

UNIVERSITY OF SÃO PAULO
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BACHELOR OF COMPUTER SCIENCE

**A Survey of Explainable AI (XAI) Methods
for Convolutional Neural Networks**

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FINAL ESSAY
MAC 499 — CAPSTONE PROJECT

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Agradecimentos

Do. Or do not. There is no try.

— Mestre Yoda

Resumo

Antonio Fernando Silva e Cruz Filho

João Gabriel Andrade de Araujo Josephik. **Title of the document: a subtitle.** Monografia (Bacharelado). Instituto de Matemática e Estatística, Universidade de São Paulo, São Paulo, 2024.

Elemento obrigatório, constituído de uma sequência de frases concisas e objetivas, em forma de texto. Deve apresentar os objetivos, métodos empregados, resultados e conclusões. O resumo deve ser redigido em parágrafo único, conter no máximo 500 palavras e ser seguido dos termos representativos do conteúdo do trabalho (palavras-chave). Deve ser precedido da referência do documento.

Palavras-chave: Palavra-chave1. Palavra-chave2. Palavra-chave3.

Abstract

Antonio Fernando Silva e Cruz Filho

João Gabriel Andrade de Araujo Josephik. **A Survey of Explainable AI (XAI) Methods for Convolutional Neural Networks.** Capstone Project Report (Bachelor). Institute of Mathematics and Statistics, University of São Paulo, São Paulo, 2024.

Keywords: Keyword1. Keyword2. Keyword3.

List of Abbreviations

AI	Artificial Intelligence
ML	Machine Learning
XAI	Explainable AI
MLP	Multilayer Perceptron
CNN	Convolutional Neural Network
Conv	Convolution
IME	Institute of Mathematics and Statistics
USP	University of São Paulo

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Introduction

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Chapter 1

Background

In this chapter, we will introduce important concepts required for the understanding of this study. We begin by introducing Explainable AI (XAI) and its principles. We also introduce Neural Networks and important concepts such as Gradient Descent and Back Propagation. Finally, we introduce Convolutional Neural Networks, the main focus of this study.

1.1 Explainable AI

With the rise of Machine Learning models in the last decade in the business and academic areas, Artificial Intelligence (AI) is becoming increasingly present in important decision-making tasks. However, as AI models have become more sophisticated, particularly with the advent of Deep Learning techniques, their internal workings have often remained opaque. Explainable AI (XAI) aims to make models and their decisions more transparent, interpretable and understandable to both experts and inexperienced users.

1.1.1 What is Explainable AI?

Defining a mathematical formalization to explainability of Machine Learning is a difficult task considering the subjective nature of what one may consider "explainable". In non-mathematical terms, Explainability in AI refers to the capacity to articulate or justify the behavior of a model, focusing on methods that explain a model's decisions after they are made.

Another important concept in the area is Interpretability, which can be defined as "the degree to which a human can understand the cause of a decision" by Miller (2017).¹ In this case, however, a model's decision is understandable entirely by its inherent transparency. In other terms, the model is simple enough to be interpretable by a human directly, without the use of external techniques.

¹ Miller, Tim. "Explanation in artificial intelligence: Insights from the social sciences." arXiv Preprint arXiv:1706.07269. (2017).

Models with low complexity whose decisions are understandable by humans are defined as *Interpretable Models*. Linear Regression, Logistic Regression and Decision Tree models are examples of models classified as *Interpretable Models*. Now, models with a level of complexity that prevents humans from directly understanding their decision-making processes are referred to as *Explainable Models*. Recently popular *Deep Learning Models* are one kind of *Explainable Models* and will be the main focus of this essay, especially *Deep Convolutional Neural Networks*, explored in section 1.3.

1.1.2 Why Explainable AI is Necessary

Creating explanations to a model's decisions can yield many advantages, including more ethical and fair decisions, correctly following regulatory compliances and easier model debugging.

To ensure ethical and fair decision-making, Machine Learning systems must provide justifiable decisions, as they often exploit discriminatory patterns to enhance accuracy, which can perpetuate harmful biases. For instance, the COMPAS algorithm, used in U.S. courts to assess recidivism risk, was analyzed by ProPublica¹ and found to exhibit significant bias against Black defendants, frequently overestimating their likelihood of reoffending compared to their actual risk.

