

Photonic Artificial Neural Networks: All-Optical Activation Function

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Introduction

The research field

Research in the field of artificial neural networks has become more and more popular in the last few decades as the computing power available to the average institution increased further. As a matter of fact, in the last two decades the calculation power which once belonged only to bulky, power hungry, and very expensive supercomputers, gradually became possible also for relatively more compact, efficient, and definitely cheaper servers and workstations.

This trend is dictated by the fact that artificial neural networks are usually simulated on conventional computers, which are on the other hand based on the von Neumann, or Princeton, architecture. Implementations of this general purpose architecture are able to do any logical computation with a series of instructions. However the downside of this suitability to generic computations is that it is often difficult to parallelize single operations and therefore certain tasks are inherently inefficient, both in energy and in time. Moreover another problem is that it requires explicit programming for each singular task

The formalization of the von Neumann architecture dates back to 1945 and is as old as the first attempt in artificial neural networks. Shortly after, research and industry efforts focused only on the development of the Princeton architecture, choosing it de facto as the primary design for computing devices.

In the recent years, help came from acceleration units known as Graphic Processing Units (GPU), which were developed for a completely different objective. These GPUs, which are nowadays being designed specifically to accelerate calculations of artificial neural network simulations, are able to carry out a restricted set of instruction compared to CPUs, but are much more efficient both in power and in time, because of their parallel execution.

Today's software implementations

Between the most famous examples of simulated neural network there certainly are the older Watson supercomputer from IBM and the more recent AlphaGo artificial intelligence. Both of them arose to popularity because they defeated us humans to our own games. Specifically the effort of Watson was aimed at prevailing the most strongest human contestants at *Jeopardy!*, an American television game, and succeeded in 2011. Watson was powered by a IBM supercomputer and used 0.22 MW of power. AlphaGO, nevertheless, is a deep neural network that in 2016, through reinforced learning algorithms, defeated Lee Sedol, considered the most formidable master of Go, a complicated board game.

Another interesting case is given by SpiNNaker, of the University of Manchester. It is composed by 10^4 neurons connected by more than 10^7 synapses, and is simulated on over 65 thousand 18-core ARM processors connected together. Unlike the other two, its main goals are to simulate several neural mechanisms, such as the operation of visual cortex.

Similarly to SpiNNaker, another effort in gaining better understanding of the human brain is *The Blue Brain Project*, which is leaded by École Polytechnique Fédérale de Lausanne in collaboration with over one hundred other international research institutions and it is funded by the European Union. This project is simulated on IBM Blue Gene supercomputers, each fitted with more than 100 thousand processors and consuming several MW of power.

Present objectives and future trends

We can distinguish two principal objectives in this research field. The first one is a technological objective and is that of developing a more powerful computational instrument. The second objective is instead a topic of more fundamental research: to use the tools of artificial neural networks to better understand biological neural networks such as our brain.

If the comprehension of the complex mechanism at the base of our brain has always been a very arduous assignment, the tasks that artificial neural network are facing become increasingly difficult while time passes. Whereas in the past years public and private research groups trained themselves with "easier" problems, such as television or board games, today they are concentrating their efforts on more challenging goals. Probably the most clear example of this tendency is given by the research on autonomous driving, where artificial neural networks are used primarily for real-time processing of data from several sensors..

A great number of businesses formed in recent years around these problems and many other will probably do the same for similar areas. It is very likely that to achieve their objectives, even more powerful and complex artificial networks will be required. Therefore more processors will be developed and put together to achieve impressive computing power. However, one cannot expect to increase indefinitely the power, without significant improvements in efficiency. Hence alternatives are sought.

Overview of current alternatives to simulations

Apart from the vast majority of the research on neural networks, which is being carried out with simulations on powerful processors, a smaller portion of research is focused on building neural network physically. Great efforts are made nowadays by many research institutions and companies to develop original computing architectures that allow artificial neural networks to run physically, instead of being simulated on conventional computers.

The reason why considerable improvements can be expected is that the path of designing hardware devices ad hoc for neural network computation has not yet been covered comprehensively. Majors performance or efficiency enhancements are coming from the fact that those new architectures are conceived with parallel execution of specific operations in mind. However, being designed for precise tasks makes them unsuitable for generic computation, which could in principle make them less easy to modify when needed, unlike simulations on conventional computers.

Many of these use the electronics framework, mainly because of the many benefits of the mature silicon technology and its CMOS-compatible production chain. Specifically footprint and efficiency of CMOS devices has not yet stopped to grow since its discovery, in the past century. However, a relatively small portion of these works also inquired simpler network fabrication with organic electronic materials, achieving remarkable results.

Among the most interesting projects which uses electronics there are TrueNorth of IBM and Neurogrid of Stanford University. The former is part of a long-term research project funded by DARPA and has the aim of building an artificial neural network with the capabilities of brains of small mammals, such as cats or mice. In 2014 IBM showed a chip, named TrueNorth, containing 1 million artificial neurons and 256 millions synapses. This network, relying on more than 5 billion transistors organized on an area of 4.3 cm^2 , consumed only 60 mW. The researcher also proved that connecting 48 of this chips together they obtained the equivalent of the brain of a mouse.

