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Big Data Computing

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Contents

1	Introduction	1
1.1	Data center	1
1.1.1	Reliability and Programmability problems	2
2	Distributed Deep Learning	3
2.1	Parallelize a DNN	3
2.2	Reducing communication overhead	4
2.3	Collective operations used in DNN	4
2.3.1	Broadcast	4
2.3.2	Allgather	6
2.3.3	Reduce-Scatter	7
2.3.4	Allreduce	7
2.3.5	All-to-All	8
3	Network topologies	10
3.1	Torus and Mesh	10
3.1.1	Ring (Torus 1D)	10
3.1.2	Mesh 2D	11
3.1.3	Torus 2D	11
3.2	Trees	11
3.2.1	Fat-tree	12
3.3	Dragonfly	13
3.3.1	Dragonfly+	13
3.4	Other topologies	14
3.5	Load balancing	14
3.5.1	In-Network Congestion Oblivious	15
3.5.2	In-Network Congestion Aware	15
3.5.3	Host-Based Congestion Aware	15
3.6	Congestion Control	15
3.6.1	Random Early Detection (RED)	15
3.6.2	Explicit Congestion Notification (ECN)	16
3.6.3	Data Center TCP (DCTCP)	16
3.6.4	Priority-Based Flow Control (PFC)	16
3.6.5	High Precision Congestion Control (HPCC)	16
3.7	In-Network compute	16
3.7.1	Map-Reduce	17
3.7.2	Allreduce	17

4 Software Infrastructure	18
4.1 Storage	18
4.1.1 Distributed File System	18
4.1.2 SQL Databases	19
4.1.3 NoSQL Data stores	19
4.1.4 RDMA	20
4.2 MapReduce Programming Model	20
4.3 Data-Flow Systems: Spark	21
4.3.1 Resilient Distributed Datasets (RDD)	21
4.3.2 DataFrame and Dataset	22
4.3.3 Spark components	22
5 Similarity	23
5.1 Introduction	23
5.1.1 Search problem	23
5.1.2 Vertical and Horizontal scaling	23
5.2 Similarity meaning	23
5.2.1 Euclidean Space	24
5.2.2 Cosine Similarity	24
5.2.3 Jaccard index	25
5.3 Curse of Dimensionality	25
5.4 Dimensionality Reduction	26
5.4.1 Principal Component Analysis (PCA)	26
5.5 Clustering	28
5.5.1 Document clustering	29
5.5.2 Clustering algorithm: Partition-Based	30
5.5.3 K-means	31
5.5.4 Similar algorithms to K-means	32
5.5.5 Clustering Quality	33
6 Recommender Systems	35
6.1 Content-based filtering	35
6.2 Collaborative filtering	36
6.2.1 User-based Neighborhood	37
6.2.2 Item-based Neighborhood	37
6.2.3 Comparison between User-based and Item-based	38
6.2.4 Latent-factor	38
6.2.5 Bias	40
6.3 Evaluation Metrics	40
6.3.1 Precision and Recall	40
6.3.2 Personalization	41

7 Analysis of Large Graph	42
7.1 Definition of graph	42
7.1.1 Representing a graph	42
7.2 Web search engine	43
7.2.1 Content similarity method	43
7.2.2 Scale-Free network	43
7.2.3 Computing node importance	44
7.3 Google PageRank	45

1

Introduction

With **Big Data** we refer to an actual phenomenon, defined by five properties called **5V**:

- **Value**: extracting knowledge from data is valuable.
- **Volume**: large amount of data.
- **Variety**: different format of data.
- **Velocity**: the speed at which the data are generated is very high.
- **Veracity**: reliability of the data used.

To do a computation on a lot of data at a high speed (like a google search), there are different problems that occurs:

- Disks are not large enough.
- Disks are not fast enough.
- CPUs are not fast enough.

1.1 Data center

To do this type of computations **data center** are used, which can upgrade their performances in two way:

- **Scale up**: buying new and more powerful components. This is a problem because the increase in the velocity required is faster than the performance increase.
- **Scale out**: buying more components and interconnect them to work in parallel.

A data center is composed by a series of rack interconnected between them with a private network. Every rack has servers inside them that contain CPUs, GPUs and memory. A server is considered a node in the network.

The major problem with the data centers is the bottleneck caused by the network. The network can't send all the data required for the computation in a fast way and this slow down the computation. Also, the increase in GPU power (that does the major part of computation in a data center) is much faster than the increase in network speed during the years.

1.1.1 Reliability and Programmability problems

Another problem is related to **reliability**, so the probability that a server, disk or network component fails. Even if the probability of a single component is very low (one fail every 10 years) a large data center with thousand of components will have failures very frequently. This mean that checkpoint must be used, and every failure reset to the last one.

Last problem is the **programmability**, so how to write an efficient parallel program being aware of all the components. Things that have to be taken in account are how to store data, how to send data efficiently, what to execute on CPU or GPU and many other.

2

Distributed Deep Learning

Deep Learning performs better with more data and parameters, but also require more computation.

The training of a Distributed Neural Network (DNN) is done by doing many iteration of 3 steps:

1. **Forward propagation (Compute the function)**: apply the model to input and produce a prediction.
2. **Backward propagation (Compute the gradient)**: calculate the error and find the parameters that contribute more to the error to correct them.
3. **Parameters update (Use the gradient)**: use the loss value to update the model parameters.

2.1 Parallelize a DNN

There are 3 ways to parallelize a DNN:

- Data parallelism: divide the training dataset in batches and compute the model on each batch in parallel. After each iteration we synchronize the parameters of the different models by doing a gradient aggregation. This is done by doing an Allreduce on the weights vector across the GPUs.
- Pipeline parallelism: divide the model by the layers, assigning each layer to a different GPU. In the forward phase the first GPU output will be the input of the second GPU and so on. In the backward phase is the opposite. The problem is that the last GPUs have a lot of idle time during the forward phase and the first GPUs in the backward phase. This idle time is called bubble. To reduce this bubble we can divide the batch of data in micro-batches and do the forward and backward computation on the micro-batches. There are also other complex methods to reduce the bubble.
- Operator (or Tensor) parallelism: if all the parameters of a layer do not fit in a single GPU we have to split the layer across multiple GPUs. An example could be splitting the matrices that we have to multiply (training require a lot of matrix multiplication) and using an Allreduce to sum the partial matrices. This requires a lot of communication.

In large models all three technique are used at the same time.

2.2 Reducing communication overhead

To reduce the communication overhead we can compute and communicate in the same time, sending the gradients of every layer as soon as they are computed, so the computation of the gradient of the previous layer can be done during the communication of the gradient of the successive layer.

Another way is to compress the gradient and reducing his data volume, this can be done in three ways to have a **lossy compression**:

- Quantization: reduce the bitwidth of the elements (for example for float32 to float16). Some models converge even with only one bit per weight (knowing only if is positive or negative).
- Sparsification: sending only the larger k values or by setting a threshold and sending only the weights bigger than it.
- Low rank: decomposes gradient into lower-rank matrices.

Lossy compression training converges with the same asymptotic rate, but in real life it requires more iterations.

Asynchronous systems could be also used to speed up the process and overcoming the gradient aggregation that acts as a barrier. This can be done with different techniques, like aggregating after not one but many iterations or not waiting the slowest GPUs.

2.3 Collective operations used in DNN

In DNN different collective operations are used:

- Allreduce for gradient aggregation in data parallelism.
- Allgather and Reduce-Scatter in sharded data parallelism.
- Allgather and Allreduce for matrix multiply in operator parallelism.
- Alltoall in expert parallelism.

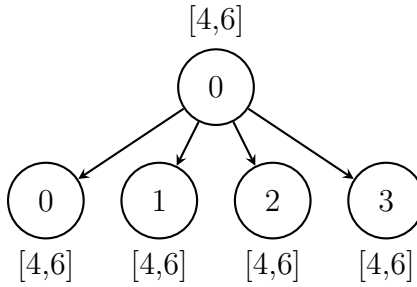
The cost model can be expressed with this formula:

$$T = k\alpha + \delta n\beta$$

In which α is the latency, a fixed cost, β is the cost per byte sent, n are the byte sent and δ is the fraction of data sent in respect to n , considered as the maximum between the input and the output.

2.3.1 Broadcast

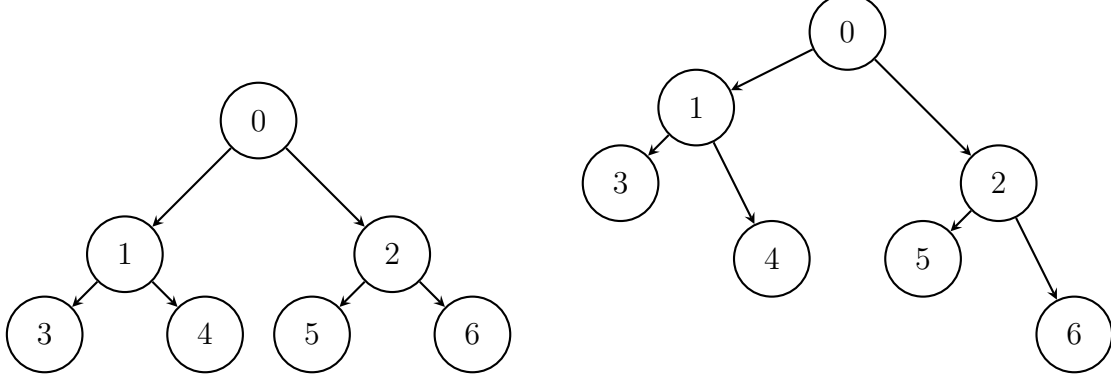
A simpler collective operation is the **Broadcast**, in which data from a process are copied to all the other processes.



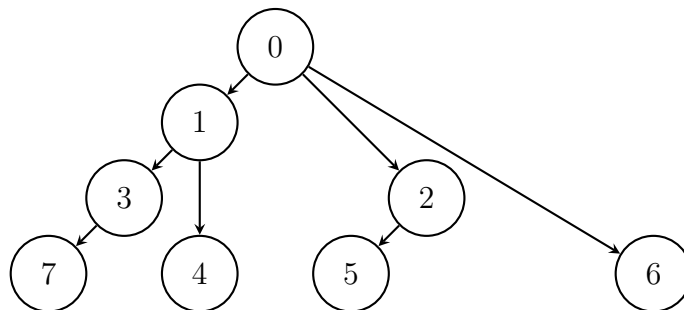
The basic algorithm to compute the Broadcast is the **chain algorithm**, in which at every step the last process to have received the data sends them to the successive. So, in step 0 the process 0 (who has the data at the start) sends them to the process 1, in the next step 1 will send them to process 2 and so on. The cost of this algorithm is:

$$T = (p - 1)(\alpha + n\beta)$$

Another algorithm involves the binary trees, in which at each step the processes that have received the data at the last step send all the data to two processes that don't have yet.



Exists also an algorithm with binomial trees, in which a process doesn't stop sending after two steps but continue to send at every step.



The cost of this algorithm is:

$$T = \log p(\alpha + n\beta)$$

The Broadcast can also be done as a Scatter (takes a vector from a process and divide it evenly between all processes) plus a Allgather. The cost of the Scatter, using the binomial trees, is

similar to the Broadcast algorithm, with the difference that the data sent halves every step, so for a step k the data sent will be $\frac{n}{2^k}$. The cost of the Scatter is:

$$T = (\log p)\alpha + n\beta$$

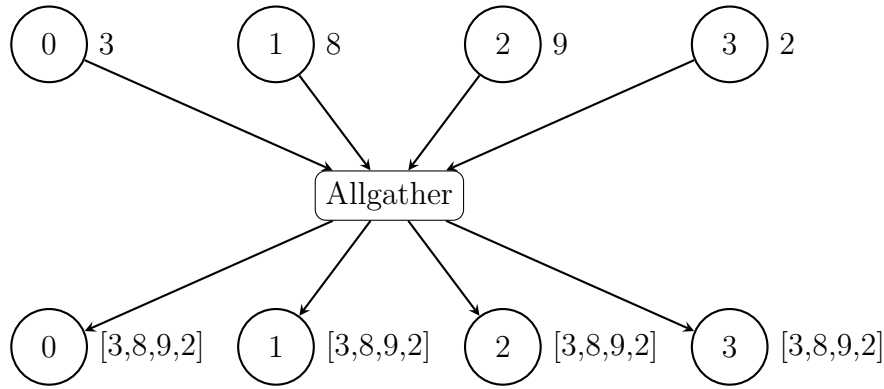
For the Allgather we use the Recursive doubling algorithm explained in Section 2.3.2.

The total cost of this approach for the Broadcast is:

$$T = \underbrace{(\log p)\alpha + n\beta}_{\text{Scatter}} + \underbrace{(\log p)\alpha + n\beta}_{\text{Allgather}} = 2[(\log p)\alpha + n\beta]$$

2.3.2 Allgather

The **Allgather** is a collective operation that takes data from every process, combine them in a vector and copies the resulting vector to all the processes.

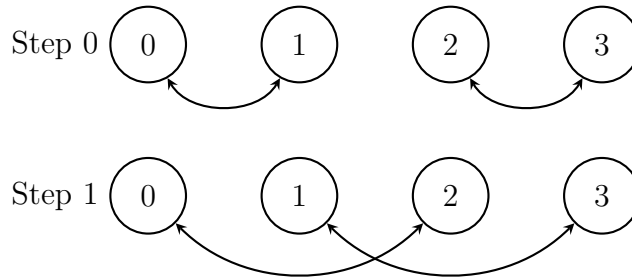


The basic algorithm to compute the Allgather consists in $p - 1$ phases in which every process sends his data to another process. The cost is:

$$T = p\alpha + n\beta$$

Another simple algorithm is the **Ring** algorithm, which use the same structure of the basic algorithm, but instead of sending the data to a different process every step, a process i will always send data to process $i+1$. The cost of the algorithm remain the same, but the difference is that the every link in the system is used by only one communication at a time.

The **Recursive doubling (Butterfly)** algorithm has a logarithmic number of phases and consist in exchanging data in a bidirectional way between process with distance that doubles at every step. In a step k , a process i will exchange all the data with a process at distance 2^k .

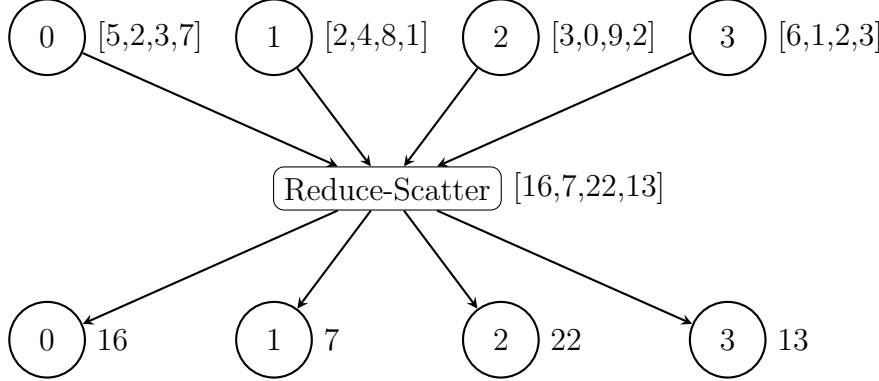


The cost is:

$$T = (\log p)\alpha + n\beta$$

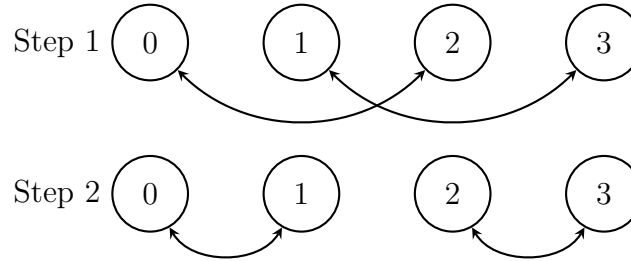
2.3.3 Reduce-Scatter

The **Reduce-Scatter** is a collective operation that takes a vector of data from every process and execute an operation on them. After calculating the resulting vector, it is splitted evenly between the processes, following the indexes.



The basic and Ring algorithm for the Reduce-Scatter are the same algorithm of the Allgather but in reversing order of communication.

Instead of Recursive doubling, for Reduce-Scatter we use the **Recursive halving (Butterfly)** algorithm, which works in a similar way but with the distance that halves instead of doubling. Also the data exchanged is more in the first stage and halves every stage.

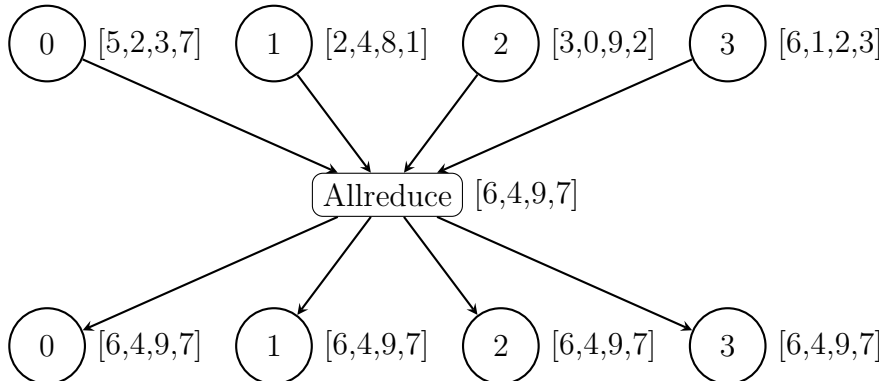


The cost is equal to the Recursive doubling algorithm of the Allgather:

$$T = (\log p)\alpha + n\beta$$

2.3.4 Allreduce

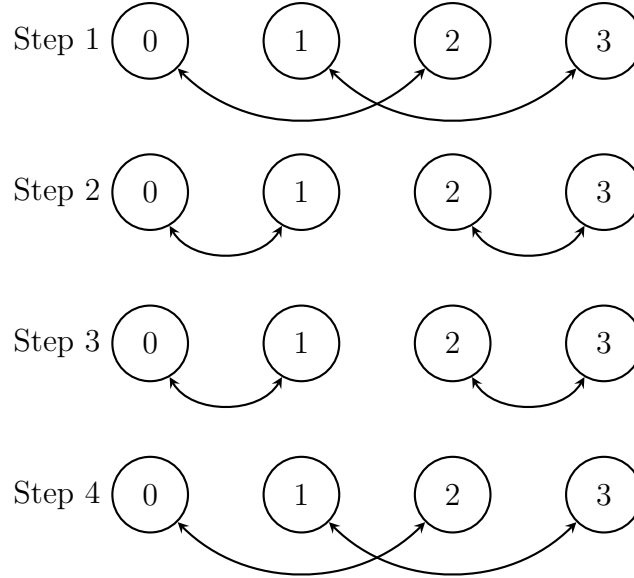
The **Allreduce** is a collective communication that takes a vector from every process, execute an operation on them and the resulting vector is copied in every process.



The Ring algorithm is done by executing a Reduce-Scatter and an Allgather with Ring algorithm, so the cost will be the sum of the two:

$$T = 2(p\alpha + n\beta)$$

The **Rabenseifner (Butterfly)** algorithm use a Reduce-Scatter with Recursive halving algorithm followed by an Allgather with Recursive doubling algorithm.



Also in this case the cost is the sum of the two algorithms:

$$T = 2[(\log p)\alpha + n\beta]$$

There is an algorithm that isn't the sum of a Reduce-Scatter and Allgather, the **Recursive doubling** algorithm. The algorithm is similar to the one to execute the Allgather but every step the processes send all the vector. The cost is:

$$T = (\log p)(\alpha + n\beta)$$

2.3.5 All-to-All

The **All-to-All** is a collective communication that takes a vector from every process and moves the data so that at the end every process i has all the data with index i in the starting vectors. If we imagine the vector as columns of a matrix, the result of the All-to-All is the transposition of the matrix.

p_0	p_1	p_2	p_3		p_0	p_1	p_2	p_3
00	10	20	30	$\xrightarrow{\text{All-to-All}}$	00	01	02	03
01	11	21	31		10	11	12	13
02	12	22	32		20	21	22	23
03	13	23	33		30	31	32	33

The basic algorithm is a linear algorithm in which in every fase k every process i sends data to process $i + k$. The cost is:

$$T = p\alpha + n\beta$$

The **Bruck (Butterfly)** has a logarithmic number of phases and works similarly to the Recursive halving algorithm for Reduce-Scatter. The difference is that half the vector is sent at every step.

p_0	p_1	p_2	p_3		p_0	p_1	p_2	p_3		p_0	p_1	p_2	p_3
00	10	20	30		00	11	22	33		00	11	22	33
01	11	21	31	Shift	01	12	23	30	Fase 1	30	01	12	23
02	12	22	32		02	13	20	31		02	13	20	31
03	13	23	33		03	10	21	32		32	03	10	21

	p_0	p_1	p_2	p_3
Fase 2	00	11	22	33
	30	01	12	23
	20	31	02	13
	10	21	32	03

So the cost of the algorithm is:

$$T = (\log p) \left(\alpha + \frac{n}{2} \beta \right)$$

3

Network topologies

The network **topology** define how the nodes in the system are connected between them. The topologies can be of different types and can be divided in categories using different parameters:

- Blocking and Non blocking: a network is non blocking if every pair of switch can be connected through disjoint path. So in an ideal case there is no congestion.
- Direct and Indirect: a network is direct if every switch is connected to at least one server.
- Regular and Irregular: a network is regular if it can be represented with a regular graph.

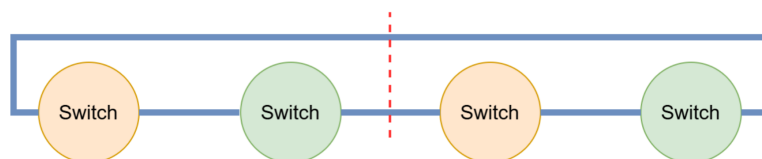
To start we need to explain some definitions:

- Switch radix: number of switch ports.
- Network diameter: maximum distance between two switches.
- Bisection cut: minimum number of links that needs to be cut to divide the network in two almost equal parts. The total bandwidth of the links cut is the Bisection bandwidth.
- All-to-All bandwidth (global bandwidth): the bandwidth the network has when running a linear All-to-All. Non blocking topologies have full global bandwidth.

3.1 Torus and Mesh

3.1.1 Ring (Torus 1D)

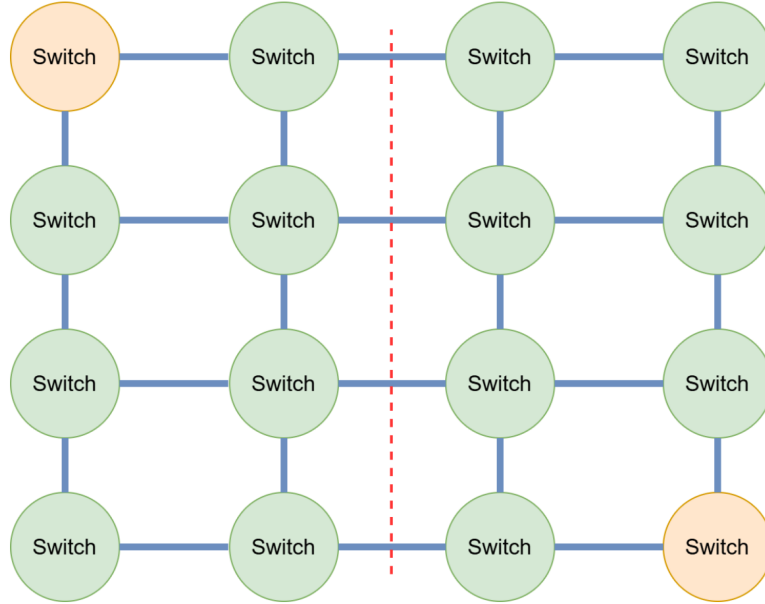
The **Ring** topology is a topology in which every server needs only two links, one to the previous server and one to the successive server.



