Unsupervised-based Distributed Machine Learning for Efficient Data Clustering and Prediction

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January 26, 2024

Abstract

Unsupervised ML-based approaches have emerged for driving critical decisions about training data samples to help solve challenges in many life critical applications. This paper proposes parallel and distributed computing unsupervised ML techniques to improve the execution time of different ML algorithms. Various unsupervised ML models are developed, implemented, and tested to demonstrate the efficiency, in terms of execution time and accuracy, of the serial methods as compared to the parallelized ones. We developed sequential, parallel, and distributed cloud computing unsupervised ML models based and determined the most efficient model through comparative analysis. As a case study, sequential, parallel, and distributed approaches of Simple K-Means, Minibatch K-means, and Fuzzy C-Means are investigated to study the developed models’ efficiency using country datasets for multiple organizations to train and test the developed model. Parallel and distributed computing models are developed utilizing could computing architect, i.e., cloud Amazon SageMaker, to study their efficiency in the execution time and model accuracy. The results show that the proposed parallel and distributed Fuzzy C-Means outperforms the other two clustering methods in terms of execution time with 0.932ms and 0.623ms with a minimal impact on the accuracy of the developed models.
Abstract—Unsupervised ML-based approaches have emerged for driving critical decisions about training data samples to help solve challenges in many life critical applications. This paper proposes parallel and distributed computing unsupervised ML techniques to improve the execution time of different ML algorithms. Various unsupervised ML models are developed, implemented, and tested to demonstrate the efficiency, in terms of execution time and accuracy, of the serial methods as compared to the parallelized ones. We developed sequential, parallel, and distributed cloud computing unsupervised ML models based and determined the most efficient model through comparative analysis. As a case study, sequential, parallel, and distributed approaches of Simple K-Means, Minibatch K-means, and Fuzzy C-Means are investigated to study the developed models’ efficiency using country datasets for multiple organizations to train and test the developed model. Parallel and distributed computing models are developed utilizing cloud computing architect, i.e., cloud Amazon SageMaker, to study their efficiency in the execution time and model accuracy. The results show that the proposed parallel and distributed Fuzzy C-Means outperforms the other two clustering methods in terms of execution time with 0.932ms and 0.623ms with a minimal impact on the accuracy of the developed models.


I. INTRODUCTION

In machine learning, one of the primary goals is to extract valuable insights or patterns from large datasets. One widely adopted technique for grouping similar data is the data clustering algorithms. These algorithms evaluate the similarity among distinct objects within a dataset and subsequently group them based on these similarities. With the growing volume of data being amassed using scientific data collection techniques, it has become increasingly imperative to ensure that machine learning algorithms are effective, efficient, and scalable. One way to improve machine learning algorithms’ performance is to parallelize them, leveraging modern computing infrastructures. However, the efficacy of parallelization is heavily contingent upon the data organization and choice of parallelization approach.

Simple K-Means algorithm is one of the first and most commonly utilized techniques. There are a few issues related to the K-Means algorithm. They are as follows: one is the initial centers initialized randomly, which is a significant issue. The other issue is that there will need to be some idea of the clusters in the dataset. Minibatch K-Means is a clustering algorithm that splits a dataset into a fixed number of groups or clusters.

Instead of processing the entire dataset, Minibatch K-Means works on smaller, randomly chosen subsets of data known as mini-batches. This approach makes it more efficient and capable of handling larger datasets that may not fit into memory. The algorithm assigns each data point to the nearest centroid and updates centroid positions based on the mean of the assigned data points. While Minibatch K-Means is faster and more scalable, it may only sometimes produce optimal results compared to the standard K-Means algorithm, mainly when the mini-batch size is too small.

Fuzzy C-Means (FCM) is an effective clustering algorithm that groups data points into distinct clusters based on their similarities. This unique algorithm enables overlapping clusters by ascribing a fuzzy membership value to each point, signifying its membership degree to each cluster. FCM continually calculates and updates the cluster centers and membership values until a predefined stopping point is reached. Many scholars have concentrated on parallelizing these techniques based on these issues.

Nevertheless, all these algorithms have a few drawbacks. One such drawback is the utilization of parallel systems with constrained programming models that automatically parallelize data line by line. This approach assumes the need to store all data in memory line by line simultaneously, which becomes highly impractical when dealing with extensive datasets containing millions of records. Consequently, a trade-off between speed and efficiency is often observed, where prioritizing one aspect may come at the expense of the other. To achieve optimal performance, it is essential to strike a balance between efficiency and accuracy.

This paper aims to investigate multiple Parallel and distributing computing for efficient unsupervised clustering. We developed different ML models for countries with funding for necessities and assistance during disasters and environmental
harm. For that, we divided the nations into categories based on the economic and medical factors that will contribute to their overall development. Organizations can determine which country needs support with the aid of analysis.

The K-Means, minibatch K-Means, Fuzzy C-Means serial approach, open MPI parallel technique, and AWS distribution strategy were all used in the investigation. We tested the speed and accuracy of the algorithms using the three different ways to make the comparison, and we then presented the findings. We used Python to implement parallelized MPI-based algorithm on AWS for computing distribution from scratch. Similarly, Minibatch K-Means, and Fuzzy C-Means utilizing the AWS cloud platform. The results shows that parallel and distributed computing MPI approaches can enhance the performance and quality of the solution of the developed ML-based models.

The reset of this paper is organized as follows. Section 2 provides background information on machine learning algorithms, Message Passing Interface and Amazon SageMaker. Section 3 related work. Section 4 focuses on parallelization strategies, describing the methods and requirements necessary for parallelization. Section 5 details the research contributions to K-Means, MiniBatch K-Means, and Fuzzy C-Means, as well as the understanding of parallelization and cloud algorithms, and the experimental setup.

Section 6 covers the evaluation matrix. Section 7 presents the results, discussions, and conclusions regarding the performance of cloud and parallel clustering distribution techniques.
versus sequential clustering. Section 8 explores potential applications. Finally, the References are included.

II. BACKGROUND

This section will cover six major topics: Machine Learning models, K-Means, Minibatch K-means, Fuzzy C-Means, Parallel Message passing interface and AWS Sage maker

A. Machine Learning

Machine learning is a subclass of artificial intelligence which allows computer systems to improve performance by learning from experiences without human instruction. It entails using large datasets to train algorithms capable of detecting patterns, making predictions, and enhancing decision-making abilities. Machine learning is classified into three main categories: supervised learning, unsupervised learning, and reinforcement learning, each having its unique techniques and areas of application.

Supervised learning deals with labeled data wherein the algorithm learns to recognize patterns, relationships, and correlations between inputs and outputs to produce accurate predictions or classifications. Unsupervised learning, on the other hand, works with unlabeled data, which means that the algorithm must identify and extract insights from the data by itself without any predefined labels. Reinforcement learning, often used in robotics and game development, relies on trial-and-error to learn the best approach to solve a particular task.

As machine learning continues to evolve, it has found applications across several industries, including healthcare, finance, retail, and transportation. It has enabled organizations to analyze diverse data sources, make informed decisions, and increase operational efficiencies. With the growing availability of data and processing power, the potential for machine learning is enormous, and its impact on various industries is expected to be transformative and here i am presenting basic model of ML Figure 1.

B. K-Means Algorithm

These grouping issues in machine learning or information science are resolved using unsupervised learning K-Means Clustering. In this chapter, we will study the K-means clustering method, how something operates, and how to apply it in Python.
Unsupervised training method K-Means Cluster divides a piece of unlabeled information into various clusters. Thus, K describes how several clusters that were not previously present should be generated due to the procedure; for instance, if K=2, the outcome will primarily be two types. If K=3, the outcome will be three clusters, and so on.

This iterative method separates the unlabeled information into k distinct clusters, each containing one dataset and sharing a set of features. It allows us both to categorize all data into different groups. It offers a workable technique for quickly and effectively determining the groups within the unlabeled dataset without having to do any training.