Similarly, the intention behind Neurogrid chip is to explore numerous hypothesis concerning mammalian brains, specifically about the inner mechanisms of operation of the cerebral cortex. Stanford's implementation reaches the line of 1 million neurons, while only requiring 5 W of power consumption. Moreover the communication lines between single neurons, the synapses, are provided by FPGAs and banks of SRAM.

Besides electronics, other kind of physical implementations are attempted from research teams. Probably the most intriguing example is given by researches in photonics, which attempts to match electronics performance by overcoming its intrinsically weaknesses, such as power consumption. Photonics has yet to reach achievements comparable to electronics, mainly due to the embryonic state of its technology.

My work

My work has two related objectives. The primary aim is to implement a proof of concept for a fundamental component of artificial neural networks: the neuron's activation function. I plan to do so by employing the framework made available by integrated photonics. The second intent is to bring together two until now distant fields of research: physics and information technology, specifically photonics and machine learning. In fact generating knowledge transversal to the two research fields is at least as much important as the primary objective

The idea at the base of our physical implementation is to employ integrated devices already manufactured to create a proof of concept for the activation function. The choice of integrated photonics is motivated by the fact that, unlike the electronic counterpart, does not suffer from the Joule effect.

Therefore in this work I will study a particular device, a microring resonator, and characterize its response to different inputs, for several working conditions. This device can process information because it has, under certain circumstances, a nonlinear response function to its inputs. Moreover, the same device might in principle be used in another important part of artificial neural network nodes: the weighted sum. However, since this last point has already been discussed in literature and it requires less effort to produce acceptable results, it will not be at the center of this study.

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Since the aim is to create a proof of concept with structures not specifically designed, I don't expect extraordinary performance both in speed and in efficiency. These aspects will be discussed in the conclusive words and will be studied to their fullest only in future works.

Furthermore, regarding the second objective, my work is part of a bigger project, BACUKP, with a wider scope and a more ambitious goal. It attempts to connect three fields of research, normally quite distant from each other: physics, computing technology, and biology. The main idea is to use the integrated photonics framework (physics) to build an artificial neural network (computing technology) on chip, where one could grow and, at least partially control, real neurons (biology). The aim is to develop a methodologies and devices that will allow us to study neural activity from a bottom-up perspective.

Structure of the thesis

The first chapter of this thesis is intended as an introduction to the world of artificial neural networks. After a brief historical summary, I describe the main aspects of neural networks and their working principles. Then I focus on the description of the simplest type of artificial networks, the feedforward network, and finally an overview on other kind of networks. Since the aim of this thesis is to tackle a specific problem, this chapter is not meant to be comprehensive foundation or compendium on the matter. Later, the chapter is concluded with an introductory description of the language Python and the library PyTorch, used in this work for artificial network simulations.

The second chapter introduces the physics which I will study. Again, after a short explanation on the basic devices used in the field, I will describe in depth the device studied for the

task. Having clarified the physics underneath, I ought to explain how I am going to use this physics to implement (part of) an artificial neural network. I will address these topics both from the theoretical point of view and from the numerical one, with simulation of the specific (simplified) system.

The third chapter is dedicated to the description of the experimental work. At the beginning the problem of the selection of the sample is discussed, along with the definition of the device chosen. After, I characterize the different behaviors with detailed data. Finally, an explanation on how the device will be used in the neural network viewpoint and the corresponding tests of operation.

Chapter 1

Artificial Neural Networks

Artificial Neural Networks (ANNs) are computational systems which elaborate information likewise biological neural networks (animal brains). These systems modify their behavior through a learning process and improve their accuracy in a certain task. The underlying idea is that biological neural networks are superior in performance to computers and extremely more efficient in difficult tasks such as classification (e.g. image recognition) and prediction (e.g. pattern recognition).

Mathematically speaking, ANNs are a collection of nodes, each of them elaborate the information and is connected to the other in a certain amount. Artificial networks can be either simulated on computers or physically built on specific hardware designed ad hoc. At this time, ANNs are mainly implemented by simulations, however the technological progress is withheld by limitation in computing power and efficiency. Training a complex artificial neural network with computers at the state of art might take even weeks. Thus research on hardware architectures is surging: digital, analog, electrical, and optical devices are being developed.

1.0.1 History

It is widely acknowledged that the opening work of this research field was made in 1943 by Warren McCulloch and Walter Pitts, a neurophysiologist and a mathematician respectively. In their research work, they described the operating mechanism of biological neurons by modeling a simple electronic circuit [1].

The following important work was made by Donald O. Hebb, who hypothesized the neural plasticity in 1949 [2]. He pointed out the fact that neural pathways are strengthened each time they are used, a concept known as *Hebbian learning*.