This topology has a diameter of $\lceil \frac{n}{2} \rceil$ and a bisection cut of 2.

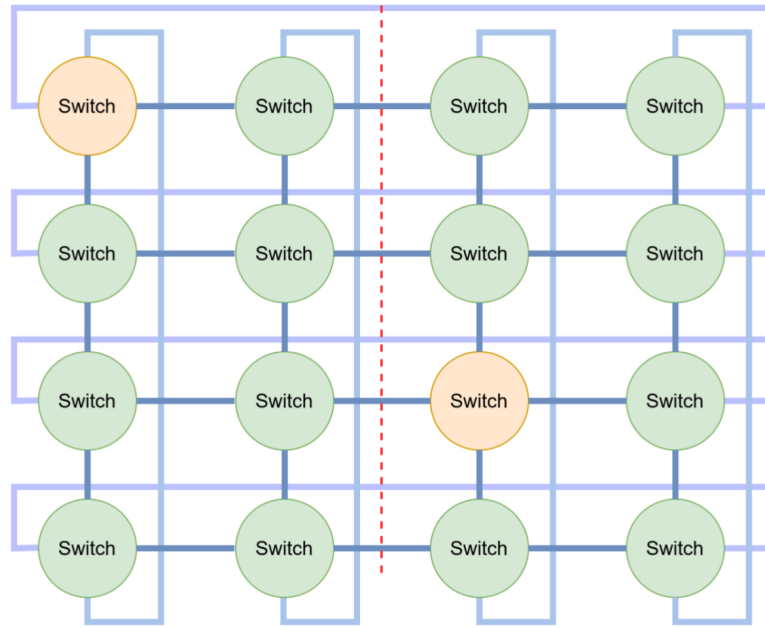
3.1.2 Mesh 2D

The **Mesh** topology is a grid and has a diameter of $2(\sqrt{n} - 1)$ and a bisection cut of \sqrt{n} .



3.1.3 Torus 2D

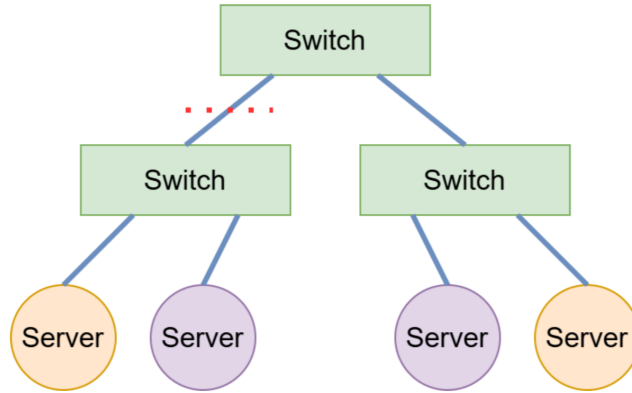
The **Torus 2D** is similar to the mesh but with the first and last switch of every row and column that are connected. The diameter is \sqrt{n} and the bisection cut is $2\sqrt{n}$.



3.2 Trees

The tree topologies are based on the switch radix r . The basic tree has a height $h = \log_{r-1}(n)$, a diameter of $2h$ and a bisection cut of 1 (doesn't make much sense if the root has more than

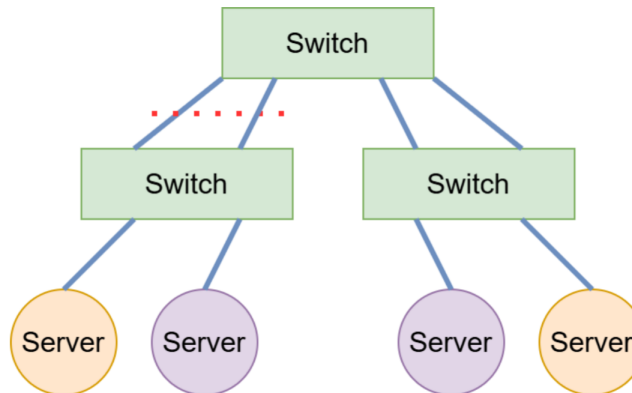
two child).



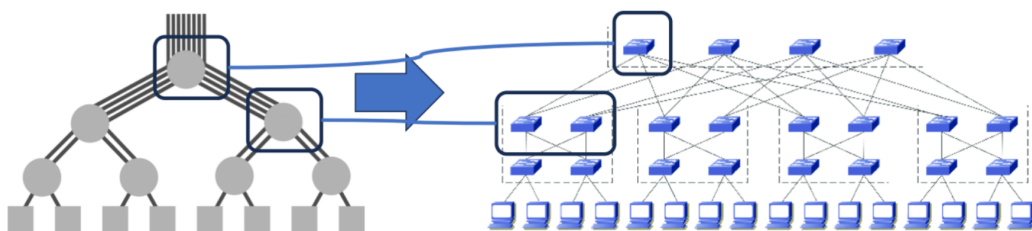
The basic tree is a blocking topology, because the switch have only one link going up.

3.2.1 Fat-tree

A **Fat-tree** is a tree in which every switch has the same number of links going up and going down. This makes this topology non blocking, providing full global bandwidth. The number of layer needed is also lower than the basic tree, having height $h = \log_2(n)$, and consequently the diameter will be smaller.



With this change the switch on different layer have different radix but in a real network every switch has the same radix. So the switch are divided in smaller ones to make every one have the same radix.



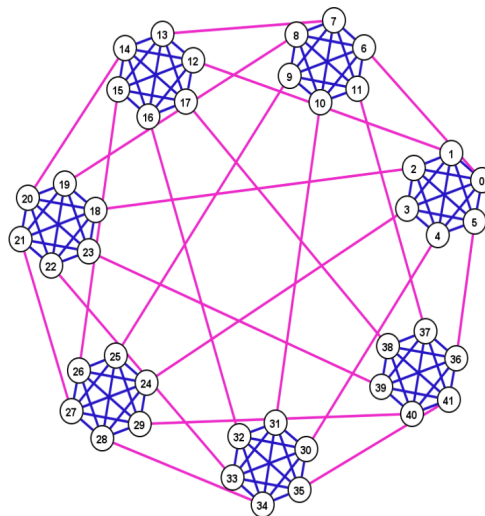
This network, also called **Folded CLOS** is the best that can be built in terms of performance but is also the most expensive one.

To reduce the cost we can make every switch have a ratio of 2:1 between links going down and links going up. This makes the topology become a blocking topology but saves money. The cost reduction is linear with the ratio.

Fat-trees don't scale well because to add nodes we have to add layers and this increase the diameter.

3.3 Dragonfly

The **Dragonfly** topology is composed of fully connected groups of switches. The groups are also fully connected between them. This topology scales better than fat-tree and has a diameter of 3 (5 if the links between switch and server are counted) that is constant.

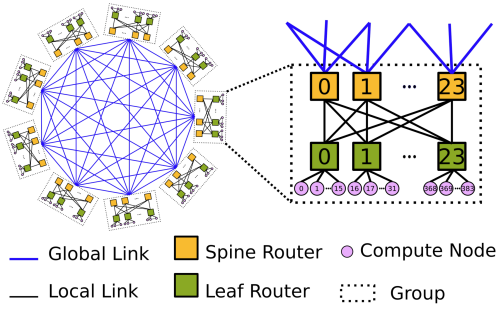


This topology has some disadvantages:

- Doesn't have full global bandwidth
- Adding new server is complicated.
- Load balancing is hard, because minimal and non-minimal path exists between two nodes.
- Can have loops, so deadlocks are possible.

3.3.1 Dragonfly+

The **Dragonfly+** is a modified version of the Dragonfly in which the groups are bipartite graphs.



The diameter is 3 like the dragonfly but can connect more server with the same switch radix.

3.4 Other topologies

- **Slimfly:** this topology is composed by two types of groups, each connected in a different way between itself. The good thing of this topology is the diameter equal to 2 but it works only for specific switch radix values.
- **HammingMesh:** this topology optimized o train Deep Learning models. Is a Mesh 2D divided into smaller meshes and in which every row and column is connected using a non blocking fat-tree.
- **Jellyfish:** this topology has no regular structure and uses a random graph so the is easy to add new servers. The problem are the management and configuration that are very hard.
- **Reconfigurable Data Center Network (RDCN):** in this topology the links are not fixed but can change over time. This can improve the performance but it also has some problems like routing and when to change the links.

3.5 Load balancing

The **load balancing** (also known as **adaptive routing**) is when there are multiple path between a pair of node and the traffic must be balanced between the different paths. The algorithms can be divided in different categories:

- Centralized: there is an external controller that makes the decisions.
- Distributed: each server makes its own decisions. Distributed algorithm can be divided based on who takes the decision:
 - In-Network: decisions are made by switches.
 - Host-Based: decisions are made by the servers.

They can also be divided based on if they choose path to improve the performance:

- Congestion Oblivious: the path is changed randomly if there is congestion.
- Congestion Aware: the path is changed to improve the performance.

3.5.1 In-Network Congestion Oblivious

The two main algorithm in this category are **Equal-cost Multi-path (ECMP)** and **Random packet spraying**.

ECMP uses an hash function to select a random output port for every message that arrives. This causes a lot of collision when multiple message get mapped to the same port.

Random packet spraying selects a random path for each packet (instead of each message). The packet might arrive out of order, that is a problem with TCP, but not with modern RDMA protocols.

3.5.2 In-Network Congestion Aware

This types of algorithms are based on **flowlet**, that are burst of packet created when the interval between two packets of the same flow is larger than a pre-set gap. This gap is set to be sure that the probability of packets arriving not in order is very low.

The first algorithm of this type is **CONGA**, in which every leaf switch keep track of the congestion on all the paths towards other leaf switches. When a flowlet arrive to the switch, it is sent to the least congested path.

Another algorithm is **Let It Flow**, in which every time a flowlet arrive to the switch, it is sent in a random path. If the path is congested, the flowlet will be smaller and will be rerouted.

3.5.3 Host-Based Congestion Aware

The algorithm of this type is **Flowbender**, that uses the ECMP hashing function to force every flow on a different path, when it detect congestion.

3.6 Congestion Control

The congestion control is needed for some problems that can't be resolved by load balancing. For example if the link congested is the one connected to an host or if a switch has two flow in input that want to exit on the same port.

Congestion control is based on detecting the congestion and adjusting the sending rate accordingly. The congestion can be detected by the sender, the switch or the receiver, but only the sender can adjust the rate.

3.6.1 Random Early Detection (RED)

The **RED** algorithm notify the congestion by dropping the packets (TCP will timeout), but instead of dropping it when the queue is full, two threshold are set, a minimum and a maximum threshold. When the average queue length is higher than the minimum threshold, the packets are dropped with a probability, if it exceed the maximum threshold the packets are always dropped.

3.6.2 Explicit Congestion Notification (ECN)

The **ECN** algorithm, uses two bits in the packet header. It follows the same logic as RED but instead of dropping the packets, it marks them.

When a packet is marked, the receiver will mark the corresponding Ack to communicate to the sender that there is congestion. The sender will then adjust the sending rate by 50%.

To avoid reporting outdated information a packet is marked when exiting the queue and not when entering.

3.6.3 Data Center TCP (DCTCP)

DCTCP works similarly to ECN but the sending rate is adjusted based on the amount of packets that are marked, if all the packets are marked the rate is reduced by 50%, but in all the other cases is reduced by a minor amount. Also, the marks control is done by using the instantaneous queue length and not the average.

3.6.4 Priority-Based Flow Control (PFC)

This algorithm assumes that the network is lossless (common in PC networks), so the switches don't drop packets.

It is based on the priority queue inside a switch, when one queue becomes too full, a message is sent to all the application sending packets on that queue to make them stop.

There are two major problems with this algorithm, the first one is deadlock, because a cycle of queues could stop each other. The second is congestion spreading, when stopping a switch stops all the ones before it and so on.

3.6.5 High Precision Congestion Control (HPCC)

The goal of the algorithm is to converge to the correct sending rate by using precise congestion signal. It works like ECN but instead of using two bits it attaches to the packet header more information: timestamp, queue length, transmitted bytes and link bandwidth. The receiver also copies these informations on the corresponding Ack.

3.7 In-Network compute

The switch can be used, in addition to simply forwarding the packets from input to output, to do complex operation.

The existing programmable switches are based on **Reconfigurable Match-Action Table (RMT)** in which the programmer declares how packets are processed.

This in-network compute can be used to do simple tasks:

- In-network telemetry: to have more details on the status of the network.
- New traffic load balancing algorithm.
- Better support for congestion control.

It also can be used to perform more complex tasks:

- In-network Map-Reduce
- In-network Allreduce

3.7.1 Map-Reduce

Map-Reduce is an approach in which the input data is splitted, mapped, shuffled and reduced to final data.

Every switch does the partial aggregation of the data and sends the result to the successive switch. At the end the last switch does the reduction on the data of only two switches. This reduce the amount of data transferred and speed ups a lot the computation.

If a switch fills its memory it sends the packets without doing the partial aggregation, in the worst case is the same as doing the Map-Reduce without in-network computation.

3.7.2 Allreduce

To do an in-network Allreduce we build a reduction tree in which the switches do the reduction. So every switch does the partial reduction of the data and sends only the results. This is now a single phase algorithm with a cost of:

$$T = \alpha + n\beta$$

It has some issues:

- Load balancing
- Reproducibility: the result could be different every run because floating point sum is not associative.
- Lack of floating-point units: floating-points are expensive in terms of area and latency so is better to use fixed-point that are composed by integer bits and fraction bits.
- Packet losses and switch failures: if an already aggregated data is lost the retransmission mechanism must be a lot complex to recover the data.

4

Software Infrastructure

4.1 Storage

Generally with storage we mean data that can't be accessed directly by the CPU, instead memory is the data addressed from the CPU.

The storage used for Big Data can have various forms:

- Distributed File System (DFS): manage large files on multiple nodes.
- NoSQL data stores: non-relational data models like key-value or graph. Have orizontal scalability e fault tolerance.
- NewSQL databases: relational databases, but scalabale and with foul tolerance.

4.1.1 Distributed File System

A famous DFS is the **Google File System (GFS)**, that is built with the assumption that there are hardware fails and multiple sequential writes of data.

Needs to store very large files, that are splitted in chunks (64 or 128 MB). The chunks are stored in chunks server and avery chunks is replicated for fault tolerance and availability. The chunks are managed by a master node that stores the file metadata and manages the operation on chunks. To guarantee availability, shadow masters take its place if the master fails.

GFS have other additional features such as:

- Data integrity: chunks are divided in 64KB blocks and fro every block a checksum is computed and checked every time data is read.
- Load balancing: chunks are balanced across chunk servers.
- API: other than traditional operation, supports two special one: snapshot (istantaneous copy) and record append (atomically append).

To write on a file GFS has a series of steps:

1. Application issues a write
2. Client translates the write position into a chunk index and sends the request to the master node
3. The master node sends the client the locations of the replicas (one of the replicas is the primary replica)

4. The client sends the data to be written to all the replicas
5. The client tells the primary replica to start the write
6. The primary replica decides in which order the data must be written in the chunk (there might be data coming from different clients in different order in the buffer)
7. The primary replica communicates the write order to the other replicas
8. The secondary replicas acknowledge the end of the write
9. The primary replica acknowledge the end of the write to the client

GFS appends data to the file at least once atomically (application must cope with possible duplicates).

Another DFS is the **Hadoop Distributed File System (HDFS)**, that is essentially a GFS clone with few differences. The most important one is the erasure coding, that replace the replicas, saving space.

Erasure coding consist in splitting the data in chunks and storing them together with m parity chunks that are used to recover the chunks in case of corruption.

4.1.2 SQL Databases

Relational Databases management promise ACID properties:

- Atomicity: either all the statements in a transaction are executed, or the transaction is aborted without affecting the databases
- Consistency: the database is consistent both before and after a transaction
- Isolation: the result of incomplete/in-progress transactions are not visible to other in-progress transactions
- Durability: Once a transaction is committed, it will remain committed even in case of a system failure

The main problem with relation databases is the scalability, because they were not designed for a distributed system. They can scale by using replication (data replicated on multiple nodes) or sharding (dividing data on multiple nodes) but either of them impact the performance.

4.1.3 NoSQL Data stores

NoSQL data stores supports flexible schema and horizontal scaling. They tradeoff reliability for higher performance. Instead of ACID properties use BASE properties:

- Basically Available: the system is available most of the time and there could exist a subsystem temporarily unavailable
- Soft state: the system might not always be consistent
- Eventually consistent: at some point, the system converges to a consistent state

NoSQL data stores are easy to scale than SQL ones and have and higher performance for large data.

4.1.4 RDMA

RDMA (Remote Direct Memory Access) is a way of handling data sends and receives without using the kernel, that occupies a CPU core.

In RDMA systems the Network Stack is divided: one part is done by the software and one by the NIC. Also, the NIC resources can be accessed directly by the user-space, bypassing the kernel.

To bypass the kernel when sending data, instead of doing the control operations (done by the kernel) for every send, all the checks are done only once and the other times are skipped.

The most used protocol for RDMA is Infiniband/RoCE that in the last version has an Infiniband transport protocol, but also uses UDP/IP for the network layer and Ethernet for the link layer to be able to communicate with a system outside that doesn't use Infiniband.

To speed up the simple operation, like receiving a message, do a simple calculation and send it back, smartNIC are used. In SmartNIC a CPU is put in the NIC to perform the simple calculation without paying the cost of sending the data to the application and back.

4.2 MapReduce Programming Model

MapReduce is a programming model used for processing big datasets with parallel algorithm on a cluster. It uses a DFS to store data on multiple nodes and guarantee fault tolerance.

The MapReduce consists of steps:

1. Input: key-value pairs $\{(k_i, v_i)^*\}$

2. Map phase: $(k_i, v_i) \rightarrow \{(k'_i, v'_i)^*\}$

Takes the input key-value pair and outputs a set of key-value pairs. The map function is called for every key-value pair in input.

3. Combine phase: $(k'_i, \{v'^*_i\}) \rightarrow (k'_i, v'_i)$

After the map task the output may contain many pairs sharing the same key, if the operation is suitable a Combine phase can be added to pre-aggregate the value before the Shuffle phase.

4. Shuffle phase: $\{(k'_i, v'_i)^*\} \rightarrow \{(k'_i, \{v'^*_i\})^*\}$

Collect all the pairs with the same key and groups the values associated.

5. Reduce phase: $(k'_i, \{v'^*_i\}) \rightarrow (k'_i, v''_i)$

The reduce function is called for every key-value pair at the end of the shuffle phase.

The MapReduce is good for problems that require many sequential data access and large batch jobs, but is not good for problems with random access to data and independent data. Also, not every application can be expressed with a MapReduce.

In MapReduce input and output are stored on the DFS, instead intermediate results of map and reduce function are stored in the local filesystem of each node.

The master node takes care of coordination, propagating information on finished tasks between mappers and reducers. It also pings nodes to detect failures and reschedule the task on other nodes if the fail occurred.

To decide how key-value pair are shuffled and sent to reducers a Partition Function is used. With R reducer nodes a key k is sent to reducer r with a function:

$$r = \text{hash}(k) \% R$$

4.3 Data-Flow Systems: Spark

In MapReduce the tasks have only two ranks: one for map and one for reduce. **Data-Flow** systems allow any number of ranks and other function than map and reduce.

The most popular Data-Flow System is **Spark**, a computing framework that provides a lot of functions, fast data sharing and execution graphs. It's also compatible with Hadoop. Spark is a fault-tolerant system with in-memory caching that provides efficient execution of multi-round algorithms.

The driver process (master in MapReduce) runs the application from a node in the cluster and its job is to maintain information about the application, respond to user input and manage the works across executors. The executor process (worker in MapReduce) run the code assigned and reports the state back to the driver. The cluster manager controls physical machines and allocates resources.

4.3.1 Resilient Distributed Datasets (RDD)

The fundamental abstraction of Spark are RDDs, a collection of elements of the same type. The works in Spark are expressed as transformation and operations on RDDs.

Each RDD is split in partitions distributed across the nodes, the program can specify the number of partition. Typically is 2 or 3 times the number of cores to be sure every core is used but the partitions are not too small.

RDDs are immutable, for multiple purpose:

- Parallelism: the data that a process is reading can't change
- Caching: there is no need for consistency
- Fault tolerance: RDD maintain a trace of the transformation done and an RDD can be re-calculated if the node fails

RDDs can be created from data storage or from the transformation of other RDDs. With the property of being recalculable the intermediate RDDs could not be saved if they are used few times. If they are used multiple times is better to save them.

On an RDD A three types of operation are possible:

- Transformation: generate another RDD B from the data of RDD A
- Action: computation on the RDD A that returns a value to the application
- Persistence: save the RDD A for later

The transformations on RDDs can require to shuffle data across nodes:

- Narrow transformation: each partition of an RDD A contributes to one partition of the RDD B , so there is no need to shuffle data.

- Wide transformation: each partition of A contributes to multiple partition of B , so a shuffle of data is needed.

Spark uses the **lazy evaluation**, means that nothing is computed until it is used in an action. When an action is triggered on an RDD, Spark builds a graph of operations that need to be executed to compute the action.

4.3.2 DataFrame and Dataset

Spark provides two interfaces to operate on structured data: DataFrame API and Dataset API. A DataFrame is a distributed collection of data organized in named columns, that allow higher abstraction than RDD. Like RDDs, DataFrames are immutable and the operations executable on DataFrames are transformations and actions. Datasets API is a more general extension of DataFrame API.

DataFrames have higher performance than RDDs due to optimizations of the query plan.

4.3.3 Spark components

Spark has different components to work with different types of data:

- Spark SQL: component for working with structured data like relational DB, CSV and JSON. Can execute queries and create SQL DataFrames.
- GraphX: component for working with graphs.
- Streaming: component that enables scalable, high-throughput, fault-tolerant stream processing of live data streams.

Spark Streaming divides data streams into micro-batches created at regular time intervals and process every batch like an RDD. The output is also pushed out in batches. The basic abstraction to work with data streams is **Discretized Stream (DStream)** that is implemented as a sequence of RDDs (so it is immutable), each one containing data from a certain interval. The DStream API is very similar to the RDD one and also the transformations are very similar.

The transformations can be stateless, if the process of each batch is independent to the previous ones, or stateful, if uses data from previous batches.

The output operations push out the DStream data to external system (like actions for RDDs, they trigger the computation).

An important type of operation when working with data streams are window operations, that apply a transformation over a sliding window of data. Two parameters are specified: the window length (duration of the window) and the sliding interval (interval at which the operation is performed). The parameters must be multiples of the batch interval.

5

Similarity

5.1 Introduction

5.1.1 Search problem

A common problem that occurs is to find a value x in a list of n items. This can be done in two ways:

- List unsorted: $O(n)$
- Sort list and binary search: $O(n \log n) + O(\log n)$

The best case scenario depends on n and on the number of search that we need to do.

Let m be the number of search, the cost of the two approach is:

- List unsorted: $O(mn)$
- Sort list and binary search: $O(n \log n) + O(m \log n)$

So, is convenient to sort if:

$$n \log n + m \log n < mn \implies m \geq \left\lceil \frac{n \log n}{n - \log n} \right\rceil$$

5.1.2 Vertical and Horizontal scaling

There are two ways in which the data can scale:

- Vertical scaling: the number of items n is very large, couldn't even fit in memory. We will assume that this happens by design.
- Horizontal scaling: a single value of the list is complex to represent, for example images. The problems shift to find the most similar image to the input one.

5.2 Similarity meaning

Similar can mean different things depending on the domain and the representation we are using. We assume data lives in a d -dimensional Euclidean space. Similarity can be compute with different metrics.

Metric and Metric Space

Given a set X , δ is a **metric** if is function $\delta : X \times X \rightarrow [0, \infty)$ where:

1. $\delta(x, y) \geq 0$ (non-negativity)
2. $\delta(x, y) = 0 \iff x = y$ (identity of indiscernibles)
3. $\delta(x, y) = \delta(y, x)$ (simmetry)
4. $\delta(x, y) \leq \delta(x, z) + \delta(z, y)$ (triangle inequality)

X is the **metric space**.

5.2.1 Euclidean Space

Given $X = \mathbb{R}^d$, for two points $x, y \in \mathbb{R}^d$ the Euclidean distance is defined as:

$$\delta(x, y) = \sqrt{\sum_{i=1}^d (x_i - y_i)^2}$$

The Euclidean distance is also referred to as the L^2 norm of the displacement vector:

$$\|x - y\|_2 = \sqrt{(x - y)(x - y)}$$

The Euclidean distance can be generalized by the **Minkowski distance** (L^p norm), that is defined as:

$$\delta_p(x, y) = \left(\sum_{i=1}^d \|x_i - y_i\|^p \right)^{\frac{1}{p}}$$

The important value of p for this formula are:

- $p = 1$ (Manhattan distance):

$$\delta_1(x, y) = \sum_{i=1}^d \|x_i - y_i\|$$

- $p = 2$ (Euclidean distance):

$$\delta_2(x, y) = \sqrt{\sum_{i=1}^d \|x_i - y_i\|^2}$$

- $p \rightarrow \infty$ (Chebyshev distance):

$$\delta_\infty(x, y) = \max_{i \in [1, d]} (\|x_i - y_i\|)$$

5.2.2 Cosine Similarity

Measures the angles between two vectors, the output has range $[-1, 1]$. It represents the orientation difference, not the magnitude.

Is defined as:

$$\cos(\theta) = \frac{xy}{\|x\|_2 \|y\|_2}$$

where θ is the angle between the two vectors.

5.2.3 Jaccard index

Measures the similarity between two finite sets. Is defined as:

$$J(A, B) = \frac{|A \cap B|}{|A \cup B|} = \frac{|A \cap B|}{|A| + |B| - |A \cap B|}$$

The output has range $[0, 1]$. The Jaccard distance is the complementary of the index:

$$\delta_J(A, B) = 1 - J(A, B) = \frac{|A \cup B| - |A \cap B|}{|A \cup B|}$$

5.3 Curse of Dimensionality

Is easy to reach a very high-dimensional space, where only few dimensions are non zero. Data tends to be more sparser as the number of dimension grows. For example, the Euclidean distance of two points is high if they are far in at least one dimension (one of the dimension $x_i - y_i$ dominates the formula). The higher the number of dimensions, the higher the probability that this occurs.

We take an unit-length (length 1 in every dimension) hypercube H with d dimensions and N points randomly in the cube. We choose a specific point p and we want to create a smaller cube h centered in p that contains the k nearest point to p . The cube has length l in every dimension and every points has distance smaller than $\frac{l}{2}\sqrt{d}$ (the diagonal of the small cube).

The volume of the smaller hypercube is $V_h = l^d$ and contains $\frac{k}{N}$ points, so:

$$l^d \simeq \frac{k}{N} \implies l \simeq \left(\frac{k}{N}\right)^{\frac{1}{d}}$$

With this formula we can clearly see that the length of the hypercube h grows exponentially as the number of dimension grows.

This is caused by the edges of the hypercube, with more dimensions the probability to not be on the edge in at least one dimension is low. Let ϵ define the edge, the probability of any point to not be on the edge with 1 dimension is:

$$P(\text{not edge}) = (1 - 2\epsilon)$$

With more dimensions the probability become smaller and with d dimensions:

$$P(\text{not edge}) = (1 - 2\epsilon)^d$$

This problem can be resolved by assuming that in the real world the data is not random, but have a pattern underneath. The **Manifold hypothesis** suggests that high-dimensional data lie on low-dimensional subspace (manifold) embedded in the high-dimensional space. For example, for a digit recognition on 20×20 images taking in account all 400 dimensions model every random patter of pixel on the image. In reality the digits cover a small fraction of all the space and can be modelled in a much simpler way.

5.4 Dimensionality Reduction

As the number of dimensions grows the number of data points must grow exponentially to keep the same density.

To work with this problem we can have three approaches:

- Feature Engineering: use specific algorithm in the domain we are working in
- Make assumptions: for example independence of dimensions (replicate the spread for each dimension), smoothness (propagate the label to neighboring region), symmetry (invariance to the number of dimensions)
- Dimensionality Reduction: a pre-processing step to represent data with fewer dimensions, must preserve as much variance in the data as possible

Dimensionality Reduction as two main approaches: feature selection in which we pick a subset of the dimensions that are the best predictors and feature extraction in which we build a new set of dimensions as a linear combination of the original ones.

5.4.1 Principal Component Analysis (PCA)

PCA defines a new set of dimensions $k < d$ as follows:

1. Take the direction of the greatest variance of data
2. Take the perpendicular direction in every of the d dimensions
3. The top k directions become the new dimensions

To take the direction of the greatest variance we want to minimize the chance that 2 points that are far end up close in the new dimensions space.

We represent each data point x_i as a list of its components $(x_{i,1}, x_{i,2}, \dots, x_{i,d})$ and we associate a random variable X to each dimension, computing the mean for each one:

$$\mu_k = E(X_k) = \frac{1}{n} \sum_{i=1}^n x_{i,k}$$

We rewrite each data point x_i as:

$$x'_i = (x'_{i,1}, x'_{i,2}, \dots, x'_{i,d})$$

in which:

$$x'_{i,k} = x_{i,k} - \mu_k$$

Now we have a set of data points that have a 0-mean in each dimension. We create a covariance matrix K in which:

$$K[i, j] = \text{Cov}(X_i, X_j) = E((\underbrace{(X_i - E(X_i))}_{=0})(\underbrace{(X_j - E(X_j))}_{=0})) = E(X_i X_j)$$

The diagonal of K will have:

$$K[i, i] = E((\underbrace{(X_i - E(X_i))}_{=0})^2) = \text{Var}(X_i)$$

If we multiply K by a random vector e_1 we will obtain another vector e_2 that is near to the direction of greatest variance. If we keep multiplying K by the resulting vector it will converge to the direction of greatest variance, this vector is called **eigenvector**. If we multiply the matrix by an eigenvector the resulting vector is equal to the eigenvector scaled by a factor λ , called **eigenvalue**.

$$Ke = \lambda e$$

We want to find the eigenvector for K with the largest eigenvalue.

To compute eigenvectors we need to solve a linear equation:

$$Ke - \lambda e = 0 \implies (K - \lambda Id)e = 0$$

in which Id is the identity matrix. Any system of this type has a trivial solution $e = 0$, to have different solutions the matrix $K - \lambda Id$ must be non-invertible, so the determinant must equal to 0. So, to find the eigenvectors we have to:

1. Find the eigenvalues λ by solving the equation:

$$\det(K - \lambda Id) = 0$$

2. Find each eigenvector e by solving the equation:

$$Ke = \lambda e$$

The result will be a ratio between the components of the eigenvector, so there are infinite solutions. By convention, we take the eigenvector for which $\|e\| = 1$.

We now need to represent each data point x in the new coordinate system defined by the eigenvectors:

$$x'' = (x'^T e_1, x'^T e_2, \dots, x'^T e_k)$$

The T means the vector x' is horizontal in respect to e that is vertical so that the product is a single number.

To choose the k dimensions we set a threshold $t \in (0, 1)$ and we take all the dimensions until they reach that threshold in respect to the total variance:

$$\frac{\sum_{i=1}^k \lambda_i}{\sum_{i=1}^d \lambda_i} \simeq t$$

PCA has problem with large numbers, means that if one dimension has a extremely large number in respect to other dimensions it will be considered the most important dimension. This can be solved by normalizing each dimension to have 0-mean and 1-standard-deviation:

$$x'_{i,k} = \frac{x_{i,k} - \mu_k}{\sigma_k}$$

PCA also doesn't work if the data live in low-dimensional space that is not linear.

Example:

We have data points with 2 dimensions and the matrix K is:

$$\begin{bmatrix} 2 & \frac{4}{5} \\ \frac{4}{5} & \frac{3}{5} \end{bmatrix}$$

To compute eigenvalues we resolve the equation:

$$\det \left(\begin{bmatrix} 2 - \lambda & \frac{4}{5} \\ \frac{4}{5} & \frac{3}{5} - \lambda \end{bmatrix} \right) = 0 \implies (2 - \lambda)(\frac{3}{5} - \lambda) - (\frac{4}{5})(\frac{4}{5}) = \lambda^2 - \frac{13}{5}\lambda + \frac{14}{25} = 0$$

The resulting eigenvalues are:

$$\lambda_1 = \frac{13 + \sqrt{113}}{10} \simeq 2.36 \quad \lambda_2 = \frac{13 - \sqrt{113}}{10} \simeq 0.24$$

We need to resolve the system of equation using λ_1 to find the eigenvector e_1 :

$$\begin{bmatrix} 2 - \lambda_1 & \frac{4}{5} \\ \frac{4}{5} & \frac{3}{5} - \lambda_1 \end{bmatrix} \begin{bmatrix} e_1 \\ e_2 \end{bmatrix} = \lambda_1 \begin{bmatrix} e_1 \\ e_2 \end{bmatrix}$$

That is:

$$\begin{cases} 2e_1 + \frac{4}{5}e_2 = 2.36e_1 \\ \frac{4}{5}e_1 + \frac{3}{5}e_2 = 2.36e_2 \end{cases} \implies e_1 \simeq 2.2e_2$$

We take the eigenvector with $\|e\| = 1$, that is:

$$\begin{cases} \sqrt{(e_1)^2 + (e_2)^2} = 1 \\ e_1 \simeq 2.2e_2 \end{cases} \implies e \simeq \begin{bmatrix} 0.91 \\ 0.41 \end{bmatrix}$$

The same can be applied to λ_2 to find e_2 .

For every data point, the new coordinates will be:

$$x = \begin{bmatrix} x^T e_1 \\ x^T e_2 \end{bmatrix} = \begin{bmatrix} (x_1 - \mu_1)e_{1,1} + (x_2 - \mu_2)e_{1,2} \\ (x_1 - \mu_1)e_{2,1} + (x_2 - \mu_2)e_{2,2} \end{bmatrix}$$

The dimension that has the greatest variance is the first so we will take it to reduce the dimensions from 2 to 1.

5.5 Clustering

Clustering is the procedure to group a set of objects into classes of similar objects. Is an unsupervised learning technique because there is no external influence other than the data. Is a method of data exploration, a way to look for patterns in data.