Every cluster has a corresponding centroid in this centroid-based approach. The principal objective of such an approach is to reduce the distance between the data points and the groups they belong.

This method starts with an intake of an unlabeled dataset, separates this into k clusters, and continues the procedure until no better clusters are found. In this method, the k value must be known in advance. Its two main functions of the k-means suggested method are:

- Determines the optimal quantity of K points or centroids via an iterative algorithm.
- Each piece of information is paired with the closest k-center. Those information fragments near a certain k-center combine to form a cluster.

Each cluster is unique and has samples that only share a small amount in a joint.

The following stages illustrate how the K-Means method functions:

- To figure out how many groups, choose K.
- Pick K locations or cluster centers randomly. (It could not be the supplied data.)
- Allocate every piece of data to its nearest centroid, which will create precisely the number of clusters that have been predetermined.
- Determine the variation, then relocate every cluster’s centroid.
- Reassign every piece of data to an updated centroid of every cluster by repeating the third section.
- Move to Phase 4 if there is a reassign; otherwise, go over to Finished.
- A finished design.

Now let us analyze the graphical plots to comprehend the steps mentioned earlier:

Below points is the explanation of graphs each point representing each graph formation Figure 2.

- Consider that there are two independent variables, P1 and P2.
- To identify the data set and divide it into various clusters, let us use a value of k for clusters or K=2. This implies that we will divide those data sets into two separate groups. To create a cluster, k points or the centroid must be chosen randomly. These coordinates may be taken from the dataset or in different locations. Since these two do not appear in the dataset, it was choosing them as the k points in this instance.

- Currently, we would identify the nearest K-point and centroid for every data point on the scatter plot. To find the distance separating two points, we will estimate it using the math we studied. To create a midpoint between both, the two centroids shall do such.

- It is evident from the previous picture that locations along the line to the left were near a K1 center, which is red, and white spots along the lines to the right are near a red centroid. For better comprehension, let us color both red and green. We will repeat the procedure by picking a new centroid to locate the nearest cluster. Let us determine those centroids centers of gravity ordered to select their fresh centroids.

- We would then reallocate every piece of data here to a new centroid. You can conduct the same procedure for locating a mean line in this, and your average should look such as the illustration of 5 one in above figure. There is one red point on the left end of the line, plus two green points on the lines. Thus, fresh centroids will be assigned to such three locations.

- Since reassign has occurred, we once more go to step 4, locating fresh centroids called K-points. Determining the exact gravitational centers of a new centroid would allow us to continue the operation.

- We will construct the average lines once more and reassign all pieces of data because we have the updated cluster centers.

- This is established due to the need for more consistent data points along both sides of the boundary.

- Now that our model is complete, we may exclude the two remaining groups and the presumptive centroids.

1) Elbow Method: The Elbow Method is widely used for determining the ideal number of clusters. A WCSS-valued idea is applied in this technique. The term "total variations inside a cluster" is abbreviated as "WCSS" and means Within Cluster Sum of Squares. This formula listed below is used to determine the number for WCSS (for 2 clusters):

The WCSS equation is:

$$WCSS = \sum_{j=1}^{k} \sum_{P_i \in C_j} (distance(P_i, C_j))^2$$  \hspace{1cm} (1)

From the formula,

In this equation, k represents the number of clusters. Cj represents the j-th cluster, and Pi represents the i-th point in the dataset.

The distance function in the equation calculates the distance between a data point Pi and the centroid of its assigned cluster Cj. The WCSS metric is important because it measures how well the data points are clustered together. The lower the WCSS value, the better the clustering solution.

The goal of the K-means clustering algorithm is to minimize the WCSS by iteratively adjusting the cluster centroids until convergence is achieved. The final result of the K-means algorithm is a set of k cluster centroids that best represent the data in the dataset.

Its Elbows technique is so named because the diagram depicts a steep bend that resembles an elbow. A graph for
the elbows approach resembles what illustration below Figure 3

C. Minibatch K-Means

Minibatch K-means is a widely used clustering algorithm that divides a given dataset into K clusters by minimizing the distances between the data points and their associated cluster centroid. In contrast to the standard K-means algorithm, Minibatch K-means processes the data’s small, randomly selected subsets, or Minibatches, to expedite convergence while maintaining comparable clustering accuracy. It is considered efficient and scalable for large datasets that exceed the available memory capacity. The algorithm assigns each data point to its closest cluster center and updates the centroids iteratively until convergence is achieved. The number of Minibatches and the batch size are hyperparameters that can be optimized to enhance the clustering performance and accuracy of the resulting partitions.

D. Fuzzy C-Means

Fuzzy C-means is a prominent unsupervised clustering approach used to group data points into distinct clusters based on their similarity. Unlike complex clustering methods such as K-means, Fuzzy C-means assigns each data point a degree of membership, represented as a fuzzy value, for every cluster. This allows a data point to be part of multiple clusters at the same time with different membership degrees. The algorithm adapts the cluster centers and their fuzzy memberships iteratively based on the data point similarity and the level of overlap among the clusters. The hyperparameter 'fuzziness' regulates the degree of blurriness of the memberships, with higher values leading to more fuzziness and an increased number of clusters. Fuzzy C-means can be utilized to address various clustering problems and provide soft classification outcomes for the data points.

E. Parallel Message Passing Interface

A standard way to communicate throughout many computers that run concurrent programs all over distributed storage is indeed the message-passing interface (MPI).

The term "node" refers to a group of processors, or even a group of core processors on a single computer, in parallel Computing. Generally, every node inside a parallel configuration focuses on a distinct aspect of an immense processing challenge. It thus becomes difficult to coordinate the activities of every simultaneous network, transfer data across nodes, or exert control over the entire parallel cluster. The message-forwarding interface defines a standardized set of commands for such activities. When a message is sent to an entity, parallel process, subroutine, function, and thread, it is usually called "passing a message." That message is again utilized to begin a different technique.

Although no formal reference implementation, such as the International Organization for Standardization or the IEEE Institute of Electrical and Electronics Engineers, had supported MPI as a benchmark, it is widely regarded as an industry norm. It serves as the base for most network systems used by parallel computing programming. Researchers also produced several MPI versions.

Fortran, C, C++, Python and Java procedures or libraries have usable syntax defined by MPI.

1) Features of the Message Passing Interface that are advantageous:

- Standardization: Previous message queue packages have been supplanted by MPI, which is now a widely recognized mainstream technology.
- A large group formulated it: Despite MPI is not a nationalized, a committee of manufacturers, integrators, and users came up with it.

2) Flexibility: Since MPI has already been developed for numerous distributed memory designs, users can move their software to other systems which the MPI standard supports without changing the source code.

- Efficiency: Usually, functionality is tailored to MPI’s device, and vendor implementations could be tuned to take advantage of built-in hardware characteristics.
- Feature set: Maximum performance on parallel processing systems and groups is a crucial feature of MPI. Furthermore, over 100 specified routines make up the fundamental MPI-1 architecture.

3) Terminology used by MPI: Key terms and instructions: A few fundamental MPI ideas or instructions are listed below Figure 4:

- Comm: This MPI communication classes link several stages. An enclosed consultation processes an individual identifier from a communication instruction, which arranges it into an orderly topology. For instance, MPI COMM WORLD is a signal for global communication.
- Color: This means that a process is assigned a color, and all activities with that coloring are housed within the same communication. MPE Make color array is a color-related instruction that modifies the offering of the product.
- Key: A key determines the ranking or ordering of a procedure in communication. The order is established by the application’s position inside the communication if two methods are assigned the same key.
Fig. 4. The rank assigned to every CPU is demonstrated using an MPI COMM process with numerous endpoints distributed across four cluster.

