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Apply Stochastic Gradient Descent algorithm on a feed forward neural network for Iris Flower classification

Date of Performance:

Date of Submission:

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Department of Artificial Intelligence & Data Science

Aim: Apply Stochastic Gradient Descent algorithm on a feed forward neural network for Iris

Flower classification.

Objective: Ability to perform optimization technique on a feed forward neural network.

Theory:

Gradient Descent is an iterative optimization process that searches for an objective function's

optimum value (Minimum/Maximum). It is one of the most used methods for changing a

model's parameters in order to reduce a cost function in machine learning projects.

The primary goal of gradient descent is to identify the model parameters that provide the

maximum accuracy on both training and test datasets. In gradient descent, the gradient is a

vector pointing in the general direction of the function's steepest rise at a particular point. The

algorithm might gradually drop towards lower values of the function by moving in the opposite

direction of the gradient, until reaching the minimum of the function.

Types of Gradient Descent:

Typically, there are three types of Gradient Descent:

• Batch Gradient Descent

Stochastic Gradient Descent

• Mini-batch Gradient Descent

Stochastic Gradient Descent (SGD):

Stochastic Gradient Descent (SGD) is a variant of the Gradient Descent algorithm that is used

for optimizing machine learning models. It addresses the computational inefficiency of

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traditional Gradient Descent methods when dealing with large datasets in machine learning

projects.

In SGD, instead of using the entire dataset for each iteration, only a single random training

example (or a small batch) is selected to calculate the gradient and update the model

parameters. This random selection introduces randomness into the optimization process, hence

the term "stochastic" in stochastic Gradient Descent

The advantage of using SGD is its computational efficiency, especially when dealing with large

datasets. By using a single example or a small batch, the computational cost per iteration is

significantly reduced compared to traditional Gradient Descent methods that require processing

the entire dataset.

Stochastic Gradient Descent Algorithm

Initialization: Randomly initialize the parameters of the model.

Set Parameters: Determine the number of iterations and the learning rate (alpha) for updating

the parameters.

Stochastic Gradient Descent Loop: Repeat the following steps until the model converges or

reaches the maximum number of iterations:

a. Shuffle the training dataset to introduce randomness.

b. Iterate over each training example (or a small batch) in the shuffled order.

c. Compute the gradient of the cost function with respect to the model parameters using the

current training.

d. Update the model parameters by taking a step in the direction of the negative gradient, scaled

by the learning rate.

e. Evaluate the convergence criteria, such as the difference in the cost function between

iterations of the gradient.



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Return Optimized Parameters: Once the convergence criteria are met or the maximum number of iterations is reached, return the optimized model parameters.

In SGD, since only one sample from the dataset is chosen at random for each iteration, the path taken by the algorithm to reach the minima is usually noisier than your typical Gradient Descent algorithm. But that doesn't matter all that much because the path taken by the algorithm does not matter, as long as we reach the minimum and with a significantly shorter training time.

Conclusion:



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Code:

```
#importing the libraries
import numpy as np
import pandas as pd
import tensorflow as tf
from sklearn.model_selection import train_test_split
from sklearn.preprocessing import StandardScaler
from sklearn.datasets import load_iris
#Loading and preprocessing the data
iris = load_iris()
X = iris.data
y = iris.target
scaler = StandardScaler()
X_scaled = scaler.fit_transform(X)
X_train, X_test, y_train, y_test = train_test_split(X_scaled, y, test_size=0.2, random_state=42)
#Building the Neural Network Model
model = tf.keras.models.Sequential([
  tf.keras.layers.Input(shape=(4,)),
  tf.keras.layers.Dense(64, activation='relu'),
  tf.keras.layers.Dense(32, activation='relu'),
  tf.keras.layers.Dense(3, activation='softmax')
])
```



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#Compiling the Model

model.compile(optimizer='sgd', loss='sparse_categorical_crossentropy', metrics=['accuracy'])

#Training the Model

 $batch_size = 32$

epochs = 100

model.fit(X_train, y_train, batch_size=batch_size, epochs=epochs, validation_split=0.1)

#Evaluating the Model

test_loss, test_accuracy = model.evaluate(X_test, y_test)

print(f"Test Loss: {test_loss:.4f}, Test Accuracy: {test_accuracy:.4f}")

Output: