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Master Spark Basics with ease



Master Spark Basics – One Post to Rule Them All!

Over the past 30 days, we've covered some of the **most essential Spark concepts** for any data engineer or big data enthusiast! Here's your **ultimate recap** to solidify your knowledge and shine in your next project or interview:

Big Data Essentials

1. **Big Data Introduction** – Why do we need big data solutions?
2. **Monolithic vs Distributed Systems** – Transition from single-node to scalable systems.
3. **Designing Big Data Systems** – Blueprint for handling massive datasets.

Hadoop Basics

4. **Introduction to Hadoop** – Spark's foundational predecessor.
5. **Hadoop Architecture** – Understand the heart of HDFS and MapReduce.
6. **How MapReduce Works** – Parallel data processing simplified.
7. **MapReduce vs Spark** – A game-changer in performance and ease of use.
8. **Core Concepts (Parts 1 & 2)** – Building blocks of distributed computing.
9. **Architecture & Execution Flow** – Dive into Spark's inner workings.
10. **DAG & Scheduler** – How Spark optimizes and executes tasks.

Advanced Insights

11. **Modes of Deployment** – Client vs Cluster modes.
13. **Memory Management** – Efficient resource utilization.
14. **Partitioning vs Bucketing** – Organize your data for efficiency.
15. **Shuffling** – The secret behind performance tuning.
16. **Lazy Evaluation** – The power of deferred execution.

RDD Mastery

17. **RDD Basics** – Spark's distributed dataset explained.
18. **Real-World RDD Examples** – Get hands-on with Spark's core API.
19. **Coalesce vs Repartition** – Data partition management simplified.
20. **Sort vs SortByKey** – Optimized sorting methods.

Functions in Action

21. **Take vs Collect** – How to retrieve just what you need.
22. **ReduceByKey vs GroupByKey** – Efficient aggregations compared.
23. **Collect, Collect_List, Collect_Set** – Transform grouped data like a pro!

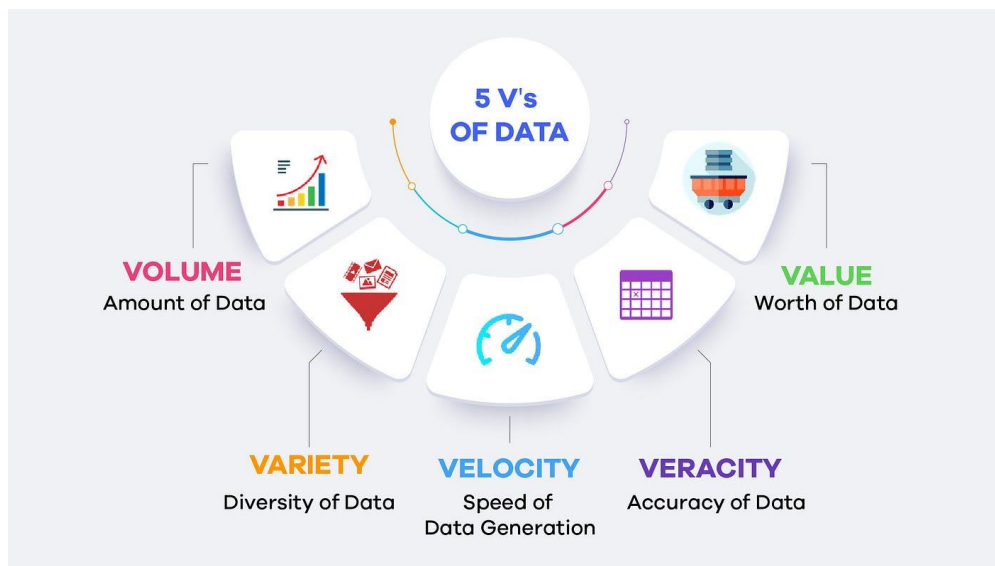


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Big Data - Definition

Big Data refers to datasets that are so large and complex that traditional data processing systems are unable to store, manage, or analyse them efficiently. These datasets are characterized by the 5V's, which include:



5V's of Big Data

1. Volume

- Massive amounts of data generated from various sources that exceed the storage capacity of conventional systems.
- Example: Social media data, sensor data in petabytes.

2. Variety

- Different formats and types of data, requiring different approaches for storage and processing:
 - Structured: Predefined schema (e.g., databases).
 - Semi-Structured: Partial schema (e.g., JSON, CSV).
 - Unstructured: No defined schema (e.g., images, videos).



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3. Velocity

- The speed at which new data is generated and needs to be processed in real-time or near real-time.
- Example: Streaming data during online sales events.

4. Veracity

- The reliability and quality of data. Systems must handle inaccuracies, inconsistencies, and incomplete data. i.e

Is the quality or correctness of Data. Data cannot always be in the right format and we should be able to handle this not so high quality data.

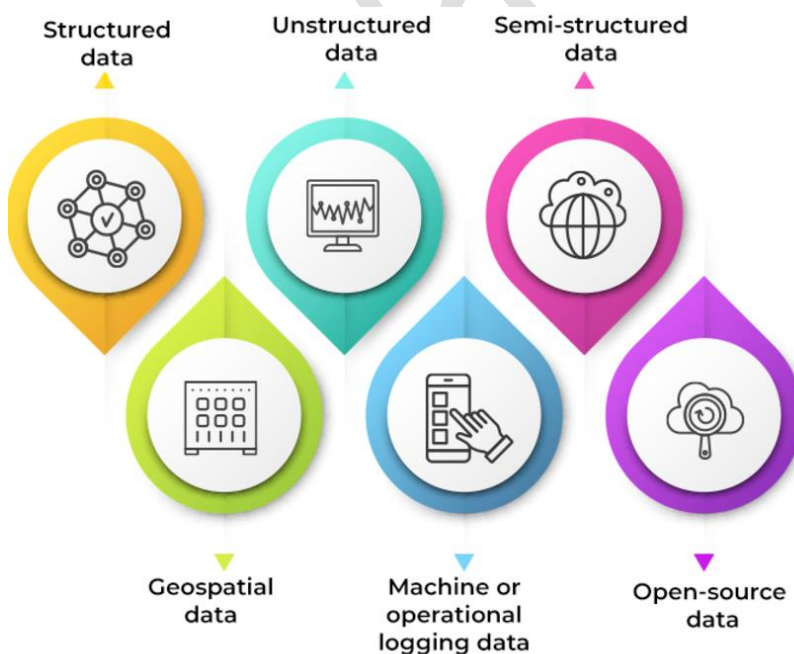
- Example: Dealing with errors in sensor readings.

5. Value

- The potential to extract meaningful insights from the data to make informed business decisions.
- Example: Analysing customer purchase patterns to improve marketing strategies.

These characteristics distinguish Big Data from traditional data and drive the need for advanced processing technologies.

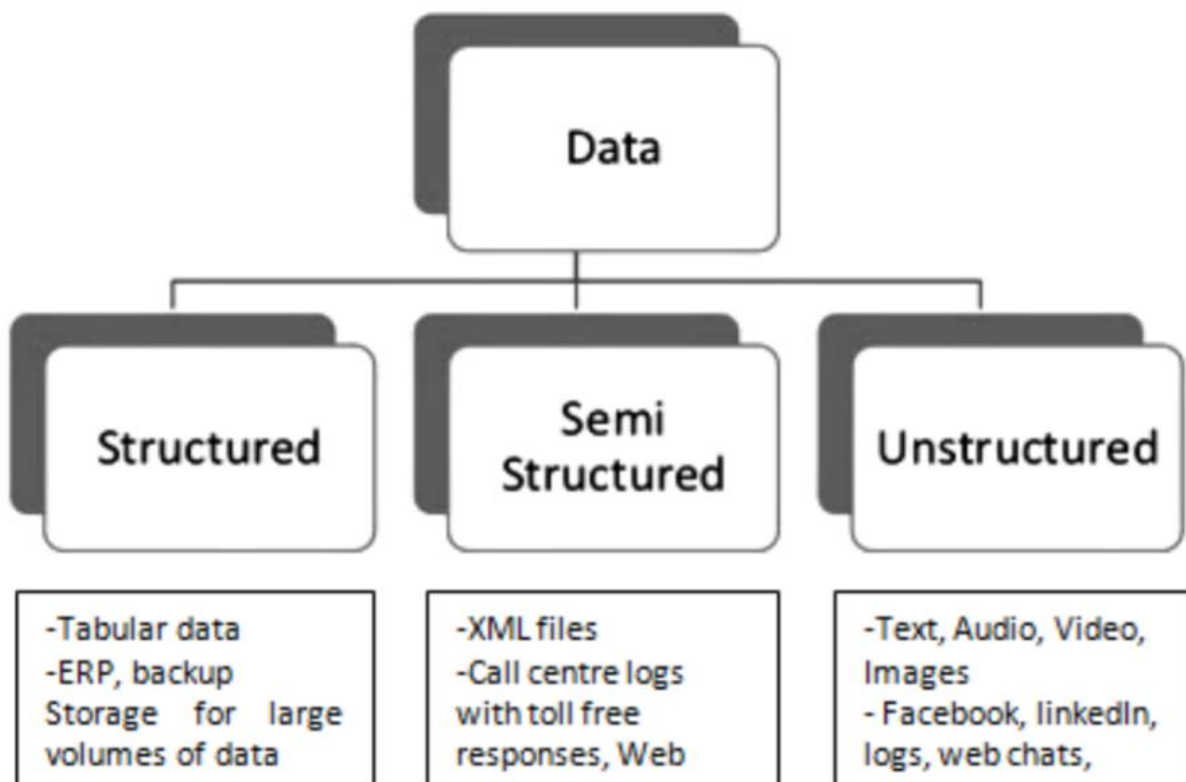
Types Of Big Data



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Various Forms of data collected across



Unstructured data

The university has 5600 students.
John's ID is number 1, he is 18 years old and already holds a B.Sc. degree.
David's ID is number 2, he is 31 years old and holds a Ph.D. degree. Robert's ID is number 3, he is 51 years old and also holds the same degree as David, a Ph.D. degree.

Semi-structured data

```
<University>
<Student ID="1">
<Name>John</Name>
<Age>18</Age>
<Degree>B.Sc.</Degree>
</Student>
<Student ID="2">
<Name>David</Name>
<Age>31</Age>
<Degree>Ph.D. </Degree>
</Student>
...
</University>
```

Structured data

ID	Name	Age	Degree
1	John	18	B.Sc.
2	David	31	Ph.D.
3	Robert	51	Ph.D.
4	Rick	26	M.Sc.
5	Michael	19	B.Sc.



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1. Structured Data

- **Definition:** Data that is organized in a predefined format (rows and columns) and stored in relational databases. It has a fixed schema.
 - **Characteristics:**
 - Easy to store, access, and analyze.
 - Follows strict data models (e.g., tables with rows and columns).
 - Queryable using SQL.
 - **Examples:**
 - Customer database:
 - Financial transactions: bank records, sales data.
 - Sensor readings (if stored in structured formats).
-

2. Unstructured Data

- **Definition:** Data that does not have a fixed format or schema and cannot be easily stored in traditional databases.
 - **Characteristics:**
 - Often large in volume and stored in files or object storage.
 - Requires special tools for processing (e.g., Natural Language Processing, computer vision).
 - Harder to query and analyze directly.
 - **Examples:**
 - Media files: Images, videos, audio recordings.
 - Text data: Social media posts, emails, PDF documents.
 - IoT data in raw formats: Logs, binary sensor outputs.
-

3. Semi-Structured Data

- **Definition:** Data that does not follow a strict tabular format but contains organizational markers (tags, keys) to separate elements. It is flexible yet partially organized.

- **Characteristics:**

- Does not conform to relational models but can be parsed with tools.
- Common in data exchange formats.
- Easier to work with compared to unstructured data.

- **Examples:**

- NoSQL databases (e.g., MongoDB, Cassandra).
- CSV files with inconsistent row structures.

4. Geospatial Data

- **Format:** Represents geographical information (coordinates, polygons).

- **Examples:**

- GPS data, satellite imagery, map boundaries (GeoJSON, Shapefiles).
-

5. Machine or Operational Log Data

- **Format:** Logs from systems, often semi-structured or unstructured.

- **Examples:**

- Server logs, IoT device logs, application error logs.
-

6. Open Source Data

- **Definition:** Freely available data for public use, often structured or semi-structured.

- **Examples:**

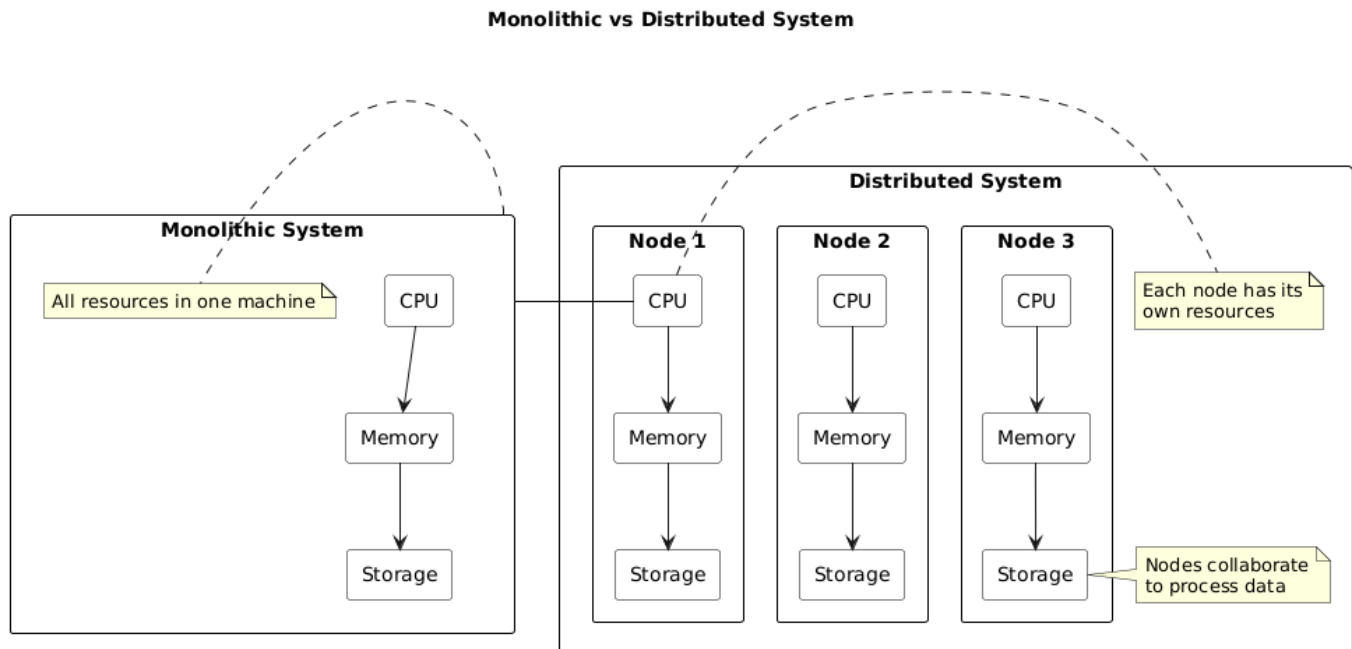
- Census data, government APIs, Kaggle datasets, GitHub repositories.



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Monolith vs. Distributed Systems in Big Data



Monolithic System

- **Architecture:**
 - Consists of a single, integrated system that contains all resources (CPU, RAM, Storage).
- **Resources:**
 - **CPU Cores:** 4
 - **RAM:** 8 GB
 - **Storage:** 1 TB
- **Scaling:**
 - **Vertical Scaling:** Increases performance by adding more resources (e.g., upgrading CPU, RAM, or storage) to the same machine.
 - **Limitations:**
 - Performance gains diminish after a certain point due to hardware constraints.



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- Eventually, hardware limitations restrict the ability to effectively scale performance in proportion to the resources added.

Distributed System

- **Architecture:**

- Composed of multiple interconnected nodes, each operating independently but contributing to a common system.

- **Node Resources** (Example of three nodes):

- **Node 1:**

- CPU Cores: 4
- RAM: 8 GB
- Storage: 1 TB

- **Node 2:**

- CPU Cores: 4
- RAM: 8 GB
- Storage: 1 TB

- **Node 3:**

- CPU Cores: 4
- RAM: 8 GB
- Storage: 1 TB

- **Scaling:**

- **Horizontal Scaling:** Increases performance by adding more nodes to the system (e.g., adding more machines).

- **Advantages:**

- Performance increases in direct proportion to the number of nodes added, achieving **true scaling**.
- Each node can independently handle its own workload, improving overall system performance and fault tolerance.



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1. Monolithic Systems

- **Definition:** In monolithic architecture, all components of a system (data storage, processing, and user interface) are tightly integrated into a single framework or application.
- **Characteristics:**
 - Centralized design with all operations performed on a single machine or tightly coupled system.
 - Easier to manage in small-scale applications.
 - Requires more resources as data grows, leading to performance bottlenecks.
 - Limited scalability, as the system can only handle the data and processing power of one machine.
- **Challenges in Big Data:**
 - Difficulty in handling large volumes of data.
 - Single point of failure: If the system goes down, the entire operation stops.
 - Harder to scale horizontally (i.e., adding more machines).
- **Example:** Traditional relational databases (like MySQL on a single server) where both storage and processing occur on the same machine.

2. Distributed Systems

- **Definition:** In distributed architecture, data storage and processing are split across multiple machines or nodes, working together as a unified system.
- **Characteristics:**
 - Data and tasks are distributed across multiple machines (nodes) that collaborate to process data efficiently.
 - Highly scalable by adding more nodes to handle increasing data volumes and processing demands.
 - Fault-tolerant: Even if one node fails, the system continues to operate using the remaining nodes.
 - Offers better performance and resilience for handling massive datasets.



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- Enables parallel processing, which significantly speeds up data analysis in Big Data environments.
- **Advantages for Big Data:**
 - Scales horizontally, making it ideal for processing large datasets.
 - Handles a variety of data types and sources efficiently.
 - Can process data in real-time, distributing workloads across multiple nodes.
 - **Example:** Apache Hadoop, Apache Spark, and other distributed systems designed for Big Data processing, where tasks are spread across multiple servers.

Key Differences:

Aspect	Monolithic System	Distributed System
Architecture	Single, tightly integrated system	Multiple nodes working together
Scalability	Limited (vertical scaling)	High (horizontal scaling by adding nodes)
Fault Tolerance	Low (single point of failure)	High (nodes can fail without affecting the system)
Processing Power	Limited to one machine	Parallel processing on multiple machines
Suitability for Big Data	Not well-suited for large datasets	Ideal for handling Big Data
Example	Traditional databases (e.g., MySQL)	Apache Hadoop, Apache Spark

Conclusion:

- **Monolithic systems** may work for smaller-scale applications but struggle with the volume, variety, and velocity of Big Data.
- **Distributed systems** are essential for efficiently processing and analyzing Big Data, offering scalability, fault tolerance, and high performance.

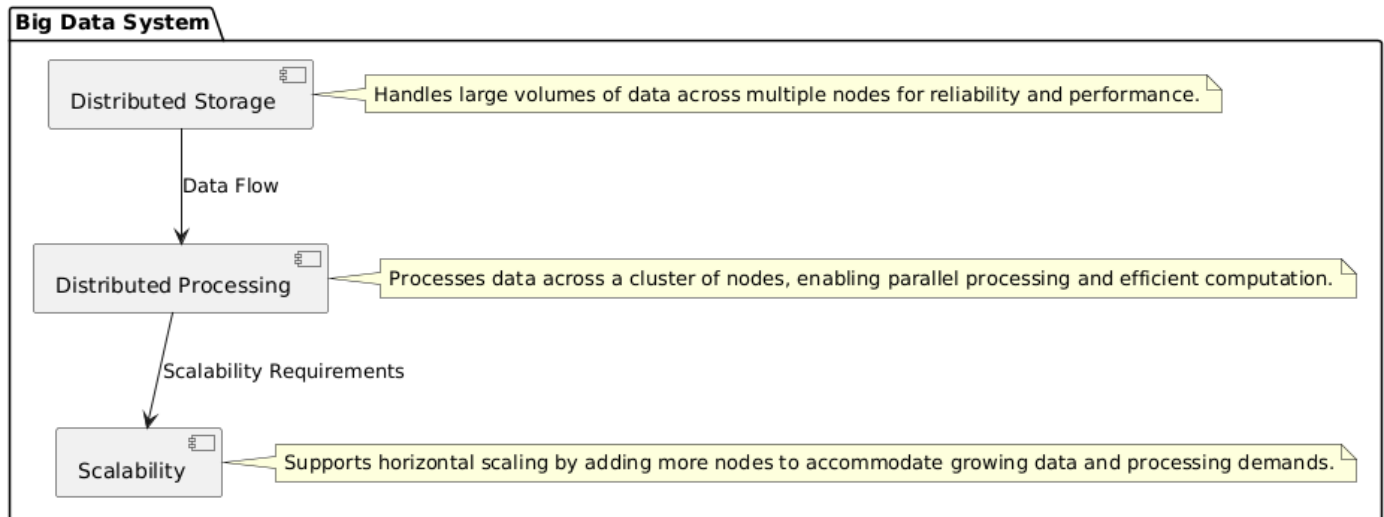


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Design the big data system

Here are the notes on the three essential factors to consider when designing a good Big Data System:



1. Storage

- **Requirement:** Big Data systems need to store massive volumes of data that traditional systems cannot handle effectively.
- **Solution:** Implement **Distributed Storage**.
 - **Definition:** A storage system that spreads data across multiple locations or nodes, allowing for better management of large datasets.
 - **Benefits:**
 - Scalability: Easily accommodates increasing data sizes by adding more storage nodes.
 - Reliability: Redundant storage across nodes enhances data durability and availability.
 - Performance: Enables faster data retrieval and processing through parallel access.

2. Processing / Computation

- **Challenge:** Traditional processing systems are designed for data residing on a single machine, which limits their capability to handle distributed data.

- **Solution:** Utilize **Distributed Processing**.
 - **Definition:** A computation model where data processing tasks are distributed across multiple nodes in a cluster.
 - **Benefits:**
 - Efficiency: Processes large volumes of data in parallel, significantly reducing processing time.
 - Flexibility: Adapts to various types of data and processing tasks without requiring significant changes to the underlying architecture.
 - Fault Tolerance: If one node fails, other nodes can continue processing, ensuring system reliability.

3. Scalability

- **Requirement:** The system must be able to adapt to increasing data volumes and processing demands.
- **Solution:** Design for **Scalability**.
 - **Definition:** The capability of a system to grow and manage increased demands efficiently.
 - **Benefits:**
 - Horizontal Scaling: Adding more nodes to the system allows for increased capacity and performance.
 - Cost-Effectiveness: Scaling out (adding more machines) is often more economical than scaling up (upgrading a single machine).
 - Future-Proofing: A scalable architecture can accommodate future growth without requiring a complete redesign.

Summary

When designing a Big Data system, it's crucial to focus on:

- **Storage:** Implement distributed storage solutions to handle massive datasets.
- **Processing:** Use distributed processing methods to efficiently compute across multiple nodes.
- **Scalability:** Ensure the system can grow to meet increasing data and processing demands, leveraging both horizontal scaling and cost-effective strategies.

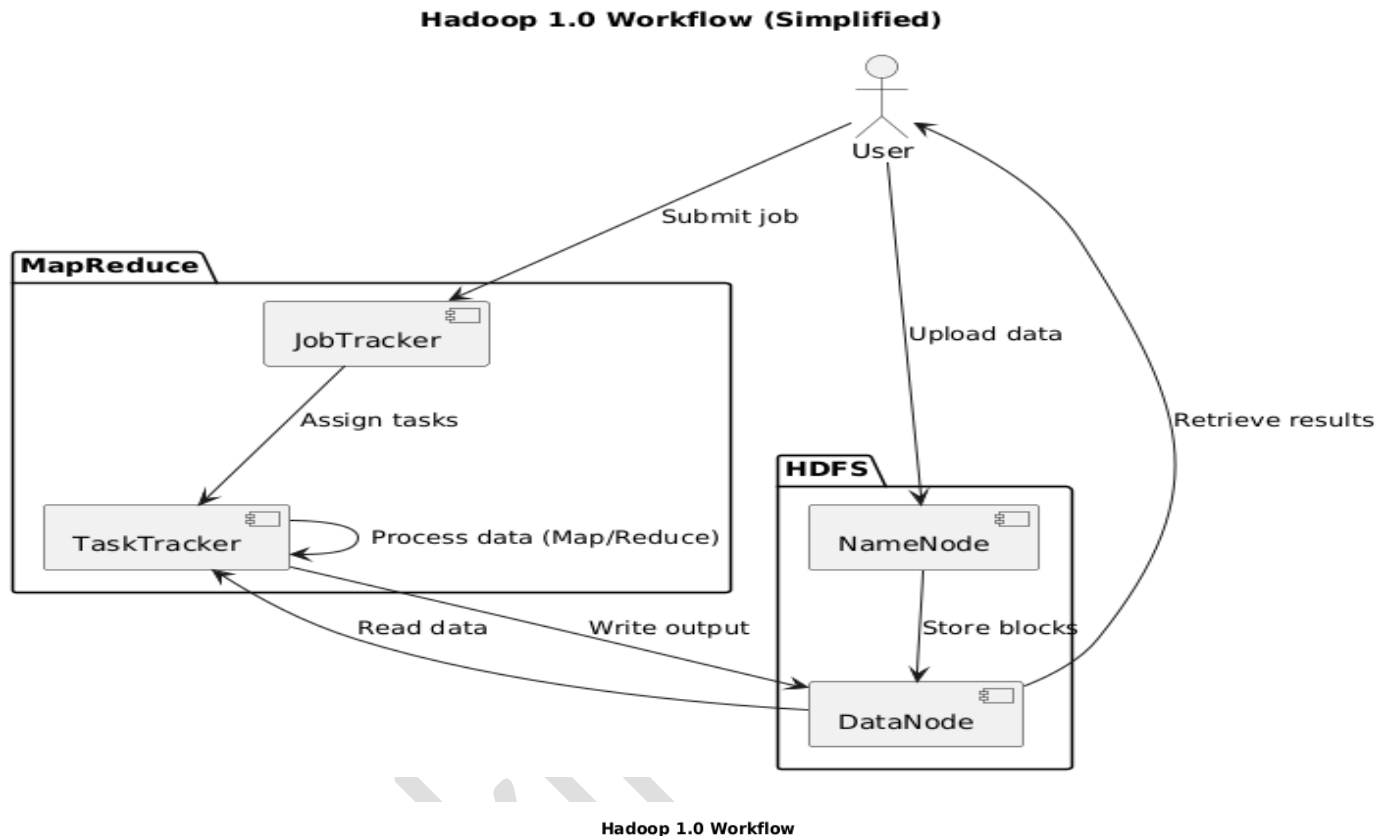


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Master Spark Concepts Zero to Big Data Hero:

Hadoop Architecture Evolution

Hadoop Architecture 1.0



Core Components:



1. HDFS (Hadoop Distributed File System):

- A distributed storage system that stores large data files across multiple nodes in the cluster.
- Data is split into blocks (default 64MB or 128MB) and stored on DataNodes.
- Key Components:
 - **NameNode (Master):** Stores metadata like file structure, block locations, and replication details.
 - **DataNode (Slave):** Stores the actual data blocks and sends periodic updates to the NameNode.

