# Auto-Scaling Techniques for Spark Streaming

Master-Thesis von Seyedmajid Azimi Gehraz aus Iran Tag der Einreichung:

1. Gutachten: Prof. Dr. rer. nat. Carsten Binnig

2. Gutachten: Dr. Thomas Heinze



Fachbereich Informatik Data Management

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Erklärung zur Master-	Th	esis
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Darmstadt, den September 25, 20	18
(Seyedmajid Azimi Gehraz)	

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#### 2 Problem Definition

#### 2.1 Introduction

Cloud computing has been on rise over the last decade. Parts of this popularity is due to its inherent features. It lets application developers to run their applications on virtual infrastructure. Virtual infrastructure lays the foundation of ondemand infrastructure. Developers acquire and release resources as required by workload. Examples of cloud providers are Amazon AWS [1], Microsoft Azure [50] and Google Cloud [24]. Today's cloud infrastructure is widely used by many customers for different purposes such as batch processing, serving static content, storage servers and alike.

As cloud environment brings up *elasticity* [32], it also introduces a new set of challenges and problems. Modern applications face fluctuating workloads. Typically, if a workload is *predictable*, resources are allocated ahead of time before load-spike starts. However, in many other scenarios predicting even near future workload is a not so easy task. Even though running an application in cloud environments helps to overcome a long standing problem of *over-provisioning*, low utilization is still one of the major problems of cloud applications. This has been confirmed by multiple studies [19] [54].

The root of the problem is originated from the fact that, most developers do not have enough insight about bottom and peak workload of their application. Thus, they fail to define an effective scaling strategy. Therefore, they end up with conservative strategies which in turn leads to low utilization. Hence, we need a system that automates the process of resource allocation. Auto-Scaling has been well studied in the context of web application. [28] [20] [33] are just a few examples. Chapter 7 explores more techniques and strategies.

#### 2.2 Objectives of Auto-Scaling Systems

The ultimate goal of an Auto-Scaling system is to automate the process of acquiring and releasing *resources* in order to minimize the *cost* with minimum violation of *service level objectives* (SLO). The definition of *resource* depends on the context. As an example, for a stateless web applications it means virtual machines or containers that run web server software. For an Auto-Scaling system to adjust required resources, it shall consider different aspects of application and environment. Additionally, the term *cost* is also defined in the context. As an example, it might mean monetary cost or just numerical value of resources. SLOs are predefined rules that shall not be violated during application runtime and are also defined in the context of application.

#### 2.3 Auto-Scaling in Data Stream Processing Systems

Data Stream Processing Systems are data processing systems that process unbounded stream of data unlike their batch-oriented counterparts. With the ever increasing adoption of IoT applications, it is critical to design stream processing systems that handles the incoming messages with high throughput and low latency. With static workloads, these problems could be solved by dominating stream processing systems like Apache Spark [7], Apache Storm [56] and Apache Flink [2]. However, the problem of low utilization also applies for stream processing systems as well. This leads us to a new generation of stream processing systems called Elastic Data Processing Systems that adopts elasticity concepts to stream processing system.

Prior to this thesis a number of studies have been performed on elastic stream processing system. [14], [29] are just a few samples. One of the dominating stream processing systems is Apache Spark which support both batch and stream processing. In order to support both workloads, Apache Spark has a unique architecture that partitions the input workload into predefined window of batches – an architecture known as micro batching. With this common architecture as a foundation, number of interesting challenges arise that need to be considered for elastic workloads.

This thesis will focus on dynamic resource allocation in the context of Apache Spark. An extensible framework will be developed based on a prior work by Kielbowicz [42]. This thesis will extend the existing prototype and implement multiple Auto-Scaling techniques for Apache Spark Streaming and will evaluate these techniques using real-world workloads. The ultimate goal is to identify how the architecture of Spark Streaming influences the performance of the different Auto-Scaling techniques.

#### 2.4 Requirements of Thesis

Since Auto-Scaling is a quite wide realm, this thesis focuses on Apache Spark Streaming with a couple of predefined requirements. Any proposed solution must adhere to the following requirements and limitations to a large extent.

- Online Decision Making. Since Spark Streaming is an online data stream processing system, any proposed solution shall apply scaling decisions in an online manner without causing any downtime on target system.
- **QoS Guarantees**. Scaling decisions shall avoid violating predefined SLOs as much as possible. Understandably, violating SLOs is an inevitable incident. However, this should be kept under an acceptable level.
- **Extendibility**. Any proposed solution shall provide some degree of extendibility such that, future modifications can be applied without any major code refactoring.
- Reconfigurability. Any proposed solution should be easily configurable by administration or *devops* team.
- Workload Independence. Any proposed solution should not impose any assumption on type of workload running under Apache Spark Streaming.
- Computationally Feasible. As mentioned in first requirement, any proposed solution should be computationally
  feasible such that it could generate results in order of seconds.
- Without Spark Core Modification. In order to make the solution extensible as much as possible, any proposed solution, is not allowed to modify spark-core or spark-streaming packages.

#### 2.5 Summary

As mentioned this thesis will focus on dynamic resource allocation in the context of Apache Spark. The thesis is organized as follows. Chapter 3 introduces and explains basics of Auto-Scaling techniques. Chapter 4 introduces the architecture of Apache Spark Streaming. Chapter 5 explains structure and design considerations of this thesis. Chapter ?? discusses implementation details and challenges faced during implementation. Chapter 6 evaluates the implementation under different workloads. Chapter 7 includes discussion of prior and related work. Finally, chapter 8 concludes.

#### 3 Introduction to Auto-Scaling

#### 3.1 Introduction

As mentioned in Chapter 2 the key characteristic of cloud environments is *elasticity* behavior. However, manually adjusting resources are is not an effective approach to exploit this feature. Hence, we need to automate this procedure with minimal human intervention. This chapter introduces foundations of Auto-Scaling techniques. Different techniques and architectures will be discussed from a high level standing point. It shall be noted that, this chapter has been heavily inspired by work done by Lorido-Botran, Miguel-Alonso, and Lozano [48].

#### 3.2 Basic Concepts

The ultimate goal of an Auto-Scaling system is to automate the process of acquiring and releasing *resources* in order to minimize the *cost* with minimum violation of *service level objectives* (SLO). However, *resource* is a broad and context-dependent term. It refers to any form processing unit – engine – that provides application developers some form of computation power. This general purpose definition is broad enough to capture different kinds. In most cases, it means virtual machines allocated by cloud provider. In more modern distributed systems, it may refer to *containers* like Google Kubernetes [25]. However, a resource might be as simple as a single process or thread.

The term *cost* refers to any form of expenditure that users pay in order to acquire a resource. It doesn't necessarily mean *monetary* cost. It can also refer to numerical values of resources, like number of virtual machines or number of running processes. Although minimizing cost is the ultimate goal of any Auto-Scaler system, not in all cases cost reduction is desirable. It should be achieved with respect to defined *Service Level Objectives*.

Service Level Objectives are any predefined rules that shall be respected during application runtime. The following, defines a couple of SLO definitions for different applications:

- 99 percentile round-trip latency of requests in a web application should be less than 150 milliseconds.
- All committed records in master database must be replicated with a maximum delay of 5 milliseconds.
- All messages pushed by a producer, should be processed by respective consumers in less than 5 minutes.
- At least 95% of images published in the last 24 hours should be served by cache servers.

Service Level Objectives are typically determined and defined by business requirements. Defining effective and meaningful SLOs is a challenge on its own. However, it is out of the scope of this thesis.

From a high level point of view, Auto-Scaling is a *trade-off* amongst cost and violation of SLOs. Resource *under-provisioning* will degrade performance and leads to SLO violations, while on the other hand, resource *over-provisioning* leads to idle resources, hence unnecessary costs. To make an effective decision, an Auto-Scaler needs to consider application and its environment:

Infrastructure pricing model. Some infrastructure providers charge customers hourly. That is, if customer
acquires a resource at 10:30AM and releases it at 11:30AM, the customer is charged for two hours. Some other
service providers might charge on minute basis. Pricing model has a huge impact on decisions made by Auto-Scaler
system, since it makes some decisions pointless.

- Service level objectives. Each application has its own set of objectives that should be adhered by Auto-Scaling system. These SLOs might be defined and applied at *soft* and *hard* levels. Violating an SLO at soft level is not critical, albeit alarming. However, violating a resource at hard level is a critical issue and is a negative point for an Auto-Scaler.
- Acquire/Release delay. Depending on type of the resource, it might take some time for the resource to become responsive and ready to process user requests. For example, booting a virtual machine typically takes couple of minutes, whereas launching a container takes time in order of seconds. An Auto-Scaler shall consider whether acquiring and releasing a heavy weight resource in a *zig-zag* manner worth the overhead or not.
- **Unit of allocation**. In some cases, it might be beneficial to allocate multiple instances of a same resource at once. This might be due to the startup and initialization overhead or it might be the case that Auto-Scaler predicted a huge load spike in near future.

Auto-Scaling can be done with different techniques and strategies. The remainder of this chapter is organized as follows. Section 3.3 defines general architecture of an Auto-Scaler. Section 3.4 clarifies which sort of actions can be applied by an Auto-Scaler. Section 3.5 classifies different techniques and briefly explains each category. Finally, section 3.6 concludes.

#### 3.3 Generic Auto-Scaler Architecture

Figure 3.1 illustrates generic architecture of an Auto-Scaler. This architecture is broad enough to capture different kinds of applications. First, responsibilities of different components shall be clarified. Table 3.1 summarizes all components of the system. An Auto-Scaling system typically consists of following sub-components.

Component	Description
Clients	Users of the application which might be applications on their own
Load Distributor	Distributes incoming requests to application instances
Application	Runs business logic defined by developer
Metrics Engine	Monitors and collects metric from application and provides it to other components
Infrastructure API	Provides API to adjust (acquire/release) resources
Auto-Scaler	Runs the auto-scaling algorithm based on metric collected by Metrics Engine

Table 3.1: Summary of Auto-Scaler Components

Clients Most kinds of applications, typically have some form of client that sends requests to the system and either waits to get a reply or operates under *fire-and-forget* strategy. It shall be noted that, a client is not necessarily an end user. In today's modern distributed applications, an application might be client of another application.

Load Distributor In order to provide some degree of *transparency*, usually clients connect to a Load Distributor component. It is the responsibility of the Load Distributer that *proxies* client request to application. Load Distributor is also a generic component and might represent different technology in read-world applications. In the context of a web application, it might be an HTTP load balancer. Even a *Message Broker* can also be represented as a form of Load Distributer. It shall be noted that Load Distributor itself can be replicated or sharded for *high availability* or *scalability* reasons.

Application The application component runs the business logic. This architecture doesn't impose any limitation on application architecture. It might be a simple stateless web application. It might access to a back-end cache or database service. It might push some messages to a message broker, as a result of client request. It might be just a simple process consuming messaging provided by a message broker. It might forward client requests to other applications for further processing.

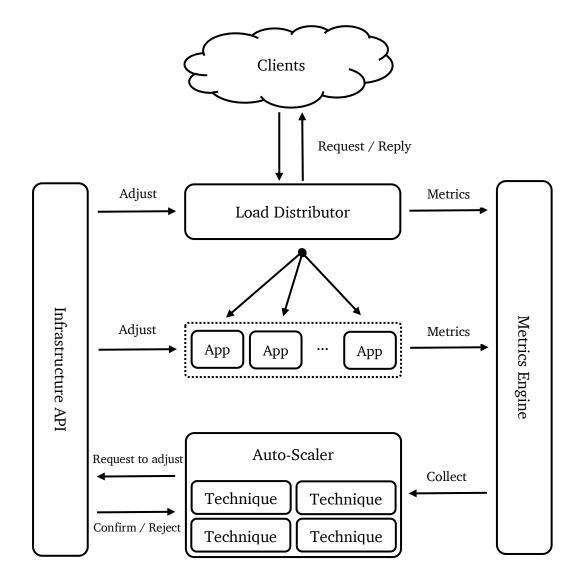


Figure 3.1: Architecture of a Generic Auto-Scaler

Infrastructure API Typically, when an Auto-Scaler system decides to take any action, it doesn't touch the application directly. In order to provide *separation of concerns*, this responsibility is handed over to infrastructure via an API provided by resource/service provider. An important issue that shall be noted here is that, service provider may schedule resource changes and execute them some time later. Thus, resource changes might not take effect immediately at the moment Auto-Scaler requests them. This fact shall be considered by Auto-Scaler system.

Metric Engine Auto-Scaler system needs to have a good insight on current status of the application and incoming requests. Metric engine – known as *monitoring engine* takes the responsibility of measuring and collecting different aspects of the application. The term *metric* refers to any form of measurable aspect of an application or its environment. Ghanbari et al. [23] has defined and proposed a list of different type of metrics that could be exploited for different purposes.

- Hardware dependent metrics such as CPU usage, disk access time, memory usage, network bandwidth usage, network latency.
- Operating System provided metrics such as CPU-time, page faults, real memory.
- Load balancer provided metrics such as size of request queue length, session rate, number of current sessions, transmitted bytes, number of denied, requests, number of errors.

- **Web server** provided metrics such as transmitted bytes and requests, number of connections in specific states (e.g. closing, sending, waiting, starting, ...).
- **Application server** provided metrics such as total threads count, active threads count, used memory, session count, processed requests, pending requests, dropped requests, response time.
- **Database server** provided metrics such as number of active threads, number of transactions in a particular state (e.g. write, commit, roll-back, ...).

Since storing and reporting metrics has its own overhead, typically metrics engine aggregates collected values at different scale depending on how fresh it is. Rationally, fresh values are more important for Auto-Scaling system. As an example, it might provide near real time values for about 15 minutes. Then, for the last 5 hours, collected values are aggregated by a window of one minute. For last two days, it is aggregated by a window of 15 minutes and finally, for any record older than last two days, it is aggregated on hourly basis. Whether these aggregated values are sufficient for Auto-Scaler to make an accurate decision is out of the scope of this thesis. However, it may be a good idea to empirically adjust this system until it fits the requirements of Auto-Scaler system.

Auto-Scaler This component is the core of Auto-Scaling system. It consists of different techniques. Typically, an Auto-Scaler works in *consecutive rounds*. First, it collects metrics from Metrics Engine and offloads them to one or multiple technique implementations. It shall be noted that, an Auto-Scaler might utilize different implementation of techniques simultaneously – for different stages of the application as an example. Even it might utilize a single implementation under different configurations. This architecture does not impose any limit on the order of techniques. Then, based on some preferences or ordering mechanism, it chooses the *final decision*. Finally, it requests the Infrastructure API to change the number of resources. Naturally the assumption is that, infrastructure provider should be able to adjust the required or released resources. However, in some case it might not be able to fulfill the task. Thus, Auto-Scaler shall wait for a response to check whether the request has been successfully fulfilled or not.

For the sake of understandability, Algorithm 1 describes the above procedure in pseudo code.

#### 3.4 Actions

In each round an Auto-Scaler decides to take an action and request that specific action to Infrastructure API. Table 3.2 summarizes feasible actions for an Auto-Scaler. This thesis, assumes three possible actions.

- Scale-In. This action implies that Auto-Scaler has decided to remove one or more resources. It doesn't necessarily means, Infrastructure API will be able to remove all the requested resources. Consequently, Infrastructure API might be able to remove any subset of resources among those that were requested to remove. In this thesis, we assume Infrastructure API offers such a behavior to notify Auto-Scaler about the actual removed resources either synchronously in response to Auto-Scaler's request or via an asynchronous API provided by Auto-Scaler.
- Scale-Out. This action implies that Auto-Scaler has decided to add one or more resources. Similar to Scale-In action, Infrastructure API might be able to fulfill the request or not. In case, it doesn't have enough resources to allocate, this incident should be reported to Auto-Scaler component. This thesis assumes this behavior. Response shall propagate back to Auto-Scaler similar to steps described in Scale-In action.
- **No-Action**. This action implies that Auto-Scaler has decided to do nothing but stay with the same number of resources as last round. This is a kind of *no-operation*.

It's noteworthy that in all cases, Auto-Scaler is allowed to store any history of actions taken so far. In fact, it is a special category of Auto-Scalers known as *stateful* Auto-Scalers. Refer to section 3.5 for further discussion and explanation on taxonomy of Auto-Scalers.

#### Algorithm 1: General Work-Flow of an Auto-Scaler

```
1 // different implementations of techniques
2 implementations ← []
3 // decision of each implementation
4 decisions ← []
5 // final decision of Auto-Scaler
6 finalDecision ← null
7 // instantiate as many techniques as required
s for i \leftarrow 0 to n do
   |implementations[i] \leftarrow InstantiateTechnique(i)
10 end
11 repeat
      // load monitoring data from metrics engine
12
      currentValue ← GetCurrentMetricsFromMetricsEngine()
13
      // initialize decisions
14
      decisions \leftarrow []
15
      for i \leftarrow 0 to n do
16
         impl \leftarrow implementations[i]
17
         decisions[i] \leftarrow GetDecision(impl, currentValue)
18
      end
19
      // calculate final decision based on some weight or ordering mechanism
20
      finalDecision ← GetFinalDecision(decisions, currentValue)
21
      // request infrstructure API to adjust resources
22
      reply ← RequestInfrastructureAPI(finalDecision)
23
      // in case infrastructure reject request, warn developers
24
25
     if reply = ReplyStatus.REJECT then
         // issue a warning
26
         LogError("request can not be fulfilled")
27
28
      end
29 until Auto-Scaler is running
```

Action	Description
Scale-In Scale-Out	Remove/Release one or more resources Acquire/Add one or more resources
No-Action	Do nothing

Table 3.2: Feasible Actions of an Auto-Scaler

Another aspect of taking an action is that, Auto-Scaler is allowed to Scale-In/Out horizontally or vertically in each round independent of previous rounds. Horizontal Scale-In/Out refers to a category of actions that acquires or releases resources in parallel to each other. For example, in the context of a web application, adding or removing one or more virtual machines is considered as a horizontal scaling action. While on the other hand, Auto-Scaler might decide to just scale by adding hardware resources. For example, it might decide to add more RAM or remove couple of CPU cores in one specific virtual machine. This kind of scaling action is considered as vertical scaling action.

Actions are not necessarily applied at a constant rate. Auto-Scaler is in full charge of taking actions at *exponential* rates. For example, in consecutive rounds, an Auto-Scaler can decide to acquire 1, 2, 4, 8, 16 virtual machines per round. This also applies for Scale-Out actions. Nothing hampers an Auto-Scaler from changing rate of scaling actions in each round. It might even decide to apply different rates for different stages of the application like *startup* phase, or *near-ending* phase, etc.

Last but not least, an Auto-Scaler might decide to apply a *grace period* after taking an action independent of previous rounds. A grace period is a time frame, in which Auto-Scaler does not take any further action in order to let cluster of resources stabilize. Similar to action rates, grace period can also be applied at different rates. For example, Auto-Scaler might wait for 10, 20, 40 seconds after taking Scale-Out action in consecutive rounds.

#### 3.5 Taxonomy of Auto-Scaling Techniques

Auto-Scalers can be modeled and classified in different categories. In a sense, it is a multi-dimensional space of features and characteristics. Table 3.3 lists and summarizes different dimensions of Auto-Scaling. In the rest of this chapter each dimension is described and explored in details.

Dimension	Description
Schedule-based versus Rule-based	Whether Auto-Scaler applies decisions manually or based on predefined rules.
Reactive versus Proactive	Whether Auto-Scaler reacts to workload changes or predict future workload ahead of time.
Execution mode	Whether Auto-Scaler is central component or operating in a distributed fashion.
Algorithm family	Which family of algorithms is applied to make the decision.

Table 3.3: Dimensions of Auto-Scaling

#### 3.5.1 Schedule-Based versus Rule-Based

Some applications have a very basic cyclic workload pattern on daily basis which could be manually modeled as *cron* style jobs. Schedule-based systems can not adopt to unplanned workload changes. Since this type of Auto-Scalers are in conflict with thesis requirements, it won't be studied in this thesis.

On the other side of the extreme, there exists *rule-based* Auto-Scalers. These Auto-Scalers take actions based on set of rules defined by application developers inspired by business requirements. Each rule is based on one or multiple *constraints*. Each constraint consists of one or a set of conditions around some variable. Variables can be defined by *application* – number of tweets per second, as an example – or by its environment – CPU utilization, for example. For example, if average network bandwidth is more than 80% of maximum available bandwidth for 10 minutes, then take Scale-Out action.

There are different criteria to calculate defined variables. Some applications might define minimum or maximum values for variables. In some other cases, average values might be useful. Furthermore, average/minimum/maximum values might be defined for a window of time or number of occurrences of specific event. Time or event windows in turn could be defined as a *static* window – where it moves with a fixed interval – or as a *sliding* window – where it smooths over elements.

#### 3.5.2 Reactive vs Proactive

In another dimension, Auto-Scalers can be partitioned into two categories. *Reactive* approaches monitor the workload in order to find a meaningful change. Thereafter, they apply some algorithm to figure out the final decision. To view different family of algorithms, refer to section 3.5.4. It's noteworthy to express that, in some applications by the time a reactive Auto-Scaler decides to take some action, it might be too late. In other words, taking an action in an already overloaded application might not be desirable in some cases. Taft et al. [58] argues that applying reactive approaches in the context of OLTP databases is not desirable. This leads us to another class of Auto-Scalers known as *proactive* Auto-Scalers.

*Proactive* Auto-Scalers try to predict *future* workload ahead of time before facing workload spikes. Hence, they are called as *predictive* approaches. Whether the term *future* is defined as near future or long term future depends on the type of Auto-Scaler. This has the advantage that, by the time load spike occurs, the required resources are already available, warmed-up and ready to respond user requests. As mentioned in section 3.2, allocating some resources takes some time to become fully initialized. For example, adding a database replica is not an immediate action. It takes some time re-replicate database records and validate the replicas. Thus, for some scenarios, even though reactive approaches might seem to be sufficient, but due to initialization latency it is not an applicable approach.

#### 3.5.3 Execution Mode

The generic architecture which is described in section 3.3 does not impose any limit on the operation and execution mode of the Auto-Scaler. In other words, Auto-Scaler itself might run as a *multi-instance* application. There are different types of proposed execution modes. But they can be classified in following generic groups without loosing generality. Table 3.4 summarizes operation modes.

<b>Execution Mode</b>	Description
Global Controller	At a specific point in time, a single controller is responsible of taking actions.
Distributed Without Coordination	Auto-Scalers operate independent of each other without performing any form of coordination.
Distributed Coordinated	Auto-Scalers perform in distributed mode, but they are allowed to communicate and cooperate with each other.
Hierarchical Controllers	A hierarchy of controllers cooperate with each other to make final decision.

Table 3.4: Execution Modes of an Auto-Scaler

Global Controller In this architecture, Auto-Scaler runs and controls the application as a *central* or *master* component. One or multiple *backup* or *slave* controllers might also accompany master node. In case master controller fails, one of the backup controllers kicks in and takes over the responsibility. In this model, it's the responsibility of master node to make decisions and execute actions. To detect failure of master controller, there exist already well established *cluster coordination* utilities like Apache Zookeeper [8] or CoreOS Etcd [17]. Figure 3.2 depicts a variation of original Auto-Scaler architecture that performs under master-slave model coordinated by a Zookeeper ensemble.

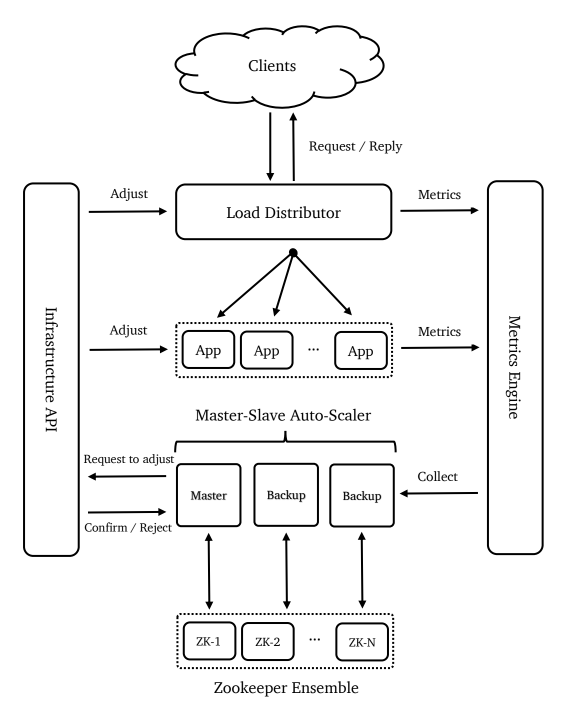


Figure 3.2: Architecture of Master-Slave or Global Controller

Distributed Without Coorindation In this architecture, typically Auto-Scaler runs alongside the application instances and each Auto-Scaler instance only *controls* the *local* application. The major difference against global controller is that, in this model each Auto-Scaler makes decision at its own will without any form of coordination with other Auto-Scalers. It's usually the most scalable solution when size of the cluster grows. However, since each Auto-Scaler instance only controls the local application, it might lead to sub-optimal decisions because local controllers lacks global view of the cluster. Figure 3.3 illustrates this architecture.

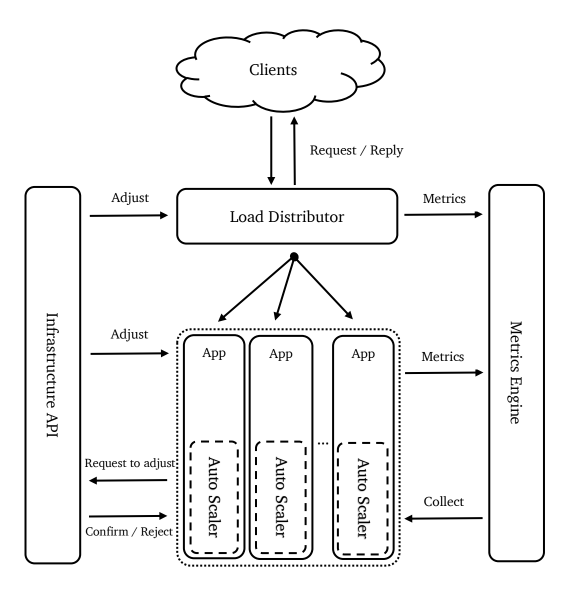


Figure 3.3: Architecture of Distributed Non-Coordinated Controller

Distributed Coordinated This architecture is similar to previous model. However the difference is that, Auto-Scaler components coordinate with each other over a communication channel to gather more information. The amount of communicated information heavily depends on underlying algorithm. In some scenarios only neighbor nodes are contacted. In other cases, all nodes might communicate with each other in order to achieve consensus. Figure 3.4 depicts this architecture.

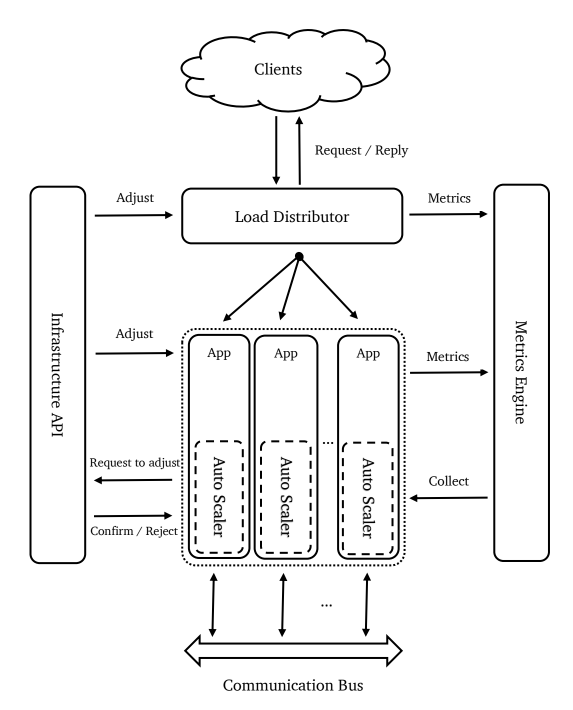


Figure 3.4: Architecture of Distributed Coordinated Controller

Hierarchical Controllers This architecture is the mixture of previous models. It's a combination of *Global Controller* and *Distributed Coordinated Architecture* model. Local Auto-Scalers communicate with a global controller in order to perform necessary actions. Local Auto-Scalers might also communicate with each other. Additionally, one or more back controller might also accompany global controller. Furthermore, there might be no limit on who actually performs the actions. Each Auto-Scaler – whether it is operating in local or global model – might perform actions independent of the others. Figure 3.5 illustrates this architecture.

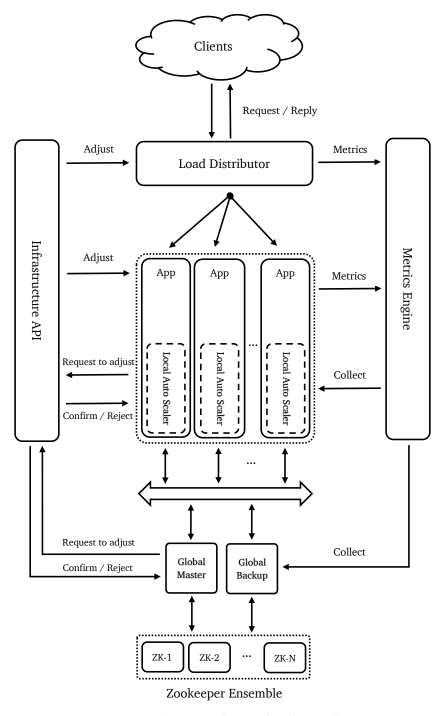


Figure 3.5: Architecture of Hierarchical Controllers

#### 3.5.4 Algorithm Family

From algorithmic point of view, Auto-Scalers can be classified into 5 different categories.

- Threshold-Based Policies
- Time-Series Analysis
- · Reinforcement Learning
- · Queuing Theory
- · Control Theory

In this section, each category is explained to some extent.

#### 3.5.4.1 Threshold-Based Policies

According to section 3.5.2, threshold-based approaches follow a purely reactive approach. It lacks any form of future workload prediction. Threshold-based techniques usually involve a set of constraints which monitors performance data gathered from Metrics Engine. In its naive form, each rule define an *upper* and/or *lower* bound plus two time periods that define how long that specific metric was above the upper threshold or bellow the lower threshold.

Algorithm 2 better describes this approach in a pseudocode. Refer to section 7 to review list of proposed solutions.

#### Algorithm 2: General Work-Flow of Threshold-Based Techniques

```
1 // upper threshold
2 UpperThreshold ← LoadFromConfig()
3 // time period that performance metric was above upper threshold
4 UpperPeriod ← LoadFromConfig()
5 // lower threshold
6 LowerThreshold ← LoadFromConfig()
7 // time period that performance metric was lower than lower threshold
8 LowerPeriod ← LoadFromConfig()
9 // number of resources to acquire in each round
10 NumberOfResourcesToAcquire ← LoadFromConfig()
11 // number of resources to release in each round
12 NumberOfResourcesToRelease ← LoadFromConfig()
13 repeat
     currentValue ← GetCurrentMetricValue()
14
     if currentValue > UpperThreshold for UpperPeriod seconds then
15
        // acquire resource
16
        AcquireResource(NumberOfResourcesToAcquire)
17
        DoNothingDuringGracePeriod()
18
     end
19
     if currentValue < LowerThreshold for LowerPeriod seconds then
20
        // release resource
21
        ReleaseResource(NumberOfResourcesToRelease)
22
        DoNothingDuringGracePeriod()
23
     end
24
25 until Auto-Scaler is running
```

#### 3.5.4.2 Time-Series Analysis

In contrast to threshold-based techniques, time-series analysis approaches are purely proactive or predictive approaches. A *time-series* is a collection of data points sampled and ordered iteratively at uniform time intervals [48]. Time-series analysis typically requires to store a history of data points. Hence, storage and computation requirements are more intensive than threshold-based approaches. Refer to section 7 for evaluation of prior works in time-series.

According to the theory of time-series analysis [11] [33], time-series can be decomposed into multiple components.

- Season The season component captures recurring patterns that are composed of one or more frequencies, e.g. daily, weekly or monthly patterns. These dominant frequencies can be determined by using a *Fast Fourier Transformation* (FFT) or by *Auto-Correlation* algorithms.
- **Trend** The trend component can be described by a *monotonically* increasing or decreasing function that can be approximated using common regression techniques.
- **Noise** The noise component is an unpredictable outliers of various frequencies with different amplitudes. The noise can be absorbed to some extent by applying smoothing techniques like *Weighted Moving Averages* (WMA), by using *Lower Sampling Frequency* or by a *Low-Pass Filter* that eliminates high frequencies.

A time-series has a couple of important characteristics that reveals more information about the values in it. The following represents some of these characteristics.

- **Burstiness** The burstiness defines the impact of fluctuations within the time series and usually calculated by the ratio of the *Maximum Observed Value* to the *Median* within a sliding window.
- **Length** The length of the time series mainly influences the accuracy of approximations for the components mentioned above. It can be modeled as *static* or *sliding* window of time.
- **Relative Monotonicity** The relative monotonicity is the maximum number of consecutive monotonic values either *increasing* or *decreasing* within a window and indirectly determines the influence of the noise and seasonal components.
- Mean, Median, Standard Deviation and Quartiles These values are important indicators for the distribution of values how values are spread in time series.
- **Frequency** The frequency of a time series represents the number of values that form a period of interest. This value is an important input as it defines the starting point to find seasonal patterns.

#### 3.5.4.3 Reinforcement Learning

Reinforcement Learning is a technique that relies on direct experience from environment. The decision maker – known as *agent* – tries to learn the best possible action for each *state* of the environment. The final goal is to maximize the returned *reward*. In the context of Auto-Scaling problem, the agent is the Auto-Scaler component that gets current system state (any group of performance metrics) and tries to minimize/maximize some aspect of the application (response times, throughput, etc.) by performing Scale-In, Scale-Out or No-Action.

In order to formally define Reinforcement Learning environments, some basic terminology shall be defined [57]. At each time step t, where t = 0, 1, 2, ... is a sequence of discrete time steps, the agent receives a representation of the environment's state  $s_t \in S$ , where S is the set of possible states, and based on that, selects an action,  $a_t \in A(s_t)$ , where  $A(s_t)$  is the set of possible available actions in state  $s_t$ . One time step later, as a consequence of performing action  $a_t$ , the agent receives a numerical reward  $r_{t+1}$  and lands in a new state  $s_{t+1}$ . At each time step, the agent implements a mapping

from states to probabilities of selecting each possible action. This mapping is called the agent's *policy* and is denoted as  $\pi_t$ , where  $\pi_t(s, a)$  is the probability of action a at state s. This procedure is depicted by Figure 3.6<sup>1</sup>.

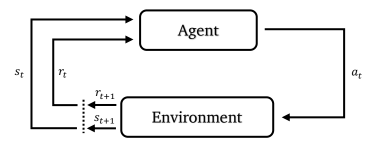


Figure 3.6: Reinforcement Learning Agent

In Reinforcement Learning environments future states can be determined only with the current state, regardless of the past history. This is known as *Markov Property*, which formally states that the probability of a moving to a new state  $s_{t+1}$  only depends on the current state  $s_t$  and the a. In other words, it is *independent* of all previous states and actions [57] [48]. Equation 3.1 defines this property.

$$P(s_{t+1} = s', r_{t+1} = r | s_t, a_t) = P(s_{t+1} = s', r_{t+1} = r | s_t, a_t, r_t, s_{t-1}, a_{t-1}, r_{t-1}, \dots, s_1, a_1, r_1, s_0, a_0)$$
(3.1)

A stochastic process that satisfies the Markov property is called a *Markov Decision Process* (MDP). It is typically represented as a 5-tuple consisting of *states*, *actions*, *transitions probabilities*, *rewards* and *policy* function.

- Set *S* which represents the state space of the environment.
- Set *A* which represents the total possible actions.
- Reward function R defined as  $S \times A \to R$  which represents the reward for each state-action pair. R(s,a) is the reward received after executing action a at state s.
- Probability distribution T defined as  $S \times A \to P(S)$  which specifies a probability distribution over the set S. P(s'|s,a) specifies the probability of landing in state s' assuming that the agent is in state s and performs action a.
- Policy function  $\pi$  defined as  $S \to A$  that maps state s to some action a. A policy function that maps state s to best possible action with highest expected reward is called *optimal* policy denoted as  $\pi^*$ .

An important aspect of reward function is that, reward must be accumulated with a *Discount Rate* denoted as  $\gamma$ . This is introduced to prevent *infinite* reward accumulation and force reward values to converge. Equation 3.2 defines reward function in combination with discount rate.

$$R_t = r_{t+1} + \gamma r_{t+2} + \gamma^2 r_{t+3} + \dots = \sum_{k=0}^{\infty} \gamma^k r_{t+k+1}$$
(3.2)

Last but not least, under policy  $\pi$ , *value-state* function  $V^{\pi}(s)$  defines an estimate value of expected reward for state s. Equation 3.3 defines value-state function where  $E_{\pi}$  defines the expected reward value. The value-state function is the core of most Reinforcement Learning techniques. In section 5, two of the techniques used in this thesis will be further discussed. Furthermore, section 7 evaluates prior work regarding Reinforcement Learning approaches.

$$V^{\pi}(s) = E_{\pi}(R_t|s_t = s) = E_{\pi}\left(\sum_{0}^{\infty} \gamma^k r_{t+k+1}|s_t = s\right)$$
(3.3)

The figure has been taken from Sutton and Barto [57]

#### 3.5.4.4 Queuing Theory

Queuing theory has been thoroughly used in the context of packet processing to estimate the average wait time until a packet could be routed. The same principle can be applied in the context of Auto-Scaling problem. Clients send requests at a *Mean Arrival Rate* denoted as  $\lambda$ . Requests are enqueued until they are processed with at a *Mean Service Rate* denoted as  $\mu$ . Figure 3.7 portrays two architecture of queuing systems. Furthermore, multi-tiered applications are modeled as multiple queues attached *serially* or in *parallel*, depending on the architecture of the application.

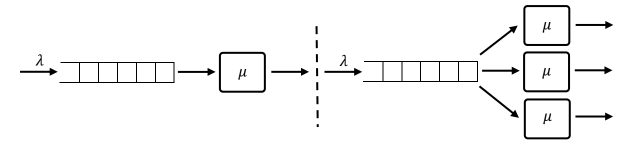


Figure 3.7: Queuing Theory with Single Processor (Left), with Multiple Processors (Right)

Queuing systems are modeled with *Kendall*'s notation [41]. A queue is modeled as A/B/C/K/N/D. Each variable is defined and further explained in the following.

- A Request inter-arrival time distribution.
- B Service time distribution.
- **C** Number of request processors.
- K Maximum length of the queue. This is an optional parameter and is set to unlimited if not specified.
- N Calling population. This is an optional parameter and is set to unlimited if not specified. This parameter controls the population of requests entering the system. In case of an *open queuing* system, this value is unlimited. While in a *closed queuing* system the population of customers is a finite value.
- **D** Priority order which defines in which order requests will be processed by request processors. This parameter is optional and is set to *First Come First Served* (FCFS) if not specified.

Queuing theory is mostly capable of modeling stationary systems with constant parameters defined above. In case of Auto-Scaling problems, these values have to periodically recomputed, since request inter-arrival distribution, service time and number of request processors are changing constantly. These parameters can be re-calculated mostly based on two approaches. First, by applying *analytic* techniques – suitable for simple queuing systems. Second, by applying *Discrete Event Simulation* (DES) techniques – suitable for more complex models.

#### 3.5.4.5 Control Theory

Control Theory is categorized as mixture of reactive and proactive systems and decomposed intro three major categories.

• Open Loop Systems These systems do not consider any feedback returned by underlying systems. They only consider current state of the system in order to make decision. The output of the system is not considered to check whether it is in stable and desired state.

<sup>&</sup>lt;sup>1</sup> The figure has been taken from Lorido-Botran, Miguel-Alonso, and Lozano [48]

- **Feedback Systems** These system monitor the output of the system in order to check whether it is complying to defined business conditions. These systems are mainly reactive systems. Figure 3.8<sup>1</sup> depicts these controllers.
- Feed-Forward System These systems pro-actively try to predict the system's future state and behavior.

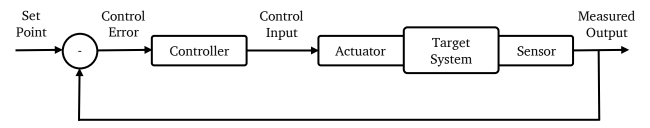


Figure 3.8: Feedback Control System

Feedback driven control systems are further divided into multiple categories [51].

- **Fixed Gain Controllers** The most simple type of feedback control systems. However, the configuration parameters of the controller are fixed during system runtime.
- Adaptive Controller These controllers allow to change some of the configuration parameters during runtime
  which makes them an applicable option for slowly-changing environments, but not for suitable for bursty workloads.
- Reconfiguring Controller In contrast to Adaptive Control controllers in which only configuration parameters can be changed at runtime controller itself does not change in this model the controller itself can also be changed at runtime. With this enhancement, Reconfiguring Controllers are more resilient to bursty workloads.
- **Model Predictive Controller** These controllers mix some *predicative* features into feedback controllers which make them even more resilient to unanticipated workloads.

#### 3.6 Conclusion

In this chapter, different architectural aspects and major design considerations of Auto-Scalers have been discussed. As mentioned, it is multi-dimensional space of features and capabilities. Many of the current proposals do not map to one category, but combine divergent set of features. Section 7 compares and evaluates prior research on different systems.

<sup>&</sup>lt;sup>1</sup> The figure has been taken from Patikirikorala and Colman [51]

#### 4 Apache Spark and Spark Streaming

#### 4.1 Introduction

Apache Spark [7] is a general purpose data processing framework that supports wide range of applications from *batch* processing, *stream* processing, *graph* processing to *machine learning*, etc. In this chapter the architecture of Apache Spark and Spark Streaming – as one of its sub projects – will be explored and discussed. Throughout the chapter, minor examples will also be presented to further simplify the concepts. This chapter is organized as follows. Section 4.2 explains the basic concepts of Apache Spark. Then, section 4.3 explains *Resilient Distributed Datasets* (RDDs) which is the most integral component of the Apache Spark. Section 4.4 explains *Discretized Streams* (DStreams) as the fundamental building block for data stream processing applications. Finally, section 4.5 concludes this chapter.

#### 4.2 Basic Concepts

Map-Reduce [18] and its derivative projects have been widely used by *data-oriented* applications to process and crunch huge datasets over last the decade. In traditional Map-Reduce environments, developers typically create *acyclic* data flow graphs to process input data. However, as confirmed by Zaharia et al. [65], there are two categories of applications that are not well suited for this architecture.

- Iterative Jobs Many machine learning applications process the same input *iteratively*. In traditional model, each iteration should be defined as a separate MapReduce job. While this is feasible, but for each job, input data has to be loaded from disk which leads to serious performance issues.
- Interactive Analysis MapReduce derivative projects like Hive [4] and Pig [6] have been extensively used to run SQL queries on top of massive datasets. Whenever a user submits different queries over the same dataset, the ideal solution would be to load all datasets into memory once and then execute different queries on top of it. However, with traditional model of execution each query shall be defined as a separate job which reads input data from disk.

Apache spark has been designed from ground up to resolve these issues. It provides a large stack of tools to facilitate processing large datasets. Figure 4.1 depicts tools provided by Spark.

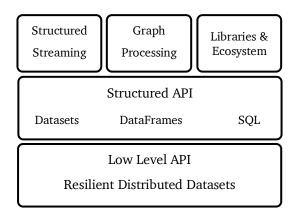


Figure 4.1: Apache Spark Stack

#### 4.2.1 Spark Runtime Architecture

Any Spark application consists of multiple components at runtime. The following lists the relevant components from a high level point of view.

Driver Process Driver It is the core component of every Spark application. It is a single process and runs on one of the nodes in the cluster. It maintains all the critical information such as user's program, input files, output files and data flow of the application. The driver process should run during the runtime the of the application. In case driver process fails, the whole application is considered as *dead* and should be restarted by any *fault-tolerant* mechanism available in the cluster.

Distributed File System It provides a shared file system accessible by any node in the cluster. There is no limitation on type of the file system but typically *Hadoop Distributed File System* (HDFS) [3] is used.

Worker Processes A collection of worker processes known as *executors* that run on cluster nodes. Executors run user defined code. During the runtime of the application each worker consume any number of input records, processes it based on user defined code and emits any number of output records. Similar to driver process, liveness of the worker processes shall by monitored. However, the role of worker processes are not as critical as driver process, even though they may fail for any reason. Workers have access to load/store data files from/to distributed file system or local memory of the machine. During lifetime of the application, status of the workers will be reported to driver process.

User Defined Code Provided by user and caries the main logic of the application.

Figure 4.2<sup>1</sup> depicts coarse grained architecture of a Spark application and table 4.1 summarizes these components.

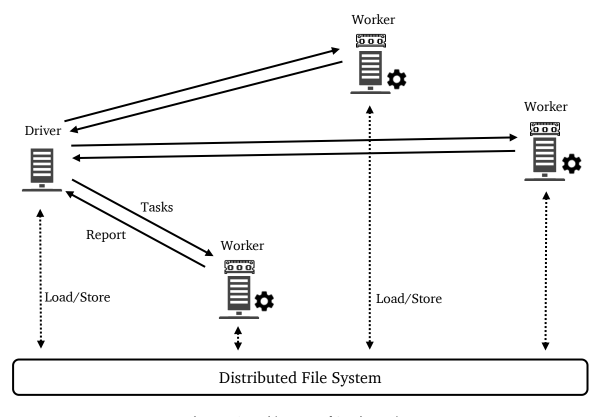


Figure 4.2: Architecture of Spark Runtime

<sup>&</sup>lt;sup>1</sup> The figure has been partially taken from Zaharia et al. [64]

Component	Description
Driver	Maintains all relevant information – including user defined code – to process input data and produce output.
Distributed File System	Provides shared file system accessible by all nodes in cluster.
Worker Process	Workforce of the cluster. Gets necessary information from driver and executes user defined code.
User Defined Code	Caries the main application logic.

Table 4.1: Summary of Spark Runtime Components

#### 4.2.2 Spark Cluster Manager

Section 4.2.1 described the coarse grained architecture of Spark. However, from a more fine grained point of view, there is missing component known as *Cluster Manager*. Cluster Manager controls the *assignment* of executors to cluster nodes. It monitors liveness of executors during lifetime of the Spark application. *Spark Session* is the entry point for all Spark applications and has the responsibility to communicate with Cluster Manager to distribute tasks and collect task progress reports from executors. It also distributes user defined code on Worker nodes. Figure 4.3<sup>1</sup> illustrates the role of Cluster Manager in Spark.

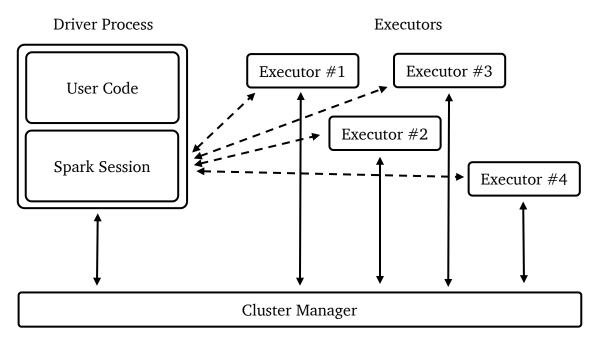


Figure 4.3: Architecture of Spark Cluster Manager

Note that, the term *executor* is a conceptual term and implementation details may differ among different resource managers. Spark supports multiple Cluster Manager implementations. The following describes the most prominent implementations. It shall be noted that this list is a growing over time due to Spark's pluggable resource manager architecture. Table 4.2 summarizes available Cluster Manager architectures.

Apache YARN It is referred as *Yet Another Resource Negotiator* [60] and is the default resource manager for modern Hadoop workloads. It has a master-slave architecture and consists of a single global *Resource Manager* – most probably accompanied by a backup as well – and several *Node Managers* that run on each worker node. The concept of executor is implemented as *containers* in YARN. Each container can hold a specific amount of node's processing power (CPU, RAM,

<sup>&</sup>lt;sup>1</sup> The figure has been partially taken from Chambers and Zaharia [15]

Cluster Manager	Description
Apache YARN	Hierarchical failure detection. Global Resource Manager monitors local Node Managers and Application Masters. Backup Resource Managers monitor global Resource Manager. Node Managers monitor local containers.
Apache Mesos	Hierarchical failure detection. Mesos Master monitors Mesos Slaves and Framework Schedulers. Backup masters monitor Mesos Master. Mesos Slaves monitor local workers.
Spark Standalone	Spark Master monitors Driver program and Spark Slaves. Backup master monitor Spark Master. Spark Slaves monitor local executors.

Table 4.2: Summary of Spark Cluster Managers

Network, etc.). In YARN, any application is modeled with two components. First, *Application Master* that maintains the necessary information to run the application – Spark driver process in this case and second, several *Workers* that run the user defined code. Both Application Master and Workers are executed in the context of YARN containers.

Fault-tolerance is provided at multiple levels. First, several backup nodes monitor the master resource manager via Zookeeper. In case master resource manager fails, one of the backup masters takes over the responsibility. Second, Applications Master reports its status to the global resource manager. In case, application master fails, the global resource manager launches a new Application Master on another node. Third, Workers provide different progress reports to Application Master and Node Managers. In case any Worker container fails, a new container will be allocated and the old one will be killed by the local Node Manager.

Apache Mesos It is another dominant cluster scheduler for Spark [35]. It is a master-slave resource manager. A global resource manager – known as *Mesos Master* – has cluster level view. On each node runs a single *Mesos Slave* process. Mesos has a pluggable architecture for different class of application schedulers. That is, a single cluster can run a mixture of Spark, MPI, etc. jobs with different prioritizes for each application type. Each *Framework Scheduler* handles corresponding jobs. For example, Spark scheduler, maintains multiple driver processes or MPI scheduler maintains multiple MPI applications. Free resources on each Mesos Slave are represented as empty *slots* – very much like containers in YARN – and are allocated to one of the currently running jobs.

Fault-tolerance is provided by a similar hierarchical approach like YARN. Mesos Master is monitored by several backups through Zookeeper. Mesos Slaves as well as Framework Schedulers are in turn monitored by Mesos Master. Each job is further monitored by corresponding Framework Scheduler. And finally, executors are monitored by Mesos Slaves.

Spark Standalone It is the default resource manager for Spark and will be used throughout this thesis. It also follows the master-slave model. *Spark Master* is the global cluster level resource manager. On each cluster node runs a *Spark Slave* process. Spark Slaves are responsible to run and monitor worker executors. It is possible to set default number of CPU cores and available memory for each executor though configuration, either *statically* – default value for all jobs – or at specific job *submission* time – per job basis.

Fault-tolerance is achieved by several counter measures. Figure 4.4<sup>1</sup> depicts all the measures. Similar to YARN and Mesos, a multi-level failure detection approach is exploited. Multiple backup nodes monitor the Spark Master via Zookeeper (Case 1). Spark Slaves and Drivers are monitored by Spark Master (Case 2). Progress of each executor – whether the assigned task is done or not – is monitored by Driver (Case 3). Liveness of each executor is further monitored by corresponding local Spark Slave process (Case 5). Initial input and final output of the jobs are stored in a distributed replicated file system like HDFS. In case any executor fails to store its final output in DFS, it will relaunched by Spark on another node (Case 4). Intermediate results are stored locally without any fault-tolerance in mind. However, *checkpoints* – stored on DFS – can be used in this case to recover from executor failures (Case 4). Refer to section 4.3 for more information on checkpointing process.

<sup>&</sup>lt;sup>1</sup> The figure has been partially taken from Chambers and Zaharia [15]

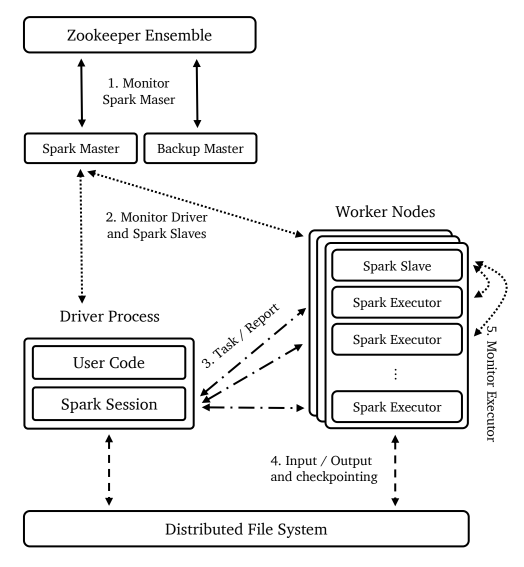


Figure 4.4: Architecture of Spark Fault-Tolerance

#### 4.3 Resilient Distributed Datasets

Resilient Distributed Dataset (RDD) [64] is the basic building block of Spark's data processing pipeline. It is inspired by the same concepts behind *shared memory* and enables *iterative algorithms* and *interactive analytics* to perform common operations directly on memory. RDDs are fault-tolerant and parallel data structures to let intermediate results of a Spark application to be stored in memory during multiple iterations. Existing distributed shared memory abstractions for cluster computing provide *fine-grained* interface – typically key/value – to modify state. Unfortunately, with this approach the only way to provide fault-tolerance is the *log shipping* approach that is already offered by distributed database systems.

Whereas, RDDs offer *coarse-grained* interface based on *immutable transformations*. Each transformation (map, filter, foreach, groupByKey, etc.) applies same application logic to all records that it contains. A dataset's *lineage graph* is the *chain – sequence* – of multiple transformations that produced the aforementioned dataset. With this approach only the transformation itself needs to be logged and shipped to other nodes to provide fault-tolerance. Since RDDs are already immutable, recomputing the lineage on other node is a trivial task. The RDD abstraction is useful in data intensive processing applications, since same operation is applied to many records. Table 4.3<sup>1</sup> summarizes the differences between RRDs and common distributed shared memory technologies.

<sup>&</sup>lt;sup>1</sup> The table has been taken from Zaharia et al. [64]

Dimension	RDD	Distributed Shared Memory
Reads	Per record or per input file	Per key/value
Writes	Coarse grained	Per key/value
Consistency	Easy (Immutable RDDs)	Application dependent
Fault-Tolerance	Lineage recomputation	Application dependent
Straggler Mitigation	Parallel backup tasks	Application dependent
Operator Placement	Based on data locality	Application dependent
Behavior if not enough memory is available	Controlled serialization to disk	OS managed swapping

Table 4.3: High-Level Comparison of RDDs and Distributed Shared Memory Systems

An RDD is a *read-only*, *partitioned* collection of records [64]. RDDs can only be created through deterministic transformations from either *stable stoage* or *other RRDs*.

- **Stable storage**. In this case dataset is typically stored on a shared file system like HDFS and accessible from all nodes of the cluster.
- Other RDDs. In this case subsequent RDDs depend on each other forming the lineage graph.

An RDD has enough information about how it was derived from other RDDs. This means it doesn't have to be materialized in every step of the pipeline. In other words, an RDD cannot reference another RDD that it cannot reconstruct after a failure. Additionally, users are able to control *persistency* and *partitioning* of RDDs.

- **Persistency**. Users can define which RDDs will be reused in next steps of the pipeline. They choose a storage strategy in-memory or disk-based to persist RDDs.
- **Partitioning**. Some transformations like groupByKey or join partition the original RDD into multiple secondary partitions. RDDs allow application developers to perform partitioning hash, range or any custom partitioning strategy such that relevant records are *co-partitioned* on same machine. This is particularly useful for *operator placement* optimizations.

In order to simplify the concept of lineage graph, listing 4.1<sup>1</sup> and figure 4.5<sup>1</sup> illustrate a simple log processing pipeline and its corresponding generated lineage graph.

Listing 4.1: Parsing Errors in Log Files From HDFS

<sup>&</sup>lt;sup>1</sup> The figure and sample code has been partially taken from [64]

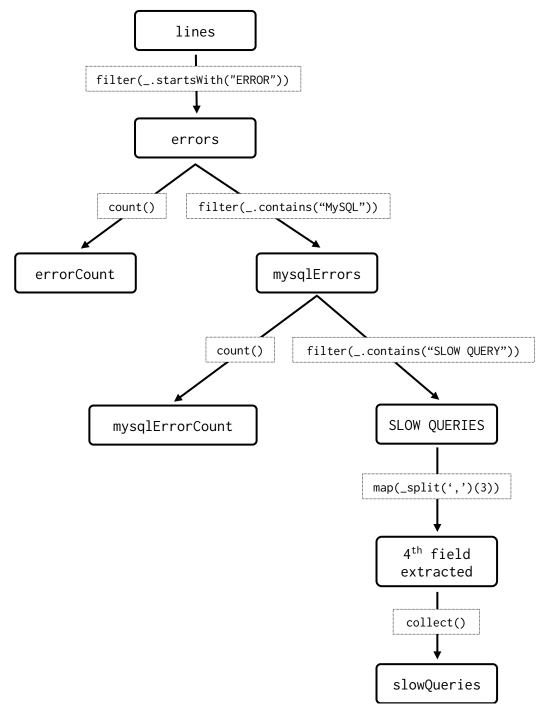


Figure 4.5: RDD Transformations and Corresponding Lineage Graph

#### 4.3.1 RDD Transformations and Dependencies

As mentioned, RDDs are the result of *lazy* transformations derived from one another. In other words, RDDs are not materialized unless it is really needed. However, not all transformations are lazy. There are three major types of transformations.

- Transformations without shuffling. These are lazy transformations that doesn't cause any sort of shuffling among different nodes. That is, transformed records can still be further processed on the same node. For example map and filter are classified in this group.
- **Transformations with shuffling**. These are also lazy transformations but they will trigger shuffling process between nodes. The behavior of the shuffling processes is influenced by the implementation of *partitioners*. As an example, groupByKey, reduceByKey and join belong to this group.
- Actions. These transformations trigger RDD materialization process. That is, the actual computation of RDDs
  don't realize until this type of transformation is met in lineage graph. For example, count, collect and save are
  classified in this group.

In a lineage graph RDDs are derived from one another. This makes each RDD depend on one or several RDDs depending on type of the transformation. There are two types of dependencies among RDDs.

- **Narrow**. When one partition of the parent RDD is used by at most one partition of the child RDD, then the dependency is considered as narrow. For example map, filter, union and joins with *co-partitioned* inputs create narrow dependencies. Narrow dependencies allow to process RDDs without triggering shuffling process a phenomenon known as *pipelined execution*. Additionally, failure recovery is easier in case of narrow dependencies because only lost parent RDDs need to be recomputed. Figure 4.6<sup>1</sup> depicts narrow dependencies.
- **Wide.** When one partition of the parent RDD is used by more than one partition of the child RDD, then the dependency is considered as wide. For example, groupByKey and joins with inputs *not co-partitioned* create wide dependencies. Wide dependencies trigger the shuffling process. In contrast to narrow dependencies, recovering from node failures are less efficient since a failed node might cause loss of some partitions from all ancestors which requires full recomputation [64]. Figure 4.7<sup>1</sup> depicts wide dependencies.

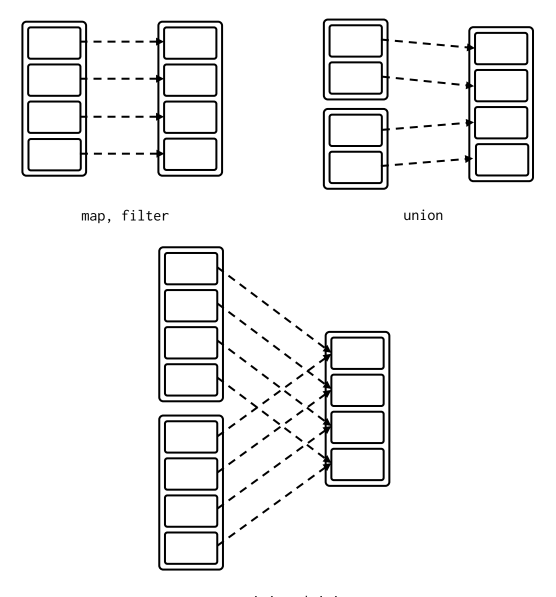
#### 4.3.2 Fault-Tolerance with Checkpointing

The idea of the lineage graph and recomputable transformations simplifies fault-recovery significantly. However, in long chains recomputing failed RDDs can be a very time/process intensive operation. Hence, in some cases it is useful to checkpoint some RDDs to stable storage – like shared file system. In general checkpointing is useful for applications with deep lineage graphs containing wide dependencies, since a node failure in parent RDDs leads to full recomputation in child RDDs. If the lineage graph is small or most dependencies are narrow, checkpointing tends to be less useful.

In order to provide full control to application developers, Spark exposes checkpointing behavior though its API – via REPLICATE flag in persist() function – and leaves the decision up to developers. There are some scenarios where automatic checkpointing without intervention of developers may seem a feasible choice. Because job scheduler is completely aware of the size all RDDs and naturally it should be able to implement more effective checkpointing strategy compared to developers. However, this feature is not currently available in Spark.

With checkpointing in-place, providing fault-tolerance becomes a fairly straightforward process. Note that, RDDs are immutable set of records. This feature makes it easy to flush them in a background process, since there is no *concurrency* and *consistency* concerns involved. Noteworthy, there is an additional fault-tolerance mechanism known as *WAL-based fault-tolerance* for streaming pipelines. This topic is further discussed in section 4.4.

<sup>&</sup>lt;sup>1</sup> The figure has been partially taken from Zaharia et al. [64]



co-partitioned join

Figure 4.6: RDD Narrow Dependencies

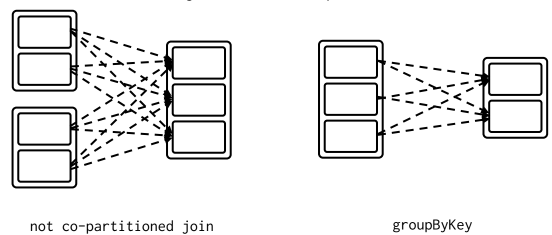


Figure 4.7: RDD Wide Dependencies

## 4.3.3 RDDs and Spark Job Stages

Spark's Job scheduler is similar to Dryad [38]. However, it takes RDD locations into account as well, since some partitions of RDDs are cached into memory or persisted to stable storage. When a job is submitted, Spark's job scheduler does the following.

- Calculating Lineage Graph. In this step, Spark calculates the lineage graph of RDDs with corresponding transformations and dependencies.
- **Defining Stages**. In this step Spark defines the stages required to run the job. From developer's point of view a job is a sequence of RDDs. However, at runtime a job is modeled as *Direct Acyclic Graph* (DAG) of stages. A *stage* is basically the longest possible chain of RDD transformations with shuffling operations in the boundaries. Additionally, different partitions of RDDs in their corresponding stages will also be computed.
- Launching Tasks. In this step, Spark launches *tasks* which is going to be run by executors. Each task process one or multiple partitions of some RDDs. Depending on the location of RDDs, tasks will be launched on machines with data-locality awareness to reduce network transmission. In case a task fails, it can be run as long as parent stages are still available. If one or more parent stages become unavailable, missing partitions from parent stages shall be recomputed according to lineage graph.

Figure 4.8 shows a sample job with logical operators and Figure 4.9 depicts its corresponding DAG with two stages at runtime.

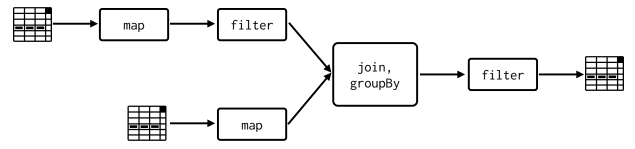


Figure 4.8: Spark Job with Logical Operators

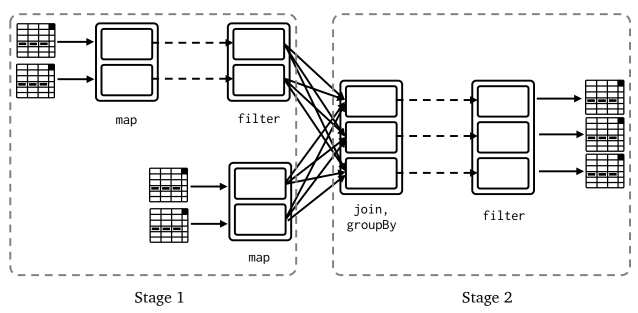


Figure 4.9: Spark Job Stages at Runtime

#### 4.4 Discretized Streams

Many modern big data applications process data in real-time as they enter the processing pipeline. Online machine learning, fraud detection, spam detection, etc. are just few examples of applications that require near real time processing of data. In order to process incoming records in near real-time, new generation of data processing frameworks has been emerged which is known as *stream processing systems*. Apache Storm [56] is an example of stream processing systems. Although these systems provide stream processing pipelines, but they also come with a couple of deficiencies. First, section 4.4.1 explores the architecture of traditional stream processing systems and its problems. Then, section 4.4.2 describes the new stream processing model known as *Discretized Streams* (D-Streams) [63] that is built on top of RDDs in order to resolve the problems of traditional model.

## 4.4.1 Continuous Operator Processing Model

Though the concept of stream processing is nothing new and there has been many proposals and implementations like [26] [56] [2], but most of these systems are based on a processing model known as *Continuous Operator Processing Model*. In this model streaming computations are divided into a set of *long-lived stateful* operators that process messages in a loop:

- Get one or more messages from previous operators.
- Apply the computation business logic on newly received messages. Query the internal state if required.
- · Update internal state if necessary.
- Produce any number of messages as the result of computation. These messages will be sent to next operators down the pipeline.

While continuous processing model minimizes latency, the stateful architecture of operators and nondeterminism that comes from record interleaving on the network, makes it hard to provide fault tolerance efficiently. In particular, recovering from a *failed* or slow node – *stragglers* – is challenging. The are usually two standard approaches to overcome these issues.

Replication In replication model which is borrowed from database systems, there are two copies processing graphs. Produced messages are sent as duplicates to down stream operators. However, just replicating messages is not sufficient. A *consensus protocol* should exist in-place to ensure that both operator replicas see incoming messages in the same order that is sent by upstream operators. Even though replication is a costly operation but it recovers from failures very fast, since both replicas are processing message online in synchronized steps.

Upstream Backup The basic idea behind this model [36] is to checkpoint the internal state every once in a while to a stable shared storage. In case an operator – or node – fails, a backup operator takes over and reloads the last successfully written checkpoint. Then backup operator rebuilds the state by replying new messages and reproducing lost messages as necessary. Although this model is more efficient in terms of replication costs, but suffers from high fail-over time. Backup node should replay newly published messages from the last checkpoint and apply them to the internal state. Depending on checkpointing interval, this might be a time consuming operation.

Besides the fault-tolerance costs of this model, dealing with straggler nodes are even more challenging. Replication model provides no solution at all, since two replicas are processing messages synchronously. The only way to resolve this issue in Upstream Backup model is to fail – kill – the slow operator and let the backup operator takes over the responsibility which will undergo slow recovery process. Figure 4.10 depicts these two approaches with additional synchronization and replication messages involved.

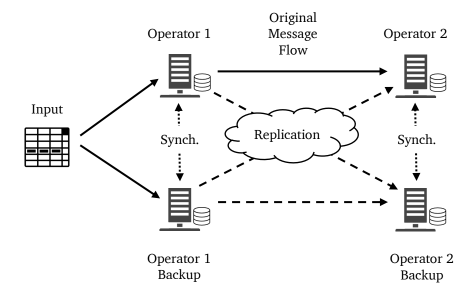


Figure 4.10: Continuous Operator Processing Model

## 4.4.2 D-Stream Processing Model

*Discretized Streams* (D-Streams) [63] is a novel technique to overcome the issues the Continuous Operator Processing Model. The basic idea behind D-Streams is to separate the computation from its state and keep the state as RDDs. This separation provides the following benefits.

- Computation of messages becomes a set of stateless and deterministic tasks operating on RDDs.
- RDDs already provide resiliency through in-memory replication and lineage recomputation. There is no need for further mechanisms to provide fault-tolerance.
- Unified processing model for batch and stream processing.

As a consequence of storing operator intermediate state as RDDs, a streaming computation can be modeled as series of *deterministic batch computations* on small intervals – known as micro-batch. Messages received in each batch interval is stored reliably across the cluster to form an input dataset for that interval. Once the time interval completes, this dataset is processed just like traditional batch processing using map, filter, groupByKey, etc. Formally, A D-Stream is a sequence of *immutable*, *partitioned* datasets (RDDs) that can be acted on by deterministic transformations [63]. These transformations produce new D-Streams, and may create intermediate state represented as RDDs. Figure 4.11 shows high-level D-Stream computation model.

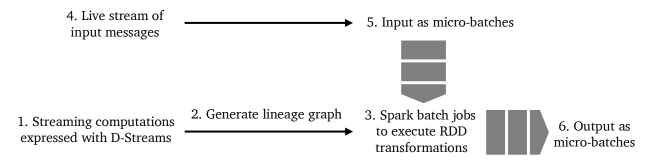


Figure 4.11: D-Stream High Level Processing Model

In order to clarify D-Stream API, listing 4.2¹ shows a simple streaming word count example. It creates a pageViews D-Stream by reading even stream over TCP, and groups them into 1-second batch intervals. Then, it transforms the event stream to a new D-Stream of (URL,1) pairs called ones. Finally, it performs a running count with a *stateful* runningReduce operator. Figure 4.12¹ depicts the corresponding lineage graph and how the streams are divided are into batch intervals. Note that smaller rectangles illustrates RDD partitions.

```
val pageViews = readStream("tcp://...", "1s")
val ones = pageViews.map(event => (event.url, 1))
val counts = ones.runningReduce((a, b) => a + b)
```

Listing 4.2: Streaming Word Count using D-Stream API

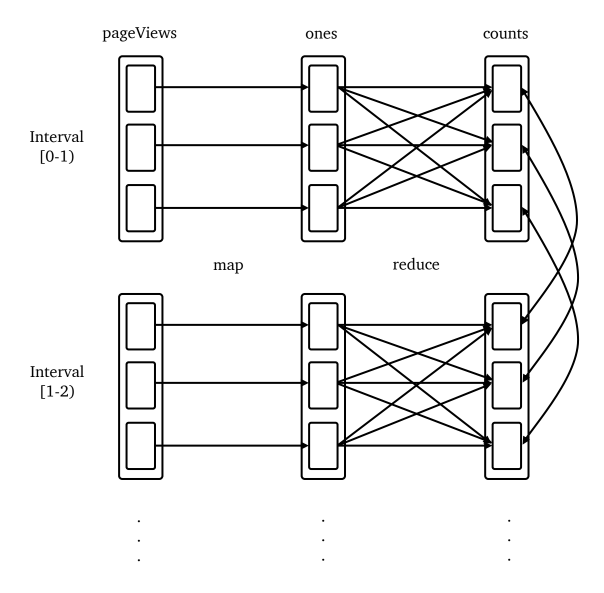


Figure 4.12: Streaming Word Count Using D-Stream API

<sup>&</sup>lt;sup>1</sup> The code and figure has been partially taken from Zaharia et al. [63]

## 4.4.3 Fault-Tolerant Message Consumption

Since RDDs provide the basic building block of Spark Streaming, achieving fault-tolerance in streaming applications is a straightforward process. In case of any node failure, failed RDD partitions can be reproduced easily by evaluating lineage graph and determining which transformations need to be recomputed. Furthermore, for stateful transformations, Spark already provides checkpointing. However, there is one case that needs more attention.

