutional neural network (CNN) models: the *difference CNN model* and the *standard CNN model*.

The *difference CNN model* processes difference images created by the `LingerImageClassifier`, which uses settings of 5 neighbors for comparison to preprocess the original CIFAR images. This model's architecture, detailed in Table

4.34, emphasizes feature extraction and computational efficiency, culminating in a regression output for predictions.

Simultaneously, the *standard CNN model* was trained directly on normalized CIFAR-100 images, utilizing a similar architecture but concluding with a softmax layer for 100-class classification. Both models were trained over 10, 30, and 100 epochs, with a batch size of 32 and a 20% validation split.

Post-training, model performance was evaluated based on accuracy, comparing predicted labels against the test set. This dual-model approach facilitates a thorough analysis of differing methodologies in image classification efficacy.

Table 4.34: Comparison of CNN Architectures for Standard and Difference Models

Layer Type	Standard Model	Difference Model
Input Layer	32x32x3 images	
Conv2D	32 filters, 3x3, padding=same, relu	32 filters, 3x3, padding=same, relu
Conv2D	32 filters, 3x3, relu	32 filters, 3x3, relu
MaxPooling2D	2x2	2x2
Dropout	0.2	0.2
Conv2D	64 filters, 3x3, padding=same, relu	64 filters, 3x3, padding=same, relu
Conv2D	64 filters, 3x3, relu	64 filters, 3x3, relu
MaxPooling2D	2x2	2x2
Dropout	0.3	0.3
Conv2D	128 filters, 3x3, padding=same, relu	128 filters, 3x3, padding=same, relu
Conv2D	128 filters, 3x3, relu	128 filters, 3x3, relu
MaxPooling2D	2x2	2x2
Dropout	0.4	0.4
Flatten	Flatten output	
Dense	512, relu	512, relu
Dropout	0.5	0.5
Output Layer	100 classes, softmax	1 output, linear

Results

Table 4.35: Regression Performance on CIFAR-100 Dataset Over Different Epochs

Model	Epochs	MSE	Epochs	MSE	Epochs	MSE
LingerImageRegressor CNN	10	0.0078	30	0.0078	100	0.0078
Standard Regressor CNN	10	0.0426	30	0.0543	100	0.0486

Analysis

The analysis of the LingerImageClassifier and Standard CNN on the CIFAR-100 dataset highlights the challenges of using a regression-based approach for classification tasks. While both models share similar convolutional architectures, they differ notably in their output layers. The Standard CNN is optimized for classification with a softmax layer, showing improved MSE from 0.0543 at 30 epochs to 0.0486 at 100 epochs. Conversely, the LingerImageClassifier uses a linear output, typical for regression, to predict label differences, not direct classes, resulting in a low MSE of 0.0078 across all epochs. This consistently low MSE does not necessarily indicate high classification accuracy but rather the model’s precision in estimating label differences, which may not effectively reflect actual class distinctions. This misalignment suggests the need for future model adaptations that better suit classification by aligning output mechanisms with standard classification metrics and better addressing the categorical nature of the dataset. Future improvements might include a modified CNN that categorizes label differences directly, potentially involving a broader range of output classes. Which includes all possible differences.

4.2.8 Experiment 8: Image Regression Using MNIST Dataset

Methodology

In this proof-of-concept study, we adapted the MNIST dataset, traditionally used for classification, to evaluate image-based regression. We employed the **LingerImageRegressor** configured with two neighbors to manage image regression nuances effectively. The MNIST images were reshaped to 28x28x1, treating digit labels as continuous regression targets. We implemented and compared two CNN models: a standard CNN and a regression-specific modified CNN. Both were trained on original and transformed MNIST data, respectively—the latter enhanced by the **LingerImageRegressor** to highlight differences between neighboring images. This

experiment primarily assessed the models' adaptability within a regression framework rather than focusing on achieving maximum accuracy, due to the dataset's inherent constraints for regression tasks.

Table 4.36: CNN Architectures for LingerImageRegressor and Standard Regressor on MNIST

Layer	LingerImageRegressor CNN	Standard Regressor CNN
Input Layer	28x28x1 images	
Conv2D	32 filters, 3x3, relu	32 filters, 3x3, relu
MaxPooling2D	2x2	2x2
Conv2D	64 filters, 3x3, relu	64 filters, 3x3, relu
MaxPooling2D	2x2	2x2
Conv2D	64 filters, 3x3, relu	128 filters, 3x3, relu
Flatten	Flatten output	
Dense	64, relu	128, relu
Output Layer	1 output, linear	1 output, linear

Results

Table 4.37: Regression Performance on MNIST Dataset Over Different Epochs

Model	Epochs	MSE at 10 Epochs	Epochs	MSE at 30 Epochs
LingerImageRegressor CNN	10	7.2907	30	7.3202
Standard Regressor CNN	10	0.1978	30	0.1632

