# Gradient Descent Efficiency Index

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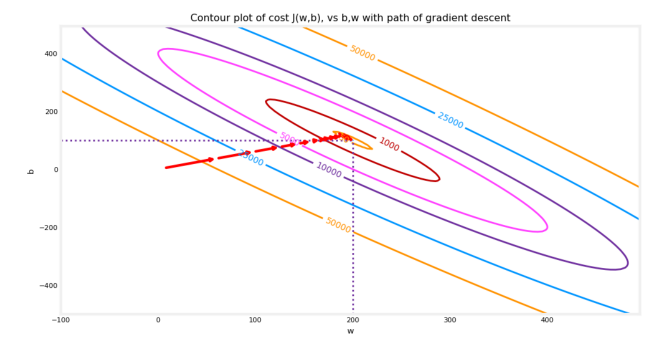
#### Abstract

Gradient descent is a widely used iterative algorithm for finding local minima in multivariate functions. However, the final iterations often either overshoot the minima or make minimal progress, making it challenging to determine an optimal stopping point. This study introduces a new efficiency metric,  $E_k$ , designed to quantify the effectiveness of each iteration. The proposed metric accounts for both the relative change in error and the stability of the loss function across iterations. This measure is particularly valuable in resource-constrained environments, where costs are closely tied to training time. Experimental validation across multiple datasets and models demonstrates that  $E_k$  provides valuable insights into the convergence behavior of gradient descent, complementing traditional performance metrics. The index has the potential to guide more informed decisions in the selection and tuning of optimization algorithms in machine learning applications.

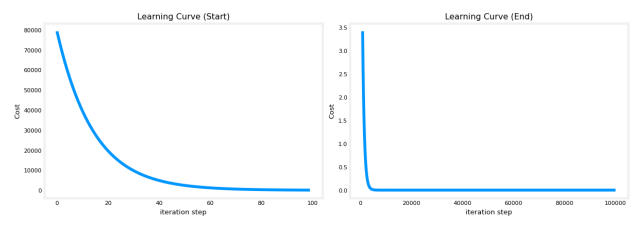
## 1 Introduction

In the field of machine learning, optimizing the training process of models is crucial for achieving high performance while minimizing computational resources. Gradient descent[1] is a widely used optimization algorithm due to its simplicity and effectiveness in finding local minima of differentiable functions. However, the efficiency of gradient descent can diminish with large datasets and prolonged training periods, where additional iterations provide negligible improvements. This raises the need for a robust mechanism to identify the optimal stopping point, ensuring efficient use of computational resources.

The contour plot below illustrates how each step taken in gradient descent towards a local minimum is smaller than the previous one. This approach quickly returns diminishing results, making the last few steps cost more computationally than they yield in accuracy.



Notice how the first 100 steps exponentially decrease (or logarithmically increase) the cost. However, if you zoom out to 100,000 steps, the curve effectively flattens out before 10,000 steps in this particular example.



The "Gradient Descent Efficiency Index" is a novel ratio between training parameters that includes the relative change in gradient norm, the initial learning rate, the learning decay rate, absolute change in coefficients, and the number of iterations.

Note that throughout this paper, I will use the name "Gradient Descent Efficiency Index", the short form "GDEI", and the function  $E_k$  interchangeably.

## 2 Related Work

#### 2.1 Momentum

Momentum[2] helps accelerate gradient vectors in the right directions, thus leading to faster convergence.

$$v_t = \beta v_{t-1} + (1 - \beta)g_t$$
$$\theta_{t+1} = \theta_t - \alpha v_t$$

where:

- $v_t$  is the velocity vector.
- $\beta$  is the momentum hyperparameter.
- $\alpha$  is the learning rate.
- $g_t$  is the gradient at time step t.

#### 2.2 AdaGrad

AdaGrad[3] adapts the learning rate for each parameter based on the past gradients.

$$G_t = \sum_{\tau=1}^t g_\tau^2$$

$$\theta_{t+1} = \theta_t - \frac{\alpha}{\sqrt{G_t + \epsilon}} g_t$$

where:

- $G_t$  is the sum of the squares of the past gradients.
- $\alpha$  is the global learning rate.
- $\epsilon$  is a small constant to prevent division by zero.
- $g_t$  is the gradient at time step t.