Given a set of data points and a notion of distance, groups data in such a way that:

- Members of a cluster are similar to each other
- Members of different clusters are dissimilar to each other

The problem with clustering is with high-dimensional spaces, where almost all pairs are at the same distance.

5.5.1 Document clustering

To group document on the same topic we work in the space of words. Document with similar words probability talk about the same topic.

There are different ways to represent document in the space of words:

1. Set of words: we only count if a word appear or not in the document
2. Bag of words: we count the multiplicity of words in the document
3. Bag of n -grams: we count the group of n consecutive words that appear

In the set of words method two document are simply described as the set of all the words that appear in the document.

In the bag of words method a vocabulary is used, that contains the list of all the possible words contained in all documents. Each document is a vector in which the index i indicates the occurrences of the i -th word in the vocabulary.

If we define:

- $D = \{d_1, \dots, d_n\}$: the collection of n documents
- $V = \{w_1, \dots, w_m\}$: the vocabulary of m words
- $d_i = (f(w_1, i), \dots, f(W_m, i))$: the vector representing d_i
- $f : V \times D \rightarrow \mathbb{R}$: the function that maps every word to a value

We can define the function f in different ways:

- One-Hot binary weighting scheme:

$$f(w_j, i) = \begin{cases} 1 & w_j \text{ appears in } d_i \\ 0 & \text{otherwise} \end{cases}$$

- Term-frequency weighting scheme:

$$f(w_j, i) = tf(w_j, i)$$

in which tf calculates the number of times w_j appear in d_i .

- TF-IDF weighting scheme:

$$f(w_j, i) = tf(w_j, i) \cdot idf(w_j)$$

in which idf is defined as:

$$idf(w_j) = \log \left(\frac{n + 1}{n_j + 1} \right)$$

in which n_j is the number of documents containing the word w_j .

Depending on the representation used the similarity method that can be used is different:

- Set of words: Jaccard distance
- One-Hot bag of words: Euclidean distance
- TF or TF-IDF bag of words: Cosine similarity

5.5.2 Clustering algorithm: Partition-Based

The clustering algorithms are divided in different categories, but we will see only the **Partition-Based** algorithms.

There are two type of clustering:

- Soft clustering: a data point can belong to different clusters with probabilities that sum up to 1.
- Hard clustering: a data point can only belong to one cluster.

Partition-Based algorithms have as input a set of N data points and a number K of clusters and as output a partition of the data points into K clusters. The goal is to find the partition which optimizes a specific criteria. These algorithms solve the problem of hard clustering. Finding the best K is also part of the problem.

Every data point is assigned to only one cluster so the partition can be defined by a vector in which:

$$C[i] = k \iff i\text{-th data point is in cluster } k$$

The possible clustering outputs are K^N , which makes this problem NP-Hard. So, is impossible to find the global optimum but there are some effective heuristics like K-means to approssimate the optimal solution.

Let $\{x_1, \dots, x_N\}$ be the set of N data points and $\{C_1, \dots, C_K\}$ the set of K clusters. We define a data point θ_k that is the representative of the cluster C_k , means that it summarise the cluster with a single data point.

We define a Loss function L :

$$L(A, \Theta) = \sum_{n=1}^N \sum_{k=1}^K \alpha_{n,k} \delta(x_n, \theta_k)$$

in which Θ is the set of cluster representatives, δ the distance function and A is a $N \times K$ matrix where:

$$\alpha_{n,k} = \begin{cases} 1 & n\text{-th data point is in cluster } k \\ 0 & \text{otherwise} \end{cases}$$

We want to find A and Θ that minimize the L function.

A greedy algorithm to solve this problem is the **Lloyd-Forgy algorithm**, an iterative algorithm that is made of two steps: assignement and update. This algorithm doesn't guarantee to find the global optimum because it may stuck on a local optimum.

In the assignement step we try to minimize L fixing Θ , only modifying A . Minimize L fixing the representatives means assigning each data point to the closest representative using δ .

$$L(A|\Theta) = L(A; \Theta) = \text{function of } A \text{ parametrized by } \Theta$$

$$\alpha_{n,k} = \begin{cases} 1 & \delta(x_n, \theta_k) = \min_{j \in [1, K]} (\delta(x_n, \theta_j)) \\ 0 & \text{otherwise} \end{cases}$$

In the update step we minimize L fixing A , only modifying Θ . To do that we take the gradient (vector of partial derivatives) of L and set it equal to 0.

$$\nabla L(\Theta; A) = \left(\frac{\partial L(\Theta; A)}{\partial \theta_1}, \dots, \frac{\partial L(\Theta; A)}{\partial \theta_K} \right)$$

To make $\nabla L = 0$ all the partial derivatives must be set to 0. For a casual θ_j :

$$\frac{\partial L}{\partial \theta_j} = \frac{\partial}{\partial \theta_j} \left[\sum_{n=1}^N \sum_{k=1}^K \alpha_{n,k} \delta(x_n, \theta_k) \right]$$

When calculating the partial derivative of θ_j all the other θ_k are treated as constant, so the inner summation can be removed.

$$\frac{\partial L}{\partial \theta_j} = \frac{\partial}{\partial \theta_j} \left[\sum_{n=1}^N \alpha_{n,j} \delta(x_n, \theta_j) \right] = 0$$

We need to solve this equation for all θ_j independently, but we need to define the distance function δ before.

5.5.3 K-means

A special case of Partition-Based algorithm is **K-means** in which each cluster is represented by its centroid, the mean of all the data points assigned to that cluster. The idea is to construct clusters minimizing the Sum of Square Distance (SSD) within the cluster.

$$\theta_k = \mu_k = \frac{1}{|C_k|} \sum_{n \in C_k} x_n$$

The loss function L is:

$$L(A, \Theta) = \sum_{n=1}^N \sum_{k=1}^K \alpha_{n,k} \underbrace{(\|x_n - \theta_k\|_2)^2}_{\delta(x_n, \theta_k)} = \sum_{n=1}^N \sum_{k=1}^K \alpha_{n,k} (x_n - \theta_k)^2$$

In the assignement step the matrix A is modified in such a way that:

$$\alpha_{n,k} = \begin{cases} 1 & (x_n - \theta_k)^2 = \min_{j \in [1, K]} ((x_n - \theta_j)^2) \\ 0 & \text{otherwise} \end{cases}$$

In the update step we set the partial derivatives as 0:

$$\frac{\partial L}{\partial \theta_j} = \frac{\partial}{\partial \theta_j} \left[\sum_{n=1}^N \alpha_{n,j} (x_n - \theta_j)^2 \right] = 0$$

This formula is in effective equal to the one that calculates the mean:

$$\begin{aligned} \frac{\partial}{\partial \theta_j} \left[\sum_{n=1}^N \alpha_{n,j} (x_n - \theta_j)^2 \right] = 0 &\iff \sum_{n=1}^N -2\alpha_{n,j} (x_n - \theta_j) = 0 \iff 2 \sum_{n=1}^N \alpha_{n,j} \theta_j = 2 \sum_{n=1}^N \alpha_{n,j} x_n \\ &\iff \theta_j \sum_{n=1}^N \alpha_{n,j} = \sum_{n=1}^N \alpha_{n,j} x_n \iff \theta_j = \frac{\sum_{n=1}^N \alpha_{n,j} x_n}{\sum_{n=1}^N \alpha_{n,j}} = \frac{1}{|C_j|} \sum_{n \in C_j} x_n \end{aligned}$$

A run of the K-means algorithm goes as follows:

1. Specify the number of clusters K
2. Select K data points at random as the initial cluster representatives
3. Assignment step
4. Update step
5. Repeat assignment and update until a stopping criterion is met, for example fixed number of iteration, cluster assignment doesn't change (with threshold) or representatives don't change (with threshold).

Using an algorithm based on Lloyd-Forgy ensure that the result will converge (not necessarily to a global optimum) at some point, because a step can't be worst than the previous one. This is an instance of a more general **Expectation maximization (EM)** that is known to converge. The complexity of this algorithm, knowing that computing a distance between two d -dimension data points is $O(d)$, is:

- Assignment step: $O(KN)$ distance computations, so $O(KNd)$
- Update step: $O(N)$ average computations, so $O(Nd)$
- Total cost: $O(RKNd)$ with R iterations. The cost is linear to the number N of data points.

The convergence rate and clustering output depends on the initial random choice of representatives, to mitigate this problem multiple run can be done with different seeds and only pick the best one.

5.5.4 Similar algorithms to K-means

A different method to choose the initial representatives is K-means++, that spreads them:

1. The first representative is chosen at random
2. For each data point x , the distance between x and the nearest representative $D(x)$ is calculated
3. Choose a random data point to be a new representative with probability proportional to $D(x)^2$
4. Repeat step 2 and 3 until K representatives are chosen

K-means++ gives an upper-bound on how much the solution is worst than the optimal one, the clusters obtained are $O(\log n)$ worst than the optimal.

If we don't want to use Euclidean distance as the δ function, other distance can also be minimized by the means like Cosine distance (Euclidean distance on normalized data points) and Correlation (Euclidean distance on standardized data points). In the case of Manhattan distance the minimizer is the median (K-medians).

For an arbitrary δ the K-medoids algorithms works, choosing as representatives the data point closer to any other in the cluster. The drawback is the expensive cost that is $O(K(N - K)^2)$.

A variant of K-means for large datasets is Bradley-Fayyad-Reina (BFR) K-means. Is better for high-dimensional Euclidean space, but has strong assumption on the clusters: normally distributed around the representative and independence between direction.

5.5.5 Clustering Quality

The clustering quality can be evaluated using **internal evaluation** based on the data that was clustered. There are a lot of methods:

- Davies-Bouldin Index:

$$DB = \frac{1}{K} \sum_{i=0}^K \max_{j \neq i} \left(\frac{\sigma_i + \sigma_j}{\delta(\mu_i, \mu_j)} \right)$$

In which μ_k is the centroid of cluster C_k and σ_k is the average distance between all elements of cluster C_k and the centroid μ_k , calculated:

$$\sigma_k = \frac{1}{|C_k|} \sum_{x \in C_k} \|x - \mu_k\|^2$$

Smaller the number, better the clustering.

- Dunn Index:

$$D = \frac{\min_{1 \leq i < j \leq K} \delta(\mu_i, \mu_j)}{\max_{k \in [1, K]} \delta'(C_k)}$$

In which $\delta'(C_k)$ is the max distance between any pair of object in cluster C_k . The higher the value, better is the clustering.

- Silhouette Coefficent: For every data point x_i , two values are calculated:

$$a(x_i) = \frac{1}{|C_i| - 1} \sum_{x_j \in C_i} \delta(x_i, x_j)$$

$$b(x_i) = \min_{k \neq i} \frac{1}{|C_k|} \sum_{x_j \in C_k} \delta(x_i, x_j)$$

The coefficent is calculated:

$$s(x_i) = \begin{cases} 1 - \frac{a(x_i)}{b(x_i)} & a(x_1) \leq b(x_1) \\ \frac{b(x_1)}{a(x_i)} - 1 & a(x_1) > b(x_i) \end{cases}$$

The higher the value, the better.

The quality can be also evaluated using **external evaluation** knowing the ground-truth (optimal clustering that we want). There are a lot of methods:

- Purity:

$$\text{purity}(C_i) = \frac{1}{|C_i|} \max_{j \in [1, J]} n_{i,j}$$

$$\text{purity} = \frac{1}{K} \sum_{i=1}^K \text{purity}(C_i)$$

In which we define L_1, \dots, L_J a set of J labels and $n_{i,j}$ the number of items with label L_j in cluster C_i . The higher value, the better, but this method has a flows because with one data point in each cluster the purity is maximized.