- New comm: Fresh communication can be created using this instruction.
- Categories of data collected: A description of the kind of data exchanged among tasks is required for MPI functions. These variables can be predefined with the help of MPI INT, MPI CHAR, or MPI DOUBLE.
- Refer: It communicates among two particular operations. Two popular delaying techniques for juncture messaging were MPI Send and MPI Recv. Block is the method by which the transmitting and receiving systems hold off on transmitting and finishing a communication until a whole signal has been appropriately transmitted and received.
- Cooperative foundations: All activities in a particular process must communicate to perform these group tasks. However, one method is MPI Bcast, which distributes information from a single node to each process in a process.
- One-sided: Usually, when this phrase refers to external communications like MPI Put, MPI Get, or MPI Accumulate. Publishing to storage, receiving to recollection, or minimizing operations on a single recollection between activities are specially mentioned.

4) MPI’s Past and Iterations: Around 1991, a tiny research group from Austria started talking about either a message-carrying interface. A Center for Studies in Parallel Computing supported a workshop about message-passing protocols inside a distributed memory system that occurred in Williamsburg, Virginia, per year afterward. Therefore To develop the standard method, a work was formed.

The prototype of MPI-1 was developed in Nov 1992, and the specification was published in 1993 just at Supercomputing ’93 conference. Around 1994, MPI version 1.0 was published after receiving more comments and modifications. Since then, MPI has been open to everyone working in the powerful computational field, with more than 40 companies currently involved.

About 115 capabilities are offered under the more dated MPI 1.3 specification, sometimes known as MPI-1, and more than 500 features and a high degree of older systems using MPI-1, a subsequent MPI 2.2 specification, or MPI-2, are available. Unfortunately, a complete MPI-2 version is not offered by all MPI implementations. Along with variable system integration and remote storage functions, MPI-2 introduced novel concurrent I/O. The Nov. introduction of the MPI-3 specification results in increased speed, multicore or cluster capabilities, and improved scalability, with availability and high interoperability. MPI 4.0 was published by The MPI Forum around 2021. It featured innovative features like Permanent Communes, segmented networking, and a new product layout. Now, MPI 5.0 is being created.

F. AWS Sage Maker

1) Overview: Most data analysts use the hosted platform to develop, train, and publish machine learning algorithms, and they were regrettably still unable to change resources as required. To go live quicker and with less expense, AWS Sage Maker makes it simpler for programmers to design and retrain models. This essay will review the capabilities, use applications, and advantages of AWS Sage Maker and Computer Vision using AWS Sage Maker.

2) AWS: A cloud-based system that offers services that are ordered over the web is known as Amazon Web Services. Every public cloud type may be developed, monitored, and
deployed using AWS services, which is where AWS Sage Maker is useful.

3) About AWS Sage Maker: Amazon manages the machine learning services offered as Sage Maker. Sage Maker enables data researchers or programmers to create machine learning models quickly, train those, and afterward instantly release them into a virtual machine that is prepared for use in operation. It usually includes methods for machine learning designed for application in a distributed scenario using extensive data sets. Sage Maker provides flexible, dispersed education options that fit existing operations Figure 5.

4) Attributes of Sage Maker: Sage Maker now features new features that Amazon has developed since its debut in 2017. Accessibility to functionality is made possible via AWS Contributory factor Studios, an Integrated Development Environment that combines all abilities. One of two methods can be utilized to build a Jupiter notebook:

- Sage Maker Studio’s internet version of the IDE.
- A machine learning server in Amazon Sage Maker that Amazon EC2 hosts.

5) Utilizing AWS Sage Maker and machine learning: Let us examine the development, testing, fine-tuning, then deployment of a model for machine learning utilizing AWS Sage Maker.

6) Builds:

- There are approximately 15 popular learning algorithm skills and making included.
- It enables us to select the configuration settings needed for the notebook instances.
- To start coding, a user may use a notebook example (for building model training tasks).
- Choose and enhance the necessary methods, such as: K-Means, Linear Regression and Logic-based Regression.
- AWS Sage Maker’s Jupiter notebook API allows programmers to alter Supervised Learning servers.

7) Tune and Test:

- All required modules should be made or imported.
- During Amazon model development, establish and control just a few configuration settings.
- The model is taught and tuned using Sage Maker.
- Sage Maker achieves a set of parameter tuning by combining some algorithm parameters.
- Sage Maker uses Amazon S3 for keeping records as it is a secure protocol.
- Sage Maker uses extensible ECR to control Container technology.
- Docker setup, management, and storage are made more accessible with ECR.
- The developed skills source is preserved in ECR, whereas the supervised learning is separated and stored in Amazon S3.
- Next, Sage Maker generates, learns, and stores a given input cluster in Amazon S3.

8) Implement:

- Upon adjusting, objects could be released to Sage Maker destinations.
- Just in the end, a forecast is made in real-time.
- To determine whether the strategy has achieved your company goals, one must evaluate it.

9) Uses of Sage Maker: Many different businesses use AWS Sage Maker. Machine learning groups use Sage Maker for the below task:

- Accessing and exchanging codes.
- Accelerate the creation of production-ready AI components.
- Enhance data interpretations and build more exact database schemas iteratively.
- Streamline data both in and out.
- Much information is to be processed.
- Codes for modeling exchange.
10) Advantages:
- We are making deep learning easier to use.
- More people will be able to innovate using machine learning thanks to integrated tools like computer scientists and also no potential input for analysts.
- Huge data preparation.
- Collect, categorize, and analyze vast quantities of organized (tabular) and unorganized (photos, videos, and audio) information via computer vision.
- Accelerate the advancement of computer vision.
- Without improved facilities, learning can be completed in just a few moments instead of days.
- Using specially designed tools, employees to comply might rise as much as ten times.
- Enhance the machine training cycle to be more efficient.
- Streamline or unify MLOps practices throughout a company to design, train, publish, and maintain algorithms at volume.

11) Modern Developments: Since the initial release of reinvent 2021, Amazon has added a feature for its Sage Maker Python SDK that offers abstracts to speed up models’ deployments, plus Model Registry, which makes it easier to connect virtualized inferences destinations and MLOps processes. One can utilize Sage Maker Cloud Hosting Deduction with high-traffic applications now that Amazon has increased the provisions requests per endpoints limitation from 50 to 200.

12) Conclusion: AWS charges every sage Maker user for Computing, memory, or information computer resources used to create, test, deploy, and log machine learning algorithms or predictions. The expenditures of S3 were related to maintaining train and prediction data sets. The sage Maker application’s design is flexible and responsive, enabling the whole lifespan of machine-learning applications through modeling creation to model implementation. This implies that the sage Maker can be used independently for the proposed model, retraining, or distribution.

III. RELATED WORK

The parallel K-Means technique, according to Mohanavalli, Jaisakthi and Aravindan, Both a distributed memory system utilizing MPI programming and a memory version utilizing OpenMP programming has a parallel K-Means model developed based. With MPI programming, a hybrid OpenMP implementation was also tested. To calculate Amdahl’s effect to evaluate the speedups achieved and the efficiency of the parallel algorithms were examined. In addition to being 50 percentage faster than MPI, the hybrid technique performed better under a balanced load [1]. They have demonstrated that this approach is added to increase the effectiveness of parallel K-Means. Both the Enslaver/Slave concept and the parallel processing method are used. The tests revealed that this technique is more effective and applicable to Yufang Zhang’s situation[2]. Showed this is suggested to use an initiation technique for K-Means parallel processing, establish the first cluster centers, that not only speeds up performance but also produces consistent results conducted by Swamy, Raghuwanshi and Gholghate[3].

Boukedhir, Lachiheb and Gouider’s research focuses on increasing K-Means’ capacity to deal with large datasets by speeding up its execution, suggesting a map Reduce-based approach. In addition we will suggest two additional algorithms. The first eliminates unnecessary, and the other dynamically chooses the starting cluster center to stabilize the outcome. The suggested technique is demonstrated to be significantly quicker than three different known algorithms from the literature when implemented just on the Hadoop framework[4]. Applied Parallel MapReduce-based K-Means clustering approach because it is a straightforward yet effective method for parallel programming. These results of experiments indicate that the suggested algorithm can process massive datasets on affordable equipment while scaling up and down effectively research evaluated by Weizhong, Huifang and Qing [5].