During the late 1950s and the early 1960s, as computers became more powerful, many promising works were published. Some simulated operation of artificial networks on calculators, e.g. Farley and Clark [3] and Rochester [4]. Other produced circuitry that implemented on hardware such networks, e.g. Rosenblatt [5], [6]. However, despite the early successes of neural networks, the traditional computing architecture (von Neumann architecture) was chosen as the preferred computing architecture. As a result, funding and therefore also research diminished drastically in the following decades.

The reason this happened, probably, was due to several concurring facts. First of all, in the same time period a research paper wrongly suggested that there could not be any multiple-layered neural network. Moreover, the early successes of some works on neural networks pointed to an overestimation of the artificial neural networks potential, also held back by the technological capacity of the time. Finally, important questions of philosophical nature come to light, such as the fear fueled debate on the impact on our society of a class of computers able to think. This very controversy, i.e. artificial intelligence (AI) problem, is still a discussed topic today [7].

Sometime during the 1980s the interest for this computing method was reinvigorated. The main stimulation was probably given by a number of works which suggested methods to implement multi-layered networks distributing pattern recognition errors through the all the layers in the network. This method is now called *backpropagation*.

In today's research and technology, artificial neural networks are used in numerous applications. However, the development is made slowly, due to technological limitation in computational power of present processors.

1.0.2 Comparison with conventional computers

1.1 Basis of Neural Networks

A neural network is a collection of processing elements, or nodes, interconnected in an arbitrary topology. From its input nodes, the network accepts information, which will propagate into the inner nodes through the interconnections and will get elaborated at each node. At the end of the network, there will be a number of output nodes, with the task of reading a portion of the inner nodes. The inner nodes are also called hidden, because they are not meant to be accessible to the external world. A generic scheme of such network is shown in Figure 1.1 on this page.

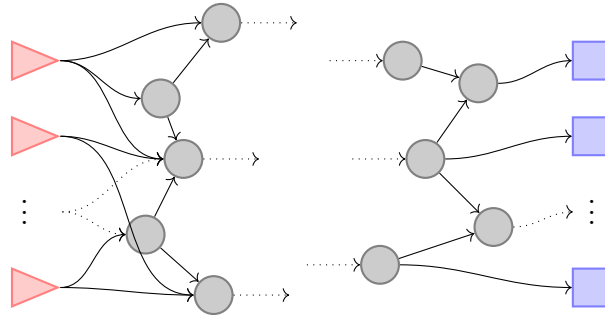


Figure 1.1: Generic scheme of a neural network. Triangles (red) are input nodes, circles (grey) are inner nodes, and squares (blue) are output nodes. Interconnections among nodes are represented by arrows: continuous when both elements are drawn, and dotted otherwise. I chose a non-standard representation to emphasize the nonlinear nodes, which will use a nonlinear activation function.

Each node can operate in the same way of the others or in a completely different manner, depending on the type of neural network. The operation of nodes resembles that of animal neurons: various input gets collected and elaborated together to obtain an output, which will become one of the many inputs for subsequent neurons/nodes. Specifically, the most used model for neurons is the McCulloch–Pitts (MCP) neuron. It is divided into two parts, as shown in Figure 1.2 on the next page: the first part is a weighted sum of the inputs, while the second part is given by the so called activation function.

The node is described mathematically by Equation 1.1

$$y = f_a \left(w_0 + \sum_{i=1}^n w_i x_i \right), \quad (1.1)$$

where f_a is the activations function, evaluated on the sum of the input x_i weighted with w_i , plus a bias w_0 .

Each node accepts values at its inputs and produces an output accordingly. However, the output depends also on the node's parameters: the weights and the bias, which are usually

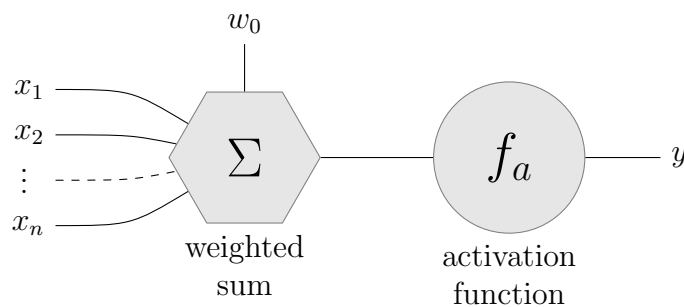


Figure 1.2: Generic node representation. x -values are inputs, y -values are outputs, w_0 is the bias.

changed outside the operative phase of the neural network, as I will explain in Section 1.2 later on the following page.

Moreover it is mandatory for the activation function $f_a(\cdot)$ to be nonlinear, because otherwise a collection of nodes will result in just a weighted sum of its inputs. Two examples of nonlinear function are shown below in Figure 1.3.