Naturally, in streaming applications, live stream of records are coming from some data sources. For example, in log processing application it may originate from web servers or in live monitoring applications it comes from sensors. If one of the RDD transformations crashes during message consumption, two questions arise:

- Are the messages that were consumed before crashing, be redelivered after relaunching failed transformations?
- In case lost messages are redelivered, will they be delivered with the same order as consumed before crashing?

The answer to these two questions leads us to three types of data sources.

Reliable Ordered With this type of data sources, the data source provides ordering and reliable message consumption capabilities. Messages will be kept in data source until its reliably confirmed – ACKed – by consumer. In case of consumer crash, messages will be delivered with the same order as before. Many of publish-subscribe message brokers such as Apache Kafka [5] provide this feature.

Reliable Unordered With this type of data sources, the data source provides reliable message store but redelivery is not guaranteed with the same order as before consumer crash. For cases where ordered message consumption is crucial for business logic of the application, Spark Streaming provides *WAL-based* consumers. With WAL-based consumers, each consumer appends the received message to a log file before starting to consume the message. These log files are stored in a distributed file system like HDFS. In case of consumer failure, another consumer re-opens the the log file and re-process the messages with same order as before.

Unreliable This type of data source does not provide any form of reliability, ordering and fault-tolerance. Hence, WAL-based message consumption shall be enabled by application developer.

## 4.5 Conclusion

In this chapter different aspects and features of Spark has been explored. Built on top of unique features of Resilient Distributed Datasets (RDDs), Spark offers a unified approach for batch and stream processing. As explained during the chapter, deficiencies and problems of traditional systems have been addressed to some extent by Spark.

#### 5 Design and Implementation Detail

## 5.1 Introduction

In this chapter design and project structure of implemented techniques will be introduced and discussed. Two techniques have been selected to be implemented. This chapter is organized as follows. Section 5.2 explains selected techniques and the reasoning behind it. In this section, selected techniques are explained at theoretical level. Section 5.3 describes the structure of the implementation for Spark Streaming in detail. Section 5.4 describes the configuration parameters required to launch the implementation. Finally, Section 5.5 concludes this chapter.

## **5.2 Choosing Auto-Scaling Techniques**

As discussed in Section 3.5.4, Auto-Scalers can be categorized into 5 groups.

Threshold-Based In this approach, one or multiple threshold values are defined [28] which specifies the behavior of the Auto-Scaler when the system load goes beyond these thresholds. However, manually defining thresholds is a tricky process [20]. Spark's default dynamic resource manager [7] and Online Parameter Optimization [30] are partially based on this approach and they will be evaluated in Section 6.

Time-Series Analysis In this approach, a history window of workload – depending on prediction accuracy – is considered to predict future workload. However, in case the workload is changing in an unpredictable way – which is not an unusual phenomena in streaming applications – then this approach becomes less useful. As confirmed by DEBS grand challenges [39] [40] [27], the general assumption for streaming applications is that the workload is unpredictable. Due to this limited applicability, no time-series technique is implemented.

Queuing Theory In this approach, the system is modeled as a queue network. Since queuing parameters – message interarrival and service rates – are static, they have to be re-calculated in a timely manner. One of the novel implementations is done by Lohrmann, Janacik, and Kao [45]. However, authors made two unconvincing assumptions. First, worker nodes shall be homogeneous in terms of processing power and network bandwidth. Second, there should be an effective partitioning strategy in place in order to load balance outgoing messages between stages. In reality both assumptions rarely occur. Large scale stream processing clusters are built incrementally. Depending on workload, data skew does exist and imperfect hash functions are widely used by software developers. Furthermore, as confirmed by Rajarshi, Tesauro, and Walsh [52], queuing models are very complicated to build and less adaptive to changing environments. As a consequence, no queuing theory technique is implemented.

Reinforcement Learning Reinforcement Learning has shown its capabilities to adapt ever changing environments. However, some of the proposed solutions are conflicting with the requirement of this thesis as discussed in Section 2. The following summarizes the reasoning why each proposal is accepted or rejected by this thesis.

- Herbst et al. [31] proposed a solution based *Bayesian Networks*. There are two major problems with this proposal. First, it needs sampling and offline training which is inapplicable for changing streaming workloads. Second, in case the model is complex, the training phase is long and computationally infeasible for streaming workloads. As mentioned, both issues are conflicting with thesis requirements.
- Tesauro et al. [59] proposes a hybrid approach to resolve performance issues of online training which consists of two components. First an online component based on queuing theory. Second, Reinforcement Learning component that is trained offline. This proposal models each node as a queue. However, the only way to apply this technique is to modify spark-core package. Thus, this solution is not implemented.

- Rao et al. [53] proposed a solution to manage virtual machine resources. However, it is also based on offline
  training and sampling which is obtained from a separate supervised training phase. Thus, it's not applicable for
  dynamic streaming workloads.
- Enda, Enda, and Jim [22] proposed a parallel architecture to Reinforcement Learning without any global controller involved. Nodes (RL agents) have two tables. Local table is trained by each node separately. Global table contains values learned by other nodes. When an agent learns anything new, it broadcasts it to other nodes. From theoretical point of view, this solution might seem feasible, since parallel learning speeds up initialization process. However, Spark has a single global controller. Applying this technique to Spark requires heavy modification to spark-core package.
- Heinze et al. [29] implemented Reinforcement Learning in the context of FUGU [26]. Each node, maintains its own Q-Table and imposes local policy without coordinating with other nodes. Although this architecture is not applicable for Spark, but its core idea *Temporal Difference* [57] algorithm— is applicable. Thus, this thesis implemented this proposal by adopting it to Spark architecture. Refer to Section 5.2.1 for theoretical background and Section 5.3 for implementation detail.
- Cardellini et al. [12] proposed a two level hierarchical architecture for resource management in Apache Storm [56]. Local controller applies local policy on each node and coordinates with the global controller for confirmation of its actions. Although, this architecture seems to be a promising approach, however it has been implemented by modifying Storm's internal components. As mentioned above, this is in conflict with thesis's requirements.
- Dutreilh et al. [21] proposed a model-based Reinforcement Learning approach for resource management which is based on a global controller. In order to overcome the slow convergence of model-free learning, authors proposed to estimate environment dynamics based on collected samples at runtime. Then it switches to *Value Iteration* [57] algorithm instead of *Temporal Difference*. This approach has also been partially adopted by this thesis. Refer to Section 5.2.2 for theoretical background and Section 5.3 for implementation detail.

Control Theory Techniques based on Control Theory are also promising for elastic data streaming. Because it monitors input and output of the application, it can respond to workload changes very fast and adapt the system if necessary. Thus it is perfectly capable of handling dynamic environments. The comparison between Reinforcement Learning and Control Theory techniques is left for future work.

As mentioned, two techniques – *Temporal Difference* and *Value Iteration* – will be implemented in this thesis. In next two sections, theoretical foundation of these algorithms will be laid out. Noteworthy to mention, these sections are heavily inspired by Sutton and Barto [57]. Furthermore, Section 7 discusses more techniques and the reasoning why they have been rejected by this thesis.

## 5.2.1 Temporal Difference

Section 3.5.4.3 explains the basics of Reinforcement Learning. Temporal Difference (TD) learning is one of the foundational algorithms of Reinforcement Learning. It can learn from applying experience without having prior knowledge about environment's dynamics. This property is potentially useful for data stream processing systems in which the incoming workload is changing without any particular pattern.

Before starting to dig into details, some formal notions shall to be explained. An *episode* is series of experiences that is taken by the agent. At each time step t, the agent moves from one *state*  $S_t$  to another state  $S_{t+1}$ . A *policy*  $\pi$  defines the action  $A_t$  that should be taken by the agent at each state. Followed by each action, the agent receives a *reward*  $R_{t+1}$  from the environment. Note that, the agent will receive reward for its corresponding action at next state  $S_{t+1}$ . Thus, an episode can be viewed as Equation 5.1.

... 
$$S_t, A_t, R_{t+1}, S_{t+1}, A_{t+1}, R_{t+2}, S_{t+2}, A_{t+2}, R_{t+3}$$
 ... (5.1)

An episode does not necessarily have a *terminal* state. In some environments – like data stream processing – an episode is a never ending sequence of experiences. Prior to taking an action, the agent has an estimate of *expected reward* V which specifies future reward if it follows the same actions returned by policy  $\pi$  from that specific state. Thus, it is referred as  $V_{\pi}$  which translates to expected future reward under policy  $\pi$ . During the episode, the agent tries to *maximize* the reward that it received from the environment. Any policy that leads to maximum possible reward is called the *optimal* policy  $\pi^*$  and its corresponding estimate of future reward is referred to as  $V_{\pi}^*$ .

Unlike Monte Carlo [57] methods that requires agent to wait until the end of episode, TD approaches only need to wait until the end of next time step to get a feedback from environment. Then, the agent updates corresponding estimate of previous state. Thus, the most simple TD approach can be formulated as Equation 5.2. Equation 5.2 contains two important parameters of TD.

$$V(S_t) \longleftarrow (1 - \alpha)V(S_t) + \alpha \left[ R_{t+1} + \gamma V(S_{t+1}) \right]$$
(5.2)

- $\alpha$  denotes *Learning Factor*. It specifies, how much the agent shall learn from new experiences. A higher  $\alpha$  means learning with a faster pace. It also leads to forgetting history faster. On the other hand, a lower  $\alpha$  leads to slower learning process which means the agent trusts its experience history more it gives more weight to history rather than new experiences.  $\alpha$  is defined as a number between (0,1].
- $\gamma$  denotes *Reward Factor* or *Discount Factor*. It specifies, whether the agent shall optimize for *future* or *immediate* reward. A higher  $\gamma$  leads to optimizing for future reward, whereas a lower  $\gamma$  leads to optimizing for immediate reward. It is defined as a number between (0,1). As defined by Equation 5.3, reward is cumulating at each time step which causes the sum to grow indefinitely. In order to prevent infinite reward problem, it is crucial to define  $\gamma$  as a number less than one to force convergence.

$$R_t = r_{t+1} + \gamma r_{t+2} + \gamma^2 r_{t+3} + \dots = \sum_{0}^{\infty} \gamma^k r_{t+k+1}$$
 (5.3)

Since expected future reward is defined per action, Equation 5.2 changes slightly and becomes as Equation 5.4. This update is applied after every transition from a nonterminal state  $S_t$ . In case  $S_{t+1}$  is terminal, then  $Q(S_{t+1}, A_{t+1})$  is set to zero. As a consequence, each experiment can be defined with a quadruple of (State, Action, Reward, State') or

## Algorithm 3: Q-Learning Work-Flow

- 1 Algorithm parameters:  $\alpha \in (0,1], \gamma \in (0,1)$
- 2 Initialize Q(s, a) for all  $s \in \{S terminal\}$ ,  $a \in A(s)$  arbitrarily and Q(terminal, a) = 0

## з repeat

- 4 | Take action *A* based on policy  $\pi$  derived from *Q* table
- 5 Observe reward R and land in state S'
- 6 Choose A' from S' using the same policy  $\pi$
- $7 \quad Q(S,A) \leftarrow (1-\alpha) Q(S,A) + \alpha \left[ R + \gamma Q(S',A') \right]$
- $S \vdash S \leftarrow S'$
- $g \mid A \leftarrow A'$
- 10 **until** End of episode or S is terminal

 $(S_t, A_t, R_{t+1}, S_{t+1})$ . This method is known as *Sarsa* algorithm and named *Q-Learning* because of the Q-Table used in the equation. Algorithm  $3^1$  describes this procedure.

$$Q(S_t, A_t) \longleftarrow (1 - \alpha)Q(S_t, A_t) + \alpha \left[R_{t+1} + \gamma Q(S_{t+1}, A_{t+1})\right]$$

$$(5.4)$$

An agent takes action based on a policy function  $\pi$  which in turn is derived – partially or completely – from the Q-Table. However, it shall be noted that in the beginning of episode the Q-Table is initialized to zero. In such situations, it takes very long time for agent to discover new states which is particularly harmful for data stream processing systems. This problem is usually solved by introducing small degree of randomness. The policy function decides randomly with a small probability  $\epsilon$  and in any other case  $(1-\epsilon)$ , it behaves based on Q-Table. Other solutions do exist to solve state discovery problem. For example, Q-Table can be initialized with a random value or based on some *heuristic* function that is partially derived from domain knowledge.

Furthermore, if two or more actions have equal expected reward, then policy function has to decide on one of them by breaking the tie. There are different strategies to solve this issue. Choosing a random action is one of them. However, in some domains this might be dangerous or harmful. Rather than choosing randomly, a simple heuristic inspired by domain knowledge can be helpful too.

## 5.2.2 Value Iteration

As mentioned in last section, in streaming workloads the common belief is that the workload is not predictable. In other words, reward values of each action and probability of landing in a specific state after taking an action – environment dynamics – is unknown. However for the sake of discussion, let us assume environment dynamics is *known*. In such situations, Reinforcement Learning can be solved using *Dynamic Programming* techniques. In this section, the problem will be solved for environments with known dynamics. Then, some of the core ideas will be applied to adopt the solution for environments with unknown dynamics.