Work targeted is clicked Plus, OpenMP and CUDA just on CPU and GPU, respectively, are used to improve its modified version for the K-Means cluster. Various data, including photos of varying data quantities, are shown, including the findings, with such a focus on rather big data. Covered are various combinations of characteristics and clusters[6]. So, to speed up K-Means, the study presents two novel methods for transmitting data. Expanded Vector is the initial method (KMMR-EV). K-Means over MapReduce utilizing Boundaries File is indeed the name of the second technique (KMMR-BF). Compared to the simple MapReduce implementation of K-Means, both methods achieve speed up had undergone considerable experimental investigation using actual and artificial data, coupled with an excess assessment to demonstrate the efficacy between both methods[7].

One goal of the researcher would be to create a K-Means method version that can run on a standard PC using NVIDIA graphics cards or tackle more extensive data sets using CUDA[8]. It concentrates upon cluster method performance problems, and a more sophisticated initial centroids core approach is used. The topic of a single CPU computer’s operational limit while dealing with large data sets also is investigated, and a parallel K-Means technique is investigated. The analysis shows that such advancements can significantly increase performance, allowing for the group of many data sets less precisely and fast analysis shown by Tian, Zhu, Zhang, and Liu[9].

Using GPU’s cluster, we demonstrate the conception and execution of an effective parallel K-Means algorithm. They dynamically use load balancing to evenly spread demand across the many GPUs inside the cluster to improve the overall cross-performance of the parallel K-Means. Additionally, they utilize software distributed shared memory to facilitate inter-node cooperation and communication. By keeping load balance on GPU clusters, the evaluation’s findings demonstrate the parallelism K-Means’ enhanced results.[10].

The latest update of K-Means separates overall issues into minor issues handled separately with one or more GPUs’ Broadcast Multiprocessors. They had developed the Graphics card K-Means (CUDA) authors[11]. It creates a buffer among collected data that did not alter their clusters over the following group, which might dramatically lower the burden of large data
sets. Its operation time was reduced by up to 70 Percentage inside the prototype system[12].

On such a CPU-GPU heterogeneous system, they provide a quick implementation of the spectral clustering technique. My method uses the multicore CPU’s processing capability and the GPU’s huge multithreading and SIMD abilities. They suggest a concurrent technique to generate a sparse refers to a network expressed in a common sparse codified form; the data points are given as input in high-dimension space. Next, using the ARPACK program, the CU SPARSE library’s backward transceiver k is ordinarily quite large—where the smallest k eigenvectors were computed by three of the Channel matrix. Additionally, we use GPU’s to create a rapid parallel processing K-Means method, and the solution is much quicker[13]. A technique known as the enhanced K-Means method was used to expand that standard K-Means strategy. Our experimental findings demonstrate that such an expanded K-Means algorithm may effectively organize the data when the number of components for each grouping has to be constrained. This method changes the number of attributes in every category by utilizing a greedy strategy by faliu and inkyu[14].

This genetic algorithm is utilized to optimize K-Means grouping to ensure that the shortcomings of K-Means can be overcome. K-Means using the GA algorithm suggested innovative products in this field of research when the outcomes of conventional K-Means clustering versus genetics K-Means grouping were contrasted and analyzed[15].

The first is MPI, and the second is shared memory utilizing OpenMP, or heterogeneous Computing utilizing CUDA-C-programmed NVIDIA GPUs. The first investigation contrasts the various strategies, while others have worked at speeding K-Means grouping. Where effectiveness of K-Means also has highly dependent on the first means used. We compare many thermally activated in serial and select the most effective one to use throughout the method. From modest (300300 pixels) to huge (11641200 pixels), I test the results on various photos. Our findings demonstrate that almost all three distributed programming paradigms resulted in speedups, with OpenMP used for small pictures and CUDA-C for bigger pictures yielding the best results by Janki, Miriam, and ningfang[16].provided a hybrid algorithm that decides when to employ the triangle inequality in an improved way. Numerous studies show that the approach performs better than the Computer or Multicore Kmeans[17].

The examination of share price pricing data from the prior yrs., with findings interpreted following intensive training using an algorithm for machine learning on CUDA, keeping in mind the time restrictions of trading. With the aid of machine learning approaches, the program’s performance has been significantly increased. In this research, a massively parallel methodology is applied to hasten the generation of findings. Compared to typical ways of employing a solitary Processor Core Unit, the execution time is significantly decreased due to the recent high achievements of CUDA parallel computing technology (CPU). By accurately anticipating stock prices in advance, it reduced computation time by a significant margin and resulted in net profits, which is the end objective of trade. Just characterized by three groups as projected utilizing algorithms, traders could choose to maintain the share, sell it, acquire new equities, or stay impartial. If a user makes a neutral choice, that indicates he must hold onto any stock he already has and refrain from purchasing any more. This suggested approach is appropriate for trading stocks based on stock price[18].

Its efficiency of a program in a concurrent environment was enhanced by adding a K-Means clustering algorithm to an MPI4py module. This report examines the effectiveness of executing the K-Means method consecutively versus using a Message Passing Interface (MPI) parallel design for grouping information in the form of operational costs and processing time[19]. This study presented a parallelization K-Means method for sparse, elevated text (PKHT). This suggested technique provides an 11x shorter runtime by utilizing both GPUs with MPI by xiaolei, yanking, and yuxin[20].

Maximum performance, flexibility, and portability are all characteristics of MPI as a message-passing type system. This led to the motivation for this research to present MKmeans, a similar K-Means clustering technique with MPI. The approach allows efficient use of the clustering algorithm inside a parallel setting. Research using experimental statistics shows that MKmeans operates well on a variety of large amounts of data with very few time resources[21]. Researchers investigate leveraging NVIDIA Graphics Processing Units (GPUs) written with CUDA C to enhance the speed of K-Means clustering. Various optimization methods are being used, including using constant memory for cluster data, shared memory, or picture data. Its outcomes were assessed on various images, ranging in size from tiny (256x256 pixels) to large (1024x1024 pixels) and with 4 to 256 clusters. For four clusters, we observe that now the serial version is typically 9x slower than the parallel version. Even as the number of clusters rises to 256, the speedup jumps to 57x[22].

We suggested a parallel implementation of the conventional K-Means algorithm to run it on the Hadoop distributed framework. The research findings show that when clustering a vast volume of data, our suggested K-Means method works better than conventional K-Means[23]. In terms of overhead expenses and execution, the overall efficiency of K-Means grouping the information is examined among parallel and sequential implementation in the Message Passing Interface Infrastructure by Ragunthar, Ashok, and Gopinath[24].

The study utilizes NVIDIA’s Compute Unified Device Architecture (CUDA) to create the GPU-based Harmony K-Means Algorithm in content clustering. During this testing, compared with CPU-based software, the GPU-based application could achieve a speedup of increase than Twenty times[25]. An approach suggested in this research is based upon the modified K-Means algorithm of a Hadoop platform; initially, a preliminary clustering is obtained using the canopy algorithm, after which the pinpoint of the exact or cluster numbers is also adapted new using the ISODATA algorithm. Lastly, integrate the MapReduce distributed computation architecture with the K-Means method to achieve the ideally planned CF-K-Means technique to improve the convergence point list and K calculated value by the Canopy method. The experiments
show that now the CI-K-Means technique resolves the issues associated with the Canopy K-means algorithm’s inconsistent result in a more excellent and also the complexity of choosing the Centre for the K-Means approach. Its correctness, overall speedup proportion, and grouping efficiency have already been vastly enhanced compared to both approaches used before improvement[26].

They examine two examples of calculating dense similarity matrices to cluster using vast data sets. By sparsifying a vector, contrast one technique using the Nyström methodology. Next, decide to sparsify the matrices while keeping their closest neighbors, plus we look into their parallelization. On dispersed computers, we parallelize both memory usage and computations. We demonstrate the practicality of our parallel approach by conducting an empirical investigation on two colossal data sets: a large-scale photo dataset with 2,121,863 examples and a document data set with 193,844 examples[27].