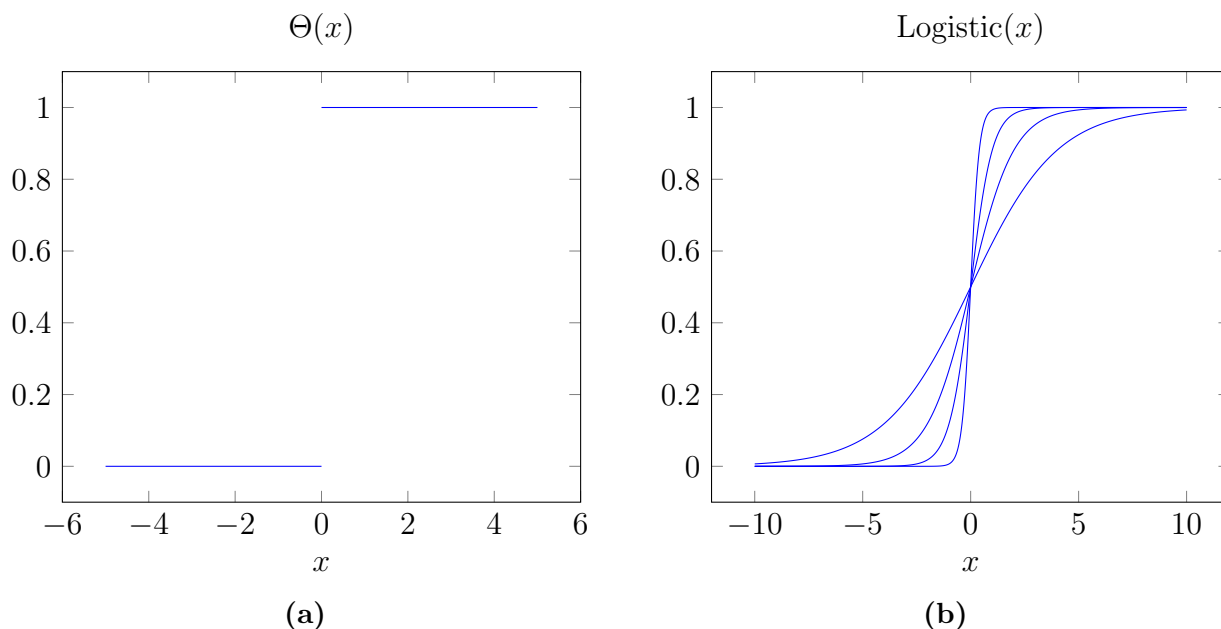


Figure 1.3: Examples of activation function: (1.3a) is the well-known step function, or Heaviside Θ , (1.3b) depicts a few functions from the family of the Logistic functions.

One can distinguish at least three type of nodes in every neural network: input, inner/hidden, and output nodes. Input nodes take one input value, from the outside of the neural network, and pass it on to the inner nodes unchanged. Inner/hidden nodes take many inputs and generate an output through the activation function. Output nodes, similarly to input nodes, take one input value, from the inside of the NN, and pass it on to the outside.

This is not the orthodox description, however it is consistent with the idea of functional *black box*, in which input and output are the only visible nodes, while the other are hidden inside. Therefore I can easily distinguish the nodes which will use a nonlinear activation function.

WHAT ARE NEURAL NETWORK GOOD FOR?

WHEN DO I TALK ABOUT THAT?

E.G. NN ARE GOOD FOR PATTERN RECOGNITION.

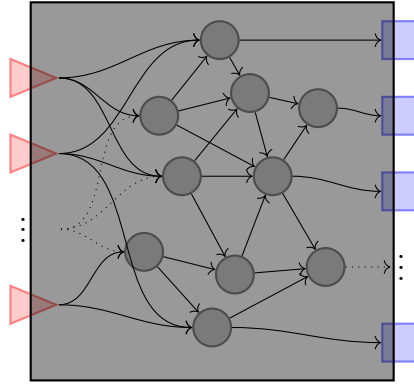


Figure 1.4: Representation of the black-box concept

1.2 Working Principles of ANNs

Because of its topology, each neural network will behave in a different manner from other neural networks with diverse, or even similar, arrangements of nodes. Moreover the same neural network will perform a certain task better or worse also depending on how inputs are weighted at each hidden node, and normally those parameters are initialized with a random value at the creation of the network. For this reason, before a neural network is considered ready to perform a task, it usually must go through three training stages: learning phase, validation phase, and testing phase. Every one of these stages is meant to prepare the network to work as required from the designer.

1.2.1 Learning Process

During the learning process the neural network is run on a set of known inputs x , each paired with its correct answer y , or target, in a second set of data. The neural network will produce at the output a third set \hat{y} , which should be as close as possible to the correct answers, when the network works properly. However this happens rarely, if ever, and a change in the way data is elaborated becomes necessary. The usual ?? way is to keep the same topology, but tweak the weights that connect the hidden nodes together. On account of this need, one needs to quantify the distance of the predicted result of the artificial neural network from the correct answer. This is made by means of the loss function.