Finding an optimal policy function, when environments dynamics is known can be *recursively* solved by applying dynamic programming. As mentioned previously, the agent is trying to maximize expected reward by taking optimal actions. Here expected reward is defined with  $\mathbb{E}$  and max selects maximum value based on specified action.

$$V^{*}(s) = \max_{a} \mathbb{E} \left[ R_{t+1} + \gamma V^{*}(S_{t+1}) | S_{t} = s, A_{t} = a \right]$$

$$= \max_{a} \sum_{s',r} p(s',r|s,a) \left[ r + \gamma V^{*}(s') \right]$$
(5.5)

The pseudocode has been taken from Sutton and Barto [57]

$$Q^{*}(s,a) = \mathbb{E}\left[R_{t+1} + \gamma \max_{a'} Q^{*}(S_{t+1}, a') | S_{t} = s, A_{t} = a\right]$$

$$= \sum_{s',r} p(s', r | s, a) \left[r + \gamma \max_{a'} Q^{*}(S_{t+1}, a')\right]$$
(5.6)

For scenarios where p and r are unknown – which is the case for this thesis – these values can estimated by collecting samples. As the agent runs for some period of time, it collects samples and then estimates p and r. Given that p and rare estimated to some degree, it is possible to initialize Q-Table with more useful values - compared to zero or random values – and speed up learning process. Whether this estimation is accurate or not – how many samples shall be collected for a precise estimation – is another question which will be evaluated in Section 6.

The following procedure is applied for collecting samples which is inspired by Dutreilh et al. [21]. As agent runs and observes a quadruple of (s, a, r, s'), the following counters are updated.

$$P[s, a, s'] = P[s, a, s'] + 1$$

$$R[s, a] = R[s, a] + r$$

$$C[s, a] = C[s, a] + 1$$
(5.7)

Given these statistics, estimators  $\overline{P}$ ,  $\overline{R}$  are calculated to replace original p, r in Equation 5.6 respectively.

$$\overline{P}[s, a, s'] = \frac{P[s, a, s']}{C[s, a]}$$

$$\overline{R}[s, a] = \frac{R[s, a]}{C[s, a]}$$
(5.8)

After estimating p, r, Algorithm  $4^1$  can be used to produce optimal policy. Not the difference between  $\max$  and argmax functions. The former returns the maximum value and the later returns the action that has the maximum value. Value Iteration that described in Algorithm 4 can be used independent of Temporal Difference. However, in this thesis it is used to get a useful initialized Q-Table. Thus, when sample collection period is over and Q-Table is initialized, normal Q-Learning approach is used thereafter. The reason is that in Value Iteration, after sampling period, Q-Table is never updated again. However, by combining Value Iteration - for initializing Q-Table - and Q-Learning, Q-Table is also updated at runtime which improves decision accuracy.

## **Algorithm 4:** Value Iteration for Estimating $\pi \approx \pi^*$

```
1 A small threshold \theta > 0 determining accuracy of estimation
 2 Initialize V(s) for all s \in \{S - terminal\} arbitrarily and V(terminal) = 0
 з repeat
         \Delta \leftarrow 0
 4
         foreach s \in S do
 5
              V(s) \leftarrow \max_{a} \sum_{s',r} p(s',r|s,a) \left[ r + \gamma V(s') \right]
 7
              \Delta \leftarrow \max(\Delta, |\nu - V(s)|)
 8
 9
10
         Output policy \pi \approx \pi^*, such that
         \pi(s) = \operatorname{argmax}_{a} \sum_{s', r} p(s', r \mid s, a) \left[ r + \gamma V(s') \right]
11
12 until \Delta < \theta
```

The pseudocode has been taken from Sutton and Barto [57]

## 5.3 Design

In this section technical design and structure of the thesis will be explained and discussed. As mentioned in previous section, two Reinforcement Learning techniques – namely Q-Learning (Temporal Difference) and Value Iteration – have been implemented. Both of these methods require defining *state space*, *policy* and *reward* functions. Typically, Reinforcement Learning agents move from one state to another by taking an action which is proposed by policy function, getting feedback – known as reward – from environment and landing into destination state. In next sections each of these components will be discussed.

#### 5.3.1 State Space

State space defines – models – the environment where the agent is living. Usually, it is combination of environment's properties. The following is list of sample state spaces that could be defined for different applications.

Web Applications For a typical web application, a sample state space could be defined as a tuple of

[Number of web servers, Total workload defined as requests per second, Average round trip delay of requests]

Virtualization Server For a Hypervisor, a sample state space can be defined as a tuple of

[Number of running virtual machines, Total CPU utilization, Total used RAM]

Stream Processing For a data stream processing system, a sample state space can be defined as a tuple of

[Number of worker nodes, Total workload defined as messages per second, Average latency of messages]

As state space contains more precise information, it gets bigger and bigger. Thus, in most systems it is *discretized* to some degree to reduce its size. State space reduction has a couple of benefits.

- Change Observation. For example, in most systems a 1% increase/decrease in CPU utilization is not considered as a noticeable change but 5% is.
- Absorbing Sudden Bursts. In some applications, there might be a sudden change in one aspect of the system
  for a slightly short time. For example 20% increase in CPU utilization for 5 seconds may not be considered as
  sustained workload increase.
- **Preventing Zig-Zag Decisions**. If state space is discretized to fine-grained states, it might lead to consecutive contradictory decisions A phenomena known as *Zig-Zag* decisions. For example, Auto-Scaler may decide to perform a scale-in action in one step and scale-out action in next step. A reasonable way to prevent this never ending loop is to discretize the state space into coarse-grained states.

However, it shall be noted that *over discretizing* the state space may have negative impacts on target system. Section 6 contains an experiment to evaluate discretization impact.

In this thesis, a combination of latency and direction of workload is incorporated in to state space.

Latency Each micro-batch is associated with two latencies. First, *Processing Latency* – time that takes to process a single micro-batch from start to end. second, *Scheduling Latency* – time that each micro-batch stays in a queue *waiting* to be processed. In this project, total latency (processing latency + scheduling latency) is stored as an average value calculated over a window of time.

Workload Direction It is reasonable to consider average number of messages that are being processed in state space. However, in order to reduce state space only its *direction* is stored. That is, instead of storing a pure number as average number of messages processed per window of time, only a boolean value indicating that incoming messages are increased to decreased – compared to last window – is stored.

As mentioned in Section 3 each state is associated with three actions – No-Action, Scale-In and Scale-Out. Each action has corresponding reward – Q-Value– attached to it. Figure 5.1 depicts this relationship.

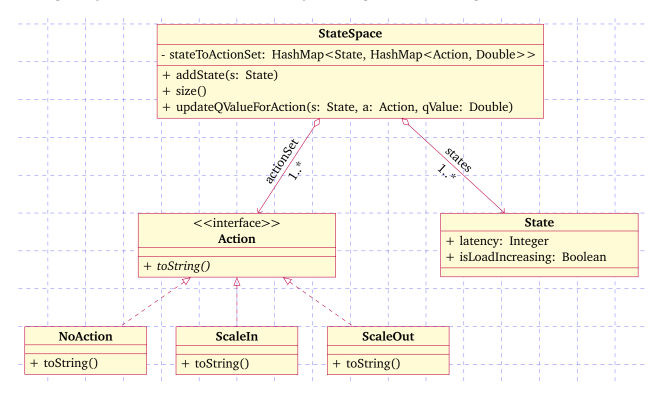


Figure 5.1: State Space Class Diagram

# 5.3.2 State Space Initialization

Typically when the agent starts, its Q-Table is empty – initialized with zero. Over time, as the agent take more actions and discovers its environment, it updates its Q-Table. Since Q-Table is initialized with zero, the agent usually takes some random actions from time to time with a small probability. This helps to discover the environment. In some applications, this behavior is sufficient. However, in fast changing environments like streaming systems this process takes very long time to converge. Thus developers usually try to initialize Q-Table beforehand to speedup the learning process. In this thesis, three different initializers have been implemented. Figure 5.2 shows class diagram of these initializers.

Zero Initializer This is the default implementation which initializes all the Q-Values – reward values – to zero.

Random Initializer This initializer assigns a random value between (-1,1) to Q-Values.

Optimized Initializer This initializer uses a *heuristic* to initialize the Q-Table. Typically in most systems, there is a well defined set of SLOs defined by users that should be respected by Auto-Scaling system. This initializer exploits a normalization function defined in Equation 5.9. If recent latency window is sufficiently large enough, workload-to-latency graph looks similar to Figure 5.3. This graph is divided into four regions. As workload increases, latency is approaching target latency (Region 1). In case the workload keeps increasing, the agent goes beyond the target latency and SLOs are violated (Region 2). Assuming the agent takes some reasonable action, latency starts to decrease (Region 3). Finally, at some point it will drop below the target latency line (Region 4). For each region, Q-Table is initialized with different values.

$$n = \frac{\text{current latency}}{\text{target latency}} \tag{5.9}$$

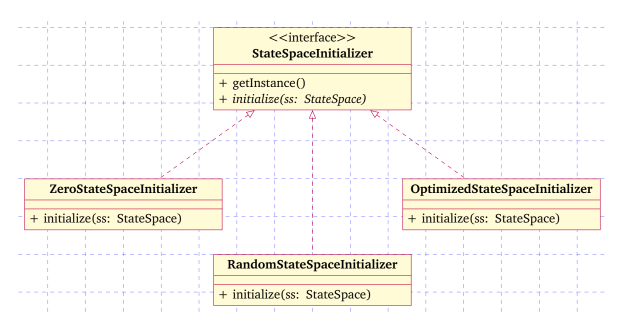


Figure 5.2: State Space Initializer

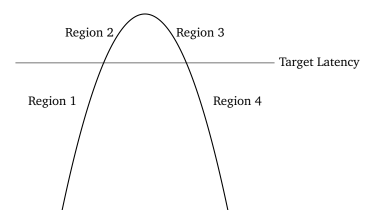


Figure 5.3: Different Regions of a Typical Workload

The following describes the way all four regions are initialized. Table 5.1 summarizes each region and describes preferred action in each region.

Region 1 In this region, latency is increasing as workload is increasing. So it is reasonable to give more reward to Scale-Out as the agent approaches target latency line. This is called *Early Scale-Out* action which helps to add worker tasks in order to prevent violating SLOs. Thus the reward for each action is initialized as Equation 5.10. In this region, Scale-Out and NoAction rewards are positive but less than one.

$$Scale-Out = \frac{current\ latency}{target\ latency} \qquad NoAction = 1 - \frac{current\ latency}{target\ latency} \qquad ScaleIn = 0$$
 (5.10)

Region 2 In this region, the agent has fall behind and latency has been increased up to the point that it is more than target latency. At this point Scale-Out is still the preferred action. It is reasonable to initialize Scale-In with a negative reward to prevent the agent from taking unjustifiable action. Reward for each action is initialized as Equation 5.11. In this region, Scale-Out reward is larger than one but NoAction reward is less than one.

$$Scale-Out = \frac{current \ latency}{target \ latency} \qquad NoAction = \frac{current \ latency}{target \ latency} - 1 \qquad ScaleIn = -1$$
 (5.11)

Region 3 In this region, either the workload is decreasing or the agent has taken a presumably reasonable action that has led latency to decrease. Usually, it is not required to take any further action here. Thus, to let latency stabilize, NoAction is a preferred action in this region. Additionally, Scale-In is still prohibited.

Scale-Out = 0 NoAction = 1 ScaleIn = 
$$-1$$
 (5.12)

Region 4 In this region, the latency has dropped below target latency and is decreasing gradually. Thus, as the agent gets farther away from target latency line, we encourage the agent to take Scale-In action. However, near target latency line, we still prefer to take NoAction, since it is still dangerous to take Scale-In.

Scale-Out = 0 NoAction = 
$$\frac{\text{current latency}}{\text{target latency}}$$
 ScaleIn =  $1 - \frac{\text{current latency}}{\text{target latency}}$  (5.13)

Region	Preferred Behavior
Region 1	Early Scale-Out is preferred to avoid hitting target latency.
Region 2	Target latency is already violated, so Scale-Out is still preferred.
Region 3	Although latency is higher than target – but decreasing, NoAction is preferred to let latency stabilize.
Region 4	Latency is stable again. If latency is sufficiently low, then Scale-In is preferred.

Table 5.1: Summary of Latency Regions

# 5.3.3 Policy Functions

Policy  $\pi$  is a function that guides the agent to take an action in each state. Different strategies can be implemented but at its core, a policy function is either extracted from Q-Table. An optimal policy  $\pi^*$  is a policy that returns actions that lead to maximum expected reward. Before getting into details about implemented policies of this thesis, *monotonicity constraint* [34] shall be explained. Monotonicity constraint mandates:

- If Scale-Out was the chosen action in last state and latency has been increased compared to last state, then it is not possible to take Scale-In in current state. The reason is that, even by applying Scale-Out, latency has been increased. So, for a even higher latency, there is no reason that Scale-In could ever help.
- If Scale-In was the chosen action in last state and latency has been decreased compared to last state, then it is not possible to take Scale-Out in current state. The reason is that, latency is decreasing without taking Scale-Out. So, for a even lower latency, there is no need for Scale-Out. Even though Scale-Out is a safe action in this case, but in Spark Streaming any Scale-In/Out action cause reshuffling of intermediate results. Thus, it is crucial to avoid unnecessary although safe actions.

As confirmed by Heinze et al. [29], monotonicity constraint improves learning process. Algorithm 5 shows the pseudocode of this procedure.

## **Algorithm 5:** Monotonicity Constraint

- 1 feasibleActions ← GetActionsFromStateSpace(currentState)
- 2 if currentState.latency > lastState.latency AND lastAction == ScaleOut then
- 3 | feasibleActions ← feasibleActions − ScaleIn
- 4 end
- 5 **if** currentState.latency < lastState.latency AND lastAction == ScaleIn **then**
- 6 | feasibleActions ← feasibleActions − ScaleOut
- 7 end

Three different policies have been implemented in this thesis. Figure 5.4 depicts class diagram of these polices. Table 5.2 summarizes the behavior of each policy.

Greedy Policy This policy takes action in two phases. First, it applies monotonicity property. After that, it selects the best possible action. Best action here means the action with largest/lowest Q-Value, depending on how reward function has been implemented.

One Minus Epsilon Policy This policy wrap any other policy and adds some degree of randomness to it. That is, with a probability of  $\epsilon$  it returns a random action and with a probability of  $1-\epsilon$  it hands over the process to underlying policy. Underlying policy is not necessarily an optimal – greedy – policy.

Decreasing One Minus Epsilon Policy This policy is similar to previous policy. The different is that randomness ( $\epsilon$ ) is decreasing over time until it reaches zero after which it unconditionally hands over to underlying policy.