- Rand Index:

$$\text{Rand} = \frac{TP + TN}{TP + TN + FP + FN}$$

In which:

- TP: true positive
- TN: true negative
- FP: false positive
- FN: false negative

This values are calculated taking every pair of object and comparing their relation in the clustering and in ground-truth:

	Same cluster in C	Different cluster in C
Same cluster in GT	TP	FN
Different cluster in GT	FP	TN

- Precisio Recall, F-measure:

$$P = \frac{TP}{TP + FP} \quad R = \frac{TP}{TP + FN}$$

$$F_1 = \frac{2 \cdot P \cdot R}{P + R}$$

We can balance the contribution of FN by using a parameter β :

$$F_\beta = \frac{(\beta + 1) \cdot P \cdot R}{\beta^2 P + R}$$

6

Recommender Systems

Online, the catalog of products or services is almost unlimited, so it is important to match the customers to the most appropriate product or service.

Recommender systems have a set of users $U = \{u_1, \dots, u_m\}$ and a set of items $I = \{i_1, \dots, i_n\}$. A user-item matrix is defined by a function $r : U \times I \rightarrow R$ in which R is a set of ordered ratings. Ratings can be discrete $\{0, 1, \dots, v\}$ or continuous $[0, 1]$.

Recommender systems have 3 problems:

- Data collection: gathering known ratings to populate the matrix. Can be done in an explicit way, asking people to rate items or in an implicit way, learning ratings from user actions.
- Rating prediction: extrapolate unknown ratings from known ones. The matrix is sparse because most people have not rated most items, also new items have no history.
- Recommendation evaluation: measure the performance of recommender methods.

Recommender systems are based on 3 approaches:

- Content-based filtering
- Collaborative filtering
- Hybrid

6.1 Content-based filtering

This method of recommendation is based on recommending items to a user u that are similar to other items rated high by u .

The first step consists in building an item profile for each item. This is done using a vector of numerical or categorical features.

The second step is to build user profiles. Let $I_u \subseteq I$ be the set of items rated by u and i_j the vector profile of item j . The simplest method could be creating a user profile using the average of the items rated:

$$u = \frac{1}{|I_u|} \sum_{i_j \in I_u} i_j$$

This doesn't take into account ratings and treat every item rated equal.

A better method is to do the mean of the ratings that the user u gave to items. For example if a user rated 5 items, each represented by a binary vector:

	Item 1	Item 2	Item 3	Item 4	Item 5
Rating	3	1	2	5	4
Vector	[1,0]	[0,1]	[0,1]	[1,0]	[0,1]

The ratings are normalized subtracting the user mean rating. In this case, the mean is 3.

	Item 1	Item 2	Item 3	Item 4	Item 5
Rating	0	-2	-1	2	1
Vector	[1,0]	[0,1]	[0,1]	[1,0]	[0,1]

The user profile will be:

$$u = \left[\frac{0+2}{2}, \frac{-2-1+1}{3} \right] = \left[1, -\frac{2}{3} \right]$$

Given a user profile u we can estimate the missing entry in the matrix for u . We recommend the top k items for which the cosine similarity between u and the item vector is higher. This is done in an iterative way:

$$\begin{aligned} A^1 &= \arg \max_i (\text{sim}(u, i) | i \in I - I_u) \\ A^2 &= \arg \max_i (\text{sim}(u, i) | i \in I - I_u - A^1) \\ &\dots \\ A^k &= \arg \max_i (\text{sim}(u, i) | i \in I - I_u - A^1 - \dots - A^{k-1}) \end{aligned}$$

The k recommended items are:

$$R_{u,k} = \bigcup_{j=1}^k A^j$$

Content-based filtering is good because it only needs the ratings of that user and doesn't have cold-start problem when an item has no ratings it can still be recommended.

It also has some cons like finding the appropriate item features, the fact that items outside user profile are never recommended, the ratings from other users are not counted and a cold-start problem for users, because there is no profile.

6.2 Collaborative filtering

The idea of this method of recommendation is based on recommending items to a user u based on preferences of users similar to u . There is no need to create user or item profiles.

Collaborative filtering has different approaches:

- Neighborhood-based (memory-based): divided also in:
 - User-based
 - Item-based
- Latent-factor-based (model-based)
- Hybrid (memory-model-based)

6.2.1 User-based Neighborhood

Evaluates a user preference for an item based on the ratings of "neighboring" users for that item.

Given a user u and an item i not rated by u , we want to estimate the rating $r(u, i)$. From the set of users we extract the k neighbours of u . Two users are similar if their ratings are similar, so we represent each user by the vector of ratings. To measure the similarity between two users we use the Pearson similarity, that is essentially the cosine similarity with the ratings normalized subtracting the average.

Let r_u be the ratings vector of user u and \bar{r}_u the average of that vector, the similarity between two users u and v is:

$$\text{Pearson}(r_u, r_v) = \frac{(r_u - \bar{r}_u) \cdot (r_v - \bar{r}_v)}{\sqrt{(r_u - \bar{r}_u)^T \cdot (r_v - \bar{r}_v)} \times \sqrt{(r_v - \bar{r}_v)^T \cdot (r_u - \bar{r}_u)}} = \text{cosine}(r_u - \bar{r}_u, r_v - \bar{r}_v)$$

The neighbours of user u are:

$$U^k = \arg \max_{U' \subseteq U - \{u\}, |U'|=k} \sum_{u' \in U'} \text{sim}(u, u')$$

We define the set of items rated by neighbours of u :

$$I^k = \{i \in I \mid \exists r_{u',i} \wedge u' \in U^k\}$$

To aggregate the neighbours ratings we can do a plain average or a weighted average:

$$\begin{aligned} r_{u,i} &= \frac{1}{k} \sum_{u' \in U^k} r_{u',i} \\ r_{u,i} &= \frac{1}{k} \sum_{u' \in U^k} r_{u',i} \cdot \text{sim}(u, u') \end{aligned}$$

The user-based recommender has 3 main issues:

- Sparsity: system performs bad when there are many items but few ratings
- Efficiency: computing similarities between all pairs of users is expensive
- Aging: user profiles change quickly and the system must be recomputed

6.2.2 Item-based Neighborhood

Systems have typically more users than items and each items have more ratings than each user, so the item rating average is more stable over time. This approach doesn't suffer from aging so the model doesn't need to be recomputed frequently.

From the set of items we extract the k neighbours of i . Two items are similar if users who rated them are similar, so we represent each item by the vector of ratings.

Let r_i be the ratings vector of item i and I_u the set of items rated by u . The neighbours of item i are:

$$I_u^k = \arg \max_{I'_u \subseteq I_u, |I'_u|=k} \sum_{i' \in I'_u} \text{sim}(i, i')$$

To aggregate the neighbours ratings we can do a plain average or a weighted average:

$$\begin{aligned} r_{u,i} &= \frac{1}{k} \sum_{i' \in I_u^k} r_{u,i'} \\ r_{u,i} &= \frac{1}{k} \sum_{i' \in I_u^k} r_{u,i'} \cdot \text{sim}(i, i') \end{aligned}$$

6.2.3 Comparison between User-based and Item-based

In general, item-based works better than user-based because the item vector is less sparse because there are more users than items.

Computing the k most similar users or items is the most expensive step, so this should be pre-computed offline. To do that we can approximate the k -nearest neighbours by using **Locality-Sensitive Hashing (LSH)**.

6.2.4 Latent-factor

This method tries to predict ratings by representing both items and users with a number of hidden factors created from observed ratings. The predicted rating of an user to an item would equal the product between the item vector and the user vector.

The most successful latent factor models are based on matrix factorization (MF). Both items and users are mapped in a latent factor d -dimensional space.

Let x_u be the vector representing user u and w_i the vector representing item i , the predicted rating of item i by user u will be:

$$r_{u,i} = \sum_{j=1}^d x_{u,j} \cdot w_{i,j}$$

The difficult part is to compute the vector for each user and item. Using a dataset D and the observed rating matrix R , to compute the latent factors representing x_u and w_i we need to minimize this loss function:

$$L(X, W) = \sum_{(u,i) \in D} (r_{u,i} - x_u \cdot w_i)^2 + \lambda \left(\sum_{u \in D} \|x_u\|^2 + \sum_{i \in D} \|w_i\|^2 \right)$$

For future simplification we multiply L by $\frac{1}{2}$.

There are two possible optimization methods, the first one is **Stochastic Gradient Descend (SGD)**. In SGD, for each instance (u, i) we compute the gradient of the loss with respect to x_u and w_i :

$$\begin{aligned} \nabla L(x_u; w_i) &= \frac{1}{2}[-2(r_{u,i} - x_u \cdot w_i)w_i + \lambda x_u] = -(r_{u,i} - x_u \cdot w_i)w_i + \lambda x_u \\ \nabla L(w_i; x_u) &= \frac{1}{2}[-2(r_{u,i} - x_u \cdot w_i)x_u + \lambda w_i] = -(r_{u,i} - x_u \cdot w_i)x_u + \lambda w_i \end{aligned}$$

We define $e_{u,i} = r_{u,i} - x_u \cdot w_i$ and at each iteration we update the value x_u and w_i as follows:

$$\begin{aligned} x_u^{t+1} &= x_u^t + \eta(e_{u,i} \cdot w_i^t - \lambda x_u^t) \\ w_i^{t+1} &= w_i^t + \eta(e_{u,i} \cdot x_u^t - \lambda w_i^t) \end{aligned}$$

SGD has different properties:

- Incremental: updates occur sample by sample
- Scales well with sparse data
- Order matter
- The learning rate (η) must be tuned carefully
- Converges to a local optimum

The main problem is the number of parameters to optimize, that is $d(m + n)$ and can get very large, requiring other optimizer.

The second method to optimize the loss function is **Alternative Least Squares (ALS)**. In ALS the two matrices X and W are alternately considered constant while updating the other one. This makes the function quadratic (convex) and can be solved optimally.

Assuming W constant the gradient in respect to the user vector is:

$$\nabla L(x_u; w_i) = - \sum_{i \in D} (r_{u,i} - x_u \cdot w_i) w_i + \lambda x_u$$

And the gradient in respect to the item vector, assuming X constant is:

$$\nabla L(w_i; x_u) = - \sum_{u \in D} (r_{u,i} - x_u \cdot w_i) x_u + \lambda w_i$$

Setting both equal to 0 we find that:

$$\begin{aligned} x_u &= (W^T W + \lambda I d)^{-1} \cdot W^T \cdot r_u \\ w_i &= (X^T X + \lambda I d)^{-1} \cdot X^T \cdot r_i \end{aligned}$$

The algorithm works as follows:

1. Initialize X and W randomly
2. Fix W and solve for X (users)
3. Fix X and solve for W (items)
4. Repeat 2 and 3 until convergence

ALS has some properties:

- Deterministic update: each step is a minimizer
- No learning rate to tune
- Parallelizable
- Converges in fewer iterations than SGD but each iteration is computationally heavier

6.2.5 Bias

The observed ratings variation depends on biases associated with users or items. Some users could rate higher than others and some items could be rated higher than other.

The bias of a rating $r_{u,i}$ is defined as follows:

$$b_{u,i} = \mu + b_u + b_i$$

In which μ is the average rating, b_u is the deviation of the user u from the average and b_i the deviation of the item i from the average.

If we include bias, the estimated rating function becomes:

$$r_{u,i} = x_u \cdot w_i + b_{u,i}$$

And the loss function to optimize becomes:

$$L(X, W) = \frac{1}{2} \left[\sum_{(u,i) \in D} (r_{u,i} - (x_u \cdot w_i + b_{u,i}))^2 + \lambda \left(\sum_{u \in D} \|x_u\|^2 + \sum_{i \in D} \|w_i\|^2 + \sum_{u \in D} (b_u)^2 + \sum_{i \in D} (b_i)^2 \right) \right]$$

This formula can still be solved with ALS.