Loss function

The loss function $L(y, \hat{y})$ evaluates the difference between the predicted and the correct answer. Usually, this quantity is linked to the geometrical distance between the predicted output and the target $|\hat{y} - y|$.

The most common loss function is the *mean-square error*. Assuming to have an input set of N examples paired with the same number of targets, and that the outputs and the targets are composed by C values, or classes, the function becomes:

$$L(y, \hat{y}) = f_{MSE}(y, \hat{y}) = \frac{1}{N} \sum_{n=1}^N \sum_{i=1}^C (\hat{y}_{n,i} - y_{n,i})^2, \quad (1.2)$$

where each example in the set is subtracted to its target and then squared. Finally the mean of all squares gives the expected result.

Another commonly used function is the cross-entropy loss (also known as negative log likelihood),

$$L(y, \hat{y}) = f_{CEL}(y, \hat{y}) = -\frac{1}{N} \sum_{n=1}^N \sum_{i=1}^C y_{n,i} \log(\hat{y}_{n,i}), \quad (1.3)$$

which expects positive values at the input. Hence the error $-y \log(\hat{y})$, quantified for each element in each example, is always a positive number. The mean over all examples in the set returns the results.

Alternatively, variations of the previous methods are given by not taking the sum of the examples in place of the mean, or by calculating a loss function for each example instead of evaluating it for the whole set.

Weights Update Process

The weights update process is a difficult task, and probably the most computationally expensive one in running a neural network. There is a variety of methods to choose from, depending on the type of artificial network and the resources available.

A widely used algorithm is the gradient descent, from its most simple version to more complex variations such as stochastic gradient descent (SGD). This method updates the weights by subtracting a value proportional to the gradient of the loss function in respect to the weights themselves times a positive factor called *learning rate*, as shown below.

$$w_i|_{n+1} = w_i|_n - lr \cdot \frac{\partial L}{\partial w_i|_n} \quad (1.4)$$

where $w_i|_n$ are the current weights, lr is the learning rate, $\frac{\partial L}{\partial w_i|_n}$ is the first derivative of the loss function in respect to the i -th weight at the current step, and $w_i|_{n+1}$ are the updated weights. This method is equivalent to minimize the error on the loss function, by following the gradient $\nabla_w L$. This vector lives in the multidimensional space of the loss function $L : \mathbb{R}^W \mapsto \mathbb{R}$, where W is the total number of parameters in the network.

The most efficient ?? algorithm is called *backpropagation*: it computes the first derivative of the loss function L in respect to all the parameters of the network, the weights, starting from the end of the artificial network and going backward toward the input, hence the name backpropagation. Since the number of connections between nodes might be even order of magnitude bigger than the number of nodes, it is simple to understand how large networks are computationally expensive to train.

Other types of learning processes are used, e.g. unsupervised/reinforced.

1.2.2 Validation Process

1.2.3 Testing Process

At the end, there is the process of testing the artificial neural network. Ideally the network is tested on a new set of data, for which the target results are known, similarly to the preceding phases. This time, however, the predicted outputs are compared to the correct answers to obtain an overall value for the correctness, often expressed in percentage.

1.2.4 Datasets

Since there are three phases of preparation for any artificial neural network, there must be an appropriate number of examples to feed to it.

HOW DO I DIVIDE A DATASET?

WHAT HAPPENS WHEN THERE ARE TOO FEW EXAMPLES?

AND WHEN THERE ARE TOO MANY?

1.3 Feedforward NN

The first and most simple type of neural network is called Feedforward. In this kind of neural network, nodes are divided into groups called *layers*. A layer is a collection of nodes that accepts inputs from a preceding group and generate as many outputs as the number of nodes in the layer. Each layer of a Feedforward neural network is connected in series with the others, except of input layer at the beginning and the output layer at the end. As for the single nodes, the inner layer are called hidden, because usually not accessible.

The information travels from the input to the output and gets elaborated from each hidden layer: there are no connection between nodes of the same layer, nor loops or feedback between layers. Depending on the topology of the network, there might be more or less layers, each composed by the same or a different number of nodes. Moreover the connection between the layers might be complete, i.e. each node in the layer accepts each input of the preceding layer, in that case the layer is said to be *fully connected*, or sparse as in the case of convolutional layers (see Section 1.3).

Perceptron

The most naive topology of a Feedforward neural network is given by the so called *Perceptron*. The Perceptron dates back to the 1957, when the homonym *Perceptron algorithm* was software implemented by Frank Rosenblatt on a computer (IBM 704) and only subsequently in hardware as the *Mark 1 perceptron*[5], [6]. The graph of a generic (single layer) perceptron ANN is shown in Figure 1.5 below.