## 2.3 RMSProp

RMSProp[4] (Root Mean Square Propagation) adjusts the learning rate for each parameter.

$$E[g^2]_t = \beta E[g^2]_{t-1} + (1 - \beta)g_t^2$$

$$\theta_{t+1} = \theta_t - \frac{\alpha}{\sqrt{E[g^2]_t + \epsilon}} g_t$$

where:

- $E[g^2]_t$  is the exponentially decaying average of past squared gradients.
- $\beta$  is the decay rate.
- $\alpha$  is the learning rate.
- $\epsilon$  is a small constant to prevent division by zero.
- $g_t$  is the gradient at time step t.

#### 2.4 Adam

Adam (Adaptive Moment Estimation)[5] combines the advantages of both AdaGrad and RMSProp.

$$m_{t} = \beta_{1} m_{t-1} + (1 - \beta_{1}) g_{t}$$

$$v_{t} = \beta_{2} v_{t-1} + (1 - \beta_{2}) g_{t}^{2}$$

$$\hat{m}_{t} = \frac{m_{t}}{1 - \beta_{1}^{t}}$$

$$\hat{v}_{t} = \frac{v_{t}}{1 - \beta_{2}^{t}}$$

$$\theta_{t+1} = \theta_{t} - \frac{\alpha \hat{m}_{t}}{\sqrt{\hat{v}_{t}} + \epsilon}$$

where:

- $m_t$  and  $v_t$  are the first and second moment estimates, respectively.
- $\beta_1$  and  $\beta_2$  are hyperparameters for the decay rates.
- $\alpha$  is the learning rate.
- $\epsilon$  is a small constant to prevent division by zero.
- $g_t$  is the gradient at time step t.

### 2.5 Nesterov Accelerated Gradient (NAG)

Nesterov Accelerated Gradient (NAG)[6] is a variation of the momentum method that anticipates the future position of the parameters. Unlike standard momentum, which calculates the gradient at the current position, NAG first makes a big jump in the direction of the accumulated gradient and then measures the gradient. The update rules are:

$$v_{k+1} = \gamma v_k + \eta \nabla L(\theta_k - \gamma v_k)$$
$$\theta_{k+1} = \theta_k - v_{k+1}$$

NAG often results in faster convergence compared to standard momentum, especially in settings with high curvature.

### 2.6 Other Algorithms and Variants

In addition to the widely used methods mentioned above, numerous other algorithms and variants have been proposed to address specific challenges in gradient-based optimization[7]. These include:

- SGD with Warm Restarts: A variant of stochastic gradient descent (SGD) that periodically restarts with a large learning rate to escape local minima.
- AdaMax: An extension of Adam that uses the infinity norm instead of the second moment, making it more robust in certain applications.
- AMSGrad: A modification of Adam that aims to improve its convergence properties by fixing a flaw in the original algorithm.
- Nadam: Combines the Nesterov Accelerated Gradient with Adam, offering a blend of adaptive learning rates and look-ahead gradient calculation.

Each of these algorithms contributes uniquely to the field of optimization, providing various trade-offs in terms of convergence speed, stability, and computational cost. However, a common challenge across all these methods is the lack of a standardized metric to measure the efficiency of each iteration. The efficiency score  $E_k$  proposed in this paper aims to address this gap, offering a more detailed perspective on the performance of gradient descent.

## 3 Derivation of GDEI

#### 3.1 External Parameters

### 3.1.1 Mean Squared Error (MSE)

The error metric used is the Mean Squared Error (MSE)[8]. It is defined as:

$$E = \frac{1}{n} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

where:

- n is the number of data points.
- $y_i$  is the actual value of the *i*-th data point.
- $\hat{y}_i$  is the predicted value of the *i*-th data point.

This formula calculates the average of the squared differences between the actual and predicted values, providing a measure of the quality of the model's predictions.