2. MapReduce:

- A distributed data processing framework that processes data in parallel.
- It involves two phases:
 - **Map Phase:** Processes and filters data, generating key-value pairs.
 - **Reduce Phase:** Aggregates the data produced by the Map phase.
- Key Components:
 - **JobTracker (Master):** Manages job scheduling and resource allocation.
 - **TaskTracker (Slave):** Executes individual tasks assigned by the JobTracker.

How Hadoop 1.0 Worked (Step-by-Step)

1. Data Ingestion and Storage in HDFS:

- Input data is uploaded to HDFS using commands or APIs.
- The file is split into fixed-size blocks (e.g., 64MB) and replicated (default replication factor: 3).
- NameNode stores the metadata about block locations, while DataNodes store the actual data blocks.

2. Job Submission:

- A user submits a MapReduce job to the JobTracker.
- The job contains:
 - Input data location in HDFS.



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- Mapper and Reducer logic.
- Output data location.

3. JobTracker Assigns Tasks:

- JobTracker splits the job into smaller tasks (Map and Reduce tasks).
- It assigns these tasks to TaskTrackers on different DataNodes.
- TaskTrackers fetch input data blocks locally (Data Locality optimization).

4. Map Phase Execution:

- TaskTrackers execute the Mapper code on their assigned data blocks.
- Each Mapper processes its block and generates intermediate key-value pairs.
- The intermediate data is temporarily stored locally.

5. Shuffling and Sorting:

- Intermediate key-value pairs are shuffled (grouped by keys) and sorted.
- Data is then sent to the appropriate Reducers.

6. Reduce Phase Execution:

- TaskTrackers run the Reducer code to process grouped key-value pairs.
- The Reducer aggregates or performs computations (e.g., summing, counting) and generates final results.

7. Output Storage in HDFS:

- Reducers write the final output data back to HDFS.
- Users can access this data through HDFS commands or APIs.

8. Monitoring and Reporting:

- JobTracker continuously monitors task execution and reassigns failed tasks to another TaskTracker (fault tolerance).
- After all tasks complete, JobTracker provides a final report to the user.



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Key Highlights of Hadoop 1.0 Workflow:

- **Data Locality:** Processing happens where the data resides to minimize network overhead.
- **Fault Tolerance:** Failed tasks are retried or reassigned using replication.
- **Batch Processing:** Suitable for large-scale, offline data processing tasks.

This process enabled reliable, parallel processing of massive datasets, although it suffered from limitations like single points of failure and scalability issues in large clusters.

Why Hadoop 1.0 Failed: Limitations

1. Single Point of Failure:

- **NameNode:** If the NameNode crashed, the entire system became unavailable since it stored all metadata.

2. Scalability Issues:

- The **JobTracker** managed job scheduling, task monitoring, and resource management, creating a bottleneck in large clusters.

3. Resource Utilization:

- **Fixed Slot Model:** Each node had fixed slots for Map and Reduce tasks, leading to inefficient resource usage. For example, idle Map slots couldn't be used for Reduce tasks and vice versa.

4. Lack of General-Purpose Computing:

- Hadoop 1.0 was designed only for **MapReduce** workloads, limiting its usability for other types of data processing like streaming or interactive queries.

5. Cluster Utilization:

- Tasks couldn't share resources dynamically, leading to underutilization of CPU, memory, and disk.

6. Latency:

- High overhead in job initialization and monitoring due to reliance on **JobTracker**.

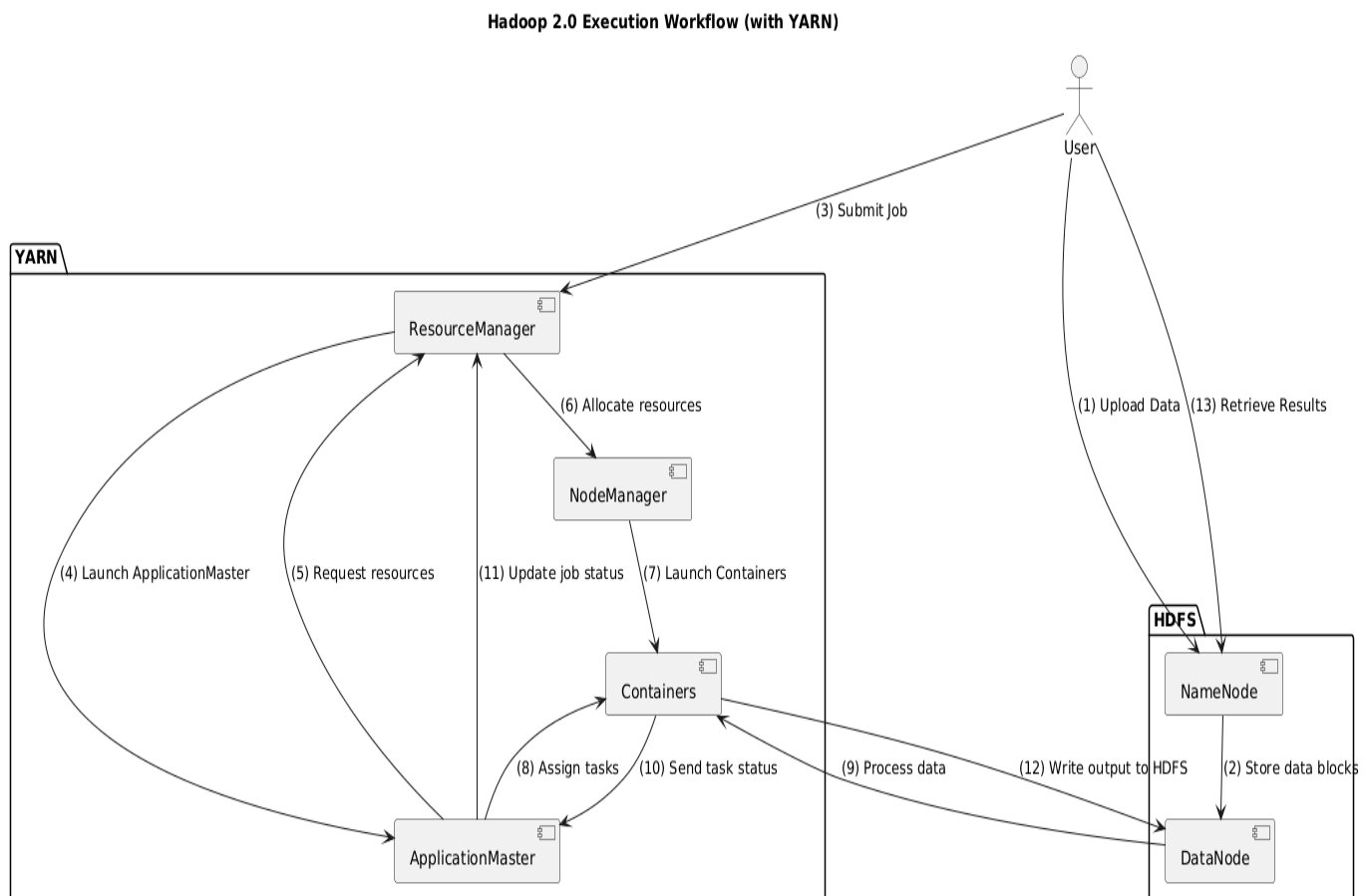


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Hadoop 2.0: Overview and Steps

Hadoop 2.0 introduced **YARN (Yet Another Resource Negotiator)** to address the limitations of Hadoop 1.0 and decouple resource management from the MapReduce framework.

Hadoop 2.0 Workflow in Steps



1. Introduction of YARN:

- YARN separates **resource management** from **job scheduling** and **monitoring**.
- Key components:
 - **ResourceManager (Master):** Manages resources across the cluster.
 - **NodeManager (Slave):** Runs on each node and reports resource usage.
 - **ApplicationMaster:** Handles application-specific task scheduling.

2. Data Storage in HDFS:

- HDFS remains the storage layer, but now it supports fault tolerance and NameNode High Availability (HA) using standby NameNodes.



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3. Resource Allocation:

- **ResourceManager** assigns resources dynamically to various applications (not just MapReduce) via containers.

4. Application Submission:

- User submits a job to the **ResourceManager**.
- The **ApplicationMaster** is launched to coordinate tasks for that specific job.

5. Task Execution:

- **NodeManagers** on individual nodes launch containers to execute tasks.
- Tasks can belong to any framework, such as MapReduce, Spark, or Tez.

6. Dynamic Resource Utilization:

- Containers dynamically allocate CPU, memory, and disk resources based on the workload, improving utilization.

7. Improved Scalability and Fault Tolerance:

- YARN allows scaling to thousands of nodes by delegating specific responsibilities to the ApplicationMaster and NodeManagers.
- NameNode HA minimizes downtime.

8. Support for Multiple Workloads:

- Beyond MapReduce, Hadoop 2.0 supports frameworks like Spark, Flink, and HBase for a variety of workloads.

Hadoop 2.0 Advantages:

- Eliminated **single point of failure** with NameNode HA.
- Improved resource utilization with dynamic allocation.
- Enabled **multi-tenancy** with support for multiple frameworks.
- Scaled efficiently for large clusters.

Hadoop 2.0 marked a major milestone in making Hadoop versatile, robust, and suitable for modern data processing needs.



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Master Spark Concept Zero to Hero:

What is Hadoop?

- **Hadoop** is an open-source framework designed for processing and storing large datasets in a distributed computing environment. It comprises a suite of tools that work together to address the challenges posed by Big Data.

Evolution of Hadoop

- **2007**: Introduction of **Hadoop 1.0**, laying the foundation for Big Data processing.
- **2012**: Release of **Hadoop 2.0**, introducing YARN for improved resource management.
- **Current Version: Hadoop 3.0**, featuring enhancements for scalability and performance.

Core Components of Hadoop

1. HDFS (Hadoop Distributed File System):

- A distributed storage system that enables the storage of large files across multiple machines, ensuring data redundancy and fault tolerance.

2. MapReduce:

- A programming model for processing vast amounts of data in parallel. It simplifies the processing of data across the HDFS, though it can be complex and requires substantial code for execution.

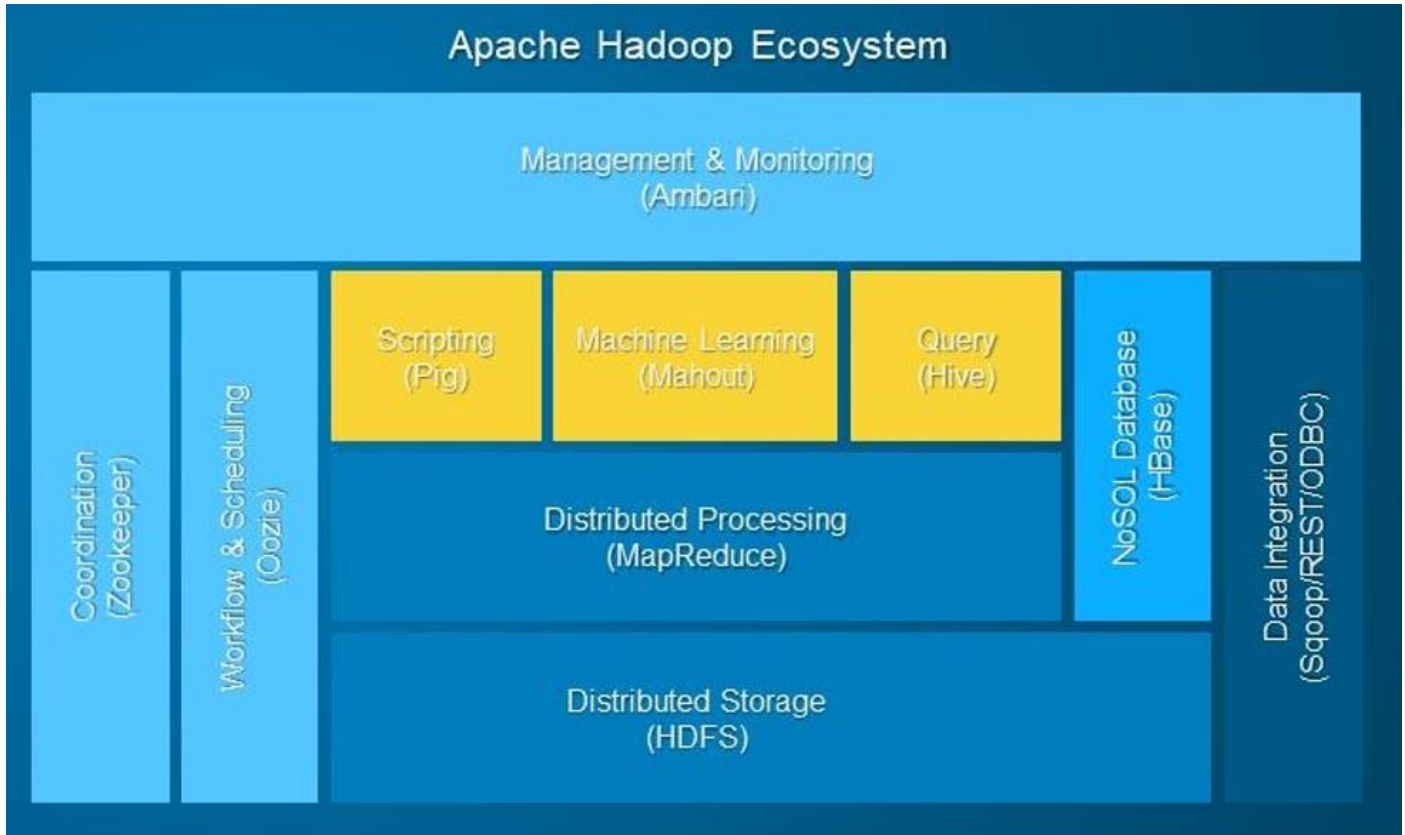
3. YARN (Yet Another Resource Negotiator):

- A resource management layer that allocates and manages system resources across the Hadoop cluster, allowing multiple applications to run simultaneously.



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Hadoop Ecosystem Technologies



- **Sqoop:**
 - Facilitates data transfer between HDFS and relational databases. It automates the import/export processes, making data movement efficient. *Cloud Alternative:* Azure Data Factory (ADF).
- **Pig:**
 - A high-level scripting language used for data cleaning and transformation, which simplifies complex data manipulation tasks. *Underlying Technology:* Uses MapReduce.
- **Hive:**
 - Provides a SQL-like interface for querying data stored in HDFS, translating queries into MapReduce jobs for execution.
- **Oozie:**
 - A workflow scheduler for managing complex data processing workflows, allowing for dependencies and scheduling of multiple tasks. *Cloud Alternative:* Azure Data Factory.



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- **HBase:**
 - A NoSQL database for quick, random access to large datasets, facilitating real-time data processing. *Cloud Alternative:* CosmosDB.

Challenges with Hadoop

1. MapReduce Complexity:

- Developing MapReduce jobs can be intricate, often requiring extensive Java coding, making simple tasks time-consuming.

2. Learning Curve:

- Users must master various components for different tasks, resulting in a steep learning curve and increased complexity in the workflow.

Conclusion

Hadoop revolutionized the way we process and store Big Data. While its core components like HDFS and YARN remain vital, the complexity of MapReduce and the necessity to learn multiple ecosystem tools present significant challenges. Despite its evolution, alternatives are emerging to simplify Big Data processing, but Hadoop's foundational role in the Big Data era is undeniable.



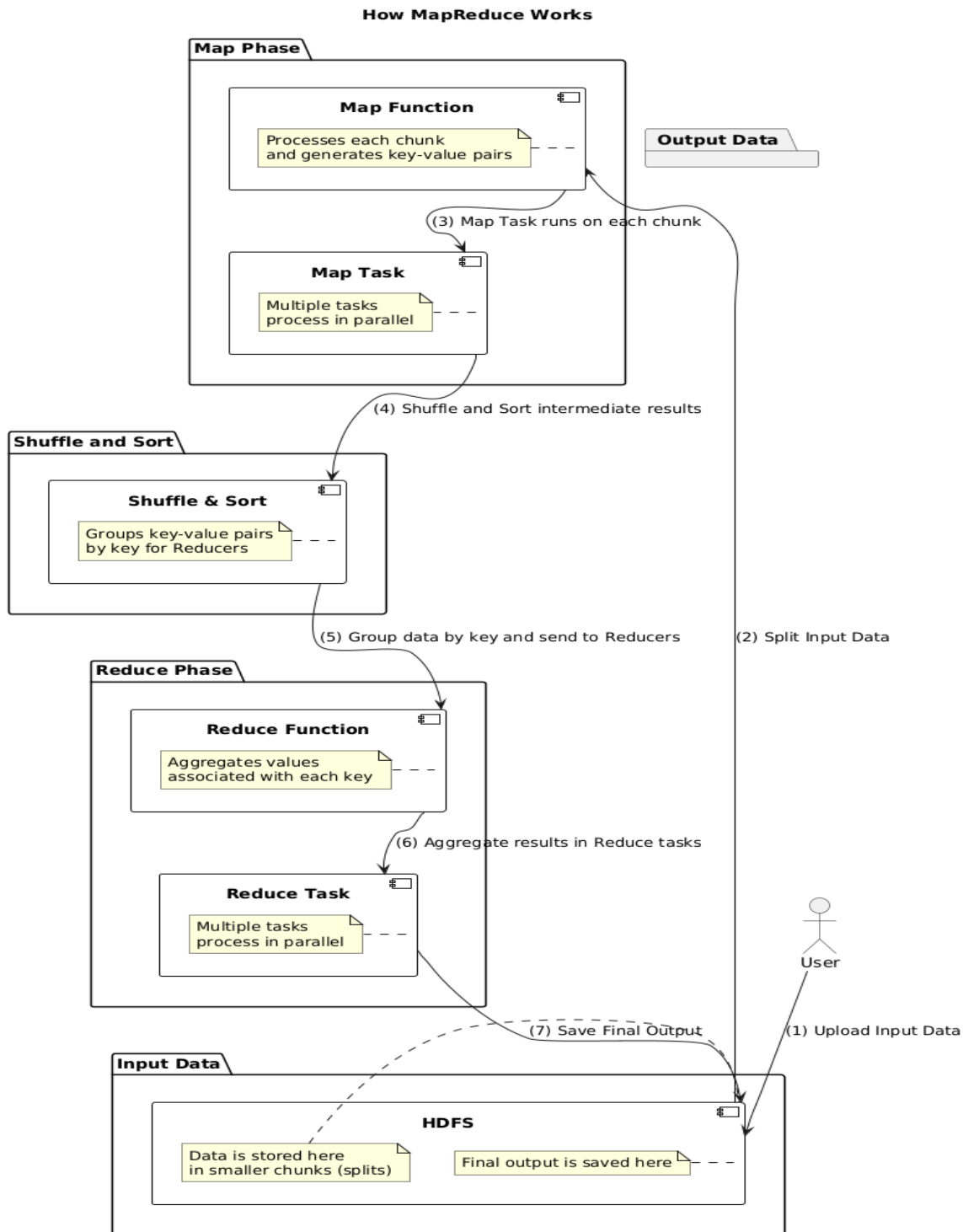
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How does Map Reduce work in Hadoop (Simplified)

MapReduce is a programming model used for processing large datasets across distributed systems like Hadoop. It divides tasks into smaller jobs, processes them in parallel, and then combines the results.

Here's a simple breakdown of how **MapReduce** works:



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1. The Basic Process:

MapReduce involves **two main steps**:

- **Map:** Process input data and generate intermediate results.
- **Reduce:** Aggregate the results to produce the final output.

2. The MapReduce Workflow in Steps:

1. Input Data:

- Data is stored in a distributed file system like **HDFS** (Hadoop Distributed File System).
- The data is divided into smaller chunks called **splits**.

2. Map Step:

- Each chunk of data is processed by a **Map function**.
- The **Map function** processes each record (e.g., a line of text) and outputs **key-value pairs**.
 - Example: For word counting, the map function will output (word, 1) for each word in a document.

3. Shuffle and Sort:

- After mapping, the system **shuffles and sorts** the data by the key (e.g., all occurrences of the word "apple" will be grouped together).
- This step ensures that all records with the same key go to the same **Reduce function**.

4. Reduce Step:

- The **Reduce function** takes each key and its list of values (e.g., for the word "apple", the values could be [1, 1, 1]).
- The Reduce function processes these values (e.g., sums them up) and produces a final result (e.g., the total count of the word "apple").

5. Output Data:



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- The final results are saved in **HDFS** for further use or analysis.
-

3. Example: Word Count

Let's say we want to count the number of occurrences of each word in a document.

Map Function:

- Input: A line of text.
- Output: Key-value pairs for each word in the line.

Input: "apple orange apple"

Output:

```
("apple", 1)
("orange", 1)
("apple", 1)
```

Reduce Function:

- Input: Key ("apple") and list of values ([1, 1]).
- Output: Total count for the word ("apple", 2).

Input:

```
("apple", [1, 1])
("orange", [1])
```

Output:

```
("apple", 2)
("orange", 1)
```

4. Summary of Steps:

1. **Map:** Process each record and generate key-value pairs.
2. **Shuffle & Sort:** Group the pairs by key.
3. **Reduce:** Aggregate the values for each key.
4. **Output:** Save the final results.



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Disadvantages of MapReduce and Why Hadoop Became Obsolete

Hadoop's **MapReduce** framework revolutionized big data processing in its early years but eventually became less favorable due to the following **disadvantages**:

1. Limitations of MapReduce

1. Complex Programming Model:

- Writing MapReduce jobs requires significant boilerplate code for even simple operations.
- Developers need to write multiple jobs for iterative tasks.

2. Batch Processing Only:

- MapReduce is designed for batch processing, making it unsuitable for real-time or streaming data processing.

3. High Latency:

- The system writes intermediate data to disk between the Map and Reduce phases, resulting in high input/output overhead and slower performance.

4. Iterative Computations Are Inefficient:

- Iterative tasks like machine learning or graph processing require multiple MapReduce jobs, with each job reading and writing to disk, causing inefficiency.

5. Lack of In-Memory Processing:

- MapReduce does not leverage in-memory computation, which is faster compared to disk-based processing.

6. Resource Utilization:

- MapReduce uses a static allocation of resources, leading to underutilization of cluster resources.

7. Not Fault-Tolerant for Iterative Tasks:

- While MapReduce can recover from node failures, the re-execution of failed tasks for iterative workloads is time-consuming.



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8. Dependency on Hadoop 1.0's Architecture:

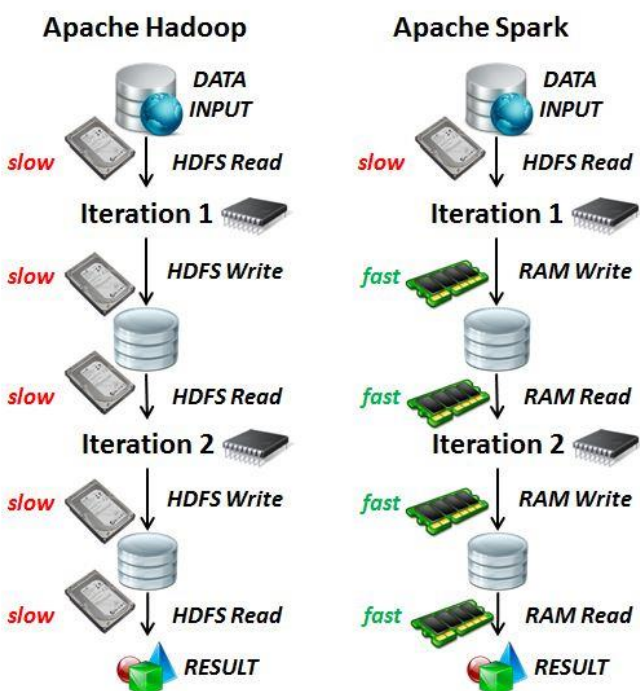
- The reliance on the **JobTracker/ TaskTracker** model caused scalability issues and made resource management inefficient.

2. Why Hadoop MapReduce Became Obsolete

- **Emergence of Alternatives:** Advanced tools like Apache Spark provided a more efficient way of processing data with better APIs and support for real-time data processing.
- **Slow Evolution:** MapReduce couldn't keep up with the growing demand for interactive queries, in-memory computation, and real-time analytics.
- **Disk I/O Bottleneck:** Repeated disk read/write operations made it slower compared to in-memory processing engines.
- **Streaming Limitations:** Hadoop MapReduce lacked native support for stream processing, a critical need for modern applications.

Advantages of Apache Spark

Apache Spark, introduced as a faster and more versatile data processing engine, addressed the limitations of MapReduce and offered several advantages:



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1. In-Memory Processing

- Spark processes data in memory, significantly reducing the time spent on disk I/O.
- This makes it much faster for iterative computations like machine learning and graph processing.

2. Unified Framework

- Supports multiple workloads:
 - **Batch Processing** (like MapReduce)
 - **Stream Processing** (real-time data)
 - **Interactive Queries** (e.g., Spark SQL)
 - **Machine Learning** (via MLlib)
 - **Graph Processing** (via GraphX)

3. Easy-to-Use APIs

- Provides high-level APIs in multiple languages like **Python**, **Scala**, **Java**, and **R**.
- Rich libraries for SQL, streaming, and machine learning simplify development.

4. Fast Performance

- Spark is **10-100x faster** than Hadoop for certain tasks, thanks to in-memory computation and optimized DAG execution.

5. Support for Real-Time Processing

- Spark Streaming enables the processing of real-time data streams, unlike Hadoop's batch-only capabilities.

6. Dynamic Resource Management

- Spark utilizes cluster managers like **YARN**, **Kubernetes**, or **Standalone**, dynamically allocating resources to improve cluster utilization.

7. Fault Tolerance

- Spark achieves fault tolerance using **RDDs (Resilient Distributed Datasets)**, which track transformations and can recompute data automatically in case of failures.

8. Scalability



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- Spark can scale efficiently from small clusters to thousands of nodes, handling petabytes of data.

9. Wide Ecosystem

- The Spark ecosystem includes libraries like:
 - **Spark SQL:** For structured data processing.
 - **Spark Streaming:** For real-time data processing.
 - **Mlib:** For machine learning.
 - **GraphX:** For graph-based computations.

10. Improved Iterative Processing

- Spark retains data in memory across iterations, making it highly efficient for algorithms like PageRank or k-means clustering.

Comparison of Hadoop MapReduce and Apache Spark

Feature	Hadoop MapReduce	Apache Spark
Processing Speed	Slower (Disk-based)	Faster (In-memory processing)
Ease of Use	Complex to code	Easy APIs for various languages
Real-Time Processing	Not Supported	Supported via Spark Streaming
Fault Tolerance	Yes, but slower recovery	Yes, with RDD-based fast recovery
Resource Utilization	Inefficient (Static)	Efficient (Dynamic allocation)
Supported Workloads	Batch processing only	Batch, Streaming, ML, Graph

Conclusion

Hadoop MapReduce played a pivotal role in big data processing during its time but became obsolete due to its inefficiency and inability to adapt to modern requirements. Apache Spark, with its fast, versatile, and easy-to-use framework, has emerged as the go-to solution for distributed data processing.



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Recap of Spark Concepts with Interview question on week 2:

Core Components of Spark

1. What are the core components of a Spark cluster?
 2. Explain the role of the Driver Node and Worker Node in Spark.
 3. What is the function of the Cluster Manager in Spark? Name a few cluster managers compatible with Spark.
 4. How are tasks assigned to worker nodes in Spark?
 5. What happens if a worker node fails during a job execution?
-

JVM, On-Heap Memory, Off-Heap Memory, and Garbage Collector

6. Explain the difference between on-heap and off-heap memory in Spark.
 7. Why is off-heap memory preferred for caching in Spark?
 8. How does the Garbage Collector affect Spark performance?
 9. What techniques can you use to optimize memory usage in Spark?
 10. How does JVM tuning impact Spark applications?
-

Spark Architecture

11. Can you explain the high-level architecture of Apache Spark?
 12. How does Spark achieve fault tolerance?
 13. What are the main differences between Spark's physical and logical plans?
 14. Explain the role of executors in Spark.
 15. What are broadcast variables, and how do they optimize Spark jobs?
-

Execution of Spark Jobs

16. Describe the execution flow of a Spark application.
17. What is the role of the SparkContext in job execution?
18. What happens when you call an action on an RDD or DataFrame in Spark?



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19.How are transformations and actions different in Spark?

20.What are the key stages of Spark execution?

Lazy Evaluation and Fault Tolerance

21.What is lazy evaluation, and why is it important in Spark?

22.How does Spark handle fault tolerance for RDDs?

23.Explain the concept of lineage graphs in Spark.

24.Why is checkpointing used in Spark, and how does it help with fault tolerance?

Directed Acyclic Graph (DAG) and Its Components

25.What is a DAG in Spark?

26.Explain the components of a DAG in Spark.

27.How is the DAG scheduler different from the task scheduler?

28.What are stages and tasks in a DAG? How are they related?

29.What triggers the creation of a new stage in a Spark job?

DAG Rescheduler

30.What is a DAG rescheduler, and when is it invoked?

31.How does the DAG scheduler recover from a failed task?

32.What is the role of speculative execution in Spark?

Job, Stages, and Tasks

33.Explain the relationship between jobs, stages, and tasks in Spark.

34.How does Spark decide the number of tasks in a stage?

35.What happens when a stage fails during execution?

36.How does Spark's locality-aware scheduling impact task execution?

37.What is the role of a shuffle in determining stages?

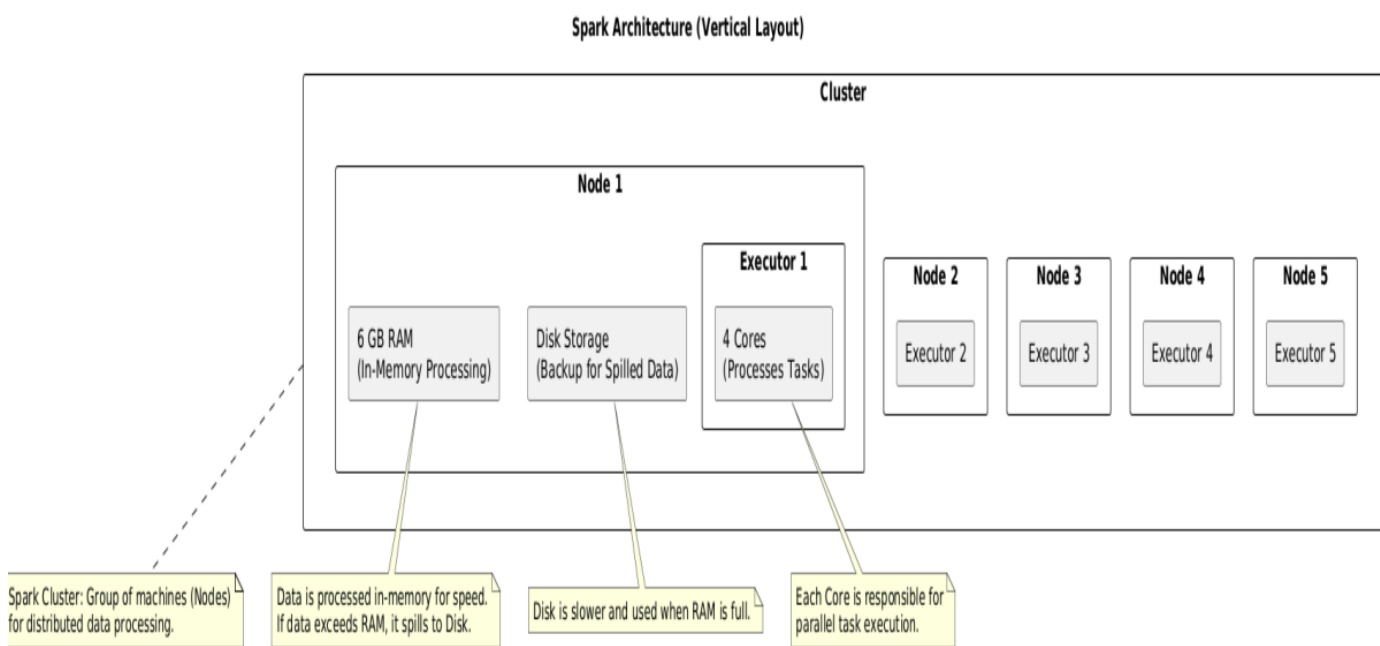


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Master Spark Concepts Zero to Big Data Hero:

Spark's Core Concepts: Cluster, Core, Executor, RAM, and Disk

Apache Spark is a powerful tool for handling big data, and understanding how it works can help optimize your workflows. Let's break down its key components:



1. The Cluster – Your Team of Workers

- Concept:** A Spark Cluster is like a team of workers (nodes) in a warehouse, where each worker is assigned a part of the warehouse (data) to work on.
- In Spark:** The cluster is a group of machines (worker nodes) that share the load of processing data. It distributes the data across these nodes to ensure no single machine is overwhelmed.

2. Core – The Brainpower of Each Worker

- Concept:** Think of each worker having several tools (cores) to perform tasks. More tools = more work done in parallel.
- In Spark:** A Core is the CPU unit responsible for handling tasks. Each worker node has one or more cores.



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- **Example:** If each node has 4 cores, it can process 4 tasks at once. This increases the speed of processing by working on multiple tasks in parallel.

3. Executor – The Workers Doing the Job

- **Concept:** The Executor is like a worker in the warehouse who actually does the work (e.g., sorting, filtering, analyzing data).
- **In Spark:** Each executor runs on a worker node and is responsible for executing the tasks of a job. Executors perform data processing tasks and store the results.
 - **Example:** Executors handle multiple tasks based on the number of cores assigned to them.

4. RAM – The Worker's High-Speed Toolbox

- **Concept:** RAM is like the worker's toolbox, holding essential tools needed for fast work. The larger the toolbox, the fewer trips to the storage room (disk) are needed.
- **In Spark:** RAM stores intermediate data during processing. If the data fits into memory, Spark can process it quickly without needing to write to disk (which is slower).
 - **Example:** If an executor has 6 GB of RAM, it can process data faster. If the data exceeds this memory, Spark will spill data to disk, slowing down the process.

5. Disk – The Worker's Slow Backup Storage

- **Concept:** Disk is like a storage room for tools. When the worker's toolbox (RAM) is full, they must run to the storage room to get more tools, which is slower.
- **In Spark:** When there's not enough RAM, Spark spills data to disk. Although this helps with large datasets, it slows down processing.
 - **Example:** If there is not enough memory (RAM), Spark will use disk storage to temporarily hold data, but accessing data from disk is slower.



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Real-World Example in Spark:

Imagine you have a **100 GB** dataset of customer orders, and you want to analyze the top-selling products:

- **Cluster:** Distributes the data across **5 worker nodes**.
- **Cores:** Each node has **4 cores**, so each node can process **4 tasks in parallel**.
- **Executors:** Each executor processes tasks such as filtering or aggregating the data.
- **RAM:** If each executor has **6 GB of RAM**, Spark will process the data in-memory, which is faster. If the data exceeds RAM capacity, Spark will spill excess data to disk, which slows down the process.

Optimization Tips:

- Increase the number of **cores per executor** to process more tasks simultaneously.
- Allocate more **RAM** to avoid spilling data to disk.
- Ensure your **cluster** has enough executors to efficiently process the data.

Summary:

- **Cluster:** A team of machines working together.
- **Core:** The CPU unit on each machine that processes tasks.
- **Executor:** The worker responsible for completing tasks on the data.
- **RAM:** High-speed memory for quick data access.
- **Disk:** Slower storage used when RAM is full.

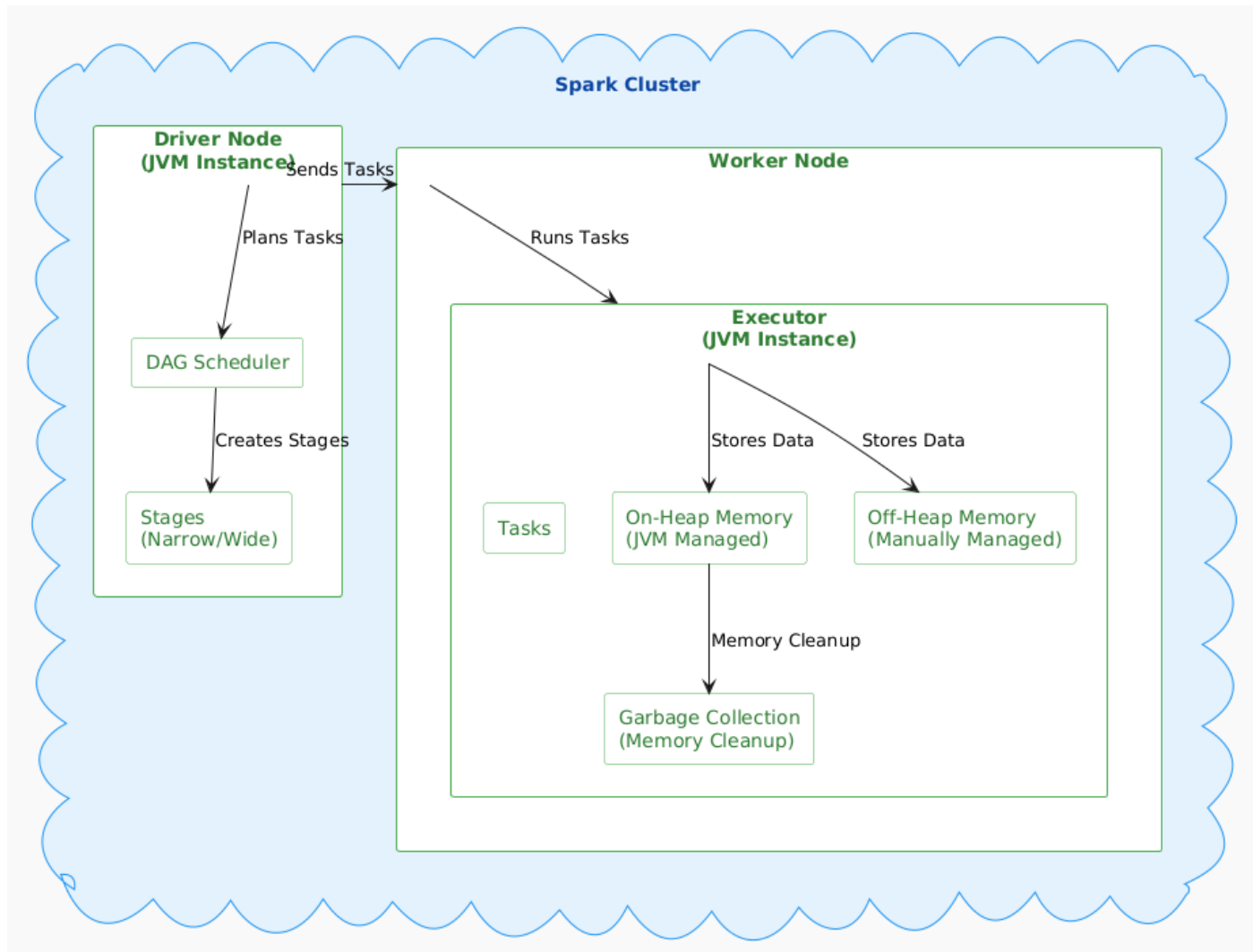
Understanding how these components work together in Spark helps you optimize your big data jobs for faster and more efficient processing.



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Master Spark Concepts Zero to Big Data Hero:

Detailed Notes on Worker Node, Executor, Task, Stages, On-Heap Memory, Off-Heap Memory, and Garbage Collection in Apache Spark



What is JVM?

The **Java Virtual Machine (JVM)** is an abstraction layer that allows Java (or other JVM languages like Scala) applications to run on any machine, regardless of hardware or operating system. It provides:

- **Memory Management:** JVM manages memory allocation, including heap and stack memory.
- **Execution:** JVM converts bytecode (compiled from Java/Scala code) into machine code.
- **Garbage Collection:** JVM automatically handles the cleanup of unused memory through garbage collection.



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JVM's Role in Spark

Apache Spark heavily relies on JVM for executing its core components:

- **Driver Program:** The driver, which coordinates the execution of jobs, runs inside a JVM instance on the driver node.
- **Executors:** Executors, which run tasks on worker nodes, are JVM processes that are responsible for task execution and data storage.

The Spark driver and executors both run as separate JVM instances, each managing its own memory and resources.

1. Worker Node

- **Role:** A **worker node** is a machine within a Spark cluster that performs the execution of tasks and handles data storage.
- **Purpose:** The worker node runs **executors** that are responsible for running tasks on data, managing intermediate results, and sending the final output back to the driver.
- **Components:**
 - **Executors:** Each worker node can have multiple executors, each handling a portion of the job.
 - **Data Storage:** Worker nodes store data either in memory or on disk during execution.

Key Points:

- Worker nodes are essentially the physical or virtual machines that process data in a distributed manner.
- They communicate with the **driver program** to receive tasks and return results.

2. Executor

- **Role:** An **executor** is a JVM process that runs on worker nodes and is responsible for executing tasks and storing data.
- **Lifecycle:** Executors are launched at the beginning of a Spark job and run for the entire duration of the job unless they fail or the job completes.
- **Task Execution:** Executors run tasks in parallel and return results to the driver.
- **Data Management:** Executors store data in-memory (or on disk if necessary) during task execution and shuffle data between nodes if required.



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Key Points:

- Executors perform computations and store intermediate data for tasks.
- Executors handle **two main responsibilities**:
 - (1) executing the tasks sent by the driver and
 - (2) providing in-memory storage for data.
- Executors are removed when the job completes.

3. Task

- **Role**: A **task** is the smallest unit of work in Spark, representing a single operation (like a map or filter) on a partition of the data.
- **Assignment**: Tasks are assigned to executors, and each executor can run multiple tasks concurrently.
- **Execution**: Tasks are generated from stages in the Directed Acyclic Graph (DAG), which defines the order of operations in a job.

Key Points:

- Tasks operate on a single partition of the data and are distributed across multiple executors for parallel processing.
- Tasks are responsible for applying transformations or actions on the data partitions.

4. Stages

- **Role**: Stages represent a logical unit of execution in Spark. Each Spark job is divided into stages by the **DAG Scheduler**, based on the **data shuffle boundaries**.
- **Types**: Stages can be categorized as **narrow** (tasks in a stage can be executed without reshuffling data) and **wide** (requires data shuffling).
- **Creation**: When an action (like count() or collect()) is triggered, Spark creates stages that represent the transformation chain.

Key Points:

- Each stage contains a set of tasks that can be executed in parallel.
- Stages are created based on the **shuffle** dependencies in the job.



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5. On-Heap Memory

- **Role:** In Spark, **on-heap memory** refers to the memory space allocated within the JVM heap for Spark's computations.
- **Usage:** Spark's operations on RDDs or DataFrames store intermediate data in the JVM heap space.
- **Garbage Collection:** On-heap memory is subject to **JVM garbage collection**, which can slow down performance if frequent or large collections occur.

Key Points:

- On-heap memory is prone to the inefficiencies of JVM garbage collection.
- The default memory management in Spark is **on-heap**.

6. Off-Heap Memory

- **Role:** **Off-heap memory** is memory managed outside of the JVM heap. Spark can use off-heap memory to store data, reducing the burden on the JVM's garbage collector.
- **Usage:** It helps in better memory utilization and reduces the likelihood of **OutOfMemoryError** due to garbage collection issues.
- **Configuration:** You can enable off-heap memory using the `spark.memory.offHeap.enabled` configuration.

Key Points:

- Off-heap memory helps optimize memory usage by bypassing the JVM's garbage collector.
- This memory is managed manually, offering more control over memory usage in high-memory workloads.

7. Garbage Collection

- **Role:** **Garbage collection (GC)** in Spark is the process of cleaning up unused objects in JVM memory to free up space for new data.
- **Types of GC:**
 - **Minor GC:** Cleans up the young generation of memory (short-lived objects).
 - **Major GC:** Cleans up the old generation (long-lived objects) and can lead to **stop-the-world** events, pausing task execution.



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- **Impact:** Frequent garbage collection, especially **major GC**, can negatively impact Spark job performance by increasing the overall execution time.

Key Points:

- Inefficient garbage collection can lead to **OutOfMemoryException** and **Driver Out of Memory** errors.
- Spark provides configurations (spark.executor.memory, spark.memory.offHeap.enabled) to optimize memory usage and reduce the impact of GC.

Summary:

- **Worker Node:** The physical or virtual machine that performs the task execution and manages storage in the Spark cluster.
- **Executor:** The process that runs on a worker node, handling task execution and data storage.
- **Task:** The smallest unit of work in Spark, operating on data partitions.
- **Stages:** Logical units of execution that divide a Spark job, categorized as narrow or wide depending on shuffle dependencies.
- **On-Heap Memory:** JVM-managed memory for storing Spark data, subject to garbage collection.
- **Off-Heap Memory:** Memory managed outside the JVM heap to avoid garbage collection delays and memory issues.
- **Garbage Collection:** A JVM process to reclaim memory, but if not optimized, it can negatively affect performance.

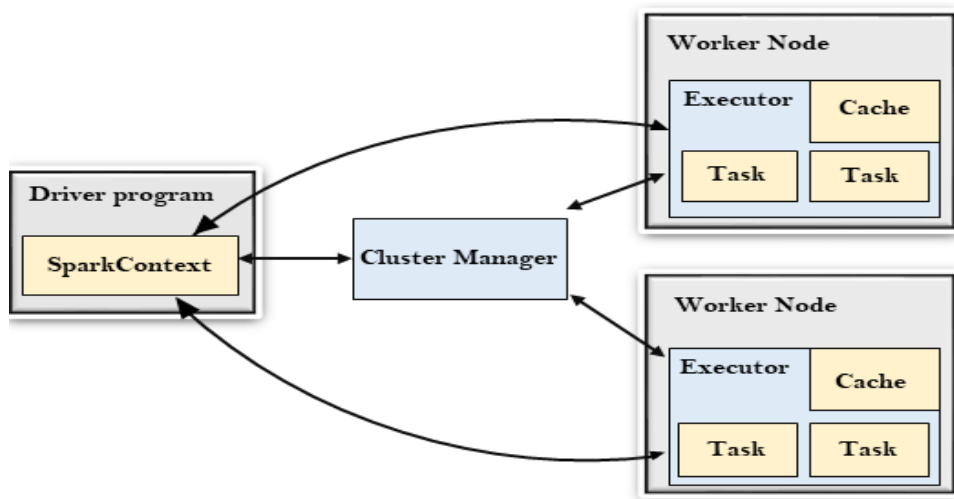


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Master Spark Concepts Zero to Big Data Hero:

Detailed Notes on Spark Architecture and Execution Flow

Apache Spark is a powerful open-source distributed computing system that enables fast and efficient data processing. Here's a quick overview of its architecture to help you understand how it works:



Key Components of Spark Architecture

- **Driver Program**

Description: The central coordinator that converts user code into tasks.

Role: Manages the execution of the Spark application and maintains information about the status of tasks.

- **Cluster Manager**

Description: Manages resources across the cluster.

Types: Standalone, YARN, Mesos, Kubernetes.

Role: Allocates resources and schedules tasks on the cluster.

- **Executors**

Description: Workers that run the tasks assigned by the driver program.

Role: Execute code and report the status of computation and storage.

- **Tasks**

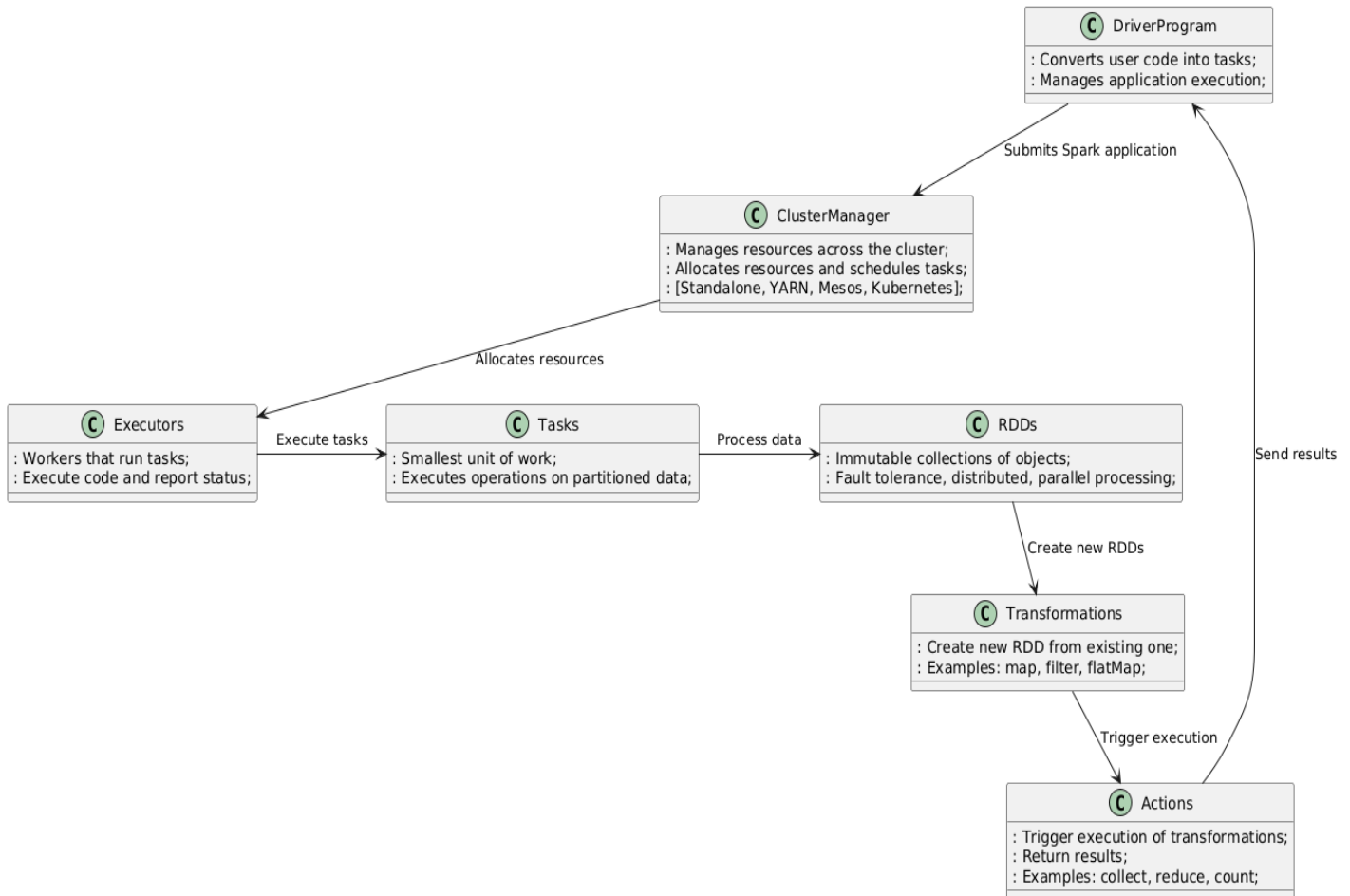
Description: The smallest unit of work in Spark.

Role: Executes individual operations on the partitioned data.



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Apache Spark Architecture



Data Processing in Spark

- **RDDs (Resilient Distributed Datasets)**

Description: Immutable collections of objects that can be processed in parallel.

Features: Fault tolerance, distributed processing, and parallel execution.

- **Transformations**

Description: Operations that create a new RDD from an existing one.

Examples: map, filter, flatMap.

- **Actions**

Description: Operations that trigger the execution of transformations and return a result.

Examples: collect, reduce, count.

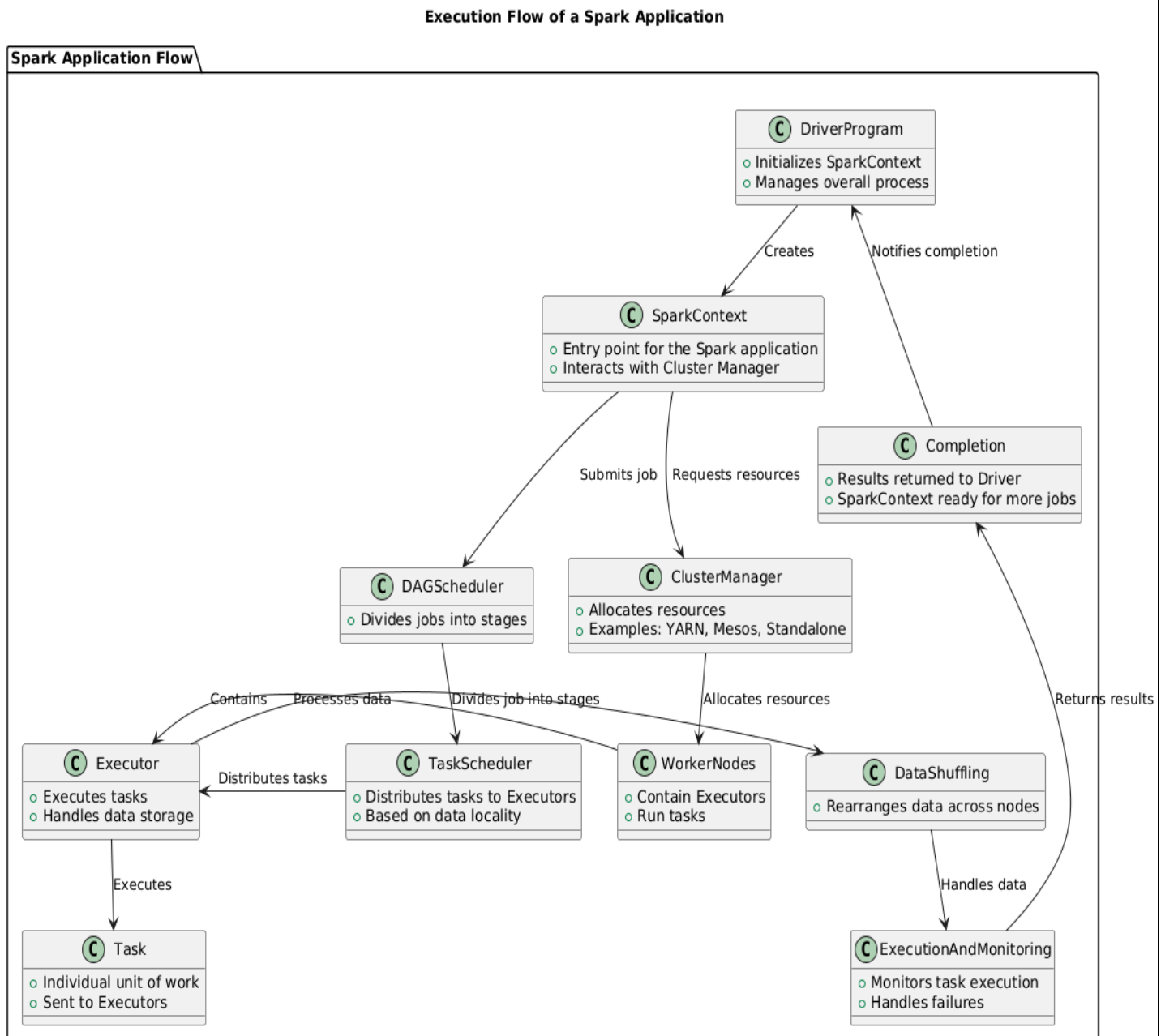


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Example Workflow

1. The Driver Program submits a Spark application.
2. The Cluster Manager allocates resources across the cluster.
3. Executors run on worker nodes, executing tasks on the data.
4. Transformations create new RDDs, and Actions trigger the execution.
5. Results are sent back to the Driver Program.

Execution Flow on Spark Application



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The execution flow in Apache Spark outlines how data processing occurs from the initial job submission to the final result collection. Here's a step-by-step breakdown:

1. SparkContext Creation

- The driver program starts and initializes the SparkContext.
- This context serves as the main entry point for Spark functionality and manages the entire Spark application.

2. Cluster Manager Interaction

- The SparkContext interacts with the Cluster Manager (e.g., YARN, Mesos, or Standalone).
- It requests resources (CPU, memory) needed for the application to run.

3. Job Submission

- The user defines transformations and actions within the Spark application.
- A job is created when an action is called (e.g., collect, count).
- At this point, Spark begins to build a logical execution plan.

4. DAG Scheduler

- The Directed Acyclic Graph (DAG) Scheduler takes the job and breaks it down into stages.
- Each stage corresponds to a set of transformations that can be executed in parallel.

5. Task Scheduler

- The Task Scheduler takes the stages defined by the DAG Scheduler and converts them into tasks.
- It distributes these tasks to the executors based on data locality, optimizing resource usage by minimizing data transfer.

6. Executor Execution

- The tasks are executed on the worker nodes by the executors.
- Executors process the data according to the tasks assigned and handle intermediate data storage as required.



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7. Data Shuffling

- If operations like reduceByKey or join are performed, data shuffling may occur.
- This involves redistributing data across the cluster, which can be resource-intensive.

8. Execution and Monitoring

- The driver program continuously monitors the execution of tasks.
- It handles any failures that may occur during processing, re-executing tasks if necessary.

9. Completion

- Once all tasks are completed, the results are sent back to the driver program.
- The SparkContext is now ready to process more jobs, maintaining a seamless workflow.

Conclusion

Apache Spark's architecture is designed to handle large-scale data processing efficiently and effectively. Understanding its components and workflow can help you leverage its full potential for your big data projects.

Important Interview Question from previous post

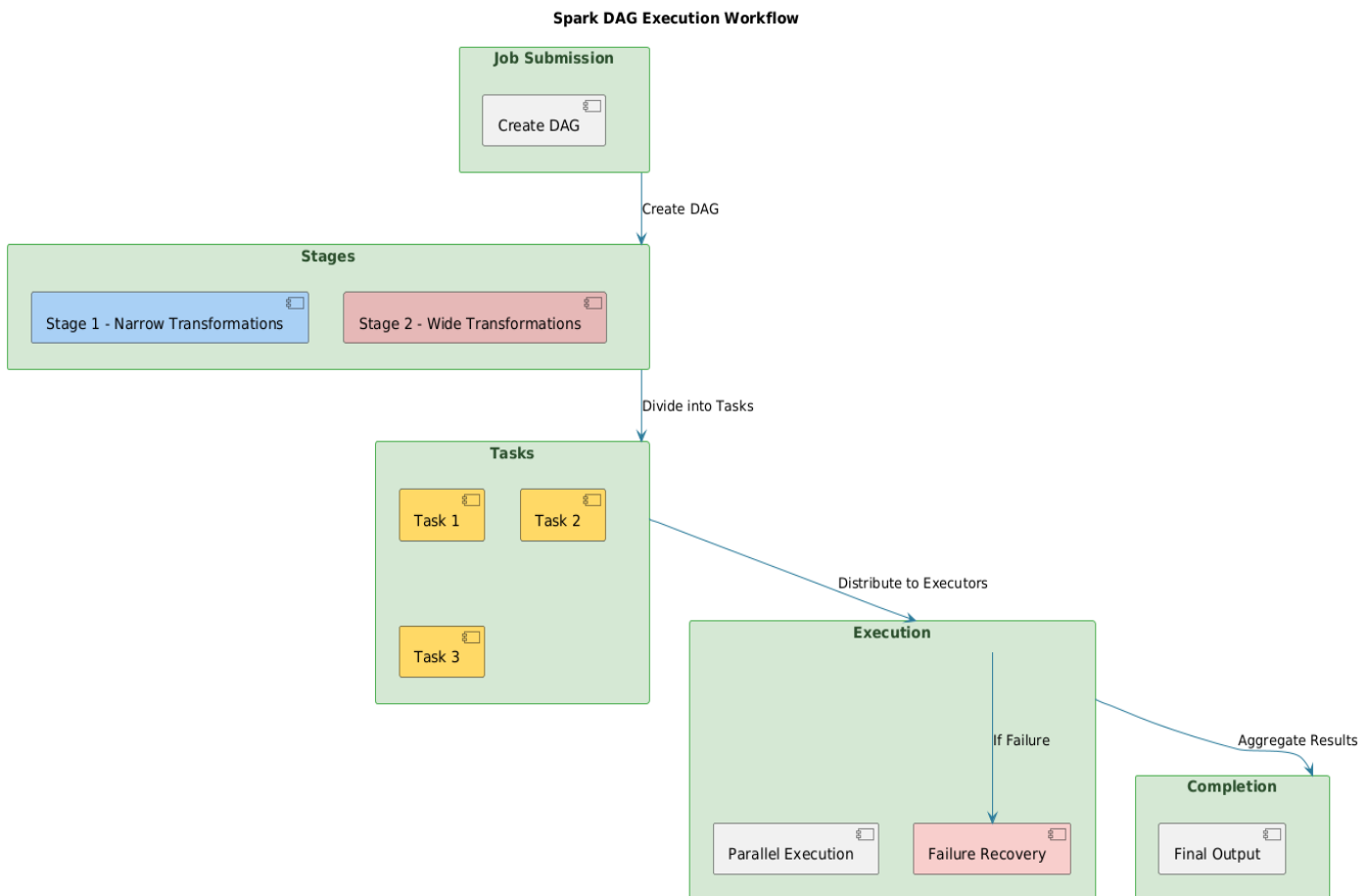
- 1. Explain Hadoop Architecture?**
- 2. How MapReduce Works?**
- 3. Difference between MapReduce and Spark?**
- 4. Why Spark is better than MapReduce?**
- 5. What are the components of Spark?**
- 6. What you mean by JVM in Spark and what are its component?**
- 7. What is use of Driver node and Work Node?**
- 8. Explain Spark Architecture?**
- 9. Explain the flow of execution in Spark?**



Master Spark Concepts Zero to Big Data Hero:

What is Spark Directed Acyclic Graph (DAG)

In Apache Spark, the Directed Acyclic Graph (DAG) is the framework that underpins how Spark optimizes and executes data processing tasks. Below is a breakdown of key concepts:



1. What is a DAG?

- **Definition:**

A DAG represents the series of operations (transformations) Spark performs on data.

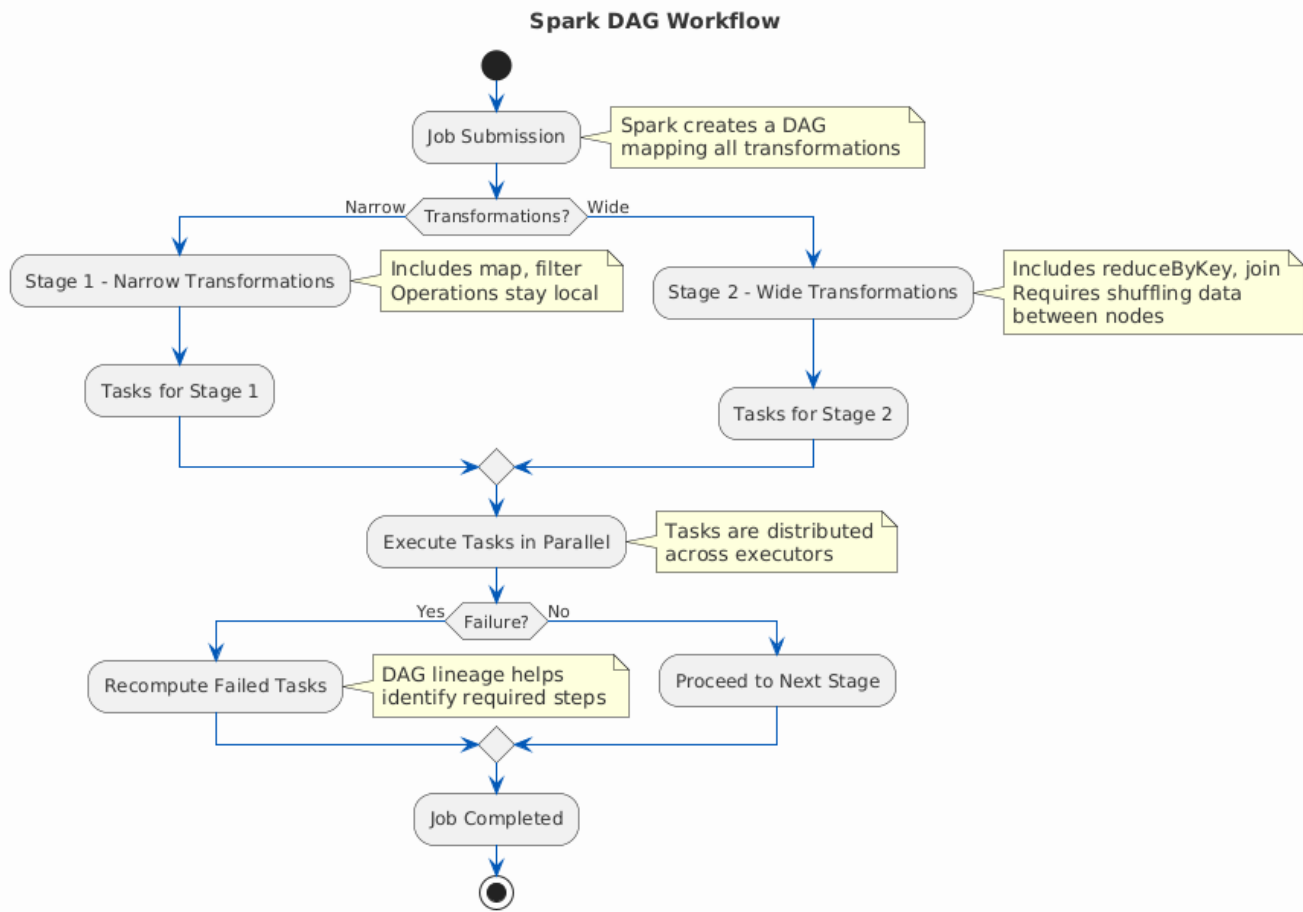
- **Components:**

- **Nodes:** Represent transformation steps (e.g., filter, map).
- **Edges:** Show the flow of data between transformations.



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2. How DAG Works in Spark



a. Job Submission

- When a Spark job is submitted, Spark creates a DAG that maps out the sequence of steps to execute the job.
- This DAG provides a visual and logical representation of how data is processed.

b. Stages

- Spark divides the DAG into multiple **stages** based on data shuffling requirements.
- **Stage Classification:**
 - **Narrow Transformations:** Operations like map and filter that don't require shuffling data between partitions.
 - Grouped within the same stage.
 - **Wide Transformations:** Operations like reduceByKey and join that require data shuffling across nodes.
 - Define boundaries between stages.



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c. Task Execution

- Each stage is further broken into **tasks**, which are distributed across nodes (executors) to execute in parallel.
- Tasks ensure that the workload is balanced for efficient execution.

d. Handling Failures

- If a task or stage fails, the DAG allows Spark to:
 - Identify the failed components.
 - Re-execute only the affected tasks or stages, saving computation time.

3. Why is DAG Important?

a. Efficiency

- DAG enables Spark to optimize task execution by:
 - Minimizing data shuffling.
 - Combining transformations to reduce redundant computations.

b. Recovery

- The DAG maintains **lineage** of operations, allowing Spark to:
 - Recompute only the necessary parts in case of a failure.

c. Speed

- By enabling parallel task execution and scheduling, DAG ensures faster data processing.

4. Key Advantages of Spark DAG

1. **Task Optimization:** Ensures efficient resource usage by structuring transformations.
2. **Parallelism:** Breaks jobs into tasks that can run in parallel across nodes.
3. **Error Handling:** Facilitates partial recomputation for failures, reducing recovery time.
4. **Transparency:** Provides a clear structure of operations, aiding debugging and analysis.

5. Key Terms to Remember

- **Transformations:** Operations performed on data (e.g., map, filter, reduceByKey).
- **Stages:** Logical segments of a DAG determined by transformations and shuffling.
- **Tasks:** Units of work derived from a stage that are executed by executors.

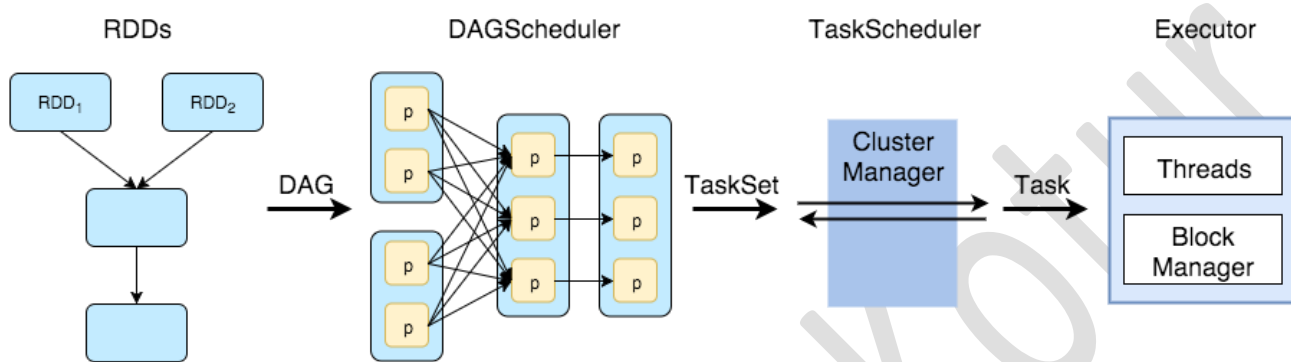


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Master Spark Concepts Zero to Big Data Hero:

How does DAG scheduler work?

The **DAG Scheduler** in Spark is responsible for **job execution planning**. It transforms a **logical execution plan** into a **physical execution plan** by dividing the work into **stages** and **tasks**. It ensures efficient task distribution, fault tolerance, and optimized execution.



Key Responsibilities of DAG Scheduler

1. Convert Logical Plan to Physical Plan:

- Spark's Catalyst optimizer generates a logical plan for transformations.
- The DAG Scheduler converts this logical plan into a physical execution plan (stages and tasks).

2. Divide Work into Stages:

- Determines **boundaries** for stages based on wide transformations (e.g., shuffle operations like `groupBy`, `join`).
- Each stage consists of tasks that can execute in parallel.

3. Manage Task Execution:

- Sends tasks to the **Task Scheduler**, which assigns them to executors.
- Ensures tasks run efficiently by using available resources.

4. Handle Failures:

- If a task fails, the DAG Scheduler recomputes only the failed tasks by tracing back the **lineage**.



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How the DAG Scheduler Chooses Jobs, Stages, and Tasks

1. Job:

- A job is created when an **action** (e.g., count, save, collect) is called on a Spark DataFrame or RDD.
- Example:
 - df.write.csv() triggers one job.

2. Stage:

- Stages are created based on **shuffle boundaries**:
 - **Narrow Transformations** (e.g., map, filter): Operations that do not require data shuffling are grouped into the same stage.
 - **Wide Transformations** (e.g., reduceByKey, groupBy): Operations requiring data shuffling (moving data across partitions) split the DAG into multiple stages.

3. Task:

- Each stage is divided into **tasks**, one for each data partition.
- Tasks are units of execution sent to executors.
- Example:
 - If there are 3 partitions in the dataset, Stage 1 will have 3 tasks.

How Stages Increase in the DAG

1. Wide Transformations Add Shuffle Boundaries:

- Wide transformations create dependencies between partitions, requiring Spark to shuffle data between nodes.
- Each shuffle creates a **new stage**.

2. Example:

- Reading and filtering: **Stage 1** (narrow transformations, no shuffle).
- Grouping data by key: **Stage 2** (wide transformation with shuffle).
- Writing results: Often merged into the final stage if it doesn't involve further shuffling.



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Detailed Example

Problem:

- Read a dataset.
- Filter rows where amount > 500.
- Group by category and calculate total sales.
- Write results to a file.

Execution Breakdown:

1. Job:

- The write action triggers **1 job**.

2. Stage Division:

- **Stage 1:** Includes the **read** and **filter** transformations. These are narrow transformations.
- **Stage 2:** Includes the **groupBy** and **write**. The **groupBy** is a wide transformation that introduces a shuffle.

3. Task Division:

- Each stage will have as many tasks as there are partitions in the dataset.
- Example: For a dataset with 3 partitions:
 - **Stage 1:** 3 tasks for filtering.
 - **Stage 2:** 3 tasks for grouped computation and writing.

Summary

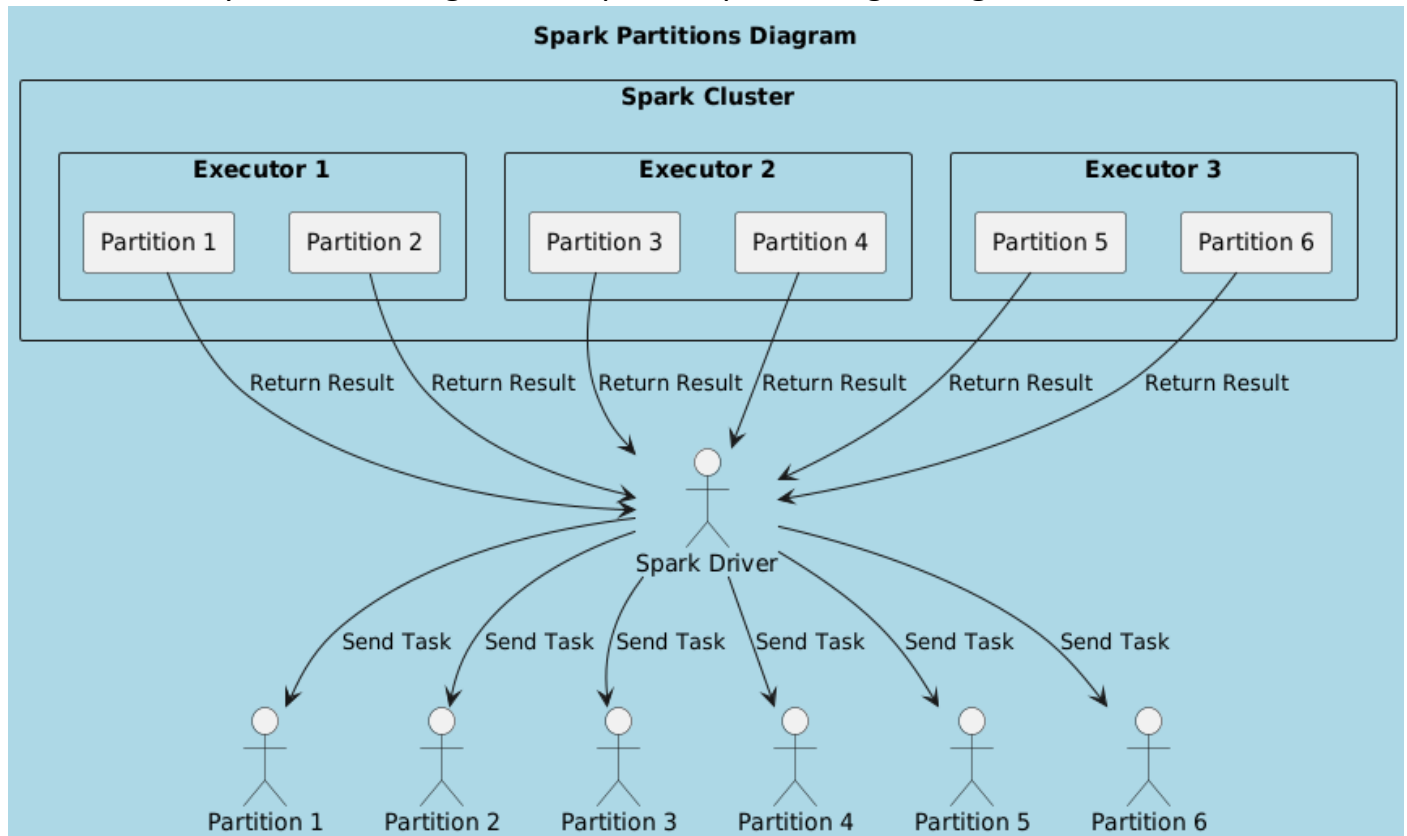
- **Job:** Represents the entire execution triggered by an action.
- **Stages:** Increase with wide transformations requiring shuffle boundaries.
- **Tasks:** Represent execution units corresponding to partitions.



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Deep Dive into Partition in Spark

A **partition** in Spark is a fundamental unit of parallelism. It represents a logical chunk of data that can be processed independently. Each partition is processed on a different node in a distributed system, enabling efficient parallel processing of large datasets.



Key Information on Partitions in Spark:

1. What is a Partition?

- A partition is a subset of the data stored in memory or on disk, allowing Spark to process large datasets in chunks across different nodes. The more partitions you have, the better the parallelism.

2. How Are Partitions Created?

- Partitions are created based on the data's source, like HDFS blocks or RDD transformations. Spark also re-partitions data when performing operations like shuffling or sorting.

3. Optimizing Partitioning:

- Proper partitioning is crucial to prevent **data skew** and ensure **balanced workload distribution**.
- Use **repartition()** to increase partitions or **coalesce()** to reduce them based on your data size and processing needs.

4. Partition Size:

- Aim for partition sizes between **128 MB to 1 GB**. Smaller partitions can cause overhead from too many tasks, while overly large partitions can result in **out-of-memory (OOM)** errors.

5. Default Partitioning:

- By default, Spark creates **200 shuffle partitions**, but this can be adjusted using `spark.sql.shuffle.partitions`.

6. Persistence and Partitioning:

- You can cache or persist DataFrames/RDDs by partitions in memory or on disk for reuse during iterative tasks, improving performance.

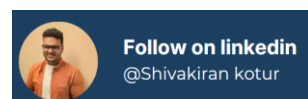
Why is Partitioning Important?

Efficient partitioning ensures:

- ✚ **Better Parallelism:** More partitions allow Spark to leverage multiple cores effectively.
- ✚ **Improved Performance:** Well-distributed partitions prevent bottlenecks from skewed data.
- ✚ **Optimized Resource Use:** Properly sized partitions reduce memory and disk overhead, speeding up operations.

By understanding and managing partitions well, you can significantly boost the performance of your Spark jobs!