Collaborative filtering has 3 main issues:

- Cold start: new items or users don't have enough data to make recommendations
- Scalability: million users and items require too much computational power
- Sparsity: most items are not rated by users

6.3 Evaluation Metrics

To evaluate the recommendation generated by the system there are a lot metrics, both online and offline.

A metric called RMSE focus on the accuracy, penalizing the diversity and not taking in account the order of the recommendations:

$$\text{RMSE} = \frac{1}{|D|} \sqrt{\sum_{(u,i) \in D} (r_{u,i} - \hat{r}_{u,i})^2}$$

In which $\hat{r}_{u,i}$ is the predicted rating while $r_{u,i}$ the real one.

6.3.1 Precision and Recall

Like the clustering evaluation we define two values P and R :

$$P = \frac{\#\text{relevant item recommended}}{\#\text{items recommended}} \quad R = \frac{\#\text{relevant item recommended}}{\#\text{items actually relevant}}$$

For example, if a system generates 5 recommendation, 3 are relevant and the system only finds 2 relevant item:

$$P = \frac{2}{5} \quad R = \frac{2}{3}$$

This metric doesn't care about order, to do that we define $P@k$ as the precision taking in account only the first k recommendations. If the system must return N items and the number of actually relevant items is $|Rel|$, we define the average precision AP as:

$$AP@N = \frac{1}{|Rel|} \sum_{k=1}^N P@k \times 1_{Rel}(k)$$

in which $1_{Rel}(k)$ is a function:

$$1_{Rel}(k) = \begin{cases} 1 & k \in Rel \\ 0 & \text{otherwise} \end{cases}$$

AP is computed on a single user, so we define the Mean Average Precision (MAP) as:

$$MAP@N = \frac{1}{|U|} \sum_{u \in U} AP@N(u)$$

6.3.2 Personalization

We need a way to know if the system is recommending many of the same items to different users. The personalization is defined as the dissimilarity between user's lists of recommendation. An high personalization indicates that the recommender system is able to provide a highly personalized experience.

As example, we have 3 users with this recommendation:

$$u_1 = [A, B, C, D] \quad u_2 = [A, B, C, E] \quad u_3 = [A, B, F, G]$$

We compute the matrix containing the cosine similarity between each pair of users:

1	0.75	0.58
0.75	1	0.58
0.58	0.58	1

The personalization score is 1 minus the average of the upper triangle:

$$\text{Personalization} = 1 - \frac{0.75 + 0.58 + 0.58}{3} = 0.36$$

7

Analysis of Large Graph

7.1 Definition of graph

We formalize the graph $G = (V, E)$ in which:

- $V = \{v_1, \dots, v_n\}$ is the set of nodes
- $E \subseteq V \times V = \{(v_i, v_j) \in V \times V | v_i \neq v_j\}$ is the set of edges

We define the degree of a node as:

$$\deg(v) = |\{u \in V | (u, v) \in E\}|$$

In the case of directed graph we can define two types of degrees:

$$\text{in_deg}(v) = |\{u \in V | (u, v) \in E\}|$$

$$\text{out_deg}(v) = |\{u \in V | (v, u) \in E\}|$$

7.1.1 Representing a graph

Graphs can be represented in different ways:

- Adjacency matrix
- Adjacency list
- Edge list

With an adjacency matrix we create a $n \times n$ matrix M in which $|V| = n$. In the matrix we have that:

$$M[i, j] = 1 \iff \exists(v_i, v_j) \in E$$

This method is intuitive and ready for mathematical manipulation but is space inefficient and hard to compute.

The adjacency list method consist in having a linked list for every node v_i containing all the other node for which exists $(v_i, v_j) \in E$. This is a compact representation and is easy to compute calculation with outgoing link but is hard to compute calculation over ingoing links.

The edge lists is simply a list of all the edges (v_i, v_j) . Is a waste of space but is the best for edge insertions.

7.2 Web search engine

7.2.1 Content similarity method

The first idea to implement a web search engine was to find relevant document using the content similarity to the query. The documents were mapped to the word space and the words were scored based on the importance in the document (using the TF-IDF method). This traditional way of Information Retrieval (IR) suffers from two main problems:

- Information overload: some queries have millions of relevant web pages that are difficult to rank.
- Trustworthiness: the similarity can be tricked by creating a lot of spam documents.

An important and trustworthy page is a page pointed by many other pages and trustworthy pages should point to each other. So we can rank the pages based on their connectivity in the web graph.

7.2.2 Scale-Free network

In the web graph the degree distribution of the nodes follows a power law where most nodes have few links and few nodes have a large number of links. Graphs with this behavior are defined Scale-Free Networks.

In a Scale-Free Network the percentage of nodes having k connections follows a power law distribution:

$$P(\deg = k) = p(k) = \alpha k^{-\gamma}$$

This power law remains unchanged, except for a multiplicative constant, independently to the scale we look at the network.

A probability distribution $p(x)$ is scale-free if:

$$\exists g(c) \implies p(cx) = g(c)p(x)$$

For the previous power law:

$$p(cx) = \alpha(cx)^{-\gamma} = c^{-\gamma} \alpha x^{-\gamma} = g(c)p(x)$$

In which $g(c) = c^{-\gamma}$.

A scale-free network can be constructed by adding node to an existing network. The links are created following a preferential attachment where the probability of the new node is linked to an existing node i is based on the number of links k_i the node already has:

$$p(\text{new node linked to } i) = \frac{k_i}{\sum_{\forall j} k_j}$$

On a log-log scale (a scale where both x and y are log based) the power law looks like a straight line.

$$\log(p(k)) = \log(\alpha k^{-\gamma}) = \log(a) + \log(k^{-\gamma}) = \underbrace{\log(a)}_{\text{constant}} + \gamma \log(k)$$

7.2.3 Computing node importance

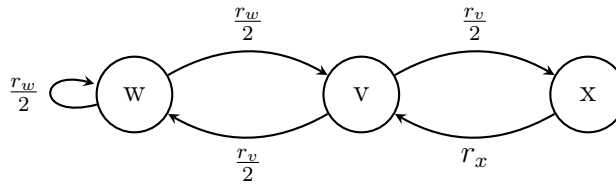
A possible link analysis approaches to compute web page importance is the PageRank. The PageRank intuition is based on a recursive definition where more incoming links a page has more important is the page and the links from important pages count more.

With the web graph $G = (V, E)$, let O_v be the set of pages linked by v and I_v the set of pages linked to v . The importance of a page v is calculated by summing the incoming links:

$$r_v = \sum_{u \in I_v} \frac{r_u}{|O_u|}$$

The interpretation can be seen in two ways, an algebrical way and a probabilistic way.

The algebrical way consist in solving a system of equations called flow equations. For example given this graph:



The system of flow equation to solve is:

$$\begin{cases} r_v = \frac{r_w}{2} + r_x \\ r_w = \frac{r_v}{2} + \frac{r_w}{2} \\ r_x = \frac{r_v}{2} \end{cases}$$

There are infinite solution to this system so we add a constraint to only have one solution, this constraint is the equation:

$$r_v + r_w + r_x = 1$$

With possibly billions of equation we need a new formulation, the matrix formulation. We use a stochastic matrix M of size $n \times n$ in which n is the number of pages. For every cell of the matrix:

$$M[w, v] = \begin{cases} \frac{1}{|O_w|} & w \text{ links to } v \\ 0 & \text{otherwise} \end{cases}$$

The matrix is column stochastic, which means that every column sums up to 1, assuming that every page has at least one outgoink link.

We define the rank vector r , a vector with an entry for each page where all the ranks sums up to 1. With the previous calculation of a rank:

$$r_v = \sum_{u \in I_v} \frac{r_u}{|O_u|}$$

The vector multiplied by the matrix must be equal to itself:

$$r = Mr$$

So r is the eigenvector of the matrix M associated with the eigenvalue 1.

The probabilistic way uses a random walk interpretation where the surfer starts in a random position and at each step follows one of the outgoing link at random. This random walk can be described as a Markov Chain M with the same definition as the stochastic matrix described before.

Let X the discrete value random variable that has $X = w$ if the server is on the page w . We can define a stochastic vector associated to X :

$$p(t) = [P(X_t = 1), \dots, P(X_t = w), \dots, P(X_t = n)]$$

The transition probability of moving in the next state depends only on the present state:

$$P(X_{t+1} = v | X_1 = x_1, X_2 = x_2, \dots, X_t = w) = P(X_{t+1} = v | X_t = w)$$

To know the probability of the surfer being on page v at time $t + 1$ we need to sum the probabilities of being on pages linked to v multiplied by the probability of going to the page v . For example if v has 2 incoming links from w and x :

$$P(X_{t+1} = v) = P(X_t = w) \cdot \frac{1}{|O_w|} + P(X_t = x) \cdot \frac{1}{|O_x|}$$

So, we can define the stochastic vector as:

$$p(t+1) = Mp(t) = M^t p(0)$$

At certain point the stochastic vector doesn't change anymore if multiplied by M , this is called the **stationary distribution**.

The two vectors p and r are equal, but we don't know if they exists and if they are unique.

Perron-Frobenius theorem

Given a column stochastic matrix M with all positive entries:

- $\lambda = 1$ is an eigenvalue of M and is the largest
- There exists a unique eigenvector r associated with eigenvalue $\lambda = 1$ with that sums up to 1

Given a column stochastic matrix M with all positive entries, there is a unique vector p such that for any $p(0)$ we have that:

$$\lim_{t \rightarrow +\infty} p(t) = \lim_{t \rightarrow +\infty} M^t p(0) = p$$

7.3 Google PageRank

The matrix M could contain some 0 making it not strictly positive to apply the theorem. There are two main issues to solve:

- Dead end: pages with no outgoing links
- Spider trap: not every page is reachable

The problem with a dead end page v is solvable creating artificial links from v to every other node, each with the same probability $\frac{1}{n}$. This solution modifies the matrix M to be column stochastic:

$$M[w, v] = \begin{cases} \frac{1}{|O_w|} & w \text{ links to } v \\ \frac{1}{n} & \sum_{\forall v} M[v, w] = 0 \\ 0 & \text{otherwise} \end{cases}$$

To solve the problem with spider traps we create artificial links from every node to every other node with probability $\frac{1-d}{n}$ in which d is the dumping factor (0.85 in the Google formulation). With this solution we can create a matrix G that is strictly positive:

$$G = dM + \frac{1-d}{n} \begin{bmatrix} 1 & \dots & 1 \\ \vdots & \ddots & \vdots \\ 1 & \dots & 1 \end{bmatrix}$$

The matrix G is very difficult to compute so we need to rearrange the equation to be manageable. Taken a single page v :

$$\begin{aligned} r_v &= \sum_{\forall w} G[v, w] \times r_w = \sum_{\forall w} \left(dM[v, w] + \frac{1-d}{n} \right) \times r_w = \sum_{\forall w} dM[v, w] \times r_w + \sum_{\forall w} \frac{1-d}{n} \times r_w \\ &= \sum_{\forall w} dM[v, w] \times r_w + \frac{1-d}{n} \underbrace{\sum_{\forall w} r_w}_1 \end{aligned}$$

So, for a single page v :

$$r_v = \sum_{\forall w} dM[v, w] \times r_w + \frac{1-d}{n}$$

And for all the vector r :

$$r = dMr + \frac{1-d}{n} \begin{bmatrix} 1 \\ \vdots \\ 1 \end{bmatrix}$$

This vector can be computed in an iterative way where in each iteration:

1. $r(t+1) = dMr(t)$
2. $r(t+1) = r(t+1) + \frac{1-d}{n} \begin{bmatrix} 1 \\ \vdots \\ 1 \end{bmatrix}$

This vector represent the rank of all the pages on the web.