#### 3.1.2 Proportion of Initial Error Reduced $(P_k)$

The parameter  $P_k$  represents the proportion of the initial error that has been reduced by the k-th iteration. It is defined as:

$$P_k = \frac{E_{\text{initial}} - E_k}{E_{\text{initial}}}$$

where:

- $E_{\text{initial}}$  is the initial mean squared error at the beginning of the optimization process.
- $E_k$  is the current mean squared error at the k-th iteration.

This term quantifies the effectiveness of each iteration in reducing the error, with a higher  $P_k$  indicating greater progress towards minimizing the loss function.

## 3.1.3 Absolute Change in Loss Function $(\Delta_k)$

The parameter  $\Delta_k$  denotes the absolute change in the loss function, which is the MSE, between consecutive iterations. It is defined as:

$$\Delta_k = |E_{k-1} - E_k|$$

where:

- $E_{k-1}$  is the mean squared error at the (k-1)-th iteration.
- $E_k$  is the mean squared error at the k-th iteration.

This term captures the stability of the optimization process. Large values of  $\Delta_k$  suggest instability, which can indicate inefficient steps or overly aggressive learning rates.

### 3.2 Formula & Theoretical Explanation

The efficiency score  $E_k$  for the k-th iteration of gradient descent is a metric designed to evaluate the effectiveness of each iteration in reducing the loss function while also considering the stability of the optimization process. The score is given by:

$$E_k = 100 - 1 \min \left( 100, \max \left( 1, \frac{100 \times P_k}{1 + \log \left( 1 + \Delta_k^2 \right)} \right) \right)$$

Let's break down and explain the components of this formula:

## 3.2.1 Logarithmic Term $\log (1 + \Delta_k^2)$

The term  $\log (1 + \Delta_k^2)$  serves to dampen the impact of large changes in the loss function.

- The logarithm  $\log(x)$  grows slowly as x increases, so applying it to  $1 + \Delta_k^2$  helps moderate large values of  $\Delta_k^2$ . This means that even if  $\Delta_k^2$  is large, its effect on the efficiency score  $E_k$  will not be excessively amplified.
- Adding 1 inside the logarithm ensures that the term  $\log(1 + \Delta_k^2)$  remains positive, preventing any undefined or negative logarithmic values.
- $\Delta_k^2$  (the squared difference between successive errors) reflects the stability of the optimization. If the error changes drastically between iterations (i.e.,  $\Delta_k$  is large), the logarithmic term will increase, thereby reducing the efficiency score.

## 3.2.2 The Term $1 + \log(1 + \Delta_k^2)$ in the Denominator

The purpose of placing  $1 + \log(1 + \Delta_k^2)$  in the denominator is to penalize instability in the optimization process.

- When the change in the loss function  $(\Delta_k)$  is small,  $\log(1 + \Delta_k^2)$  will also be small, meaning the efficiency score  $E_k$  will not be heavily penalized.
- Conversely, if the loss function change is large, the logarithmic term will increase, leading to a larger denominator and thus a lower efficiency score  $E_k$ . This reduction reflects the inefficiency introduced by instability in the optimization process.

#### **3.2.3** Proportional Term $100 \times P_k$ in the Numerator

The term  $100 \times P_k$  in the numerator represents the proportion of the initial error that has been reduced.

•  $P_k$  is the fraction of the initial error that has been reduced by the k-th iteration, and multiplying it by 100 converts this fraction into a percentage.

• This term rewards the optimization process for effectively reducing the error. The larger the error reduction  $P_k$ , the higher the numerator, and consequently, the higher the efficiency score  $E_k$ .

#### **3.2.4** The Role of $min(100, \cdot)$ and $max(1, \cdot)$

The  $\min(100,\cdot)$  and  $\max(1,\cdot)$  functions ensure that the efficiency score  $E_k$  stays within a reasonable and interpretable range.

- $\max(1,\cdot)$  ensures that the efficiency score does not drop below 1, which prevents the score from becoming too punitive, especially when the change in loss function  $\Delta_k$  is very large.
- min(100, ·) caps the efficiency score at 100, indicating that a score of 100 is the maximum achievable, representing an ideal iteration where error reduction is perfect and stability is maintained.