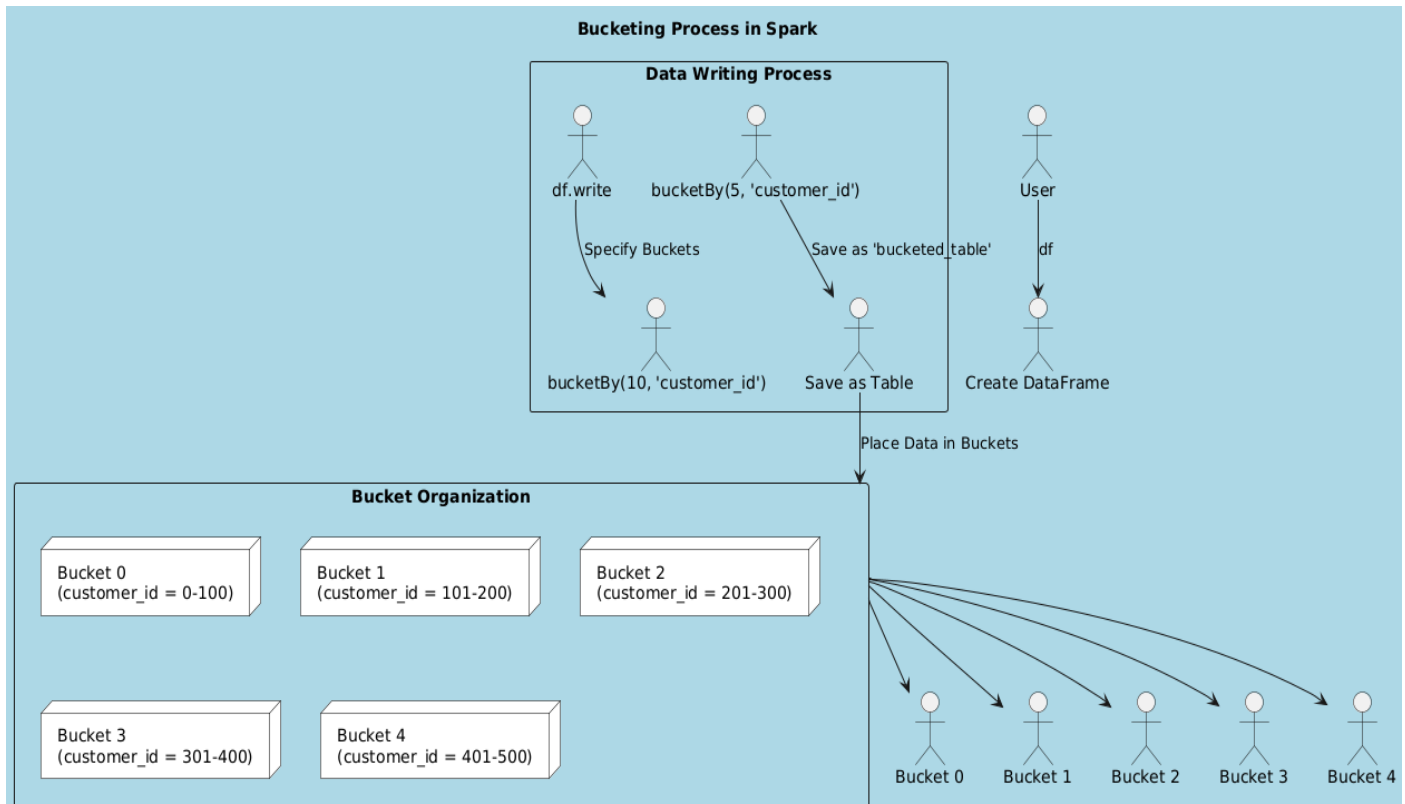
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Deep Dive into Bucketing in Spark

Bucketing in Spark: Key Concepts

Bucketing is a technique in Spark used to optimize query performance by physically dividing the data into fixed-size buckets based on the values of one or more columns. This helps reduce the amount of data shuffled during join or aggregation operations.



Key Concepts of Bucketing:

1. What is Bucketing?

- Bucketing distributes data into a predefined number of equal-sized buckets based on the values of specific columns (bucket keys). It's particularly useful for optimizing queries involving joins and aggregations on large datasets.

2. How is it Different from Partitioning?

- **Partitioning** splits data into different directories based on column values, while **bucketing** divides data into fixed-number buckets within each partition. Bucketing allows for more fine-grained control over how data is divided.
- In partitioning, there are as many partitions as there are distinct values, whereas in bucketing, the number of buckets is predefined.



3. Use Cases for Bucketing:

- Bucketing is ideal for **joining large tables or performing aggregations** on columns with a lot of distinct values.
- For example, if you often join tables on a customer_id column, bucketing both tables by customer_id can significantly reduce shuffling and speed up joins.

4. Creating Buckets:

- You can create a bucketed table in Spark by specifying the number of buckets and the column(s) to bucket on:

```
df.write.bucketBy(4,  
"customer_id").sortBy("customer_id").saveAsTable("bucketed_table")
```

5. Advantages of Bucketing:

- **Reduced Shuffling:** Bucketing minimizes data movement during joins by ensuring that data with the same key is placed in the same bucket.
- **Optimized Queries:** Bucketing enables faster query performance, especially for joins and aggregations.

6. Limitations of Bucketing:

- Once the data is bucketed, it cannot be dynamically adjusted. You need to determine the correct number of buckets ahead of time.
- Bucketing is a more static optimization compared to **Adaptive Query Execution (AQE)**, which dynamically adjusts partitions at runtime.

Can You Use Bucketing in Databricks?

Yes, bucketing is supported in **Databricks** and can be very effective for large-scale data processing, especially when you're frequently joining large datasets. Databricks allows you to leverage Spark's bucketing mechanism to optimize performance for workloads involving heavy joins and aggregations.

In Databricks, you can:

- Use **bucketing** to optimize the performance of your jobs by reducing shuffles during joins or aggregations.
- Combine **bucketing** with **Delta Lake** to optimize large datasets, ensuring better read performance and reducing execution time for analytical queries.

Example of bucketing in Databricks:

```
# Enable bucketing and sort
spark.conf.set("spark.sql.bucketing.enabled", "true")

# Create a bucketed table
df.write.bucketBy(10,
"customer_id").sortBy("customer_id").saveAsTable("bucketed_table")
```

- + **df.write**: Initiates the process of writing the DataFrame df to a table.
- + **.bucketBy(10, "customer_id")**:
 - **10**: Specifies the number of buckets to create.
 - **"customer_id"**: Indicates the column used for bucketing. All records with the same customer_id will be placed in the same bucket.
- + **.sortBy("customer_id")**: Sorts the data within each bucket by customer_id, optimizing query performance for future reads.
- + **.saveAsTable("bucketed_table")**: Saves the bucketed and sorted data as a table named bucketed_table for later querying.

Best Practices for Bucketing in Databricks:

- Choose an appropriate number of buckets based on your data volume and query patterns.
- Make sure to use the same number of buckets for both tables in joins to avoid shuffle operations.
- Combine with **partitioning** and **Delta Lake** for better performance in complex workflows.

By using bucketing in Databricks, you can ensure more efficient data processing, especially for large-scale joins and aggregations!

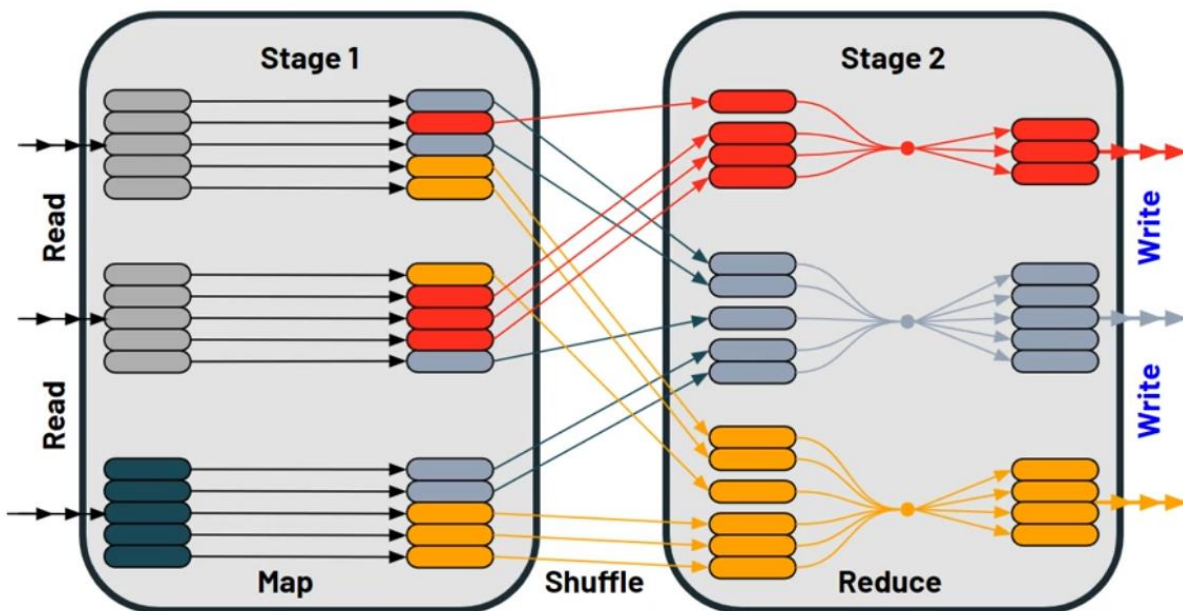
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Deep Dive into Shuffling in Spark

What is Shuffling?

Shuffling in Spark is the process of **redistributing data** across different partitions on various nodes to meet the needs of certain transformations (like `groupByKey()`, `join()`, `distinct()`, etc.). It occurs when data in one partition is required in another, often due to operations that aggregate, reorder, or repartition data.



How Shuffling Works:

- Data Redistribution:** For example, if you're summing up values by a key in a `groupByKey()`, Spark must collect all the data with the same key from different partitions and bring them together in one partition.
- Disk and Network Transfer:** When Spark performs a shuffle, the data is written to disk, serialized, and transferred across the network to another partition on another node. This introduces disk I/O and network traffic, which slows down the processing.

Key Challenges of Shuffling:

- I/O and Network Bottlenecks:** Since shuffling requires data to be written to disk and transferred between nodes, it leads to high disk I/O and network usage, slowing down performance.
- Increased Garbage Collection:** Large shuffles can cause high memory usage, leading to frequent garbage collection pauses, which impact overall processing efficiency.

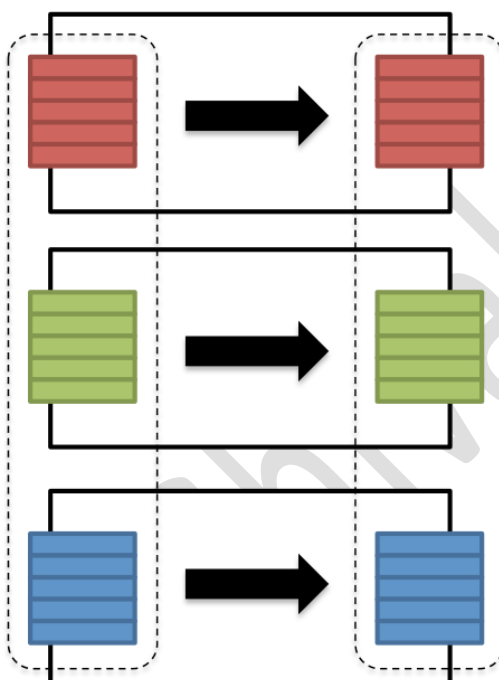
- **Stragglers:** Skewed data can cause certain partitions to be much larger than others, causing some tasks to take much longer, creating performance imbalances, often referred to as "stragglers."

When Does Shuffling Occur?

- **groupByKey():** Requires all values for the same key to be shuffled to the same node.
- **join():** Data from two DataFrames/RDDs needs to be aligned across partitions based on keys, causing a shuffle.
- **repartition():** Explicitly redistributes data into a different number of partitions.
- **distinct():** Needs to shuffle data to ensure that duplicate records are removed.

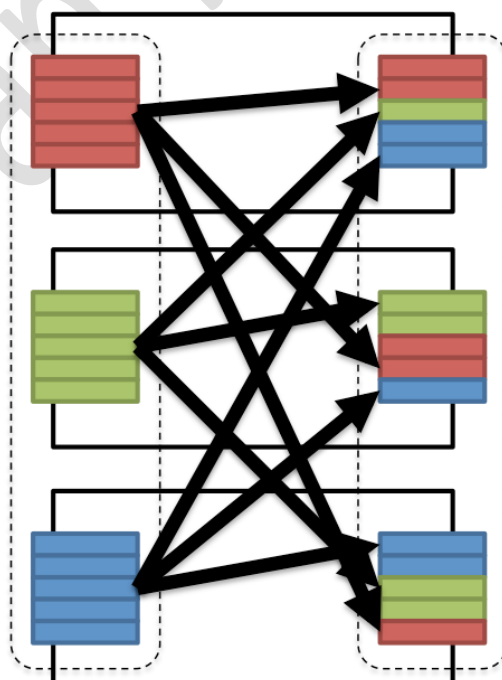
Narrow transformation

- Input and output stays in same partition
- No data movement is needed



Wide transformation

- Input from other partitions are required
- Data shuffling is needed before processing



Why Shuffling Slows Down Processing:

- **Breaks In-Memory Processing:** Spark typically tries to keep data in memory to process faster, but during a shuffle, it has to **write to disk and transfer over the network**, causing a performance hit.

- **Large Shuffle Data:** If the shuffle results in large volumes of data being moved, it can lead to **longer network transfers** and **disk write times**, impacting the speed and causing **out-of-memory (OOM)** errors.

Optimizing Shuffling:

1. Avoid Unnecessary Shuffles:

- Prefer **reduceByKey()** over **groupByKey()**, since it reduces data before the shuffle, minimizing the data size.
- Use **mapPartitions()** or **combineByKey()** to operate on data within partitions to reduce the need for shuffles.

2. Tune Shuffle Partitions:

- The number of shuffle partitions can be controlled using **spark.sql.shuffle.partitions** (default is 200). Increasing this value for larger datasets can reduce the size of each shuffle partition, which may help distribute work more evenly across nodes.
- For smaller datasets, reducing this value can avoid too many small tasks being created, leading to better parallelism.

Check or Set Shuffle Partitions:

```
# Check the current shuffle partitions setting
print(spark.conf.get('spark.sql.shuffle.partitions'))

# Set a new value for shuffle partitions
spark.conf.set("spark.sql.shuffle.partitions", "500")
```

3. Manage File Partition Sizes:

- Control how large a single partition file can be before being split using **spark.sql.files.maxPartitionBytes**. For example:

```
print(spark.conf.get('spark.sql.files.maxPartitionBytes'))
```

4. Data Skew Handling:

- Spark's **Adaptive Query Execution (AQE)** can automatically handle skewed data by splitting large partitions, reducing the impact of uneven workloads.
- **Skewed join optimizations** can be applied to detect and redistribute skewed partitions.

Key Metrics to Monitor Shuffling:

- **Shuffle Write and Read Time:** In Spark UI, you can track how much time is spent writing shuffle data to disk and reading it back. Long shuffle times are a good indicator that your application is being bottlenecked by shuffling.
- **Task Execution Time:** Look for tasks that take disproportionately longer than others (stragglers), which can often be linked to shuffle bottlenecks or data skew.

Practical Example:

Consider a scenario where you're joining two large datasets in Spark. If one dataset is heavily skewed, meaning one partition holds most of the data, Spark will shuffle the data, and that skewed partition will take much longer to process. By enabling AQE and letting it adjust shuffle partitions dynamically, you can significantly speed up the join by balancing the workload more efficiently.

Summary:

- **Shuffling** is necessary when Spark needs to **redistribute data** between partitions.
- It can cause performance bottlenecks due to **disk I/O** and **network transfers**.
- Optimizing shuffling involves **reducing unnecessary shuffles**, **tuning partition sizes**, and leveraging **AQE** to handle skewed data efficiently.

By managing shuffles effectively, you can **boost Spark performance** and ensure that large-scale data processing tasks run smoothly and efficiently.

Master Spark Concepts Zero to Big data Hero:

Lazy Evaluation in PySpark: Unlocking Performance Optimizations

Lazy evaluation is one of the core features of **PySpark** that makes it incredibly efficient and fast, even when handling huge datasets. But what exactly is it? Let's break it down with a real-world example.

What is Lazy Evaluation?

In PySpark, transformations like `filter()`, `map()`, and `groupBy()` **don't execute immediately**. Instead, they are added to a logical plan, and the computations only happen when an **action** like `show()` or `save()` is called.

Example: Word Count from a Text File

Let's count the words longer than 3 characters in a text file, which is a great example of lazy evaluation in action.

```
from pyspark.sql import SparkSession

# Initialize SparkSession spark =
SparkSession.builder \
.appName("Word Count Example") \
.getOrCreate()

# Load data from a text file data =
spark.read.text("hdfs://path/to/file.txt")

# Split lines into words, filter words longer than three characters, and create
pairs word_counts = (data.selectExpr("explode(split(value, ' ')) as word")
    .filter("length(word) > 3")
    .groupBy("word")
    .count())

# Show the word counts word_counts.show()

# Save the result to a text file
word_counts.write.mode("overwrite").csv("hdfs://path/to/output")
# Stop the Spark session
#spark.stop()
```



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How Lazy Evaluation Works:

1. **Data Loading:** When we load the data with `spark.read.text()`, Spark just **builds a logical plan**. No data is read at this point. No computation happens yet!
2. **Transformations:** We define transformations (`split`, `explode`, `filter`, `groupBy`). They **don't execute immediately** but are stored in the logical plan for later.
3. **Action Triggers Execution:** The first action (`word_counts.show()`) kicks off the execution. This is where Spark evaluates the transformations and optimizes the plan.
4. **Optimization:** Spark combines transformations, minimizes data shuffling, and reorders operations to **execute efficiently**. This could involve pushing down filters or reducing intermediate steps.
5. **Execution:** After optimization, Spark processes the data step by step based on the logical plan, fetching and transforming the data in the most efficient way possible.
6. **Subsequent Actions:** Any following actions (like `write()`) will follow the **optimized execution plan**, ensuring everything runs as efficiently as possible.

Why Lazy Evaluation is So Powerful:

- **Efficiency:** Only the necessary computations are executed, cutting down on unnecessary resource usage.
- **Fault Tolerance:** If a failure happens, Spark can **recompute** the lost data by re-running the logical plan.
- **Optimization:** By delaying execution, Spark optimizes the entire sequence, reducing time-consuming tasks like shuffling and redundant transformations.

A Real-Life Analogy:

Think of lazy evaluation like cooking a meal. Instead of cooking each ingredient immediately, you **plan out the steps** first, and only start cooking when you have everything in place. This ensures that you're not wasting time or energy on unnecessary steps!

"In PySpark, transformations such as `filter()` and `map()` are lazy, meaning they don't execute until an action is performed. This allows Spark to optimize queries by building a Directed Acyclic Graph (DAG) of operations."

 **Takeaway:** Lazy evaluation in PySpark is a game changer. It optimizes the computation flow and ensures efficient execution, even with large datasets.

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Introduction to RDD:

What is RDD?

RDD stands for Resilient Distributed Dataset. RDDs are the core data structure in Apache Spark, designed for fault-tolerant, distributed processing. They represent an immutable, distributed collection of objects that allows users to perform transformations and actions on data across multiple nodes in a Spark cluster.

RDDs allow parallel processing of data, which is critical for handling large datasets efficiently. Spark provides a programmer's interface (API) to work with RDDs through simple functions.

Key Characteristics of RDDs

- Immutable:** Once created, RDDs cannot be modified, only transformed into new RDDs.
- Partitioned:** Data in RDDs is automatically split across different nodes in the cluster, allowing distributed data processing.
- Fault-Tolerant:** RDDs are designed to handle node failures gracefully. If data is lost, RDDs automatically recover it using lineage information.
- Lazy Evaluation:** Transformations on RDDs are not executed until an action is called, which optimizes performance by minimizing data movement.
- In-Memory Computation:** Spark processes data in-memory, leading to faster computation speeds compared to traditional data processing frameworks.

RDD Operations

RDD operations are divided into two main categories:

- Transformations:** These operations are used to create a new RDD from an existing RDD. They are *lazy* operations, meaning they don't execute until an action is called.
 - Examples: `map`, `filter`, `flatMap`, `groupByKey`, `reduceByKey`
- Actions:** These operations trigger the computation of transformations and produce results, either by returning values to the driver program or by saving output.
 - Examples: `collect`, `count`, `take`, `saveAsTextFile`

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Important Terms

Term	Definition
RDD	Resilient Distributed Dataset
Transformation	Operation that produces a new RDD
Action	Operation that produces a result or local object
Spark Job	Sequence of transformations with a final action

Creating an RDD

You can create an RDD in Spark using one of the following methods:

1. From an In-Memory Collection:

```
# Create RDD from a list  
data = [1, 2, 3, 4, 5]  
rdd = sc.parallelize(data)
```

2. From an External Data Source:

```
# Create RDD from a text file  
rdd = sc.textFile("path/to/sales_data.txt")
```

RDD Transformations

- **Transformations:** These create a new RDD by applying a function to elements of an existing RDD. They are *lazy*, meaning they are not computed until an action is performed.

1. **map:** Applies a function to each element in the RDD, producing a new RDD with transformed values.

Example: squared_rdd squares each element of rdd.

2. **filter:** Selects elements that match a condition, returning a new RDD with elements that satisfy the condition. Example: filtered_rdd keeps only even numbers.



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3. **flatMap**: Similar to map, but allows returning multiple elements per input element, making it useful for tasks like tokenizing text. Example: words_rdd splits lines into words.
4. **reduceByKey**: Aggregates data for each key, making it ideal for counting occurrences or summing values by key. Example: counts_rdd counts occurrences of each word.
5. **groupByKey**: Groups elements by key, resulting in pairs of (key, iterable of values). Example: grouped_rdd groups data based on keys in rdd.

```
# Applying map transformation to square each element
squared_rdd = rdd.map(lambda x: x ** 2)

# Applying filter transformation to keep even numbers
filtered_rdd = rdd.filter(lambda x: x % 2 == 0)

# Applying flatMap transformation to split lines into words
words_rdd = lines_rdd.flatMap(lambda line: line.split(" "))

# Using map and reduceByKey to count occurrences of each word
counts_rdd = words_rdd.map(lambda word: (word,
1)).reduceByKey(lambda a, b: a + b)

# Applying groupByKey to group elements by their key
grouped_rdd = rdd.groupByKey()
```

RDD Actions

- **Actions**: Unlike transformations, actions trigger the execution of the RDD's computations, returning a result to the driver or saving it externally.
1. **collect**: Retrieves all elements in the RDD as a list. Ideal for small datasets, as it brings data to the driver.
 - Example: data contains all elements in rdd.
 2. **count**: Returns the total number of elements in the RDD.
 - Example: total_elements gives the count of items in rdd.
 3. **take**: Fetches the first n elements of the RDD.
 - Example: first_elements retrieves the first 3 elements of rdd.



4. **saveAsTextFile**: Saves the RDD content to a specified file path, writing each partition as a separate text file.
 - Example: rdd.saveAsTextFile stores rdd data to the given path.
5. **reduce**: Aggregates the elements of the RDD using a specified binary function, commonly used for summing or multiplying elements.
 - Example: sum computes the total by summing all elements in rdd.

```
# Collects all elements of the RDD into a list
data = rdd.collect()

# Counts the number of elements in the RDD
total_elements = rdd.count()

# Takes the first 3 elements of the RDD
first_elements = rdd.take(3)

# Saves the RDD as a text file at the specified path
rdd.saveAsTextFile("path/to/output")

# Reduces the RDD elements by summing them up
sum = rdd.reduce(lambda x, y: x + y)
```

Summary

- **RDDs** are immutable, fault-tolerant, and partitioned collections that support distributed processing.
- **Transformations** create new RDDs but do not compute until an action is called.
- **Actions** trigger computation and produce results.
- RDDs allow Spark to handle large datasets across distributed clusters effectively, enabling parallel processing.

By understanding and using RDDs effectively, you can harness the full power of Spark for large-scale data processing.

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RDD part 2:

Example: Sales Data Analysis with RDDs

Suppose we're analyzing a dataset of sales records, where each record includes transaction details: transaction_id, product, price.

```
# Initialize SparkContext (only needed if not using an existing
# Spark session)
from pyspark import SparkContext

# Define data as a list of tuples
data = [
    (1, "Apple", 2.50),
    (2, "Banana", 1.00),
    (3, "Apple", 2.50),
    (4, "Orange", 3.00),
    (5, "Banana", 1.00)
]

# Create an RDD using parallelize
sales_rdd = sc.parallelize(data)

# Display the RDD's content
for record in sales_rdd.collect():
    print(record)
```

```
(1, 'Apple', 2.5)
(2, 'Banana', 1.0)
(3, 'Apple', 2.5)
(4, 'Orange', 3.0)
(5, 'Banana', 1.0)
```

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Explanation

- **sc.parallelize(data)**: This command creates an RDD from the given data list, which represents sales records.
- **sales_rdd.collect()**: This action collects all elements of the RDD and returns them as a list, allowing you to print and inspect the RDD contents.

```
# Sample data as a list of strings, simulating lines in a text file
sales_data = [
    "1,Apple,2.50",
    "2,Banana,1.00",
    "3,Apple,2.50",
    "4,Orange,3.00",
    "5,Banana,1.00"
]

# Step 1: Create an RDD using sc.parallelize
rdd = sc.parallelize(sales_data)

# Print original RDD
print("Original RDD:")
print(rdd.collect())
```

```
Original RDD:
['1,Apple,2.50', '2,Banana,1.00', '3,Apple,2.50', '4,Orange,3.00', '5,Banana,1.00']
```

```
# Step 2: Parse each line into a tuple of (product, price)
product_price_rdd = rdd.map(lambda line: line.split(","))
                           .map(lambda parts: (parts[1],
float(parts[2])))

# Output after parsing
print("\nParsed Product and Price Data:")
print(product_price_rdd.collect())
```

```
Parsed Product and Price Data:
[('Apple', 2.5), ('Banana', 1.0), ('Apple', 2.5), ('Orange', 3.0), ('Banana', 1.0)]
```

```
# Step 3: Filter Records - Only "Apple" Sales
apple_sales_rdd = product_price_rdd.filter(lambda record: record[0]
== "Apple")
```

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```
# Output after filtering
print("\nApple Sales Records:")
print(apple_sales_rdd.collect())
```

```
Apple Sales Records:
[('Apple', 2.5), ('Apple', 2.5)]
```

```
# Step 4: Aggregate Data - Calculate Total and Average Sales for
Each Product
```

```
# Map each product with (price, 1) for counting
product_price_count_rdd = product_price_rdd.map(lambda record:
(record[0], (record[1], 1)))
```

```
# Output after counting
```

```
print("\nProduct Price Count RDD:")
print(product_price_count_rdd.collect())
```

```
Product Price Count RDD:
```

```
[('Apple', (2.5, 1)), ('Banana', (1.0, 1)), ('Apple', (2.5, 1)), ('Orange', (3.0, 1)), ('Banana', (1.0, 1))]
```

```
# Calculate total sales and count for each product
```

```
product_totals_rdd = product_price_count_rdd.reduceByKey(lambda a,
b: (a[0] + b[0], a[1] + b[1]))
```

```
# Output after reduction
```

```
print("\nTotal Sales and Count by Product:")
print(product_totals_rdd.collect())
```

```
Total Sales and Count by Product:
```

```
[('Orange', (3.0, 1)), ('Apple', (5.0, 2)), ('Banana', (2.0, 2))]
```

```
# Calculate average price for each product
```

```
average_price_rdd = product_totals_rdd.mapValues(lambda total:
total[0] / total[1])
```

```
# Output after calculating average prices
```

```
print("\nAverage Sales Price by Product:")
print(average_price_rdd.collect())
```

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```
Average Sales Price by Product:
```

```
[('Orange', 3.0), ('Apple', 2.5), ('Banana', 1.0)]
```

```
# Step 5: Actions to Display Results
```

```
# Total sales revenue by product
```

```
print("\nFinal Sales Revenue by Product:")
for product, totals in product_totals_rdd.collect():
    print(f'{product}: Total Revenue = ${totals[0]:.2f}, Count = {totals[1]}')
```

```
Final Sales Revenue by Product:
```

```
Orange: Total Revenue = $3.00, Count = 1
```

```
Apple: Total Revenue = $5.00, Count = 2
```

```
Banana: Total Revenue = $2.00, Count = 2
```

```
# Average sales price by product
```

```
print("\nFinal Average Sales Price by Product:")
for product, avg_price in average_price_rdd.collect():
    print(f'{product}: Average Price = ${avg_price:.2f}')
```

```
Final Average Sales Price by Product:
```

```
Orange: Average Price = $3.00
```

```
Apple: Average Price = $2.50
```

```
Banana: Average Price = $1.00
```

Transformations Overview

1. sc.parallelize(sales_data):

- **Description:** Creates an RDD from a list of strings representing sales records. This RDD is distributed across the nodes in the Spark cluster.
- **Purpose:** To initialize the dataset for processing.