Explainable AI (XAI) is sometimes a mandatory requirement, particularly under regulations like the United Kingdom's General Data Protection Regulation (GDPR). The GDPR mandates that organizations must provide clear and understandable explanations for decisions that significantly impact individuals, especially those made by automated systems commonly powered by Machine Learning algorithms. Without XAI, high-stakes decisions cannot leverage such models in the United Kingdom, highlighting the crucial role of explainability in enabling the broader adoption of Machine Learning for real-world applications while ensuring compliance and fairness.

When debugging Machine Learning models, their behavior can often be unpredictable, revealing biases that may not have been initially apparent to humans. These biases can result in high performance on training, validation, or even test datasets but lead to poor performance in real-world deployment. For instance, consider training an image classifier to differentiate between dog and cat images. The model may achieve impressive accuracy on images of dogs in green fields. However, upon examining the regions of the image the model relies on for its predictions, researchers might discover that it focuses on the background rather than the animals themselves. This happens because dog owners are more likely to photograph their pets outdoors, leading to an unintended association between dogs and green backgrounds. Techniques from Explainable AI, such as Grad-CAM ([SELVARAJU *et al.*, 2019](#)) and Gradient Saliency methods, enable researchers to visualize these image regions, providing critical insights into model behavior and helping to address such biases.

¹ "How We Analyzed the COMPAS Recidivism Algorithm", by ProPublica: <https://www.propublica.org/article/how-we-analyzed-the-compas-recidivism-algorithm>

1.2 Gradient Descent

Let $f : A \rightarrow B$ where $A \subseteq \mathbb{R}^n$ for $n \in \mathbb{N}$ and $B \subseteq \mathbb{R}^+$. Suppose we want to find the solution to the optimization problem

$$\operatorname{argmin}_{x \in A} f(x) \quad (1.1)$$

when $\frac{\partial f}{\partial x}$ is known for any value of x . Considering that the vector $\frac{\partial f}{\partial x}$ points to the direction of the steepest ascent of the function, the vector $-\frac{\partial f}{\partial x}$ will point to the steepest descent from the given point x . Therefore, one can define an initial random value for x and update x using $-\frac{\partial f}{\partial x}$ and a scaling factor η in order to find a local minimum of f and an approximation to the solution of given optimization problem.

We can define such method using the following formula, where x_t represents the value of x at iteration t of the algorithm:

$$x_{t+1} = x_t - \eta \frac{\partial f(x_t)}{\partial x_t}. \quad (1.2)$$

The term η is often called the *learning rate* used in the Gradient Descent method and is often defined manually by the user.

The Gradient Descent method can be used to optimize a neural network's parameters to solve a given problem using a *loss function*.

1.3 Neural Networks

Neural Networks are proven to be universal approximators.² That means that Neural Networks are Machine Learning models capable of representing any continuous function, therefore making Neural networks adept at modeling a range of different complex problems. This class of models have seen a growing presence across both academic and industry landscapes. However, given the architecture of multiple hidden layers of Neural Networks creating complex internal patterns, such models are classified as Explainable Models.

In this section, the inner workings of Neural Networks will be explained, starting with the *Perceptron*, considered the fundamental building block of Neural Networks.

1.3.1 Perceptron

A Perceptron is a Machine Learning model inspired by how biological neurons work. It is a simple binary linear classifier that defines its parameters by linear combinations of points in the dataset. The Perceptron model can be described by Figure 1.1.

² Hornik, K., Stinchcombe, M., White, H. Multilayer feedforward networks are universal approximators. [https://doi.org/10.1016/0893-6080\(89\)90020-8](https://doi.org/10.1016/0893-6080(89)90020-8)

³ <https://towardsdatascience.com/what-the-hell-is-perceptron-626217814f53>

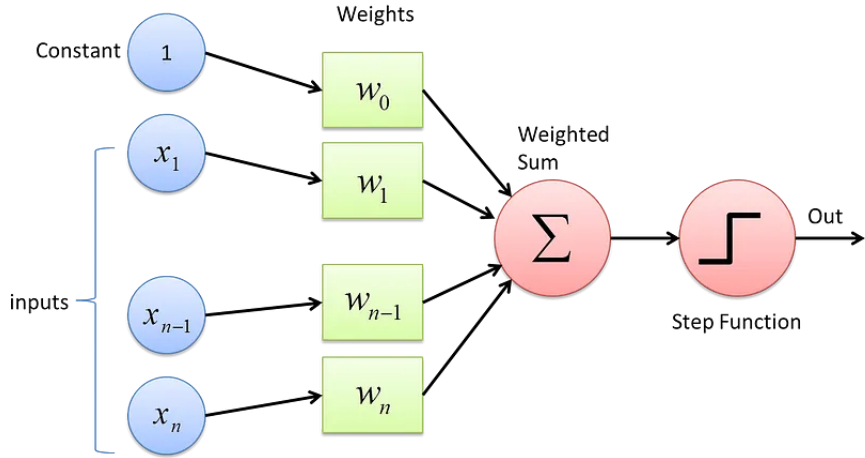


Figure 1.1: Perceptron Architecture. Font: Towards Data Science³.