The subsequent work has developed a probabilistic theory for grouping using a random point scientific process. This approach perfectly mimics Bayes decision theory regarding classification: Bayes clustering procedures available with the least predicted errors provided known underlying processes and a defined objective function. As a result, clustering becomes an ongoing method rather than a subjective one. In this study, we start to comprehend such an algorithm by presenting the circumstances where the given opportunity used in traditional K-Means clustering becomes efficient in the novel Bayes clustering concept[28].

The description of methodological improvements may allow for high computation savings in average squared data clustering. A parallel statistics program P-CLUSTER, which runs on a desktop system, now includes additional upgrades. Unsupervised categorization of photos with a familiar texture was the subject of investigations. A 96 decreasing trend in the calculation was made for some data sources[29].

The K-Means technique is suggested in this study as a single special instruction multiple data (SIMD) architecture processors (GPUs) driven algorithm. Both objects and the reconfiguration of k-centroids are assigned to the GPU concurrently inside this approach to minimize the computationally costly parts of the entire unit. With the most current development of GPUs with computational integrated system architectures, they have built such Graphics K-Means algorithm (CUDA). The numerical tests showed that the efficiency of GPU-based K-Means might be up to 40 times quicker than that of CPU-based K-Means[30]. Furthermore, for this research, we embarked on an in-depth exploration of many existing studies including [32], [33], [34], [35], [36], [37], employing a comprehensive approach that encompassed [research methodologies/tools].

IV. PARALLELIZATION STRATEGIES

The parallel method for data mining uses a parallel architecture. Moreover, for implementation purpose parallel algorithm is used. A standard parallel clustering procedure includes at least the three phases listed. The first phase is Partition. What happens is like during this phase, we will divide the information into smaller datasets. The second phase is Computing. Here we run the clustering method on each processor’s local dataset. Each processor’s clustering method may be different.

The third phase is Integration. To get the overall result, we have merged all the information, which are clusters obtained from each processor. There are two data techniques which are as follows. Finding reasons for the information set’s highly changeable components, or locating and explaining outliers, is the aim in several applications. Other applications aim to comprehend the variances of the overwhelming bulk of the given dataset components without any concern for the extremes. The first type of data mining is primarily used in science, while the second type is business applications.

In first-class applications, it appears that Computing is necessary. Because we are still determining how beneficial sampling from a vast dataset will be in implementations of the latter type. Thus, parallel Computing has a big future as a platform for data processing. However, it has yet to be apparent if this is the direction data mining will take. We can understand how the technique is already employed by looking at the applications in which the most extensive data centers are used for data gathering.

The parallelization of data involves a variety of techniques. The first method is data parallelism, and the aim is to distribute the data appropriately and divide it into processors, each of which computes the given data in parallel. Data parallelisms are classified into two types: record-based and attribute-based.

The second method is the parallelization of tasks. Each record’s likelihood of performing a data mining strategy is the same when redistributed, making equally distributing information achievable. Data partitioning based on records is commonly used in real-world parallel data mining applications. If the characteristic Partition is incorrect, the information connection breaks, and the processing quality decreases.

The second strategy is Task Parallel. It breaks the entire solution method into many sub-procedures and sends them to separate processors. It also includes two ways for achieving task parallelism. One uses a divide-and-conquer technique, assigning the work and the subtask to a designated processor. The other is based on a scheduler, which dynamically assigns data to available processors.

The execution method involves utilizing many CPUs to perform the same job and several CPUs to complete a separate one. The one picked is determined based on the area we are implementing and the data structure. Regardless of the parallel method, the load-balancing problem must be addressed. If data is misallocated, the burden may be unequal.

The third alternative is a hybrid; they need help resolving the issue. Upon properly splitting the data and distributing it among each processing unit, jobs are assigned to a designated processor in this approach, according to the data properties assigned to each processing unit.

After properly breaking down information and transmitting data to each processing, tasks are assigned to a designated processor by the attributes assigned to each processing unit. The data generated by the chain is combined when each job executing on each processor is finished. Task and data parallel techniques are used in this situation, and data interchange occurs before and after job execution, which is advantageous.
Algorithm 1: K-Means Algorithm

1: \( K \) is cluster
2: \( N \) is data objects
3: clusters\([K]\) cluster centers in array
4: objects\([N]\) data objects in array
5: membership\([N]\) objects memberships in array
6: Set threshold \( \Theta \)
7: repeat
8: \( G \) ← 0
9: for \( a = 0 \) to \( N - 1 \) do
10:   for \( b = 0 \) to \( K - 1 \) do
11:     distances ← objects\([a]\) − clusters\([b]\)
12:     if distances < \( d_{\text{min}} \) then
13:       \( d_{\text{min}} \) ← distances
14:       \( n \) ← \( b \)
15:     end if
16:   end for
17:   if membership\([a]\) ≠ \( n \) then
18:     \( G \) ← \( G + 1 \)
19:     membership\([a]\) ← \( n \)
20: end if
21: newcluster\([n]\) ← newcluster\([n]\) + objects\([a]\)
22: newclustersize\([n]\) ← newclustersize\([n]\) + 1
23: end for
24: for \( b = 0 \) to \( K - 1 \) do
25:   clusters\([b]\)[*] ← newcluster\([b]\)[*]/newclustersize\([b]\)
26:   newcluster\([b]\)[*] ← 0
27:   newclustersize\([b]\) ← 0
28: end for
29: until \( G/N \leq \Theta \)

In addition to minimizing transmission costs, the job is not interrupted.

V. Methodology

This section discusses a study focused on improving the quality of the information in a country data set using K-Means, Minibatch K-Means, and Fuzzy C-Means methods. The study uses parallelization techniques to speed up the execution times of all methods and obtain more accurate results. With the help of the elbow method, the optimal number of clusters is perceived using this method.

The study discusses different techniques for parallelizing: sequential K-Means, Minibatch K-Means, and fuzzy C-Means, parallelization using MPI (Message Passing Interface), and a cloud-based method. These techniques are implemented from scratch in Python, a popular programming language for machine learning and data processing. The results of this study may be helpful for organizations interested in aiding impoverished countries, as it provides more accurate information about the country data set.

A. Sequential Process of K-Means

Calculating centroids and cluster formation is done progressively with sequential K-means. Here is a detailed guide and breakdown of how K-Means works.

The code is a Python implementation of the K-Means clustering technique. The objective of the method is to divide a collection of \( N \) objects into \( K \) clusters according to how identical they are, where \( K \) is the number of clusters Algorithm 1.

The K-Means algorithm uses this code to cluster data. It initializes the cluster centers and places data objects into clusters. Then, it refines the cluster centers and data object membership iteratively until reaching a threshold value that terminates the algorithm. To manage data, it uses variables to handle cluster centers, data objects, and membership information. Cluster centers are an array of size \( K \), while data objects and membership data are arrays of size \( N \).

The code computes the distances between objects and each cluster center during each iteration, assigning objects to the nearest cluster based on these calculations. If the object is assigned to a new cluster, a variable counter increments to reflect changes in its membership data. The new cluster center updates by summing its data object values and dividing by the cluster size.

Finally, the algorithm checks if the termination threshold is met. If not, it updates the cluster centers and repeats the process.
Algorithm 2 Minibatch K-Means

1: Begin
2: Initialize: $k$, mini-batch size $b$, iterations $k$, data collection $X$
3: For $j \leftarrow 1$ to $k$ do
4: $a \leftarrow 0$
5: $M \leftarrow \{x \in X\}$
6: $d[x] \leftarrow f(C, x)$
7: $s \leftarrow d[x]$ for $x \in M$
8: $a[s] \leftarrow a[s] + 1$
9: $n \leftarrow 1/a[s]$
10: $s \leftarrow (1 - n)s + nx$
11: end for
12: end for
13: Stop

B. Minibatch K-Means

Here is an explanation of the code, step by step Algorithm 2:

For each point $x$ in dataset $X$, a centroid $s$ is initialized randomly from the set of possible centroids $C$. The algorithm then runs for $k$ iterations, each with the following steps.