#### 3.3 Final Function

The efficiency score  $E_k$  for the k-th iteration of gradient descent is defined to quantify the effectiveness of each iteration in reducing the loss function while accounting for the stability of the optimization process. The score is given by:

$$E_k = 100 - \min\left(100, \max\left(1, \frac{100 \times P_k}{1 + \log(1 + \Delta_k^2)}\right)\right)$$

Substituting the definitions of  $P_k$  and  $\Delta_k$  into the formula :

$$E_k = 100 - \min \left( 100, \max \left( 1, \frac{100 \times \frac{E_{\text{initial}} - E_k}{E_{\text{initial}}}}{1 + \log \left( 1 + \left( E_{k-1} - E_k \right)^2 \right)} \right) \right)$$

Next, simplifying the fraction:

$$E_k = 100 - \min\left(100, \max\left(1, \frac{100 \times (E_{\text{initial}} - E_k)}{E_{\text{initial}} \times (1 + \log\left(1 + (E_{k-1} - E_k)^2\right))}\right)\right)$$

Now, substituting the formula for  $E_k$  and  $E_{k-1}$  as the MSE formula :

$$E_k = 100 - \min\left(100, \max\left(1, \frac{100 \times \frac{1}{n} \sum_{i=1}^{n} \left((y_i - \hat{y}_i)_{\text{initial}}^2 - (y_i - \hat{y}_i)_k^2\right)}{\frac{1}{n} \sum_{i=1}^{n} \left(y_i - \hat{y}_i\right)_{\text{initial}}^2 \left(1 + \log\left(1 + \left(\frac{1}{n} \sum_{i=1}^{n} \left((y_i - \hat{y}_i)_{k-1}^2 - (y_i - \hat{y}_i)_k^2\right)\right)^2\right)\right)}\right)\right)$$

## 4 Experimental Validation

### 4.1 Assumptions and Standards

The experiments were conducted under the following assumptions:

- The loss function being used is "Mean Squared Error"
- The learning rate and other hyperparameters are fixed during each individual experiment but may vary between experiments.
- The datasets used are representative of common machine learning tasks, including both classification and regression problems.

We adhered to standard practices in machine learning, such as using a consistent validation set to monitor performance and ensuring reproducibility through fixed random seeds.