2. `rdd.map(lambda line: line.split(",")):`

- **Description:** Transforms each line of the RDD into a list of strings, splitting on commas.
- **Purpose:** To break down each sales record into its individual components (transaction_id, product, price).

3. `.map(lambda parts: (parts[1], float(parts[2]))):`

- **Description:** Maps the split lines to tuples of (product, price), converting the price to a float.
- **Purpose:** To create a structured format for further analysis, focusing on the product and its corresponding price.

4. `.filter(lambda record: record[0] == "Apple"):`

- **Description:** Filters the RDD to include only records where the product is "Apple".
- **Purpose:** To isolate sales records for the specific product being analyzed.

5. `product_price_rdd.map(lambda record: (record[0], (record[1], 1))):`

- **Description:** Maps each record to a tuple of (product, (price, count)), where count is initialized to 1.
- **Purpose:** To prepare for aggregation by counting occurrences of each product.

6. `.reduceByKey(lambda a, b: (a[0] + b[0], a[1] + b[1])):`

- **Description:** Aggregates the price and count for each product by summing the prices and counts.
- **Purpose:** To compute total sales revenue and the number of sales for each product.

7. `.mapValues(lambda total: total[0] / total[1]):`

- **Description:** Maps the totals to calculate the average price by dividing total price by count.
- **Purpose:** To determine the average sales price for each product based on the aggregated results.

These transformations collectively allow for an effective analysis of sales data, facilitating filtering, aggregation, and calculation of average values in a distributed manner using Apache Spark RDDs.

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RDD part 3: Map, FlatMap, Filter

1. Map Transformation

- **Definition:** The map transformation applies a specified function to each element of the RDD, producing a new RDD containing the results.
- **Characteristics:**
 - It maintains the same number of elements in the output as in the input.
 - The output elements can be of a different type than the input.
 - It is a **lazy transformation**, meaning the execution does not happen until an action is called.
- **Use Cases:** Use map when you want to perform an operation on each element, such as mathematical computations, data formatting, or any transformation that produces a one-to-one correspondence of elements.
- **Example:**

```
rdd.map(lambda x: x * 2)    # Doubles each element in the RDD
```

2. FlatMap Transformation

- **Definition:** The flatMap transformation also applies a specified function to each element of the RDD but can return multiple output values for each input element. The results are then flattened into a single RDD.
- **Characteristics:**
 - The number of output elements can vary; it can produce zero or more elements for each input.
 - It is also a **lazy transformation**.
- **Use Cases:** Use flatMap when you need to split elements or return multiple results per input, such as tokenizing sentences into words or expanding lists.
- **Example:**

```
rdd.flatMap(lambda x: (x, x + 1))    # Produces each element and its successor
```

3. Filter Transformation

- **Definition:** The filter transformation selects elements from the RDD that meet a specified condition, returning a new RDD containing only the elements that satisfy the condition.
- **Characteristics:**
 - The number of elements in the output may be less than or equal to the input.
 - It is a **lazy transformation**.
- **Use Cases:** Use filter when you need to extract elements based on specific criteria, such as removing outliers or selecting records that match certain conditions.
- **Example:**

```
rdd.filter(lambda x: x > 10)      # Returns elements greater than 10
```

Summary of Differences

Transformation	Purpose	Output Characteristics
map	Applies a function to each element	Same number of elements as input
flatMap	Applies a function that returns multiple elements	Can produce zero or more elements per input; flattened output
filter	Selects elements that meet a condition	Fewer or equal elements compared to input

Conclusion

- + Understanding the differences between map, flatMap, and filter transformations is essential for effective data manipulation in Spark.
- + These transformations are foundational tools for processing and analyzing large datasets in a distributed environment, allowing developers to efficiently transform data as needed for various applications.

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How Flatmap works in RDD

In PySpark, the flatMap transformation is used to process each element in an RDD, applying a function that returns zero, one, or multiple elements. This is useful for breaking down nested structures or splitting complex data into individual elements.

Function Definition in PySpark:

flatMap(f)

- f: A function that takes an element and returns an iterable (e.g., list, set, generator).

Key Points:

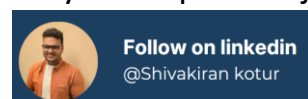
- **Purpose:** flatMap flattens nested data structures (like lists, tuples, or maps) into a single-level RDD.
- **Return Value:** A new RDD containing the flattened elements. The size of this RDD can vary depending on the function logic.
- **Processing:** Each element in the RDD is processed one at a time.

How flatMap Works:

1. Iterates through each element in the original RDD.
2. Applies the function: The user-defined function can return:
 - A single element.
 - Multiple elements (e.g., a list or set).
 - No elements (filtered out).
3. Flattens the results into a single RDD.

Common Use Cases:

- **Tokenizing Strings:** Splitting strings into individual words.
- **Flattening Nested Data:** Extracting individual elements from lists, arrays, or nested collections.
- **Expanding Key-Value Pairs:** Breaking down maps into key-value pairs or just values.



Examples:

1. Flattening Nested Lists

```
data = [[1, 2, 3], [4, 5]]  
rdd = sc.parallelize(data)  
flat_rdd = rdd.flatMap(lambda x: x)  
print(flat_rdd.collect()) # Output: [1, 2, 3, 4, 5]
```

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```
[1, 2, 3, 4, 5]
```

flatMap is used here to flatten each nested list into individual elements in a single RDD.

The expression **lambda x: x** is a simple lambda function in Python, and its purpose in the context of the code is to return the input value as it is, without any modification. Let's break it down:

Lambda Function:

In Python, a lambda function is an anonymous function defined using the `lambda` keyword. The general syntax is:

lambda arguments: expression

- **arguments:** The input parameters the function accepts (in this case, `x`).
- **expression:** The operation or expression that gets evaluated and returned when the function is called.

Breaking Down **lambda x: x:**

- **x:** This is the argument that the lambda function takes as input.
- **x (the return value):** The function then returns `x` as it is, meaning no change is made to the input value.

So, `lambda x: x` is a function that simply returns whatever is passed to it.

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Context in the Code:

Here's the relevant line in your code:

```
flat_rdd = rdd.flatMap(lambda x: x)
```

- The RDD rdd contains nested lists (e.g., [[1, 2, 3], [4, 5]]).
- When flatMap applies the lambda x: x function to each element in the RDD:
 - The input to the lambda (x) is one of the sublists ([1, 2, 3] or [4, 5]).
 - The lambda x: x function simply returns that sublist as it is, but the key part is that **flatMap will flatten these sublists** into individual elements.

Example Walkthrough:

When flatMap(lambda x: x) is applied:

1. For the first element [1, 2, 3], x is [1, 2, 3], and lambda x: x just returns [1, 2, 3].
2. For the second element [4, 5], x is [4, 5], and lambda x: x returns [4, 5].

However, flatMap **flattens** these sublists into individual elements, so the final result is [1, 2, 3, 4, 5] in a single-level RDD.

Example:

```
# Sample data: List of words
data = ["hello", "world", "spark"]

# Parallelize the data into an RDD
rdd = sc.parallelize(data)

# Use flatMap to flatten each word into individual characters
flat_rdd = rdd.flatMap(lambda word: list(word))

# Collect and print the result
print(flat_rdd.collect()) # Output: ['h', 'e', 'l', 'l', 'o', 'w', 'o', 'r', 'l', 'd', 's', 'p', 'a', 'r', 'k']
```

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```
['h', 'e', 'l', 'l', 'o', 'w', 'o', 'r', 'l', 'd', 's', 'p', 'a', 'r', 'k']
```

Explanation:

1. **Input:** The data is a list of words: ["hello", "world", "spark"].
2. **Transformation:** We apply flatMap with the lambda function lambda word: list(word):
 - For each word (e.g., "hello"), list(word) converts the word into a list of its characters.
 - flatMap then flattens all these individual characters into a single RDD.



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3. **Result:** The RDD contains the individual characters of the words: ['h', 'e', 'l', 'l', 'o', 'w', 'o', 'r', 'l', 'd', 's', 'p', 'a', 'r', 'k'].

2. Splitting Strings

```
lines = ["hello world", "how are you"]
rdd = sc.parallelize(lines)
words_rdd = rdd.flatMap(lambda line: line.split(" "))
print(words_rdd.collect()) # Output: ["hello", "world", "how", "are", "you"]
```

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```
['hello', 'world', 'how', 'are', 'you']
```

flatMap splits each line into words and flattens them into a single RDD of individual words.

3. Exploding Key-Value Pairs

```
data = {"a": [1, 2], "b": [3, 4]}
rdd = sc.parallelize(data.items())

# Option 1: Flatten to key-value pairs
exploded_kv = rdd.flatMap(lambda kv: [(kv[0], v) for v in kv[1]])
print(exploded_kv.collect()) # Output: [("a", 1), ("a", 2), ("b", 3), ("b", 4)]

# Option 2: Flatten only values
flattened_values = rdd.flatMap(lambda kv: kv[1])
print(flattened_values.collect()) # Output: [1, 2, 3, 4]
```

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```
[('a', 1), ('a', 2), ('b', 3), ('b', 4)]
[1, 2, 3, 4]
```

Option 1: Flatten to Key-Value Pairs

```
exploded_kv = rdd.flatMap(lambda kv: [(kv[0], v) for v in kv[1]])
print(exploded_kv.collect()) # Output: [("a", 1), ("a", 2), ("b", 3), ("b", 4)]
```



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1. **flatMap Transformation:** flatMap applies the lambda function to each element of the RDD and flattens the results into a single RDD.
2. **Lambda Function Explanation:** For each key-value pair (kv) in the RDD:
 - kv[0] is the key (e.g., "a" or "b").
 - kv[1] is the list of values (e.g., [1, 2] or [3, 4]).
 - [(kv[0], v) for v in kv[1]] generates a list of tuples by pairing the key with each element of the list. So ("a", [1, 2]) becomes [("a", 1), ("a", 2)], and ("b", [3, 4]) becomes [("b", 3), ("b", 4)].
3. **Result:** After flatMap flattens the results, we get [("a", 1), ("a", 2), ("b", 3), ("b", 4)].

Option 2: Flatten Only Values

```
flattened_values = rdd.flatMap(lambda kv: kv[1])
print(flattened_values.collect()) # Output: [1, 2, 3, 4]
```

1. **flatMap Transformation:** Here, flatMap applies a lambda function that simply returns the values list (kv[1]) for each key-value pair.
2. **Lambda Function Explanation:** For each key-value pair (kv) in the RDD:
 - kv[1] is the list of values (e.g., [1, 2] or [3, 4]).
3. **Result:** flatMap flattens these lists of values from each key-value pair, resulting in a single RDD of values: [1, 2, 3, 4].

In summary:

- **Option 1** pairs each key with each of its values.
- **Option 2** extracts and flattens only the values.

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RDD part 3: Map, FlatMap, Filter example

Summary of Differences

Transformation	Purpose	Output Characteristics
map	Applies a function to each element	Same number of elements as input
flatMap	Applies a function that returns multiple elements	Can produce zero or more elements per input; flattened output
filter	Selects elements that meet a condition	Fewer or equal elements compared to input



```
from pyspark import SparkContext

# Initialize Spark Context
sc = SparkContext.getOrCreate()

# Step 1: Create an RDD from a list of integers
data = [1, 2, 3, 4, 5]
rdd = sc.parallelize(data)

# Step 2: Apply Transformations

# Map Transformation: Add 10 and 20 to each element
map_result = rdd.map(lambda a: (a + 10, a + 20)).collect()
print('Map Transformation:', map_result)

# Expected Output: Map Transformation: [(11, 21), (12, 22), (13, 23), (14, 24), (15, 25)]

# FlatMap Transformation: Add 10 and 20 to each element, returning flattened results
flat_map_result = rdd.flatMap(lambda a: (a + 10, a + 20)).collect()
print('FlatMap Transformation:', flat_map_result)

# Expected Output: FlatMap Transformation: [11, 21, 12, 22, 13, 23, 14, 24, 15, 25]
```

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```
# Filter Transformation: Get only even numbers
filter_result = rdd.filter(lambda a: a % 2 == 0).collect()
print('Filter Transformation:', filter_result)

# Expected Output: Filter Transformation: [2, 4]
```

```
Map Transformation: [(11, 21), (12, 22), (13, 23), (14, 24), (15, 25)]
FlatMap Transformation: [11, 21, 12, 22, 13, 23, 14, 24, 15, 25]
Filter Transformation: [2, 4]
```

Explanation of the Code

1. **Spark Context Initialization:** A Spark context is created or retrieved, which is necessary to work with Spark functionalities.
2. **Creating RDD:** The parallelize method is used to create an RDD from a list of integers (1 to 5).
3. **Transformations:**
 - o **Map Transformation:** Each element is transformed by adding 10 and 20, resulting in a new RDD of tuples.
 - o **FlatMap Transformation:** Each element is transformed to produce two new elements (adding 10 and 20), and the results are flattened into a single list.
 - o **Filter Transformation:** The RDD is filtered to keep only even numbers.
4. **Output:** Each transformation result is collected and printed.

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RDD part 5: operations

1. Creating RDDs:

- o RDDs are created using `sc.parallelize()`, which distributes data across multiple nodes in the cluster. The second parameter specifies the number of partitions.

2. Input Data:

- o The initial RDD (`rdd1`) contains duplicate integers. Using `glom()`, you can visualize the contents of each partition.

3. Distinct Transformation:

- o The `distinct()` operation filters out duplicate values in the RDD, resulting in a new RDD with unique elements. It is useful for deduplication in datasets.

4. Union Operation:

- o The `union()` operation combines the elements of two RDDs. The result includes all elements from both RDDs, allowing for data merging.

5. Intersection Operation:

- o The `intersection()` operation finds common elements present in both RDDs, useful for filtering data based on shared characteristics.

6. Subtract Operation:

- o The `subtract()` operation returns elements in the first RDD that are not present in the second RDD. This is often used to filter out unwanted data.

7. Partitions:

- o RDDs can have multiple partitions, allowing for distributed processing. The number of partitions can be obtained using `getNumPartitions()`.

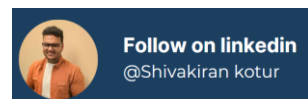
8. Coalesce Operation:

- o `coalesce(n)` reduces the number of partitions in an RDD to n without performing a full shuffle, making it a narrow transformation. This is useful for optimizing performance when reducing the amount of data.

9. Repartition Operation:

- o `repartition(n)` can increase or decrease the number of partitions in an RDD and performs a full shuffle of data. This is a wide transformation and can lead to empty partitions if not enough data is available.

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These operations and concepts are foundational for effective data manipulation and processing using RDDs in Spark, enabling efficient distributed data analytics.

```
# Creating RDDs
rdd1 = sc.parallelize([1, 1, 2, 2, 2, 3, 4, 4, 5, 5, 6], 3) # RDD with 3 partitions
rdd2 = sc.parallelize([4, 5, 6, 7, 8, 9], 3) # RDD with 3 partitions

# Display the input RDD
print('Input : ', rdd1.glom().collect())

# Distinct Transformation: Returns unique elements from rdd1
print('Distinct : ', rdd1.distinct().glom().collect())

# Union Operation: Combines rdd1 and rdd2
print('Union : ', rdd1.union(rdd2).collect())

# Intersection Operation: Returns common elements in rdd1 and rdd2
print('Intersection : ', rdd1.intersection(rdd2).collect())

# Subtract Operation: Returns elements in rdd1 that are not in rdd2
print('Subtract : ', rdd1.subtract(rdd2).collect())
```

```
Input : [[1, 1, 2], [2, 2, 3], [4, 4, 5, 5, 6]]
Distinct : [[3, 6], [1, 4], [2, 5]]
Union : [1, 1, 2, 2, 2, 3, 4, 4, 5, 5, 6, 4, 5, 6, 7, 8, 9]
Intersection : [6, 4, 5]
Subtract : [1, 1, 2, 2, 3]
```



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RDD part 6: Coalesce and repartition in RDD

Understanding Coalesce and Repartition in Apache Spark

In Apache Spark, managing the number of partitions of RDDs (Resilient Distributed Datasets) is crucial for optimizing performance. Two primary operations for managing partitions are **coalesce** and **repartition**. While both are used to adjust the number of partitions, they operate differently and serve different purposes.

1. Coalesce

Definition:

- The coalesce(n) function reduces the number of partitions in an RDD to n. This operation is typically used to decrease the number of partitions and does so without requiring a full shuffle of data.

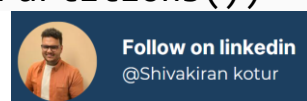
How It Works:

- **Narrow Transformation:** coalesce is considered a narrow transformation because it only combines partitions without moving data between them. It essentially combines existing partitions without redistributing the data across the cluster.
- **Efficiency:** Since it avoids a full shuffle, coalesce is generally more efficient than repartition when reducing partitions, leading to lower overhead and faster performance.
- **Use Case:** It is often used when filtering data has led to fewer records, and you want to optimize resource usage by reducing the number of partitions. For example, if you have a dataset with many partitions but only a small amount of data left after some transformations, coalescing to fewer partitions can help with performance.

```
# Creating another RDD with 50 elements and 10 partitions
rdd = sc.parallelize(range(50), 10)

# Display number of partitions
print('Number of Partitions : ', rdd.getNumPartitions())
```

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```
# Display partition-wise data
print('Partition Wise Data : ', rdd.glom().collect())

# Coalesce Operation: Reduces the number of partitions to 5
rdd_c = rdd.coalesce(5)
print('Coalesce : ', rdd_c.glom().collect()) # Narrows down the
number of partitions
```

2. Repartition

Definition:

- The repartition(n) function can increase or decrease the number of partitions in an RDD to n. This operation involves a full shuffle of data across all partitions.

How It Works:

- Wide Transformation:** repartition is considered a wide transformation because it involves reshuffling data between partitions. It redistributes the data in such a way that partitions are evenly filled, which can lead to better parallelism during computation.
- Shuffling Data:** Since repartition requires a shuffle, it can be more resource-intensive and slower than coalesce, especially when increasing the number of partitions.
- Use Case:** This operation is beneficial when you need to increase the number of partitions for better parallel processing or when the data is imbalanced across partitions. For example, if certain partitions have too much data, repartitioning can help ensure even distribution across the cluster.

```
# Repartition Operation: Changes the number of partitions to 5 and
can reshuffle data
rdd_r = rdd.repartition(5)
print('Repartition : ', rdd_r.glom().collect()) # Can create new
partitions and potentially empty ones
```

```

Number of Partitions : 10
Partition Wise Data : [[0, 1, 2, 3, 4], [5, 6, 7, 8, 9], [10, 11, 12, 13, 14], [15, 16, 17, 18, 19], [20, 21, 22, 23, 24], [25, 26, 27, 28, 29], [30, 31, 32, 33, 34], [35, 36, 37, 38, 39], [40, 41, 42, 43, 44], [45, 46, 47, 48, 49]]
Coalesce : [[0, 1, 2, 3, 4, 5, 6, 7, 8, 9], [10, 11, 12, 13, 14, 15, 16, 17, 18, 19], [20, 21, 22, 23, 24, 25, 26, 27, 28, 29], [30, 31, 32, 33, 34, 35, 36, 37, 38, 39], [40, 41, 42, 43, 44, 45, 46, 47, 48, 49]]
Repartition : [[], [25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44], [], [15, 16, 17, 18, 19], [0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 20, 21, 22, 23, 24, 45, 46, 47, 48, 49]]

```

Key Differences Between Coalesce and Repartition

Aspect	Coalesce	Repartition
Purpose	Reduce the number of partitions	Increase or decrease partitions
Data Shuffle	No shuffle (narrow transformation)	Requires a full shuffle (wide transformation)
Performance	More efficient for reducing partitions	Less efficient due to shuffling
Use Case	After filtering data, to optimize resource usage	When needing balanced partitioning or increasing parallelism

Conclusion

- + Choosing between coalesce and repartition depends on the specific needs of your Spark application.
- + Use coalesce for efficient reduction of partitions without shuffling, and repartition when you need to change the partitioning scheme for better data distribution or parallel processing.
- + Understanding these concepts helps optimize Spark applications for performance and resource management.

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Coalesce and repartition in Dataframe

Initial DataFrame Creation:

- The code starts by creating a DataFrame with 100 records (numbers 0 to 99) and repartitions it into 5 partitions.

Initial Partition Information:

- It checks and prints the number of partitions and the data within each partition before any transformations.

Coalesce Operation:

- The DataFrame is then coalesced to reduce the number of partitions to 3. This operation combines existing partitions without a full shuffle, which is more efficient for reducing partitions.

Post-Coalesce Data Check:

- The number of partitions and the data in each partition after coalescing are printed.

Repartition Operation:

- The DataFrame is repartitioned back to 4 partitions. This operation can involve a full shuffle of the data, redistributing it across the specified number of partitions.

Post-Repartition Data Check:

- Finally, it checks and prints the number of partitions and the data in each partition after repartitioning.

Partition Size Calculation:

- The sizes of each partition after both coalesce and repartition operations are calculated and displayed.

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```

from pyspark.sql import SparkSession

# Create a Spark session
spark = SparkSession.builder.appName("Partition Check with Coalesce and Repartition").getOrCreate()

# Create a DataFrame with some sample data
data = [(i,) for i in range(100)]
df = spark.createDataFrame(data, ["numbers"]).repartition(5) # creating 5 partitions

# Initial partition information
print("Initial number of partitions:", df.rdd.getNumPartitions())

# Show data in each partition before any transformations
print("Initial data in each partition:")
initial_partitions_data = df.rdd.glom().collect()
for i, partition in enumerate(initial_partitions_data):
    print(f"Partition {i}: {partition}")

Initial number of partitions: 5
Initial data in each partition:
Partition 0: [Row(numbers=3), Row(numbers=6), Row(numbers=1), Row(numbers=22), Row(numbers=23), Row(numbers=17), Row(numbers=28), Row(numbers=30), Row(numbers=27), Row(numbers=38), Row(numbers=45), Row(numbers=50), Row(numbers=56), Row(numbers=59), Row(numbers=60), Row(numbers=63), Row(numbers=72), Row(numbers=76), Row(numbers=98), Row(numbers=88), Row(numbers=96)]
Partition 1: [Row(numbers=7), Row(numbers=8), Row(numbers=14), Row(numbers=16), Row(numbers=33), Row(numbers=32), Row(numbers=44), Row(numbers=36), Row(numbers=54), Row(numbers=53), Row(numbers=67), Row(numbers=64), Row(numbers=71), Row(numbers=74), Row(numbers=83), Row(numbers=77), Row(numbers=95), Row(numbers=91), Row(numbers=92), Row(numbers=89)]
Partition 2: [Row(numbers=0), Row(numbers=2), Row(numbers=20), Row(numbers=21), Row(numbers=35), Row(numbers=34), Row(numbers=39), Row(numbers=41), Row(numbers=49), Row(numbers=57), Row(numbers=61), Row(numbers=70), Row(numbers=69), Row(numbers=73), Row(numbers=80), Row(numbers=81), Row(numbers=86), Row(numbers=93)]
Partition 3: [Row(numbers=10), Row(numbers=9), Row(numbers=13), Row(numbers=18), Row(numbers=25), Row(numbers=29), Row(numbers=43), Row(numbers=37), Row(numbers=40), Row(numbers=55), Row(numbers=58), Row(numbers=66), Row(numbers=68), Row(numbers=79), Row(numbers=82), Row(numbers=87), Row(numbers=94), Row(numbers=84)]
Partition 4: [Row(numbers=11), Row(numbers=5), Row(numbers=4), Row(numbers=19), Row(numbers=15), Row(numbers=12), Row(numbers=26), Row(numbers=31), Row(numbers=24), Row(numbers=46), Row(numbers=42), Row(numbers=47), Row(numbers=52), Row(numbers=51), Row(numbers=48), Row(numbers=65), Row(numbers=62), Row(numbers=75), Row(numbers=78), Row(numbers=97), Row(numbers=85), Row(numbers=99)]

# Coalesce to reduce the number of partitions (to 3)
df_coalesced = df.coalesce(3)
print("\nNumber of partitions after coalesce():",
df_coalesced.rdd.getNumPartitions())

# Show data in each partition after coalescing
print("Data in each partition after coalesce:")
coalesced_partitions_data = df_coalesced.rdd.glom().collect()
for i, partition in enumerate(coalesced_partitions_data):
    print(f"Partition {i}: {partition}")

```



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```

Number of partitions after coalesce(): 3
Data in each partition after coalesce:
Partition 0: [Row(numbers=3), Row(numbers=6), Row(numbers=1), Row(numbers=22), Row(numbers=23), Row(numbers=17), Row(numbers=28), Row(numbers=30), Row(numbers=27), Row(numbers=38), Row(numbers=45), Row(numbers=50), Row(numbers=56), Row(numbers=59), Row(numbers=60), Row(numbers=63), Row(numbers=72), Row(numbers=76), Row(numbers=98), Row(numbers=88), Row(numbers=96)]
Partition 1: [Row(numbers=7), Row(numbers=8), Row(numbers=14), Row(numbers=16), Row(numbers=33), Row(numbers=32), Row(numbers=44), Row(numbers=36), Row(numbers=54), Row(numbers=53), Row(numbers=67), Row(numbers=64), Row(numbers=71), Row(numbers=74), Row(numbers=83), Row(numbers=77), Row(numbers=95), Row(numbers=91), Row(numbers=92), Row(numbers=89), Row(numbers=0), Row(numbers=2), Row(numbers=20), Row(numbers=21), Row(numbers=35), Row(numbers=34), Row(numbers=39), Row(numbers=41), Row(numbers=49), Row(numbers=57), Row(numbers=61), Row(numbers=70), Row(numbers=69), Row(numbers=73), Row(numbers=80), Row(numbers=81), Row(numbers=90), Row(numbers=86), Row(numbers=93)]
Partition 2: [Row(numbers=10), Row(numbers=9), Row(numbers=13), Row(numbers=18), Row(numbers=25), Row(numbers=29), Row(numbers=43), Row(numbers=37), Row(numbers=40), Row(numbers=55), Row(numbers=58), Row(numbers=66), Row(numbers=68), Row(numbers=79), Row(numbers=82), Row(numbers=87), Row(numbers=94), Row(numbers=84), Row(numbers=11), Row(numbers=5), Row(numbers=4), Row(numbers=19), Row(numbers=15), Row(numbers=12), Row(numbers=26), Row(numbers=31), Row(numbers=24), Row(numbers=46), Row(numbers=42), Row(numbers=47), Row(numbers=52), Row(numbers=51), Row(numbers=48), Row(numbers=65), Row(numbers=62), Row(numbers=75), Row(numbers=78), Row(numbers=97), Row(numbers=85), Row(numbers=99)]
# Repartition to increase the number of partitions (to 4)
df_repartitioned = df_coalesced.repartition(4)
print("\nNumber of partitions after repartition():",
df_repartitioned.rdd.getNumPartitions())

# Show data in each partition after repartitioning
print("Data in each partition after repartition:")
repartitioned_partitions_data =
df_repartitioned.rdd.glom().collect()
for i, partition in enumerate(repartitioned_partitions_data):
    print(f"Partition {i}: {partition}")