- A mini-batch $M$ of $b$ examples is randomly sampled from the dataset $X$.
- For each example $x$ in the mini-batch $M$, the distance between $x$ and each centroid in $C$ is computed using the function $f$.
- For each example $x$ in the mini-batch $M$, the closest centroid $s$ is identified based on the computed distances, and the count of examples assigned to that centroid $a[s]$ is incremented.
- For each example $x$ in the mini-batch $M$, the position of the assigned centroid $s$ is updated using a weighted average of its previous position and the new example $x$. The weight given to the previous position is determined by the number of examples already assigned to that centroid, $s \leftarrow (1-n)s+nx$, where $n = 1/a[s]$.
- At the end of $k$ iterations, the final positions of the centroids are returned.

The main idea behind this code is to iteratively improve the positions of the centroids by adjusting their position based on the examples in each mini-batch. By randomly sampling the examples and updating the centroids incrementally, the algorithm can scale to large datasets while producing high-quality clustering results.

C. Fuzzy C-Means

Here is an explanation of the code, step by step Algorithm 3:

To perform Fuzzy C-Means clustering, we first initialize the membership matrix, $U$, by assigning random values to it or by setting it to a predefined initial value.

- At each iteration, denoted by the $k$-step, we calculate the center vectors of the clusters, representing the centroids of the clusters. The center vectors are calculated using the membership matrix $U$ from the previous iteration.
- The center vectors are calculated using the membership matrix $U$ from the previous iteration. Specifically, we calculate the center vectors, $C_i$, by taking a weighted average of the data points, $x_j$, according to their membership values, $u_{ij}$.
- A higher membership value for a data point implies a more prominent weight for that data point. Once we have calculated the center vectors, we update the membership matrix for the next iteration by recalculating the membership values, $u_{ij}$, for each data point, $x_i$, based on their distance from the center vectors, $C_j$.
- The new membership value, $u_{ij}$, depends on the sum of the distances from the data point $x_i$ to all the center vectors, $C_j$, raised to the power of $2/(m-1)$, where $m$ is a fuzziness parameter that controls the level of fuzziness or overlapping among clusters.
- We then check if the difference between the membership matrices at the $k$-step and the $(k+1)$-step is lower than a certain threshold, $\epsilon$. If the difference is smaller than $\epsilon$, the algorithm terminates and returns the final membership matrix and center vectors.
- Otherwise, the algorithm returns to step 2 to recalculate the center vectors and update the membership matrix for the next iteration. The algorithm repeats steps 2 to 7 until convergence, which implies that the difference between the membership matrices at two consecutive iterations is less than or equal to $\epsilon$.

D. Parallel Techniques using Message Passing Interface

MPI is one of the best methods for implementing parallel computing. The apparent reason for that is because of the standard libraries it has. The motivation is to aim for exactness and efficacy. With the help of open MPI, we can parallelize the K-Means, Minibatch K-Means, and Fuzzy C-Means. MPI is supported in many languages. In this, the datasets were divided equally among many of the MPI processes, and Using OpenMP instructions, data labeling for each local data within the process was carried out.
Each attribute has its dimension in the result and is created in the space based on the number of attributes. The result generated using this methodology will be assimilated with the basic version of the technology to determine whether the approach is superior and how much performance measurements, such as time for completion, are generated. This method was evaluated on the same dataset, termed "Country Data Set." The information we will validate is based on the testing dataset, and we broke that complete information into train and test for training and testing the model. Here we used eighty for training and twenty for testing, i.e., 80:20.

To implement this, we use Jupiter Notebook. Where the "divide and combine" scheme for parallel K-Means, Minibatch K-Means, and Fuzzy C-Means clustering using the Message Passing Interface (MPI) library, the process can be divided into several steps. Firstly, the data points should be divided into smaller subsets that different processes can independently process. Each process would then compute the centroid of a subset of data points using all algorithms until it converges to a stable value. After that, the partial centroids obtained from each process must be combined to obtain the final cluster of centroids. This can be achieved using the MPI reduce() function, which combines the partial centroids to ensure that all processes receive the same result Figure 7.

Finally, the results should be verified by comparing the final centroids with the expected values or by measuring the clustering quality using a performance metric such as the Sum of Squared Errors (SSE). The divide and combine scheme in parallel clustering using MPI provides an efficient way to process large datasets by dividing them into smaller subsets and processing them in parallel, resulting in faster execution times and increased scalability.

1) Steps involved in Parallel Techniques using MPI: The algorithm starts by choosing the number of clusters, k, and assigning k cluster centers to the nodes. It is crucial to remember that the clusters must be distributed evenly among the nodes, i.e., k/n, where n is the total number of nodes.

After assigning the cluster centers, each node actively computes the mean centers of the k/n groups, disseminates the centers to all nodes using MPI (Message Passing Interface), receives the cores from other nodes, computes the distance between each data point and its nearest collection. When the cluster assignments stop changing, it actively achieves convergence. Finally, the process is actively repeated until reaching convergence.

A crucial element of the clustering algorithms is convergence. It ensures the algorithms terminate once the cluster assignments have stabilized and no more adjustments are required. This enhances the clustering process’ effectiveness and precision. In conclusion, a clustering algorithm is a valuable tool for grouping data into groups, and convergence is essential to the algorithm’s precision and effectiveness.

2) MPI methodology consists of several stages:
- We first determine the centroid based on the number of nodes.
- At this point, the master component sends data chunks based on centroid to various nodes. These broadcasts are...
made possible by comm = MPI.COMM_WORLD. And communication bias.

- The means of each node is used to calculate centroids, which are then broadcast to other nodes.
- Data points are dispersed using communication scatter. This process is repeated until convergence occurs.

**E. Amazon SageMaker**

Amazon SageMaker is a cloud-based machine learning platform that enables developers to build, train, and deploy machine learning models. It offers a range of tools for data labeling, training, experimentation, model deployment, and monitoring. Amazon SageMaker uses a pay-per-use pricing model, which makes it more cost effective for small-scale projects and large-scale enterprises.

1) **Working of parallel clustering techniques using MPI in AWS:** Parallel clustering techniques using MPI in AWS SageMaker work by distributing the clustering algorithms across multiple compute nodes using the Message Passing Interface (MPI) protocol.

The input data is divided into various subsets. Each subset is given its own compute node when a training job is started in SageMaker using the clustering algorithms and MPI. Then, using MPI, the clustering algorithms are applied parallel to each subset to process the data across numerous nodes.

During the parallel processing, each computed node calculates its own local centroids and assigns data points to the nearest centroid. Then, the intermediate results are merged to obtain the final clustering solution.

The results are typically stored in an S3 bucket, which can be accessed by other AWS services or downloaded for further analysis. Using SageMaker for parallel clustering with MPI allows for the efficient processing of large datasets and significantly reduces the time required for clustering tasks.

Overall, parallel clustering using MPI in AWS SageMaker enables clustering in the context of distributed computing of big datasets and provides a scalable and efficient way to process data using multiple compute nodes.

A managed service machine learning (ML) Amazon’s Elastic Compute Cloud (Amazon EC2) notebook example, which continues to run Jupiter Software, is an Amazon SageMaker notebook example. This same notebook example is used to build and manage Jupiter notebooks for information extraction, training, and deploying machine learning models.

To begin the analysis, an Exploratory Data Analysis (EDA) of the chosen datasets will be conducted, followed by the training and creation of a model using AWS Sagemaker’s built-in version of clustering. The resulting recommendation engine can then be deployed using an AWS Sagemaker inference endpoint and integrated into an application via an API. These steps were performed within an AWS Sagemaker-hosted Jupyter Notebook instance running on an ml.t2.xlarge instance.