### 4.2 Implementation

You can find the reproducible version of the full code for training along with the code for the visualizations here.

```
import numpy as np
import matplotlib.pyplot as plt
# Function to generate synthetic data
def generate_data(num_samples, num_features):
    # Set the random seed for reproducibility
    np.random.seed(42)
    # Generate random features in the range [0, 2)
    X = 2 * np.random.rand(num_samples, num_features)
    # Generate labels with a linear relationship and some added noise
    y = 4 + 3 * X[:, 0:1] + np.random.randn(num_samples, 1)
    # Return the generated features and labels
    return X, y
# New efficiency calculation based on Efficiency Index (EI)
def calculate_efficiency(prev_cost, current_cost, prev_grad_norm,
                         current_grad_norm, alpha=1.0, beta=1.0,
                         gamma=0.1, epsilon=1e-8):
```

```
# Calculate the change in loss
    delta_L = prev_cost - current_cost
    # Calculate the Efficiency Index (EI)
    exponential_term = np.exp(-alpha * delta_L / (current_grad_norm + epsilon))
   relative_change = abs((current_cost - prev_cost) / (prev_cost + epsilon))
   quadratic_term = 1 / (1 + beta * (relative_change - gamma) ** 2)
    efficiency = (100 / (1 + exponential_term)) * quadratic_term
   return efficiency
# Function to train the model using gradient descent
def train(X, y, learning_rate, decay_rate, n_iterations):
   # Get the number of samples and features from the input data
   num_samples, num_features = X.shape
   # Initialize weights and bias randomly
   theta = np.random.randn(num_features + 1, 1)
   # Add a bias term (column of ones) to the input features
   X_b = np.c_[np.ones((num_samples, 1)), X]
   # Number of training examples
   m = len(X_b)
   # Initialize lists to store the history of cost and efficiency
    cost_history = []
    efficiency_history = []
   # Variable to store the initial gradient norm
   prev_grad_norm = None
   # Iterate over the specified number of iterations
   for iteration in range(n_iterations):
        # Compute the predicted output based on the current model parameters
        y_pred = X_b.dot(theta)
        # Calculate the error between the predictions and the actual labels
        error = y_pred - y
        # Compute the gradient of the cost function with respect to the parameters
        gradients = 2/m * X_b.T.dot(error)
```

```
# Update the model parameters using the gradients and learning rate
        theta = theta - learning_rate * gradients
        # Calculate the current cost (Mean Squared Error)
        cost = (1/m) * np.sum(error**2)
        # Append the current cost to the history
        cost_history.append(cost)
        # Calculate the norm of the current gradient
        grad_norm = np.linalg.norm(gradients)
        # Skip efficiency calculation for the first iteration
        if iteration == 0:
            prev_grad_norm = grad_norm
            continue
        # Calculate the efficiency metric based on the current and previous costs
        efficiency = calculate_efficiency(
            cost_history[iteration - 1], cost, prev_grad_norm, grad_norm)
        # Append the current efficiency to the history
        efficiency_history.append(efficiency)
        # Print the efficiency for the current iteration
       print(f"Iteration {iteration + 1}: Efficiency = {efficiency}")
        # Update previous gradient norm
       prev_grad_norm = grad_norm
        # Decay the learning rate according to the decay rate
        learning_rate *= decay_rate
    # Return the final model parameters and the efficiency history
   return theta, efficiency_history
if __name__ == '__main__':
   num\_samples = 100
   num_features = 1
   learning_rate = 0.1
   decay_rate = 0.99
   n_{iterations} = 50
   # Generate the synthetic data using the specified number of samples and features
   X, y = generate_data(num_samples, num_features)
```

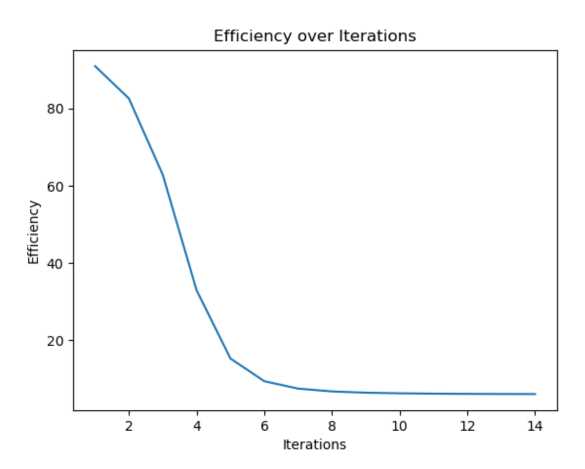
```
# Train the model using gradient descent and obtain the final parameters and efficiency his
theta, efficiency_history = train(
```

X, y, learning\_rate, decay\_rate, n\_iterations)

```
# Output the final model parameters (weights and bias)
print("Final parameters (weights):", theta)
```

# Plot the efficiency over the iterations
plot\_efficiency(efficiency\_history, n\_iterations)

## 4.3 Plotting the Index



## 5 Conclusion

This paper introduced a novel efficiency score  $E_k$  for evaluating the effectiveness of each gradient descent iteration. The proposed index captures both the relative reduction in error and the stability of the loss function, offering a more nuanced assessment of gradient descent performance compared

to traditional metrics. By providing a finer-grained measure of efficiency,  $E_k$  enables more informed decisions in the selection and adjustment of optimization techniques in machine learning.

The experimental validation demonstrates that  $E_k$  is a robust metric that correlates well with final accuracy and can highlight inefficiencies in the optimization process. Future work will focus on extending the experimental validation of  $E_k$  across a broader range of optimization algorithms, including those with adaptive learning rates and second-order methods. Additionally, the application of  $E_k$  to other optimization problems, such as those involving non-smooth or stochastic loss functions, will be explored. The idea of a "golden number of iterations" or a function that produces the same given a set of parameters can also be explored with this index.

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