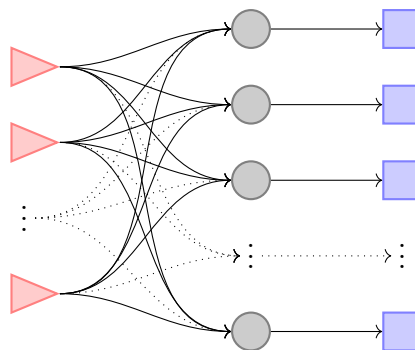


Figure 1.5: Perceptron type neural network: in this representation the perceptron has n inputs and m outputs as well as a hidden layer with m nodes. Colors, shape and styles are the same as in Figure 1.1 on page 6.

By adding more than one hidden perceptron layer to the neural network, one obtain the so called *Multi-Layer Perceptron* (MLP). This allows for more computational complexity ???. When the total number of layer is more than two, the network is called *deep*. A deep MLP is shown in Figure 1.6 on the next page. In principle any shape is possible, i.e. each layer could have a different number of nodes, however often the layers at the beginning are wider than

the layer at the end of the network ???. Besides the shape, in literature a perceptron is almost always considered fully connected ???.

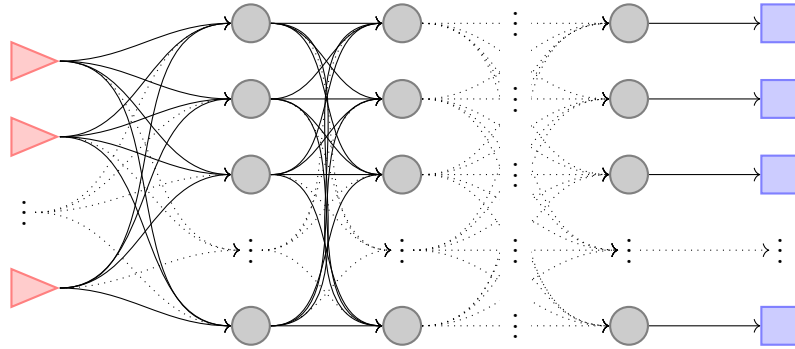


Figure 1.6: Deep Multi-Layer Perceptron (MLP), fully connected.

Other Feedforward NNs

Autoencoder neural networks are feedforward networks in which the output nodes are as many as the input nodes. The purpose of this kind of network is to reconstruct its own inputs.

Probabilistic

Time delay

Convolutional networks are inspired to the visual cortex, in which neurons are not fully connected their inputs but only to a restricted region. Convolutional neural networks are a type of feedforward network conceived to recognize images without being misled by distortions such as translation, skewing, or scaling. In this kind of network the input is often represented by a 2D matrix, instead of a 1D vector. It is usually composed by many layers, the prevalent kind is the convolutional one. This layer performs a two-dimensional convolution over the input matrix of a second 2D matrix of weights, called *feature map*. Thus, each node of the layer operates on a restricted region to understand if a feature is present or not. Commonly the operating regions are overlapping and the feature map is shared among the nodes in the same layer.

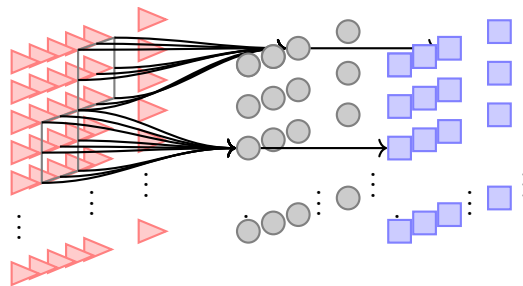


Figure 1.7: image/scheme of the conv NN

1.3.1 Other Types of NNs

By changing the topology of the nodes distribution and their connections, one obtain other networks that cannot be catalogued under the class of feedforward networks. Moreover, those

different types of network are not a niche, but are widely ?? studied as a different approach to the same or additional problems.

Recurrent NN

Recurrent neural network are a kind of network in which a portion of the input of nodes depends on the (past) output of the same nodes or nodes of subsequent layers. That is information does not propagates only forward like in the feedforward networks, but can propagate also backward, for example in loops or in feedbacks.

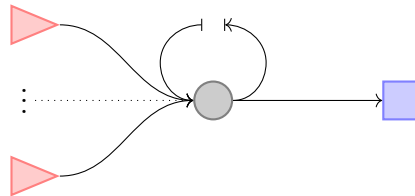


Figure 1.8

Reservoir NN

Reservoir neural networks differ from feedforward and recurrent networks in the learning approach. In fact, the topology of a reservoir network could be exactly the same as that of a deep multi-layer perceptron or that of a recurrent network. However, the reservoir computing approach claims that it is not necessary to learn all the weights of the network, but only for the last (perceptron) layer.