2) **Steps to generate a SageMaker notebook example:**

- Go to https://console.aws.amazon.com/sagemaker/ to access an Amazon SageMaker console.
- Select Notebook cases, then start creating notebook examples.
- I will provide additional data just on the Generate Note-
F. Implementation

1) Data Collection: We utilized Python programming language and machine learning techniques to implement algorithms on the KAGGLE country dataset, which consists of nine key factors. From those nine key factors, we use only four key factors to predict what organizations can use to determine financial assistance to nations. These nine attributes include Child Mortality, Exports, Health, Imports, Income, GDP, Inflation, Life Expectancy, and Total Fertility.

To perform the analysis, we used an Apple M1 processor with a memory of 16GB. Our primary objective was to use these algorithms to identify countries that may require financial aid based on their performance in these four factors.

2) Data Pre-Processing: First, the code sorts the DataFrame data by the values in the gdpp column in ascending order using the sort values function and then resets the index of the sorted DataFrame using the reset index function with the argument drop=True. This creates a new DataFrame with the same data as data but with the rows sorted by GDP per capital in ascending order and a new index.

Then, the code creates a scatter plot using the plot scatter function with the x values as the index of the sorted DataFrame and the y values as the gdpp column of the sorted DataFrame. The plot xlabel, plot ylabel, and plot title functions are used to add axis labels and a title to the plot, and plot show is used to display the plot.

Next, the code computes the quartiles of the 'gdpp' column using the quantile() function with the argument [0, 0.33, 0.67, 1] to get the lower quartile (0-33), middle quartile (33-67), and upper quartile (67-100) values. After that, the code defines the category labels as ['Low', 'Medium', 'High'].

Then, the code adds a new column to the DataFrame called GDP Category using the pd cut function with the arguments bins=quantiles and labels=categories. This function creates categories for the gdpp values based on the quantiles computed earlier and assigns the category label to each gdpp value in the new GDP Category column.

Finally, the code drops the columns health, life expec, total fer, and 'child mort' from the DataFrame using the drop function with the arguments columns, inplace=True, and axis=1. This removes the specified columns from the DataFrame, without creating a new DataFrame.

In summary, this code performs data cleaning and visualization tasks on a DataFrame that contains information about various countries. It sorts the DataFrame by GDP per capital, creates a scatter plot of the GDP per capital values, categorizes the GDP per capital values into low, medium, and high categories based on quartiles, and drops some columns from the DataFrame that are not needed for further analysis.

3) Data Transformation: Since data transformation entails transforming raw data into a similar and standardized style, it is essential to clustering. Data transformation’s primary goal is to reduce the influence of different scales, ranges, and skewness in the variables because these factors can significantly affect the results of clustering. Normalization, standardization, PCA transformation, and log transformation are a few examples of the data transformation methods frequently used in clustering. The best data conversion technique must be
chosen depending on the issue domain and the data’s nature. If the appropriate data transformation methods are applied, clustering may yield precise and insightful results.

4) Implementation of Clustering Algorithms:

1) Sequential K-Means: The code provided reads a processed data file containing information about different countries. It initializes an empty list called sse to store the sum of squared errors (SSE) for different values of k. Then it iterates through the range of k values from 2 to 6. A K-Means object is initialized with the current value of k, and the algorithm is fitted to the dataset. The SSE for the fitted model is then appended to the sse list. The next block of code selects the input column for the K-Means technique and plots the SSE for different values of k. The plot shows that the optimal number of clusters is likely 4, as the SSE begins to level off at that point.

The code initializes a K-Means object with k=4 and fits it to the dataset. The appropriate prediction method is then used to predict the cluster assignments for each data point. These predicted cluster assignments are then added as a new column to the DataFrame. Finally, the code prints out the centroid coordinates for each cluster.

In summary, the K-Means technique clusters similar data points into k clusters. The code provided reads a file of processed data, calculates the SSE for different values of k, and selects the optimal number of clusters based on the plot. It then predicts cluster assignments for each data point and adds them to the DataFrame. Finally, it prints the centroid coordinates for each cluster.

2) Sequential Minibatch K-Means: This code performs clustering on a dataset using the Mini Batch K-Means algorithm. The dataset is loaded from a CSV file, and the columns for clustering are specified. The range of k values to try is defined, and an empty list is initialized to store the sum of squared errors (sse).

A loop is then executed over each k value in the defined range. A Mini Batch K-Means model is initialized with the current k value and fitted to the dataset in each iteration. The inertia attribute of the model is then appended to the sse list. This indicates how well the data is being clustered, with a lower sum of squared errors indicating a better fit.

Once the loop completes, the results are plotted as a line graph with k on the x-axis and the sum of squared errors on the y-axis. This plot can be used to identify the optimal number of clusters. This code sets the number of clusters to 4, and the model is fitted to the dataset using the prediction method.

The predicted clusters and the original dataset are then plotted as a scatter plot, with each point colored according to its cluster assignment. The cluster centers are also plotted as red circles. The resulting plot shows how the data has been partitioned into distinct clusters based on the specified features.

3) Sequential Fuzzy C-Means: This code performs fuzzy c-means clustering on a dataset of country data. First, the data is loaded from a CSV file, and non-numeric columns are removed. The remaining numeric columns are then normalized using StandardScaler.

The number of clusters is set to 4, and the fuzzy exponent parameter (m) and the maximum number of iterations are also defined. The fuzzy c-means clustering algorithm is then initialized with the normalized data, number of clusters, m, and maxiter values.

The algorithm is run, and the predicted cluster membership for each data point is obtained by taking the argmax of the u matrix. This assigns each data point to a cluster based on the degree of membership in each cluster. The predicted clusters are added to the original dataframe, and the number of countries in each cluster is printed to the console. This indicates how well the data has been clustered and can help with further analysis or decision-making. Fuzzy c-means clustering is functional when data points can belong to multiple clusters simultaneously, allowing for more nuanced clustering than traditional k-means clustering.

4) Parallel K-Means Minibatch K-Means and Fuzzy C-Means Using MPI: This code is an implementation of the K-means clustering algorithm using MPI (Message Passing Interface) for parallelization. The code is divided into two parts: the master part (rank 0) and the worker part (other ranks).

In the master part, the code reads a CSV file containing data and initializes the number of clusters (num clusters) and other necessary variables. The data is divided into chunks, and each chunk is sent to a worker using MPI’s scatter operation. The initial centroids are also broadcasted to all workers.

In the worker part, each worker receives its chunk of data and calculates the distance between each data point and the centroids. It assigns each data point to the cluster with the closest centroid. It then counts the number of samples in each cluster and sends the count to the master using MPI’s gather operation. The worker also receives the updated centroids from the master using MPI’s all reduce operation.

This process of calculating distances, updating clusters, and centroids is repeated until the centroids no longer change. The final cluster assignments from all workers are gathered at the master, and the adjusted Rand score is calculated to compare the results with the labels obtained from running KMeans from scikit-learn (kmeans).

Finally, the code visualizes the clusters and centroids using matplotlib. It also performs some analysis, such as printing the frequency of clusters for different GDP categories, replacing cluster labels to match the original labels, computing a confusion matrix, and printing classification metrics.

Overall, this code demonstrates parallelizing the K-means algorithm using MPI and compares the results with scikit-learn’s KMeans implementation. It also provides visualizations and analysis of the clustering results. We will replace the KMeans function with MiniBatchKMeans and FuzzyCMeans and run the same code.

5) Parallel K-Means, Minibatch K-Means, and Fuzzy C-
Means using MPI in Cloud: As discussed above, we use the same code to run in AWS SageMaker, which employs parallel processing with MPI and all algorithms to cluster data.

5) Pattern Information: In Clustering, data points are grouped according to how alike they are using pattern information. The method finds patterns in the data by repeatedly assigning each data point to the closest centroid and adjusting the centroid by averaging all the data points given to it. Up until the centroids stop moving or the maximum number of repetitions has been achieved, this procedure is continued. The ensuing clusters show collections of data points with comparable patterns or traits. By comparing new data points to existing clusters, these clusters may be utilized to predict future data points and understand the underlying patterns within the data.