Where the weights w_i for $i \in \{0, 1, \dots, n\}$ are trainable parameters and the step function can be defined as $\sigma : \mathbb{R} \rightarrow \{0, 1\}$ such that

$$\sigma(x) = \begin{cases} 1 & \text{if } x \geq 0 \\ 0 & \text{if } x < 0. \end{cases} \quad (1.3)$$

Therefore, the Perceptron model can be defined as the function⁴ $f : \mathbb{R}^n \rightarrow \{0, 1\}$ where

$$f(x) = \sigma(w_0 + \sum_{i=1}^n w_i x_i). \quad (1.4)$$

The Perceptron model updates its parameters using each sample (x, y) of the dataset with the rule

$$w_i^{t+1} = w_i^t + \eta (y - f(x)) x_i \quad (1.5)$$

for $i \in \{1, \dots, n\}$ and

$$w_0^{t+1} = w_0^t + \eta (y - f(x)), \quad (1.6)$$

where η is the *learning rate* hyperparameter and t is the update iteration number.

As a linear model, the Perceptron can only model linear problems, which only represent a small subset of real world problems. As a solution, researchers started combining Perceptrons in a layered structure, called Multilayer Perceptron, also famously known as *Neural Networks*.

⁴ The independent term w_0 is usually called the *bias* of the Perceptron (or neuron)

1.3.2 Multilayer Perceptron (MLP)

By stacking multiple Perceptrons into multiple layers, one can build more complex decision boundaries and model more complex functions. For example, by using the Multilayer Perceptron, one can model a XOR function:

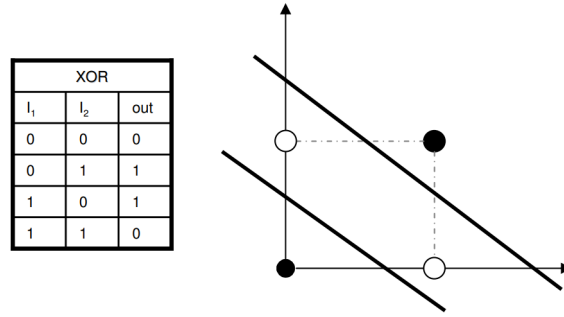


Figure 1.2: The XOR Problem Font: Kevin Swingler Lecture Notes.

The XOR problem can not be solved by using a single Perceptron, since it is not a linear problem. However, by using two Perceptrons (which corresponds to the two lines in the figure), one can model the non-linear XOR problem. In this specific example, one can define the uppermost line as the decision boundary of the Perceptron $f_1(x) = \sigma(-x_1 - x_2 + 1.5)$ and the lowermost line as the decision boundary of the Perceptron $f_2(x) = \sigma(x_1 + x_2 - 0.5)$. Considering the Perceptrons f_1 and f_2 , we can create a new Perceptron that receives the outputs of those Perceptrons and returns the result of the XOR function. For example, we can define the Perceptron $g(x) = \sigma(x_1 + x_2 - 1.5)$, generating the following results:

x_1	x_2	$f_1(x)$	$f_2(x)$	$g(f_1(x), f_2(x))$
0	0	1	0	0
0	1	1	1	1
1	0	1	1	1
1	1	0	1	0

Table 1.1: Perceptron output for binary combinations of x_1 and x_2 .

Showing that the non-linear problem can be successfully be solved by the Multilayer Perceptron.

Although the Multilayer Perceptron has the perk of being able to model complex functions, we are still limited by how to model is trained, since it cannot be trained by using the update rule of the traditional Perceptron.

By using a technique known as Backpropagation, the Multilayer Perceptron can be trained using gradient-based update rules, like the Gradient Descent.

1.3.3 Backpropagation

In order to find the weight's gradients of our MLP, one can use Backpropagation, a technique that involves computing those gradients by performing a *Forward Pass* and a *Backward Pass* on the Neural Network.

Forward Pass

The Forward Pass basically consists in computing the input through the network and comparing the output with the expected value using a loss function \mathcal{L} . During the forward pass, the output of each layer is stored in memory to be later used in the Backward Pass.