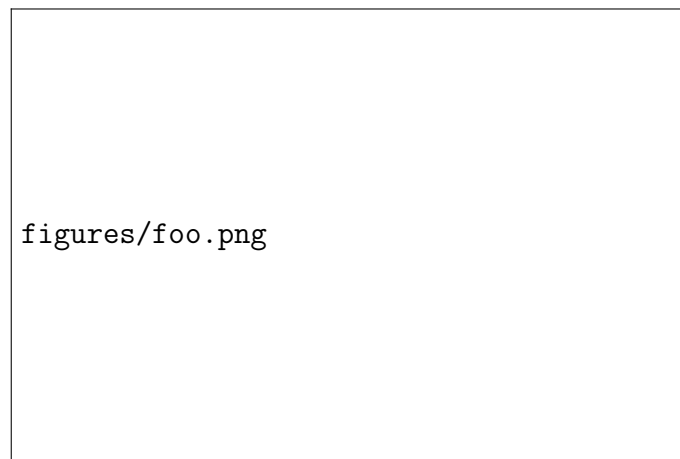


Figure 1.9: image/scheme of the reservoir NN

These kind of networks, then, can be trained much faster than their respective counterparts, i.e. feedforward and recurrent,

Modular NN

Spiking NN

Spiking artificial networks are the most different kind in respect to all the other networks until now described. In this class of ANNs, information is not coded only in the intensity of the signal, but also in the rate of signals, e.g. a high value will be encoded as a signal with high

repetition rate, whereas a low value as a signal with low repetition rate. This way of encoding information is more alike the mechanism of biological neural networks, such as our brain ??.

LOOK INTO other type of neuron model: Hodgkin–Huxley (H-H) model https://en.wikipedia.org/wiki/Binding_neuron

1.4 Real-Life Examples

? SHOULD I KEEP THIS SECTION ?

1.5 ANN Simulation

The resources and the time available allowed me to implement physically only the activation function of one node. Hence, to test this hardware implementation as I will show in Section 3.4 of Chapter 3, I had first to simulate and train *offline* a specific neural network. To do so, I chose a programming language, *Python*, and a library, *PyTorch*, which helped me in this task.

1.5.1 PyTorch

PyTorch is a python package that provides high-level features such as a deep learning research platform [8]. It is based on a backpropagation algorithm called *Reverse-mode auto-differentiation*, which is fast and flexible. Moreover integrates acceleration libraries that allow fast and lean operation.

To summarize, the library was chosen for its language and its flexibility, even though it is also powerful. This because our need was to simulate a little network to test the activation function.

Chapter 2

Integrated Photonics

The physics on which i want to build my hardware ANN is optics/photonics. Specifically integrated photonics tries to mix the good points of the integrated electronics (CMOS compatibility) with the good parts of optics.

How do I intend to create a hardware photonic ANN node?

2.1 Waveguides

In conventional optics, light is handled with bulky mirrors and lenses toward the desired position. In integrated photonics the most important device to control light is the waveguide. A waveguide is a path inside a medium, whose volume is defined by a certain number of interfaces between different materials, in which light remains confined and ideally travels with negligible losses.

One can distinguish two main types of waveguides: metallic and dielectric. The former are based on the reflection of the electromagnetic field by the metallic surface. However, for visible and infrared frequencies, metallic waveguides are not possible, due to too high absorption of the metals, The latter are based on the phenomenon of total internal reflection (TIR), and are nowadays widely used ?? in the industry and various research fields.

Total Internal Reflection and Dielectric Waveguides

Total internal reflection is the well-known phenomenon in which light coming from a material with high refractive index n_H gets reflected at the interface with a material with low refractive index n_L . For this to happen, incident light must be at an angle greater than the given by the

$$\theta_C = \arcsin\left(\frac{n_L}{n_H}\right)$$

Waveguides have many shapes, however one can initially discriminate them by the dimensionality of the confinement of light. The simplest one is called *slab* and it is composed by two interfaces, which divides the material in three volumes, as shown in Figure 2.1 below. This type is classified as a 1D-waveguide, because light is constrained in only one dimension. A generic representation is shown in the Figure 2.1 below.

2D-waveguides on the other hand are used more, since they allow to confine light in a straight path and conduct it to the designed spot. A few types are shown in Figure 2.2 below.

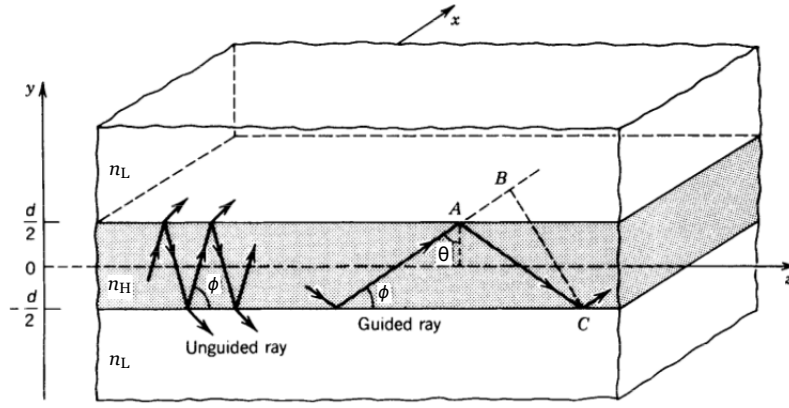


Figure 2.1: Scheme of a dielectric slab waveguide, made by two materials with refractive indexes n_H and n_L . Two rays are shown: the unguided one is incident on the interface at an angle smaller than the critical angle. Conversely the guided ray is incident at a greater angle and is therefore totally reflected inside the waveguide [9].