6) Comparison of various Techniques: We will compare all three methods of execution times, including in terms of execution speed, accuracy, precision, recall, and f1 slope and support.

G. Setup of Exploration

We tested and implemented all algorithms using Jupyter Notebook on my computer, and we also utilized Amazon SageMaker Jupiter Notebook for cloud services. We used a computer with an Apple M1 processor and 16 GB of RAM to run our algorithms during testing. To supplement our methodology, we used a KAGGLE dataset that contained 167 rows and ten columns, totaling 13.2 kb.

We utilized machine learning techniques and various Python libraries to implement our methodology. Our approach involved thorough data analysis and preprocessing, then implementing and testing various algorithms on the cleansed data. We then analyzed the results to determine the best approach for accurately predicting financial aid requirements for different nations.

VI. RESULT AND CONCLUSION

A. Elbow Method

The Elbow method is a practical approach for determining the optimal no of clusters in a model. Applying this method makes it possible to identify the most appropriate number of distinct clusters for a given dataset Figure 11.

We can create a new model with the optimal number of clusters identified, which is 4, and then examine the cluster assignments for each observation. Essentially, we would assign each data point to one of the 4 clusters established on their characteristics or similarities. This process can provide valuable insights into patterns and relationships within the data, potentially aiding in making informed decisions or predictions and the graphs below.

The structure of the clusters is displayed in the graph, and it includes the following algorithms Fig. 15 16 17 18 19 20 21 22 23

2. Parallel K-Means using MPI.

5. Parallel Minibatch K-Means using MPI.
7. Sequential Fuzzy C-Means.
8. Parallel Fuzzy C-Means using MPI.

The confusion matrix is a tool that helps to summarize the performance of a model by displaying the number of correct and incorrect predictions made for each class in the dataset. It consists of three columns and three rows, corresponding to the three classes in our data. The true values are represented on the y-axis while the predicted values are represented on the x-axis. Using the values obtained from the confusion matrix, precision, recall and F1 scores can be calculated for the model.

The confusion matrices for three different clustering algorithms; Sequential K-Means Figure 12, Sequential Minibatch K-Means Figure 13 and Sequential Fuzzy C-Means Figure 14.

| Sequential Evaluation Matrix of Sequential Clustering |
|--------------------------|----------------|-------|-------|-------|
| K-Means                  | 0.45           | 0.68  | 0.66  | 0.67  |
| Minibatch K-Means        | 0.48           | 0.75  | 0.72  | 0.71  |
| Fuzzy C-Means            | 0.46           | 0.69  | 0.65  | 0.63  |
The table shows that Minibatch K-Means performs the best out of the three algorithms in all four metrics. It has higher sequential accuracy, precision, recall, and F1-score than K-Means and Fuzzy C-Means. This indicates that Minibatch K-Means is the most effective algorithm in accurately grouping the data points into clusters Table I. Also, we presented a graphical comparison representation of Figure 24.

<table>
<thead>
<tr>
<th>Techniques</th>
<th>Execution Time (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequential K-Means</td>
<td>2.853</td>
</tr>
<tr>
<td>Parallel K-Means using MPI</td>
<td>1.050</td>
</tr>
<tr>
<td>Parallel K-Means using Cloud</td>
<td>0.828</td>
</tr>
<tr>
<td>Sequential Minibatch K-Means</td>
<td>2.310</td>
</tr>
<tr>
<td>Parallel Minibatch K-Means using MPI</td>
<td>1.215</td>
</tr>
<tr>
<td>Parallel Minibatch K-Means using Cloud</td>
<td>0.921</td>
</tr>
<tr>
<td>Sequential Fuzzy C-Means</td>
<td>1.285</td>
</tr>
<tr>
<td>Parallel Fuzzy C-Means using MPI</td>
<td>0.932</td>
</tr>
<tr>
<td>Parallel Fuzzy C-Means using Cloud</td>
<td>0.623</td>
</tr>
</tbody>
</table>

Overall Considering these findings, cloud computing platforms like AWS Sage Maker have the potential to offer substantial advantages for data processing in parallelized clusters. Also, you can see the graphical representation of Figure 27.

B. Discussion

The independent features are imports, exports, health, and inflation. Based on these four features, clusters are formed. We determine the model accuracy using the GDP feature from the dataset and the clusters. The GDP values are categorized as low, medium, and high. Based on the GDP category, we determine whether a country requires financial assistance or

<table>
<thead>
<tr>
<th>Techniques</th>
<th>Execution Time (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequential K-Means</td>
<td>2.853</td>
</tr>
<tr>
<td>Parallel K-Means using MPI</td>
<td>1.050</td>
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<tr>
<td>Parallel K-Means using Cloud</td>
<td>0.828</td>
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<tr>
<td>Sequential Minibatch K-Means</td>
<td>2.310</td>
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<td>1.215</td>
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<td>Parallel Minibatch K-Means using Cloud</td>
<td>0.921</td>
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<tr>
<td>Sequential Fuzzy C-Means</td>
<td>1.285</td>
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<tr>
<td>Parallel Fuzzy C-Means using MPI</td>
<td>0.932</td>
</tr>
<tr>
<td>Parallel Fuzzy C-Means using Cloud</td>
<td>0.623</td>
</tr>
</tbody>
</table>

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B. Discussion

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not. If the GDP is categorized as low, it signifies that the country needs help. For medium GDP, the country may or may not require assistance, and for high GDP, the country does not require help.

From Figure 28, the green color indicates help is not required, the red color indicates help need, and the yellow might need help and shows the countries in need of assistance and those that may require assistance. As a result, an NGO or organizations can quickly analyze data and assist countries.

The utilization of parallel clustering techniques results in numerous advantages over sequential clustering approaches, including enhanced efficiency, flexibility, scalability, fault tolerance, resource utilization, and real-time capabilities. These benefits make parallel clustering techniques the preferred option for dealing with large-scale datasets and time-sensitive clustering tasks.

The managed and user-friendly environment provided by Amazon SageMaker is ideal for developing and deploying machine learning models that include clustering algorithms. It offers scalability, managed infrastructure, integration with AWS services, flexibility, monitoring capabilities, and cost-effectiveness advantages that make it a leading choice over manual setup and management of parallel clustering techniques that utilize MPI.

C. Conclusion

Machine learning (ML) is a field devoted to understanding and building methods that let machines leverage data to enhance the performance and accuracy of different applications. A subset of machine learning is closely related to data clustering, which focuses on grouping data samples into various groups for data prediction, detection, identification, etc.

Unsupervised ML algorithms are the most effective methods for data clustering. This work investigates various approaches and identifies factors influencing the algorithm’s efficiency. Among them are the effects of the starting cluster on accuracy and the way the data is parallelized on efficiency. Finding the best approach based on the available data takes time and effort. We concentrated on country data and analyzed it using various parallel approaches and cloud computing. The analysis is based on four different country factors, and this analysis will assist various organizations that are prepared to assist the country.

The results demonstrate that implementing these models on a cloud platform significantly improves their performance in terms of execution time. By evaluating accuracy scores and confusion matrix values, we were able to compare the effectiveness of the three models. The findings indicated that when utilizing the sequential process, Minibatch K-Means achieves better performance compared to K-Means and Fuzzy C-Means. However, in cases where the parallel process was used, K-Means produced better results than the other two models.

D. Future Work

Our future work will concentrate on analyzing data using MapReduce and Multiprocessing, as well as developing an application that receives data as input and recommends the most efficient method for speedy execution. We aim to enhance the accuracy, precision, recall, and F1-score of our results. Additionally, we will experiment with different clustering algorithms for cluster initialization and construct an algorithm to assess the dataset and determine the most appropriate parallelization technique.

REFERENCES

Fig. 28. Help needed per country (World).


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