Backward Pass

In the Backward Pass, the gradients of the loss with respect to the weights are calculated to update the network. By using the chain rule, one can start off by calculating the gradients of the last layer of the network, and then use the result to calculate the gradients of the weights of the previous layer, *going on the opposite direction of the Forward Pass*.

Updating Weights

With the gradients of the loss with respect to the weights calculated, now the weights are updated in an iterative process, until a satisfiable loss/accuracy is achieved or new model/dataset tuning is necessary.

1.4 Convolutional Neural Networks

1.4.1 Convolutions

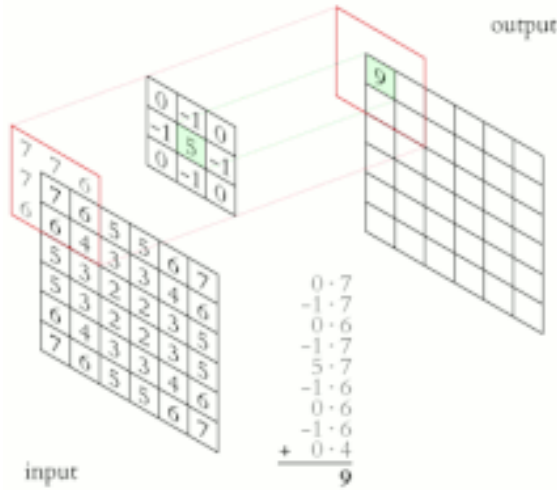
First, it is important to define what a convolution is. Given two discrete one-dimensional signals f and g , their convolution $f * g$ is defined as:

$$(f * g)[n] = \sum_{i=-\infty}^{+\infty} f[n]g[n-i]$$

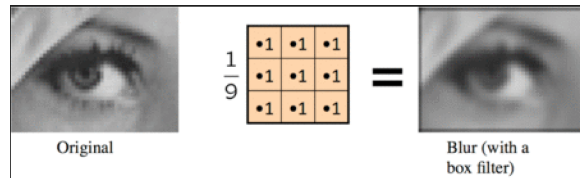
Given two discrete two-dimensional signals f and g , their convolution $f * g$ is calculated as:

$$(f * g)[m][n] = \sum_{i=-\infty}^{+\infty} \sum_{j=-\infty}^{+\infty} f[m][n] \cdot g[m-i][n-j]$$

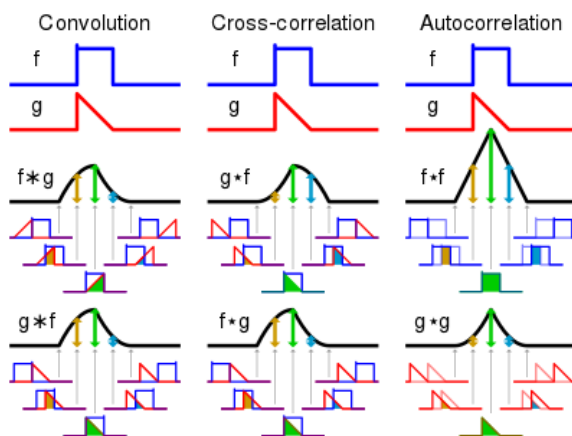
In practice, the signal g is represented by a window (or kernel), usually square and of odd size. Thus, we can abstract convolution as the multiplication of a sliding window. The picture below illustrates this process. It is important to note that the window needs to be flipped during the convolution, although this is not illustrated in the picture.

**Figure 1.3: 2D Convolution**

This process can be used to apply different filters to images. For example: using a 3×3 window with all weights equal to $\frac{1}{9}$, we can generate a filter that blurs the image (moving average). Below is an example of applying this filter:

**Figure 1.4: Filtered Image (3x3 Mean Filter)**

It is also important to note that convolution is commonly implemented in machine learning contexts as "cross-correlation," which is a very similar operation but without the flipping of the window. Note that, since the weights are learned in our case, there is no difference. Therefore, in our context, convolution and cross-correlation are synonymous.

**Figure 1.5: Comparison with Cross-Correlation**

A pertinent question that can be asked is what happens at the edges of the image.

When the window is sliding over them, what happens to the missing pixels? The process of filling in these pixels is called padding. Padding can be done with zeros, the nearest pixel, or not be done at all. Note that when there is no padding, the image decreases in size after convolution.

Another important hyperparameter that can be adjusted is the "stride." This defines how many positions the window is moved at a time. That is: a stride value different from 1 also implies a decrease in image size after convolution.