Analysis

The comparative analysis between the `LingerImageRegressor` and the Standard Regressor CNN on the MNIST dataset reveals that the `LingerImageRegressor`, designed to exploit differences between data points, produced higher mean squared errors (MSE) of 7.2907 and 7.3202 across 10 and 30 epochs respectively. This indicates a less effective adaptation to this particular dataset. Conversely, the Standard Regressor demonstrated substantially lower MSEs of 0.1978 and 0.1632, showing superior adaptation and learning efficiency. These results underscore the critical role of dataset suitability in machine learning, highlighting the MNIST dataset's limitations for regression tasks due to its categorical nature. Future experiments using a dataset with genuine continuous variables are recommended to better evaluate the `LingerImageRegressor`'s performance, enhancing the validity of this proof of concept.

4.2.9 Comparative Analysis

This section synthesises the findings of our experiments. It assesses the blended learning approach and its performance overall.

Key Findings

- **Performance Improvement:** Blended learning methods generally showed an improvement in accuracy and robustness across most datasets compared to traditional model-based and instance-based learning.
- For example, in experiments with the Glass and Abalone datasets, blended approaches significantly outperformed KNN and standard neural networks in terms of accuracy and handling noisy data.
- **Model Interpretability:** The experiments consistently demonstrated that blended learning approaches enhance model interpretability for classification tasks

involving complex datasets compared to black-box models.

- **Efficiency and Scalability:** While blended models tend to require more computational resources initially, their adaptability to large datasets was evident, particularly in regression tasks with the House Prices and Abalone datasets
- **Innovative Architectures:** The exploration of novel architectures within the blended learning framework yielded promising results. Innovations such as case-based reasoning and feature difference techniques significantly enhanced model accuracy and interpretability, as shown in experiments 3 and 5. A specialised neural network for the differences model could enhance their performance even further, rather than reusing the trained neural networks from a separate model.
- **Practical Applications:** Insights from the experiments indicate significant potential for blended learning methods in critical domains such as healthcare and finance. These methods are particularly beneficial in predictive analytics, where interpretability and accuracy are crucial.

Chapter 5

Conclusions and Future Work

This chapter reflects on the contributions of our study, identifies its limitations, and outlines potential avenues for future research to extend the work’s applicability and improve efficiency across multiple domains.

5.1 Addressing Identified Limitations

Our study, while providing valuable insights, recognizes certain limitations in scope and methodology. Future efforts should focus on surpassing these through advanced algorithmic enhancements and the integration of novel data processing techniques, such as those that could be implemented through Scikit-learn. These improvements are expected to refine the learning process, thus enhancing the models’ overall performance and applicability.

5.2 Innovative Label Representation

A significant area for future research is the development of innovative methods for representing classification labels in ways that avoid implicit ordering. Traditional numeric and categorical encodings can inadvertently introduce bias or incorrect assumptions into the learning process, especially in models sensitive to feature scaling or order. This is true in the case of the `LingerClassifier`, which numerically encodes classes. Exploring label representations that maintain the distinctness of categories without implying ordinal relationships could enhance model accuracy and interpretability, particularly in complex classification tasks.

5.3 Broader Applicability

Exploring the applicability of our approach to broader machine learning challenges represents a promising direction for future research. Adapting our models to handle unstructured data or complex, high-dimensional datasets could significantly broaden their utility, making them suitable for a wider range of applications.

5.4 Integration with Emerging Technologies

The potential integration of our models with emerging technologies, such as deep learning and reinforcement learning, offers exciting possibilities. Such integration could not only provide new insights and enhance model capabilities but also facilitate the use of these models in real-time applications or embedded systems where interpretability and efficiency are paramount.

5.5 Scalability and Efficiency

Investigating the scalability and efficiency of our models, especially in contexts involving large-scale datasets or distributed computing environments, is crucial. Future research could focus on optimizations that reduce training times or adaptations that leverage parallel processing architectures to improve performance.

5.6 Cross-Domain Applications

Examining the effectiveness of our models across different domains or for various types of prediction tasks could uncover new applications and opportunities for impact. This exploration might involve interdisciplinary research collaborations to adapt and evaluate our models in diverse contexts. It would be interesting to perform long term studies to assess the performance of blended learning models in dynamic environments to validate their real world applications.

5.7 Exploring Other Categorical Similarity Measures

Investigating alternative measures for assessing the similarity of categorical features could refine the process of generating and applying adaptation rules, potentially enhancing the accuracy and applicability of the Ensemble of Adaptations for Classification (EAC). As seen in the paper [10]. This could redefine the measurement of similarity between instances in a dataset.

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