```

```

Number of partitions after repartition(): 4
Data in each partition after repartition:
Partition 0: [Row(numbers=0), Row(numbers=8), Row(numbers=1), Row(numbers=20), Row(numbers=16), Row(numbers=17), Row(numbers=26), Row(numbers=25), Row(numbers=34), Row(numbers=46), Row(numbers=37), Row(numbers=41), Row(numbers=50), Row(numbers=51), Row(numbers=58), Row(numbers=67), Row(numbers=60), Row(numbers=62), Row(numbers=74), Row(numbers=72), Row(numbers=78), Row(numbers=87), Row(numbers=86), Row(numbers=92), Row(numbers=96)]
Partition 1: [Row(numbers=11), Row(numbers=10), Row(numbers=2), Row(numbers=19), Row(numbers=13), Row(numbers=21), Row(numbers=28), Row(numbers=31), Row(numbers=29), Row(numbers=38), Row(numbers=42), Row(numbers=40), Row(numbers=54), Row(numbers=56), Row(numbers=48), Row(numbers=61), Row(numbers=64), Row(numbers=63), Row(numbers=73), Row(numbers=83), Row(numbers=76), Row(numbers=97), Row(numbers=94), Row(numbers=93), Row(numbers=89)]
Partition 2: [Row(numbers=3), Row(numbers=5), Row(numbers=9), Row(numbers=22), Row(numbers=15), Row(numbers=18), Row(numbers=33), Row(numbers=30), Row(numbers=24), Row(numbers=44), Row(numbers=45), Row(numbers=47), Row(numbers=49), Row(numbers=53), Row(numbers=59), Row(numbers=66), Row(numbers=70), Row(numbers=71), Row(numbers=79), Row(numbers=80), Row(numbers=77), Row(numbers=95), Row(numbers=98), Row(numbers=85), Row(numbers=84)]
Partition 3: [Row(numbers=7), Row(numbers=6), Row(numbers=4), Row(numbers=14), Row(numbers=23), Row(numbers=12), Row(numbers=35), Row(numbers=32), Row(numbers=27), Row(numbers=43), Row(numbers=39), Row(numbers=36), Row(numbers=52), Row(numbers=55), Row(numbers=57), Row(numbers=65), Row(numbers=68), Row(numbers=69), Row(numbers=75), Row(numbers=82), Row(numbers=81), Row(numbers=90), Row(numbers=91), Row(numbers=88), Row(numbers=99)]

```

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```
# Calculate and show the size of each partition after coalesce and  
repartition  
coalesce_partition_sizes =  
df_coalesced.rdd.glom().map(len).collect()  
print("\nSize of each partition after coalesce:",  
coalesce_partition_sizes)  
  
repartition_partition_sizes =  
df_repartitioned.rdd.glom().map(len).collect()  
print("Size of each partition after repartition:",  
repartition_partition_sizes)
```



Size of each partition after coalesce: [21, 39, 40]

Size of each partition after repartition: [25, 25, 25, 25]

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ReduceByKey vs GroupByKey in RDD

Let's take a scenario with a Spark cluster that has 4 nodes (each node can be thought of as a worker with multiple cores) and a dataset with key-value pairs. I'll explain how reduceByKey and groupByKey operate on these nodes to highlight their key differences.

Assume we have the following key-value pairs across the nodes:

Node	Key-Value Pairs
1	(A, 2), (B, 3), (A, 5), (C, 4)
2	(A, 1), (B, 2), (C, 6), (C, 5)
3	(B, 7), (C, 3), (A, 4), (D, 5)
4	(A, 6), (B, 3), (D, 8), (C, 2)

How reduceByKey Works

- Initial Partitioning:** Spark divides the data across the 4 nodes in the cluster.
- Local Reduction on Each Node:** reduceByKey applies the specified reduction function (e.g., summing values) locally within each partition first.
 - For instance:
 - On Node 1: $(A, 2) + (A, 5) = (A, 7)$, (B, 3), (C, 4)
 - On Node 2: (A, 1), (B, 2), (C, 11)
 - On Node 3: (A, 4), (B, 7), (C, 3), (D, 5)
 - On Node 4: (A, 6), (B, 3), (D, 8), (C, 2)
- Shuffle and Aggregate:** The reduced values for each key are then shuffled to the appropriate nodes to further aggregate them.
 - For example, all A pairs are shuffled together, all B pairs together, and so on.
 - The final reduction takes place, summing up values across nodes.
 - Example results:
 - (A, 18), (B, 15), (C, 20), (D, 13)
- Result:** The final result is an RDD with each unique key and its aggregated value. By reducing data before shuffling, reduceByKey minimizes data transfer and memory usage.

How groupByKey Works

1. **Initial Partitioning:** Spark distributes the data across the 4 nodes in the cluster (same as reduceByKey).
2. **Shuffle All Key-Value Pairs:** Unlike reduceByKey, groupByKey shuffles all data for each key to one node without any local aggregation. This means:
 - o Every value associated with a particular key (A, B, C, D) is sent to the same node.
 - o For example, all key-value pairs with key A will be transferred to a single node.
3. **Group Values by Key:** After shuffling, the values are grouped under each key.
 - o Example result:
 - (A, [2, 5, 1, 4, 6]), (B, [3, 2, 7, 3]), (C, [4, 6, 3, 5, 2]), (D, [5, 8])
4. **Result:** The final result is an RDD where each key has a list of all its values. Since all key-value pairs are shuffled without any reduction, groupByKey uses more memory and network resources.

```
# Sample data with key-value pairs
data = [
    ("A", 2), ("B", 3), ("A", 5), ("C", 4),
    ("A", 1), ("B", 2), ("C", 6), ("C", 5),
    ("B", 7), ("C", 3), ("A", 4), ("D", 5),
    ("A", 6), ("B", 3), ("D", 8), ("C", 2)
]

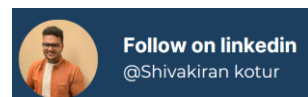
# Parallelize data to create an RDD
rdd = spark.sparkContext.parallelize(data)

# Using reduceByKey
reduce_by_key_rdd = rdd.reduceByKey(lambda x, y: x + y)
reduce_by_key_result = reduce_by_key_rdd.collect()
print("Result of reduceByKey:", reduce_by_key_result)

# Using groupByKey
group_by_key_rdd = rdd.groupByKey().mapValues(list)
group_by_key_result = group_by_key_rdd.collect()
print("Result of groupByKey:", group_by_key_result)
```

The main difference between reduceByKey and groupByKey in Apache Spark lies in their efficiency and use cases:

- **reduceByKey** performs a combination and reduction operation at the map stage, which minimizes data shuffling across the network. It applies the specified function (e.g., sum, max) directly to the values of each key in each partition, then combines the intermediate



results. This makes `reduceByKey` more efficient in terms of memory and network usage, especially for large datasets, as it reduces data before shuffling it across partitions.

- **groupByKey**, on the other hand, shuffles all key-value pairs across the network to group the values for each key on a single partition. After shuffling, it combines all values for each key without performing any aggregation. This can lead to higher memory and network costs, making `groupByKey` slower for large datasets and prone to out-of-memory errors when working with a large number of values per key.

In summary: Use `reduceByKey` when you want to perform aggregation, as it is more efficient. Use `groupByKey` only if you need to retain all values per key without aggregating.

Summary: Key Differences in Operation

Aspect	<code>reduceByKey</code>	<code>groupByKey</code>
Purpose	Aggregates values for each key (e.g., sum, count)	Groups all values for each key without aggregation
Local Aggregation	Performs local aggregation on each partition before shuffling, reducing data volume	No local aggregation; all data is shuffled, leading to potentially high network and memory usage
Shuffle Cost	Lower shuffle cost due to reduced data transfer (only partially aggregated values are shuffled)	Higher shuffle cost as all key-value pairs are transferred without reduction
Memory Usage	More memory-efficient; only aggregated values per key are kept in memory during shuffling	Higher memory usage, especially if there are many values per key
Performance	Faster and more efficient for large datasets with aggregation needs	Slower and may cause out-of-memory errors for large datasets due to shuffling all values
Use Case	Best for operations that aggregate data (e.g., summing or averaging values by key)	Use only when you need all values for each key (e.g., for further processing without reducing values)
Example Result	(A, 18), (B, 15), (C, 20), (D, 13) (each key with a single aggregated value)	(A, [2, 5, 1, 4, 6]), (B, [3, 2, 7, 3]), (C, [4, 6, 3, 5, 2]), (D, [5, 8]) (each key with all values)

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SortBy vs SortByKey in RDD

1. sortBy Operation

The sortBy operation sorts elements of an RDD based on a specified attribute or criteria. It can be used on RDDs containing any data type (not just key-value pairs), and the user specifies a function that determines the sorting criteria.

Example:

```
# Sample data
data = [5, 2, 8, 1, 3]

# Creating an RDD
rdd = sc.parallelize(data)

# Sorting in ascending order using sortBy
sorted_rdd = rdd.sortBy(lambda x: x)

# Collecting and printing the result
print(sorted_rdd.collect())
# Output: [1, 2, 3, 5, 8]
```

In this example, each element is sorted in ascending order. The lambda `x: x` function simply returns the element itself, so the entire RDD is sorted based on the element values.

Behind the Scenes:

Here's a breakdown of what happens when you use sortBy on an RDD:

1. Partitioning:

- If the RDD has multiple partitions, Spark first performs local sorting within each partition independently.
- By default, Spark uses **Timsort** (a highly efficient, adaptive sorting algorithm) to handle the sorting within each partition.

2. Sorting Within Partitions:

- Spark applies Timsort to sort the data within each partition based on the function specified in sortBy.



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- This step is parallelized, meaning each partition is sorted independently on different executor nodes in a distributed manner, which increases efficiency.

3. Merging Partitions:

- Once the data within each partition is sorted, Spark merges the sorted partitions.
- The merging process also uses Timsort to maintain the global sorted order across partitions.
- Spark efficiently combines the partitions, producing a final RDD that is sorted based on the criteria specified.

2. sortByKey Operation

The sortByKey operation is specifically used for sorting key-value pair RDDs by key. It's a common operation for RDDs with data structured as tuples (e.g., (key, value) pairs), and sorts based on the key while keeping the corresponding values.

Example:

```
# Sample data as key-value pairs
data = [(5, 'five'), (2, 'two'), (8, 'eight'), (1, 'one'), (3, 'three')]

# Creating an RDD
rdd = sc.parallelize(data)

# Sorting by key in ascending order using sortByKey
sorted_rdd = rdd.sortByKey()

# Collecting and printing the result
print(sorted_rdd.collect())
# Output: [(1, 'one'), (2, 'two'), (3, 'three'), (5, 'five'), (8, 'eight')]
```

In this example, each tuple (key-value pair) is sorted based on the key. The result is a new RDD with the tuples ordered by key in ascending order.

Behind the Scenes:

The sortByKey operation follows a similar process to sortBy, but with some additional steps related to handling key-value pairs.



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1. Partitioning and Key Extraction:

- If the RDD has multiple partitions, Spark partitions the data based on keys.
- Each partition is locally sorted by the keys using Timsort, which arranges the key-value pairs within each partition.

2. Sorting Key-Value Pairs:

- Within each partition, Spark applies Timsort to sort key-value pairs based on the keys. The corresponding values are rearranged based on the sorted keys.

3. Merging Partitions:

- Once the key-value pairs within each partition are sorted, Spark merges the partitions to produce the final sorted RDD.
- The merging process ensures that the key order is maintained across partitions.

Key Differences Between sortBy and sortByKey

Aspect	sortBy	sortByKey
Use Case	General sorting based on a specified attribute or function	Sorting key-value pair RDDs based on keys
Data Type	Works on any data type	Specifically designed for key-value pair RDDs
Example Function	sortBy(lambda x: x.some_attr)	sortByKey() (no function needed)
Partition Handling	Sorts each partition independently, then merges	Partitions and sorts by key, then merges sorted partitions

Performance Considerations

- **Partitioning:** Both sortBy and sortByKey operations may involve shuffling data across partitions to ensure a globally sorted result. This shuffle can impact performance, especially for large datasets.
- **Parallelism:** Sorting within each partition is parallelized, making both sortBy and sortByKey efficient for distributed sorting.
- **Data Skew:** If the data is skewed (uneven distribution of keys across partitions), performance may degrade as some partitions will have significantly more data to sort than others.

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Choosing Between sortBy and sortByKey

- Use sortBy when you need to sort based on a specific attribute or transformation on a non-key-value RDD.
- Use sortByKey when working with key-value pairs and sorting based on the keys without additional transformations.

These sorting operations help prepare data for downstream tasks such as joins, aggregations, or output formatting, making them essential for many data processing workflows in Spark.

```
# Sample data
data = [("apple", 3), ("banana", 1), ("orange", 2), ("grape", 5)]

# Creating an RDD
rdd = sc.parallelize(data)

# 1. Using sortBy to sort by the second element in each tuple (the
# integer value)
sorted_by_value = rdd.sortBy(lambda x: x[1])
print("Sorted by value (using sortBy):", sorted_by_value.collect())
# Output: [('banana', 1), ('orange', 2), ('apple', 3), ('grape', 5)]

# 2. Using sortByKey to sort by the first element in each tuple
# (the string key)
sorted_by_key = rdd.sortByKey()
print("Sorted by key (using sortByKey):", sorted_by_key.collect())
# Output: [('apple', 3), ('banana', 1), ('grape', 5), ('orange', 2)]
```

```
Sorted by value (using sortBy): [('banana', 1), ('orange', 2), ('apple', 3), ('grape', 5)]
Sorted by key (using sortByKey): [('apple', 3), ('banana', 1), ('grape', 5), ('orange', 2)]
```

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Take vs collect in RDD

Both take and collect are actions in Spark used to retrieve data from an RDD (Resilient Distributed Dataset), but they serve different purposes and are designed for different use cases.

1. take Action

- **Return Type:**
 - take returns an array with the first N elements of the RDD, where N is specified by the user.
- **Use Case:**
 - take is typically used to retrieve a small sample of elements from the RDD for inspection or quick testing, without bringing the entire dataset into the driver's memory.
- **Example:**

```
# Sample data
data = [1, 2, 3, 4, 5]
# Create an RDD
rdd = sc.parallelize(data)

# Retrieve the first three elements of the RDD
result = rdd.take(3)

print(result) #return as list #[1, 2, 3]
print(", ".join(map(str, result))) # Output: 1, 2, 3
```

- **Key Considerations:**
 - **Performance:** take is more efficient than collect when you only need a small subset of data, as it minimizes the amount of data transferred from the executors to the driver.
 - **Memory Usage:** Since it retrieves only a specified number of elements, take is safer to use on large datasets without overwhelming the driver's memory.

2. collect Action

- **Return Type:**

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- collect returns an array containing all elements of the RDD, bringing the entire dataset to the driver program.

- **Use Case:**

- collect is used when the entire RDD is required in the driver program for further processing, often for saving as a local data structure like an array or list.

- **Example:**

```
# Sample data
data = [1, 2, 3, 4, 5]
# Create an RDD
rdd = sc.parallelize(data)

# Retrieve all elements of the RDD
result = rdd.collect()

print(result) #output in list [1, 2, 3, 4, 5]
print(", ".join(map(str, result))) # Output: 1, 2, 3, 4, 5
```

- **Key Considerations:**

- **Data Size:** collect should only be used when the dataset is small enough to fit in the driver's memory, as large datasets can cause out-of-memory errors.
- **Performance:** Since collect transfers all data to the driver, it incurs a higher network overhead and can significantly affect performance on large datasets.

Summary Table: take vs collect

Feature	take	collect
Purpose	Retrieves the first N elements of the RDD.	Retrieves all elements of the RDD.
Return Type	Array of the first N elements.	Array of all elements.
Use Case	Use when a sample is sufficient for inspection or testing.	Use when the entire dataset is needed on the driver for processing.
Performance	More efficient, as it only brings a subset of data.	Can be resource-intensive and cause memory issues if the dataset is large.
Memory Usage	Requires minimal memory on the driver, as it only loads a few elements.	Requires sufficient memory on the driver to store the entire dataset.
Risk	Low risk of out-of-memory errors.	High risk of out-of-memory errors with large datasets.
Network Overhead	Minimal, as only a subset of data is transferred to the driver.	Higher network overhead, as it transfers all data to the driver.

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Collect(), Collect_list() , Collect_set()

In PySpark, collect, collect_list, and collect_set are functions used to retrieve or aggregate data in different ways from Spark DataFrames or RDDs. Here's a breakdown of each function, with code examples:

1. collect()

The collect() function gathers all elements of a DataFrame (or RDD) and returns them as a list on the driver node. While it's useful for small datasets, using collect() with large datasets can overwhelm the driver memory, so it should be used with caution.

Example:

```
from pyspark.sql import SparkSession

# Initialize Spark session
spark =
SparkSession.builder.appName("CollectExample").getOrCreate()

# Sample DataFrame
data = [("Alice", 25), ("Bob", 30), ("Cathy", 28)]
df = spark.createDataFrame(data, ["Name", "Age"])

# Using collect() to gather all rows
collected_data = df.collect()

print ("collect() output as below:")
print(collected_data)

print ("access each row from collect:")
# Display collected data
for row in collected_data:
    print(row)

collect() output as below:
[Row(Name='Alice', Age=25), Row(Name='Bob', Age=30), Row(Name='Cathy', Age=28)]
access each row from collect:
Row(Name='Alice', Age=25)
Row(Name='Bob', Age=30)
Row(Name='Cathy', Age=28)
```

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2. collect_list()

The `collect_list()` function is an aggregation function used with `groupBy()` in DataFrames. It gathers all values of a specified column for each group into a list, allowing duplicates.

Example:

```
from pyspark.sql import functions as F

# Sample DataFrame
data = [("Alice", "Math"), ("Alice", "Science"), ("Bob", "Math"),
        ("Cathy", "Science")]
df = spark.createDataFrame(data, ["Name", "Subject"])

+-----+
| Name|Subject|
+-----+
|Alice|  Math|
|Alice|Science|
| Bob|  Math|
|Cathy|Science|
+-----+

# Using collect_list() to gather subjects for each person
result =
df.groupBy("Name").agg(F.collect_list("Subject").alias("Subjects"))

# Show the result
result.show()
```

3. collect_set()

The `collect_set()` function is similar to `collect_list()`, but it removes duplicate values within each group, returning a unique set of values for each group.

Example:

```
# Sample DataFrame
data = [("Alice", "Math"), ("Alice", "Math"), ("Alice", "Science"),
        ("Bob", "Math")]
df = spark.createDataFrame(data, ["Name", "Subject"])

df.show()
+---+-----+
| Name|Subject|
+---+-----+
|Alice|  Math|
|Alice|  Math|
|Alice|Science|
| Bob|  Math|
+---+-----+

# Using collect_set() to gather unique subjects for each person
result =
df.groupBy("Name").agg(F.collect_set("Subject").alias("UniqueSubjects"))

# Show the result
result.show()
+---+-----+
| Name| UniqueSubjects|
+---+-----+
|Alice|[Math, Science]|
| Bob|      [Math]|
+---+-----+
```

Key Differences:

- **collect()**: Collects all elements from a DataFrame or RDD to the driver node.
- **collect_list()**: Collects values of a column for each group as a list, allowing duplicates.
- **collect_set()**: Similar to collect_list(), but removes duplicates within each group.

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Collect(), Collect_list() , Collect_set() part2 problem

Problem Statement

Suppose we have a dataset of student records with each student's Name, Subject, Marks, and Exam_Date. Our goal is to analyze the data and generate insights:

1. For each student, find all the subjects they have appeared in, allowing duplicates
2. For each student, find the unique subjects they have appeared in, removing any duplicates.
3. Find the overall list of all student records.

```
from pyspark.sql import SparkSession
from pyspark.sql import functions as F

# Initialize Spark session
spark =
SparkSession.builder.appName("StudentAnalysis").getOrCreate()

# Sample DataFrame creation
data = [
    ("Alice", "Math", 85, "2024-01-10"),
    ("Alice", "Science", 78, "2024-01-15"),
    ("Alice", "Math", 88, "2024-02-10"),
    ("Bob", "Math", 90, "2024-01-12"),
    ("Bob", "History", 85, "2024-01-20"),
    ("Cathy", "Science", 95, "2024-01-15"),
    ("Cathy", "Math", 70, "2024-02-05"),
    ("Cathy", "Math", 75, "2024-02-20"),
]
df = spark.createDataFrame(data, ["Name", "Subject", "Marks",
"Exam_Date"])

# 1. Find all subjects each student has appeared in (allowing
# duplicates)
all_subjects = df.groupBy("Name").agg(
    F.collect_list("Subject").alias("AllSubjectsTaken")
)
```



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	Name	AllSubjectsTaken
1	Alice	> ["Math", "Science", "Math"]
2	Bob	> ["Math", "History"]
3	Cathy	> ["Science", "Math", "Math"]

```
# 2. Find unique subjects each student has appeared in (removing duplicates)
```

```
unique_subjects = df.groupBy("Name").agg(
    F.collect_set("Subject").alias("UniqueSubjectsTaken")
)
```

```
print("Unique Subjects Taken by Each Student (Without Duplicates):")
unique_subjects.display()
```

	Name	UniqueSubjectsTaken
1	Alice	> ["Math", "Science"]
2	Bob	> ["Math", "History"]
3	Cathy	> ["Math", "Science"]

```
# 3. Collect the entire DataFrame to the driver
collected_data = df.collect()
```

```
print("Collected Data:")
for row in collected_data:
    print(row)
```

Collected Data:

```
Row(Name='Alice', Subject='Math', Marks=85, Exam_Date='2024-01-10')
Row(Name='Alice', Subject='Science', Marks=78, Exam_Date='2024-01-15')
Row(Name='Alice', Subject='Math', Marks=88, Exam_Date='2024-02-10')
Row(Name='Bob', Subject='Math', Marks=90, Exam_Date='2024-01-12')
Row(Name='Bob', Subject='History', Marks=85, Exam_Date='2024-01-20')
Row(Name='Cathy', Subject='Science', Marks=95, Exam_Date='2024-01-15')
Row(Name='Cathy', Subject='Math', Marks=70, Exam_Date='2024-02-05')
Row(Name='Cathy', Subject='Math', Marks=75, Exam_Date='2024-02-20')
```



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Master Spark Concepts Zero to Big data Hero:

What is Spark Submit?

Spark-submit is a command-line tool used to deploy Spark applications to a cluster. It allows users to:

1. **Specify application configurations** such as memory, cores, and dependencies.
2. **Submit Spark jobs** in different deployment modes.
3. **Interact with resource managers** like YARN, Mesos, or Kubernetes.

Key Features of Spark-submit:

- Enables the execution of distributed applications.
- Supports multiple languages like Scala, Python, Java, and R.
- Offers flexibility with deployment modes.

Deployment Modes in Spark

When submitting a Spark job using spark-submit, you must choose a **deploy mode** to define where the driver program (main application logic) will run.

Cluster Mode

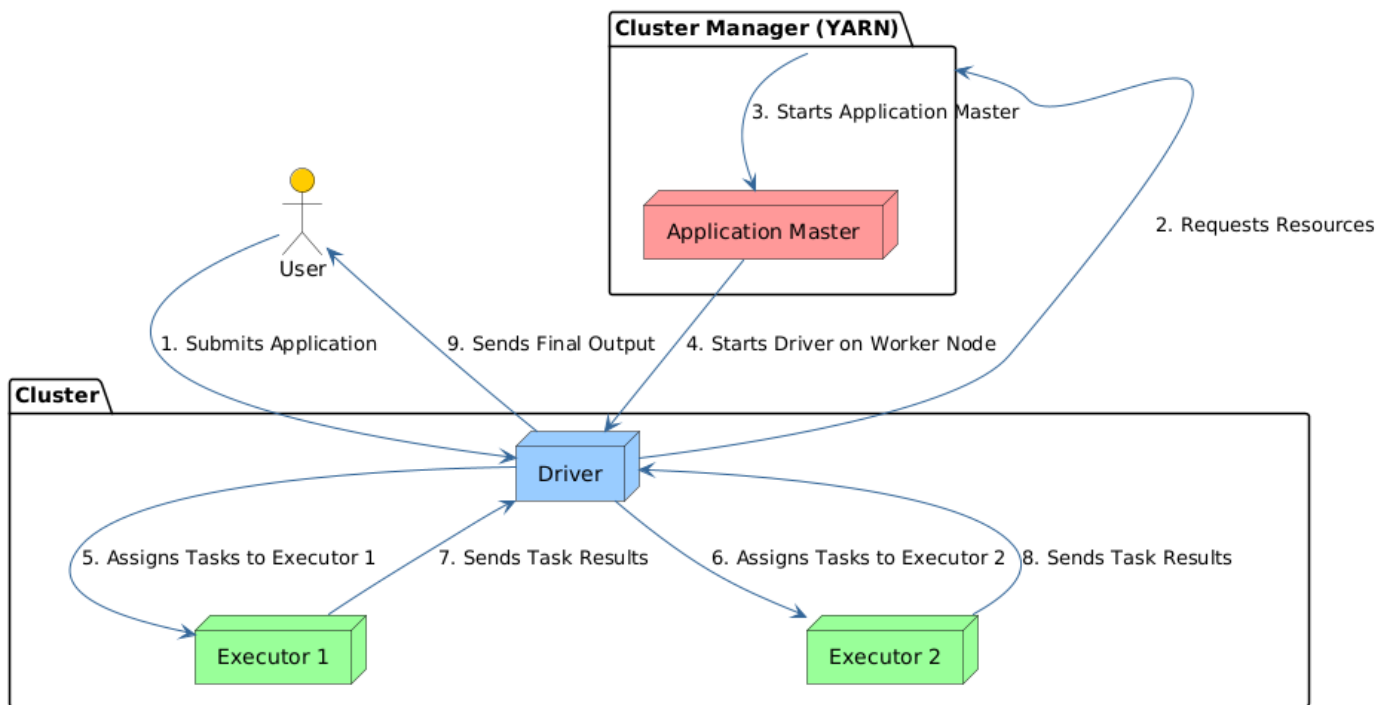
In Cluster Mode, the **Driver** runs within the cluster on one of the worker nodes, and the cluster manager allocates resources, including the Driver and Executors, to handle the application's execution.

Use Cluster Mode for **production applications** or long-running jobs, where the driver runs within the cluster for better resource management and fault tolerance.

1. User submits the Spark application to the Driver.
2. Driver communicates with the Cluster Manager (YARN) to obtain resources.
3. Cluster Manager starts the Application Master, which initializes the driver.
4. Driver assigns tasks to Executor 1 and Executor 2 for processing.
5. Executors carry out the tasks and return results to the Driver.
6. Driver aggregates all the results and sends the final output back to the User.



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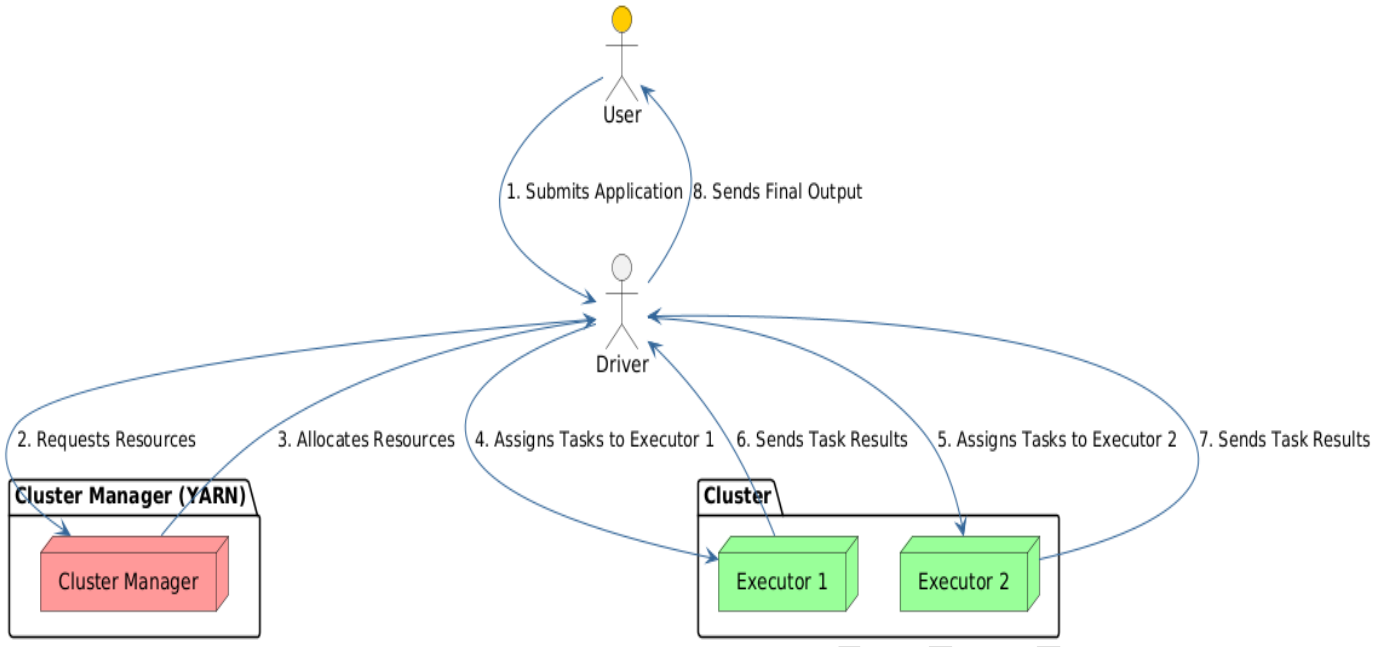


Client Mode

Client Mode: In Client Mode, the **Driver** runs on the client machine, and it directly interacts with the cluster manager to request resources and assign tasks to the worker nodes for execution.

Client Mode: Use Client Mode for **interactive applications or development and testing**, where the driver needs to run on the local machine and interact directly with the user.

1. User submits the Spark application to the Driver.
2. Driver (acting as the Application Master) requests resources from the Cluster Manager (YARN).
3. Cluster Manager allocates the required resources and returns them to the Driver.
4. Driver assigns tasks to Executor 1 for data processing.
5. Driver assigns tasks to Executor 2 for data processing.
6. Executor 1 sends task results back to the Driver.
7. Executor 2 sends task results back to the Driver.
8. Driver sends the final output back to the User.



Key Differences Between Client Mode and Cluster Mode

Aspect	Client Mode	Cluster Mode
Driver Location	Runs on the client machine	Runs on a worker node in the cluster
Dependency	Requires an active client connection	Operates independently of the client
Best For	Development and testing	Production and large-scale applications
Execution Control	Managed by the client	Managed by the cluster

How Databricks Overcomes These Limitations?

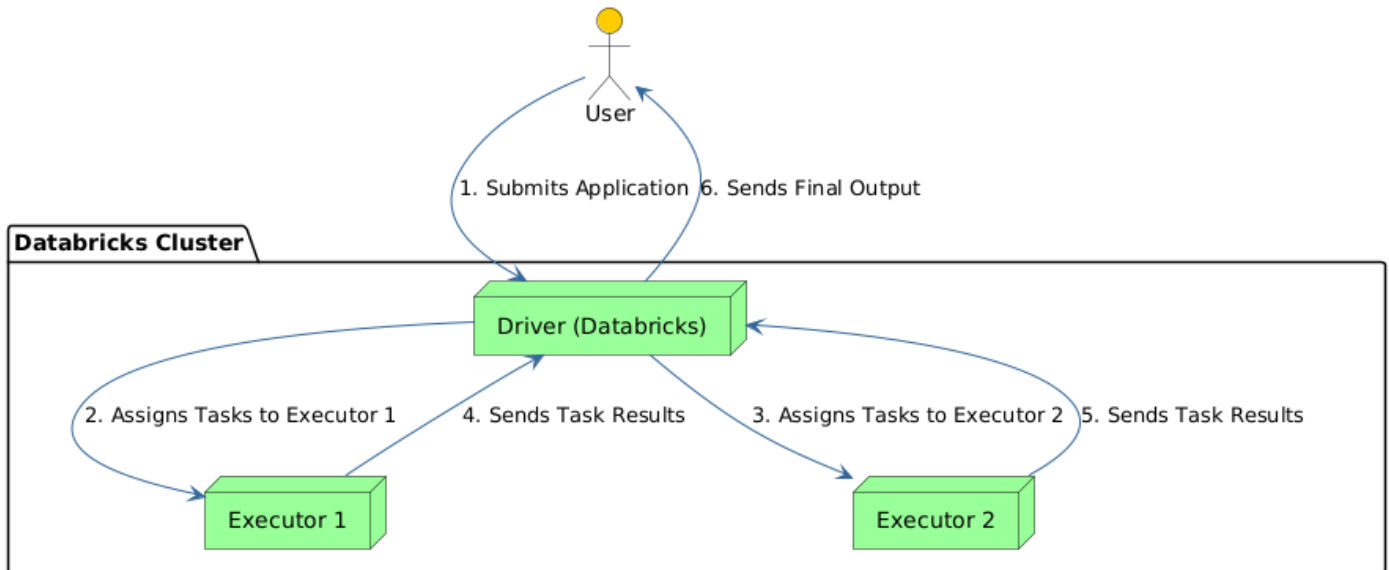
Databricks simplifies the deployment of Spark applications and abstracts the complexities of deployment modes. Here's how:

1. Unified Environment:

- Databricks combines the benefits of both **Client Mode** and **Cluster Mode** by managing the driver program and executors within its environment.



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2. Interactive Notebooks:

- Provides a seamless **notebook interface** for interactive development, similar to Client Mode, but hosted entirely on the Databricks platform.

3. Job Clusters:

- For production workloads, Databricks uses **job clusters**, ensuring stability and independence from user sessions, akin to Cluster Mode.

4. Enhanced Reliability:

- Databricks automates resource allocation and handles disconnections gracefully, allowing users to focus on development without worrying about deployment configurations.

5. Scalability and Optimization:

- Databricks clusters dynamically scale resources based on workload needs, offering better efficiency and performance compared to traditional deployment modes.

Conclusion

- Spark-submit gives flexibility to choose between Client Mode for development and Cluster Mode for production.
- Databricks takes this flexibility further by unifying and automating deployment, making Spark applications easier to develop, test, and deploy.

Master Spark Concepts Zero to Big data Hero:

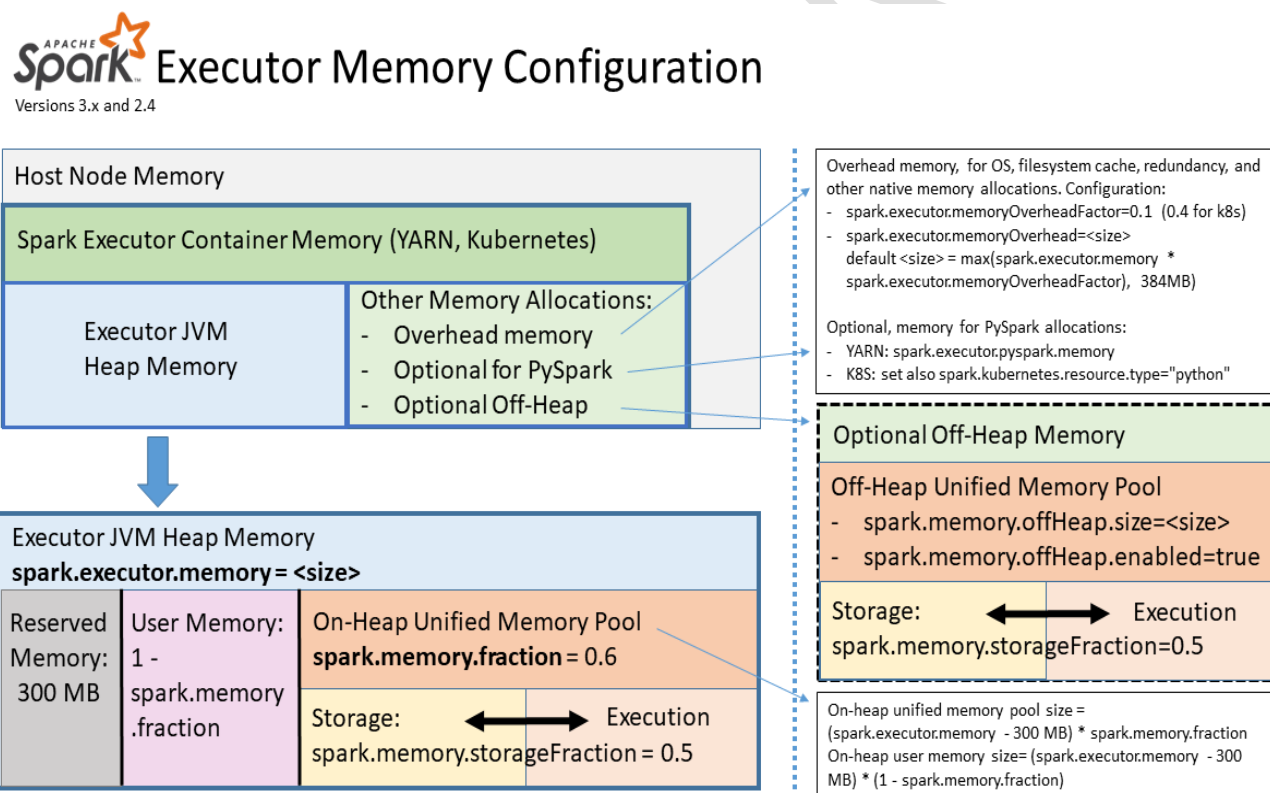
Introduction to Memory Management in Spark

Memory management in Spark is vital for optimizing performance and resource allocation. In Spark, memory is allocated at the **executor level** and involves three key areas:

1. **On-Heap Memory** – Managed by the JVM.
2. **Off-Heap Memory** – Managed outside the JVM.
3. **Overhead Memory** – Used for internal system operations.

Understanding how Spark handles these areas is crucial to avoid memory-related issues like out-of-memory errors.

Executor Memory Layout



An **executor** in Spark consists of several types of memory:

1. On-Heap Memory (JVM Managed):

- o **Execution Memory:** Used for operations like joins, shuffles, sorts, and aggregations.



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- **Storage Memory:** Used for caching (RDDs or DataFrames) and storing broadcast variables.
- **User Memory:** Stores user-defined objects, collections, and UDFs (User Defined Functions).
- **Reserved Memory:** Memory for internal system objects Spark needs to manage, like task metadata.

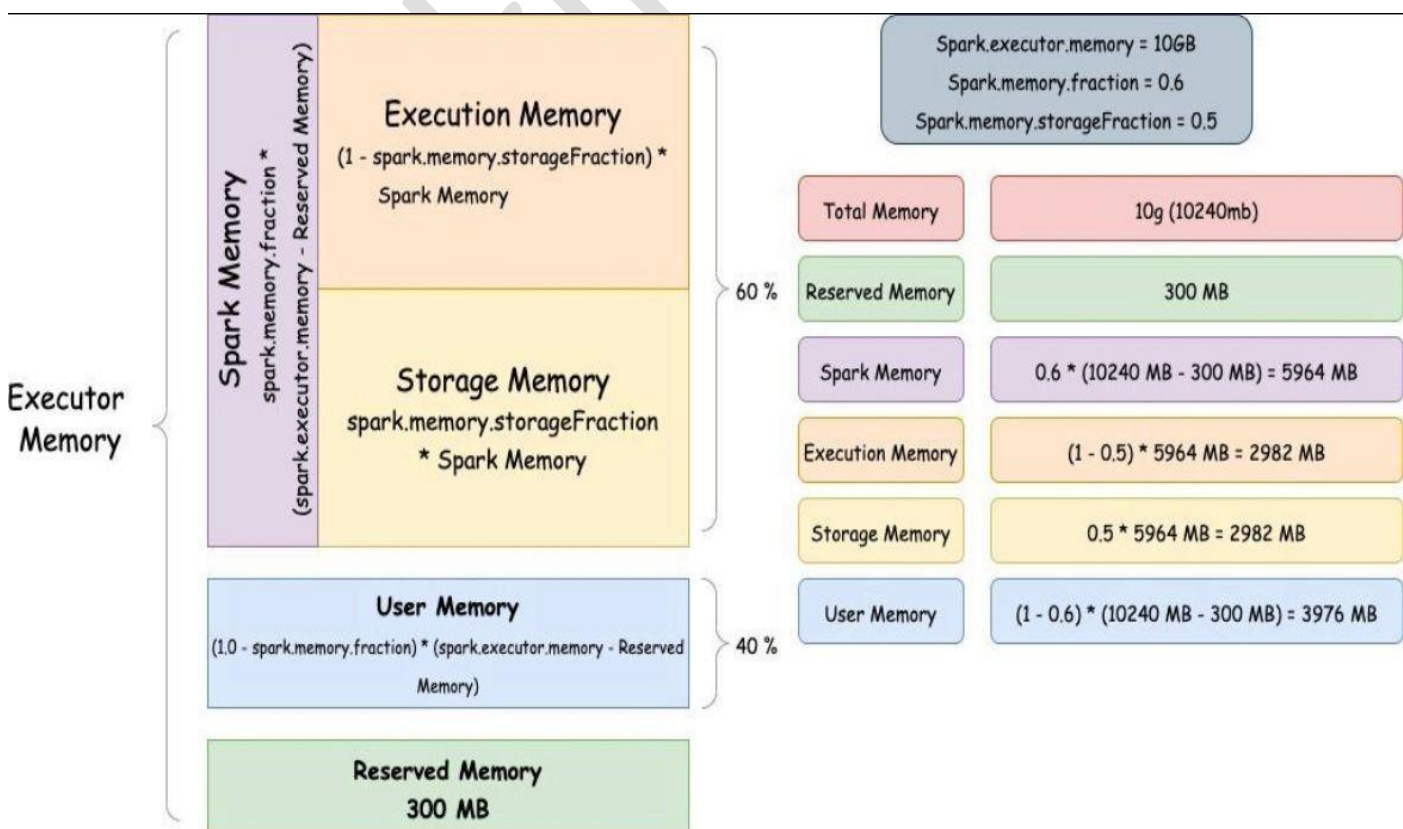
2. Off-Heap Memory (External to JVM):

- This memory is not directly managed by the JVM and can be used for certain optimizations like storing large objects or minimizing garbage collection impact. However, it must be manually enabled.

3. Overhead Memory:

- Memory required for Spark's internal operations. It is calculated as the **maximum of 384 MB or 10% of executor memory** (whichever is greater).
- Example: If the executor memory is set to 10 GB, the overhead memory would be **1 GB**.

Memory Calculation Example



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Let's consider an executor memory of **10 GB**. Here's how memory is typically divided:

- **Execution Memory:** This is for operations like sorting, shuffling, etc. It's calculated based on the configuration parameter `spark.memory.fraction`. This is typically **60% of the total executor memory**.
 - So for a 10 GB executor, **6 GB** is allocated to execution memory.
- **Storage Memory:** This is for caching RDDs and DataFrames. It's calculated based on the `spark.memory.storageFraction` parameter, which is typically set to **50%** of execution memory.
 - For a 10 GB executor, **3 GB** is allocated to storage memory.
- **User and Reserved Memory:** The remaining **3 GB** (from the 6 GB allocated to execution memory) is split into user-defined memory and reserved memory (default 300 MB).
- **Overhead Memory:** Spark will use **1 GB** of overhead memory (since 10% of 10 GB is 1 GB, which is greater than 384 MB).

Total Memory Request:

For an executor with 10 GB of memory, Spark requests **11 GB** in total (10 GB of executor memory + 1 GB of overhead memory).

Unified Memory Management (Post-Spark 1.6)

Spark's **unified memory management** is an important feature that allows the allocation of both execution and storage memory to share space dynamically. Before Spark 1.6, these memory areas were fixed and couldn't grow or shrink based on the workload, leading to inefficient memory use.

How Unified Memory Works?

- **Execution Memory:** When Spark executes transformations (like joins, shuffles, or aggregations), it uses execution memory.
- **Storage Memory:** When Spark needs to cache data (like RDDs or DataFrames) or store broadcast variables, it uses storage memory.

With **unified memory**, the execution and storage memory areas can dynamically borrow space from each other:

- **If execution memory is fully utilized**, unused storage memory can be used to fulfill execution requirements.



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- If storage memory is needed, Spark can evict cached data using the **LRU (Least Recently Used)** algorithm to free up space.
-

Off-Heap Memory

Off-Heap Memory refers to memory allocated outside the JVM heap. This is particularly useful in scenarios where:

- **Minimizing garbage collection overhead** is important.
- Storing large objects in memory without triggering JVM's garbage collection.

To enable off-heap memory, you need to explicitly configure the `spark.memory.offHeap.size` parameter. You can allocate a fixed size for off-heap memory, and Spark will use it for execution and storage just like on-heap memory.

For example:

- If off-heap memory is enabled, Spark might use **1 GB** of off-heap memory for execution and **2 GB** for storage, in addition to the on-heap memory.
-

Memory Allocation and Dynamic Behavior

- **Before Spark 1.6:** The memory for execution and storage was fixed, and Spark couldn't adapt the allocation dynamically, leading to possible memory wastage or performance issues.
- **Post-Spark 1.6:** The introduction of **dynamic memory allocation** allows Spark to allocate more memory to execution or storage depending on the workload:
 - **Execution Priority:** Execution memory always has higher priority than storage memory. This means if execution memory is needed, Spark will borrow from storage memory.
 - **Eviction:** When storage memory is needed, Spark evicts the least recently used (LRU) blocks to free up space.

Key Takeaways:

1. **On-Heap Memory:** Managed by the JVM and split into execution, storage, user, and reserved memory.



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2. **Off-Heap Memory:** External to the JVM and useful for reducing garbage collection overhead.
 3. **Overhead Memory:** Internal memory used by Spark's system-level operations.
 4. **Unified Memory:** Allows execution and storage memory to dynamically share space based on workload demands (introduced in Spark 1.6).
 5. **Dynamic Memory Allocation:** Post-Spark 1.6, memory can be adjusted between execution and storage memory, optimizing memory usage.
 6. **LRU Eviction:** Used to free up memory for execution or storage needs.
-

How Spark Requests Memory

When an executor is launched, Spark requests memory from the cluster manager (e.g., YARN):

- **Total Memory Request = Executor Memory + Overhead Memory.**
- If **off-heap memory** is enabled, it will be added to this request.

Example: For a 10 GB executor and 1 GB overhead, Spark requests **11 GB** of memory.

Conclusion

Understanding Spark's memory management system helps optimize performance by properly allocating resources and avoiding out-of-memory errors. With features like **unified memory**, **dynamic memory allocation**, and **off-heap memory**, Spark can efficiently manage memory based on workload demands, ensuring faster processing and reduced memory overhead.

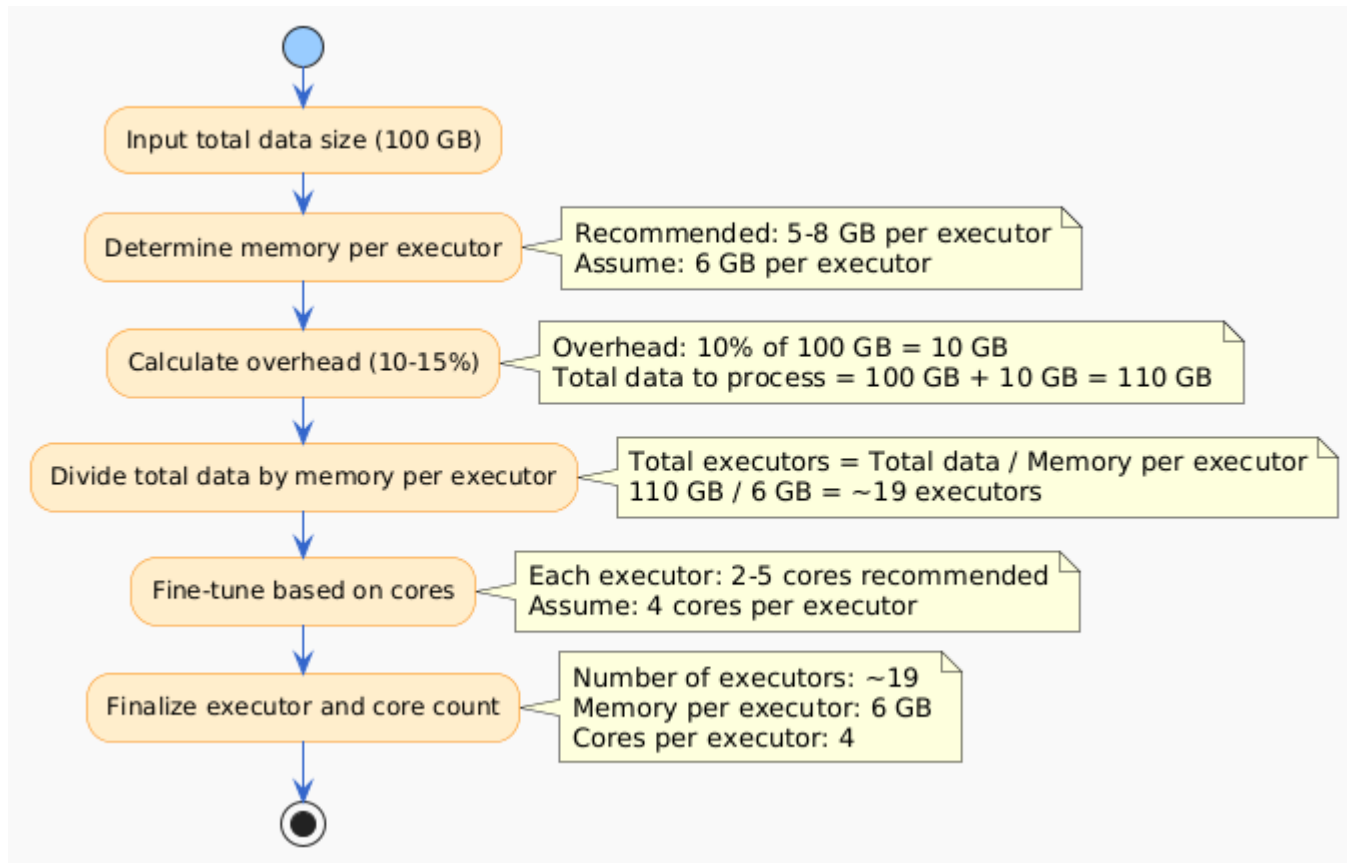


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Master Spark Concepts Zero to Big data Hero:

Calculate the Number of Executors Required to Process 100 GB

To calculate the number of executors required to process 100 GB of data in Spark, you need to consider several factors like the executor memory, overhead, core count, and how Spark partitions the data. Here's a step-by-step breakdown:



1. Determine the Memory per Executor:

The amount of memory allocated to each executor determines how much data it can handle. A general rule is to allocate around **5-8 GB per executor**, depending on the workload.

For example, let's assume **6 GB per executor** for your calculation.

2. Estimate the Data Overhead:

Spark incurs overhead for tasks like serialization, shuffling, etc. A typical overhead estimate is around **10-15% of the data size**.

For **100 GB of data**, this means:

- Overhead: **10% of 100 GB = 10 GB**
 - Total data to process: **100 GB + 10 GB overhead = 110 GB**
-

3. Estimate Data per Executor:

Next, divide the total data by the memory per executor to estimate how much data each executor will process.

For example:

- Total data: **110 GB**
 - Memory per executor: **6 GB**
-

4. Fine-tune Based on Cores:

Each executor should have at least **2-5 cores** for optimal performance. Let's assume **4 cores per executor** as a common practice. You'll need to divide the data processing load by the number of cores.

For example, if your cluster has **4 cores per executor**, the number of parallel tasks Spark can handle per executor increases, allowing for better performance and efficiency.

Summary:

To process **100 GB of data**:

- Memory per executor: **6 GB**
- Estimated overhead: **10 GB** (approx.)
- Total executors: **~19 executors**
- Each executor should have **4 cores**.

This is a general guide. For production workloads, always test and adjust based on your data's specific characteristics (like skewness, partitioning, and complexity of transformations).



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