Policy	Behavior
Greedy	Always stick to action with highest/lowest Q-Value.
One Minus Epsilon	Take random action with a probability of $\epsilon$ , otherwise stick to underlying policy.
Decreasing One Minus Epsilon	Same as $1-\epsilon$ policy, but $\epsilon$ is decreasing over time.

Table 5.2: Summary of Policy Functions

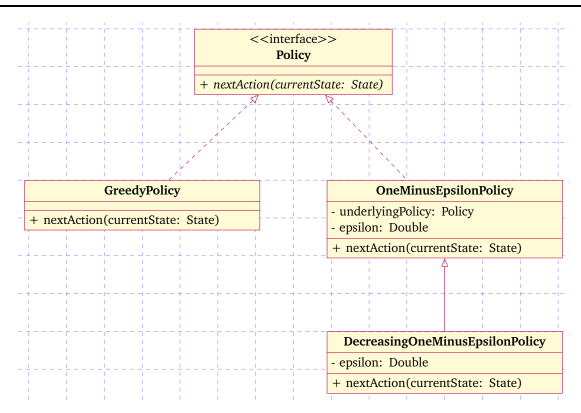


Figure 5.4: Policy Functions

# 5.3.4 Reward Function

5.4 Configuration			

6 E	/aluation
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#### 7 Related Work

#### 7.1 Introduction

Dynamic resource allocation in cloud environments has been studied extensively in literature. In this chapter prior work will be discussed and explored. It is organized as follows. Section 7.2 delves into threshold-based techniques. Section 7.3 investigates techniques based on time-series analysis. Section 7.4 analyses techniques based on queuing theory. Section 7.5 explores Reinforcement Learning techniques comprehensively. Finally, section 7.6 concludes.

## 7.2 Threshold-Based Techniques

Hasan et al. [28] proposed four thresholds and two time periods. ThrUpper defines upper bound. ThrBelowUpper is slightly below ThrUpper. Similarly, ThrLower defines lower bound and ThrAboveLower is slightly above the lower bound. In case, system utilization stays between ThrUpper and ThrBelowUpper for a specific duration, then cluster controller decides to take a scale-out action, by adding resources. On the other hand, if system utilization stays between ThrLower and ThrAboveLower for a specified duration, then the central controller decides to take scale-in action. Furthermore, in order to prevent making oscillating decisions, grace period is enforced. During this period, no scaling decision is made. Defining two levels of thresholds helps to detect workload persistency and avoids making immature scaling decision. However, defining thresholds is a tricky and manual process, and needs to be carefully done [20]. It shall be noted that, computation overhead of this approach is very low.

*RightScale* [55] applies voting algorithm among nodes to make scaling decisions. In order for a specific action to be decided, majority of nodes should vote in favor of that specific action. Otherwise, no-action is elected as a default action. Afterwards, nodes apply grace period to stabilize the cluster. The complexity of the voting process in trusted environments is in the order of  $O(n^2)$ , which leads to heavy network traffic among participants when cluster size grows. This approach also suffers from the same issue – accurately adjusting threshold values – as other threshold-based approaches.

Heinze et al. [30] proposed a novel threshold-based solution in the context of FUGU [26] – a data stream processing framework. This techniques uses an adaptive window [9] to monitor the recent changes in workload pattern. In case a change in workload is detected, optimization component is activated and fed with recent short-term utilization history. Thereafter, the optimization component determines monetary cost of current system configuration and then simulates the cost of different scaling decisions. The *latency-aware* cost function has the responsibility to calculate monetary cost of system configuration. The search function is an implementation of *Recursive Random Search* [62] algorithm which consists of two phases. First, in *exploration* phase, the complete parameter space is explored to find a solution with minimum cost. In second phase – *exploitation phase* – only specific parts of the parameter space which has been discovered in first phase, will be investigated. Kielbowicz [42] has implemented this techniques in the context of Spark Streaming. Thus, It is considered in evaluation scenarios.

#### 7.3 Time-Series Analysis Techniques

Herbst et al. [33] surveys different auto-scaling techniques based on time-series analysis in order to forecast *trends* and *seasons*. *Moving Average Method* takes the average over a sliding window and smooths out minor noise level. Its computational overhead is proportional to size of the window. *Simple Exponential Smoothing*(SES) goes further than just taking average. It gives more weight to more recent values in sliding window by an exponential factor. Although it is more computationally intensive compared to moving average, it is still negligible. SES is capable of detecting short-term trends but fails at predicting seasons. These approaches are more specific instances of *ARIMA* (Auto-Regressive Integrated

Moving Average) which is a general purpose framework to calculate moving averages. However, time-series analysis is only suitable for stationary problems consist of recurring workload patterns such as web applications. Additionally, more advanced forms of time-series analysis which are capable of forecasting seasons (such as *tBATS Innovation State Space Modeling Framework* [44], *ARIMA Stochastic Process Modeling Framework* [37]) are computationally infeasible for streaming workloads.

Taft et al. [58] applied time-series analysis in the context of OLTP databases. The authors argue that reactive approaches don't fit to database world. By the time, auto-scaler component decides to scale-out, it is already too late for a database system. This premise comes from the fact that taking scaling actions in a database doesn't take place in timely manner. The database system has to replicate some of the records which is an additional burden on a heavily loaded system. Thus, database system must take proactive approach and take scaling decisions ahead of time. While this is convincing argument, the auto-scaler module depends on a couple of parameters that are hard to calculate in heterogeneous public cloud environments. First, target throughput of a single server. Second, shortest time to move all database records with single sender-receiver thread. While this might be feasible in some scenarios, on today's cloud environments with virtual machines hosted on heterogeneous physical nodes, getting a near-precise number is unconvincing. It worth noting that author assumed an approximately uniform workload distribution for all database nodes – each database shard serves a fairly equal portion of total workload which is a questionable assumption.

## 7.4 Queuing Theory Techniques

Lohrmann, Janacik, and Kao [45] proposed a solution based on queuing theory. The solution is designed for *Nephele* [46] streaming engine which has a master-worker style architecture. Similar to Spark Streaming, a job is modeled as a DAG. It utilizes *adaptive output batching* [61] – which is essentially a buffer with variable size – to buffer outgoing messages emitted from one stage to the other. Each task – an executer that runs user defined function (UDF) – is modeled as a G/G/1 queue. That is, the probability distributions of message inter-arrival and service time are unknown. In order to approximate these distributions, a formula proposed by Kingman [43] is used. From a bird's eye view, this solution seems promising. However, authors made two unconvincing assumptions that led us to abandon the proposal. First, worker nodes shall be homogeneous in terms of processing power and network bandwidth. Second, there should be an effective partitioning strategy in place in order to load balance outgoing messages between stages. In reality both assumptions rarely occur. Large scale stream processing clusters are built incrementally. Depending on workload, data skew does exist and imperfect hash functions are widely used by software developers.

Zhang, Cherkasova, and Smirni [66] proposed a solution for multi-tiered enterprise applications based on regression techniques. Regression based models can absorb some level of uncertainty and noise by compacting samples. Each tier is modeled as G/G/1 queue and scaled differently compared to other tiers. The system has fixed number of users – a principle known as *closed-loop queuing network*. In order to calculate system workload – incoming message rate – and service time which is required by queuing models, the authors proposed to use Mean Value Analysis [49]. In order to simplify the queuing network, the system is modeled as a *transaction-based* system with independent requests coming from clients. However, It is widely believed that multi-tiered enterprise applications are *session-based* systems [16]. Each request from the same client depends on her previous request during a specific session.

#### 7.5 Reinforcement Learning Techniques

Herbst et al. [31] surveys on state of the art techniques to predict future workload. It includes workload forecasting based on *Bayesian Networks* (BN) and *Neural Networks*. There are several issues with each of them that makes them unsuitable for streaming workloads. As an example, there is no universally applicable method to construct a BN. Furthermore, it requires collecting data and training the model offline. Neural networks suffer from the same issues. That is, it requires collecting samples and periodically training the model. For complex models, training phase is typically computationally infeasible which is conflicting with requirements of thesis.

Tesauro et al. [59] proposes a hybrid approach to overcome poor performance of online training. The system consists of two components: an online component based on queuing system combined with Reinforcement Learning component that is trained offline. The offline component is based on *neural networks*. The authors model the data center as multiple applications managed under a single resource manager. Modeling streaming workloads as a queuing system has two problems. First, modeling is a complicated process and determining probability distributions requires domain knowledge. Second, it requires access to each node (so it can be modeled as a queue) which is currently not possible without modifying spark-core package. Since, it was one the requirements to provide a solution without making any modification to spark-core, this work has been abandoned.

Rao et al. [53] proposed to use Reinforcement Learning to manage resources consumed by virtual machines. It employs standard model-free learning, which is known as *Temporal Difference* [57] or *Sarsa* algorithm. The state space consists of metrics collected from virtual machines (CPU, RAM, Network IO, . . .). There is no global controller and each node decides based on its own Q-Table. As mentioned in literature, standard temporal difference has a slow convergence speed. In order to speedup bootstrap phase, Q-Table is initialized by values that were obtained during separate supervised training. Since this approach also relies on offline training, it wasn't adopted by this thesis.

Enda, and Jim [22] proposed a parallel architecture to Reinforcement Learning. Standard model-free learning (Temporal Difference) is used. No global controller is involved and each node decides locally. In order to speed up learning, all nodes maintain two Q-Tables (local and global tables). Local table is learned and updated by each node. Whenever, an agent learns a new value for a specific state, it broadcasts it to other agents. The global table contains values received from other agents. Additionally, agent prioritize local and global tables by assigning weights to each table. Weights are factors that are defined by application developers. The final decision is the outcome of combining local and global tables. Although each node learns some part of the state space (which may overlap with other nodes), it is not applicable in the context of Spark Streaming. The assumption in this architecture is that, each node is operating autonomously without intervention from other nodes (such as web servers). In contrast, Spark is a centrally managed system. That is, all nodes running Spark jobs are supervised by a single master node (probably with couple of backup masters).

Heinze et al. [29] implemented Reinforcement Learning in the context of FUGU [26] and compared it to threshold-based approaches. Each node, maintains its own Q-Table and imposes local policy without coordinating other nodes. This architecture can not be applied in the context of spark streaming, since Spark abstracts away individual nodes from the perspective of application developer. In order to decrease state space, the author applied two techniques. First, only system utilization is considered. Second, system utilization is discretized using coarse grained steps. To remedy slow convergence, the controller enforces a *monotonicity constraint* [34]. That is, if the controller decides to take scale-out action for a specific utilization, it may not decide scale-in for even worse system utilization. This feature has been adopted by this thesis.

Cardellini et al. [12] proposed a two level hierarchical architecture for resource management in Apache Storm [56]. There is a local controller on each node which is cooperating with the global controller. The local controller monitors each operator using different policies (threshold-based or Reinforcement Learning using temporal difference). In case, local controller decides to scale in or out an specific operator, it contacts the global controller and informs it about its decision. Then it waits to receive confirmation from the global controller. The global controller operates using a token-bucket-based policy [13] and has global view of cluster. It ranks requests coming from local controllers and either confirms or rejects their decisions. Although, this architecture seems to be a promising approach, however it has been implemented by modifying Storm's internal components. As mentioned above, this is in conflict with thesis's requirements.

In order to mitigate the problem of large state space in Reinforcement Learning, Lolos et al. [47] proposed to start the agent from small number of coarse grained states. As more metrics are collected (and stored as historical records), agent will discover *outlier* parameters (those parameters that are affecting agent more, CPU rather than IO as an example). Then, it partitions the affected state into two states and *re-trains* newly added states using historical records. Both Temporal Difference and Value Iteration methods can be used as learning algorithm. Gradually, agent only focuses on some specific parts of the state space, since all parameters are not equally important. This approach, effectively reduces

the size of state space. However, the trade-off is the storage cost in which historical metrics need to be stored. It worth noting that from the context of paper, storage cost (whether it is in-memory or on-disk and the duration of storing historical metrics) is unclear. Thus, this approach has been abandoned due to uncertainty.

Dutreilh et al. [21] proposed a model-based Reinforcement Learning approach for resource management of cloud applications. All virtual machines are supervised by a single global controller. Slow convergence is the bottleneck of model-free learning, in contrast to model-based learning. However, environment dynamics are not available at the time of modeling. Authors proposed to estimate these parameters as more metrics are collected and then switch to *Value Iteration* [57] algorithm instead of *Temporal Difference*. In short, statistical metrics are stored and updated for each visit of (old state, action, reward, new state) quadruple. As more samples are collected, statistical metrics become more accurate and can be directly used in *Bellman* equation. Until enough measurements get collected, a separate initial reward function is used which is essentially the original reward function but with penalty costs removed. Furthermore, In order to reduce the state space – tuple of [request/sec, number of VMs, average response time] – there exists a predefined upper and lower bound for state variables and average response time is measured at granularity of seconds. This approach has been partially adopted by this thesis.

Dutreilh et al. [20] proposed a model-free Reinforcement Learning approach (*Temporal difference* algorithm) with modified *exploration* policy. The standard exploration policy for Q-Learning is  $1-\epsilon$ . Under this policy, the agent performs a random action with probably of  $\epsilon$  and with probability of  $1-\epsilon$ , it adheres to an action proposed by optimal policy. Although the random action is necessary to explore unknown states, but it has sever consequences under streaming workloads. In some cases, it leads to unsafe states where SLOs are severely violated. Since streaming is heavily latency sensitive, this property is undesirable. Thus, author sought toward a heuristic-based policy proposed by Bodik et al. [10]. This policy is based on couple of key observations which has been adopted by this thesis:

- · It must quickly explore different states.
- It should collect accurate data as fast as possible, to speedup training.
- During exploration phase, the policy should be careful not to violate SLOs.

# 7.6 Summary

In this chapter prior work on auto-scaling scaling has been discussed and evaluated. First, threshold-based approaches are investigated. Simple threshold-based approaches are intuitive and simple to understand by application developers and are widely supported by cloud providers. However, adjusting thresholds is a tricky and error-prone process. Then, time-series analysis techniques are explored. As confirmed by other authors, advanced seasonal forecasting is a computationally intensive process, which makes it less suitable for streaming workloads. Queuing theory approaches are suitable for stationary networks with a known probability distribution for workload and service time. Reinforcement Learning techniques has the benefit that it requires zero knowledge about the environment which helps to gradually adapt to changes in environment.

8	Conclusion

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