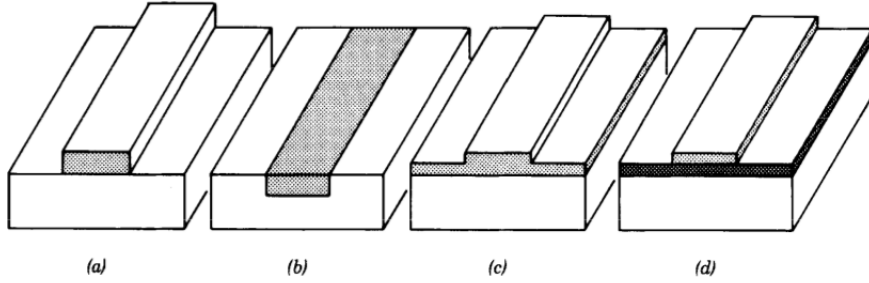


Figure 2.2: Representation of a few types of 2D dielectric waveguides. Light is constrained in two direction and can only move forward or backward in the third direction. (a) strip, (b) embedded strip, (c) rib, (d) strip loaded [9].

2.2 Microring Optical Cavity

In integrated photonics the microring is an optical cavity made by bending a waveguide on itself.

2.2.1 Nonlinear Perturbations

2.3 Photonics applied to ANNs

2.3.1 Weighted sum of inputs

This has already been demonstrated and integrated widely, so it will not be the focus of this work.

2.3.2 Nonlinear Activation Function

As opposed to the mechanism for weighted sum, an phenomenon for the activation function in an integrated photonic circuit has yet to be proposed.

2.3.3 Simulations

Chapter 3

Samples, setup and experiments

All the experiments have been made on samples manufactured for the IRIS project. This is due to the fact that in the time frame of this work there would have not been enough time to design and produce an ad hoc device. Moreover, as already stated, the aim of this thesis is to produce a proof of concept for an all-optical implementation of an activation function, rather than to construct a complete prototype.

3.1 The samples

The IRIS project studied the design and the implementation of an integrated reconfigurable silicon photonics switch matrix, a routing device, as a replacement for electronic devices used in the telecommunication industry. The completed integrated photonic circuit consists of a matrix of waveguides crossing each other and linked by couples of racetrack resonators, thermally-controlled. At the ends of the waveguides other structures, interleavers and AWGs, allowed many signals at different wavelengths to be multiplexed/demultiplexed on/from the same waveguide.

The complexity of such photonic circuit required the fabrication, for testing purposes, of each and every structure of which it is composed, with various production parameters. The collection of all these devices on a chip was produced in many samples. Isolated from the others, but also grouped together with few other elements. Specifically these test structures were disposed on a single chip and were accessible via grating couplers.

Since this work is at the beginning of the project, my choice was a system of intermediate complexity, so that it could be used both to test the activation function and the weighted sum.

The structure selected is the following: a simple waveguide, coupled to eight drop channels by a single or a couple of ring resonators each, nicknamed *mini-matrix*. In this family of devices, there were those built with single microrings, double microrings, single racetracks, or double racetracks. The final choice was to study the *mini-matrix* in which the coupling mechanism was provided by single ring resonators, because of its simpler transfer function in respect to double microrings or double racetracks.

Moreover, these samples were manufactured with thermo-electric ?? pads to heat the rings and effectively tune their resonance.

3.2 Setup

3.3 Characterization of the Activation Function

Characterization of the activation function

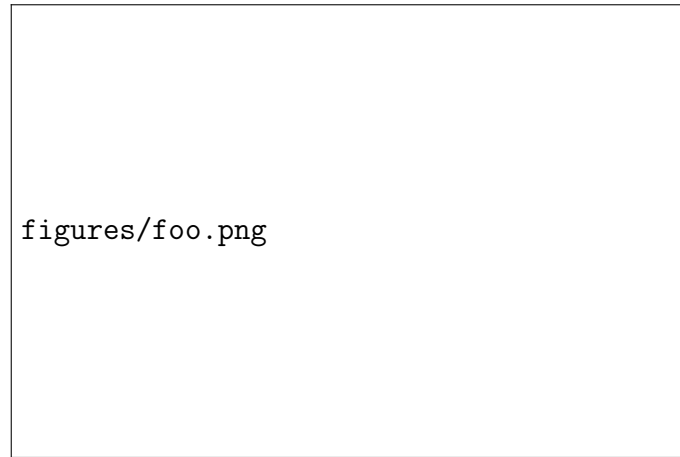


Figure 3.1: image/scheme of the minimatrix

3.4 Test of a Trained ANN

Test of the activation function

Conclusion

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