1.4.2 Convolutional Layer

In order to understand the need for convolutional networks, we have to understand why it's impractical to use fully-connected networks to process images. Let's walk through an example to see that.

Consider a color image with dimensions 512x512. If we were to process this image with a conventional neural network, the input layer would have $3 \cdot 512 \cdot 512 = 786432$ dimensions. Assume a hidden layer with only 128 neurons (which is relatively small). Just between these two layers, there would be 100663296 parameters! This is highly inefficient.

The solution to this problem is to extract features from the image, which will serve as input to the network. These features could include various aspects such as symmetry, black levels, contrast, presence or absence of patterns, etc. All of these features will serve as input to the network. As a result, we can reduce the input layer's dimensionality from several hundred thousand to just a few dozen.

However, a challenge still remains: how do we select these features? We can apply convolutions to the image to calculate interesting features, and these convolutions can be learned alongside the rest of the network! It is important to understand some essential details about these networks before proceeding. Each convolutional layer has three dimensions: height, width, and the number of channels. The input layer typically has one channel for black and white images, or three channels for color images.

Each channel in each convolutional layer combines all the channels from the previous layer. In other words, the "windows" used have weights for all the channels. These windows slide over the data from the previous layer to generate **one channel** in the next layer.

Let us consider an example. Suppose we have a network that processes color images of size 128×128 pixels. This network has 3 convolutional layers with 16, 32, and 64 channels per layer, respectively. Assume a window size of 3 for all layers. In this case, we have:

- First layer: window size is $3 \times 3 \times 3$. With 16 output channels, we will have

$$16 \cdot 3 \cdot 3 \cdot 3 = 432 \text{ parameters.}$$

- Second layer: window size is $3 \times 3 \times 16$. With 32 output channels, we will have

$$32 \cdot 3 \cdot 3 \cdot 16 = 4608 \text{ parameters.}$$

- Third layer: window size is $3 \times 3 \times 32$. With 64 output channels, we will have

$$64 \cdot 3 \cdot 3 \cdot 32 = 18432 \text{ parameters.}$$

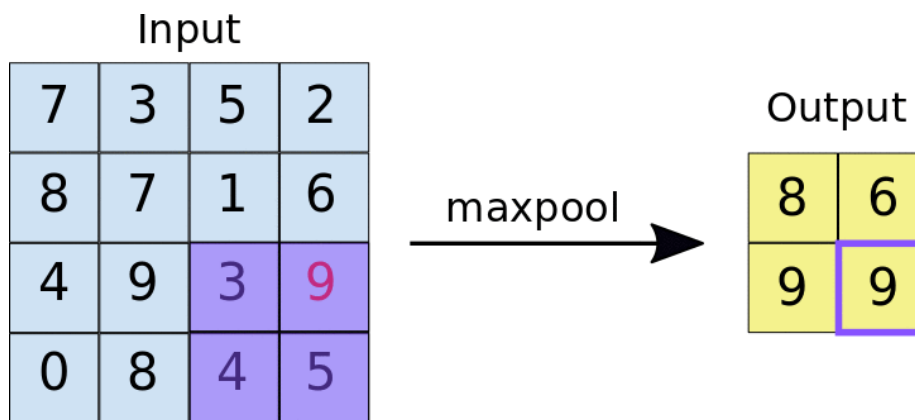
Another important detail is the output dimension of each layer. This depends on whether or not **padding** is used. **Padding** refers to how the layer behaves at the image edges. We can complete the image with zeros, the nearest pixel value, or the pixel value from the opposite edge of the image. If padding is not used, the output dimensions will decrease by $2\lfloor \frac{W}{2} \rfloor$, where W is the window size. For instance, with a window size of 3, each layer will reduce the image size by 2 pixels. If the input is 128×128 , the output of the first layer will be 126×126 , the second layer will output 124×124 , and so on.

1.4.3 Pooling

Remember, each convolution extracts a feature from the image. Therefore, when we perform another convolution using the outputs from the previous layer, we are combining features extracted from the image to compute new features. As a result, deeper layers extract more complex features from the image. For instance, the first layer may extract features like the presence of vertical straight lines, while the tenth layer may extract features like "presence of dog snouts."

Thus, the features involved gradually become less localized and more global (pertaining to the entire image). This is why it is useful to summarize information into smaller dimensions as the network deepens.

To accomplish this, we use "pooling" layers. These layers work similarly to convolutions: windows slide over the data and compute an output based on nearby pixels. However, this time, a function is used to aggregate these data. Common functions include "max" (maximum value) and "avg" (average value).

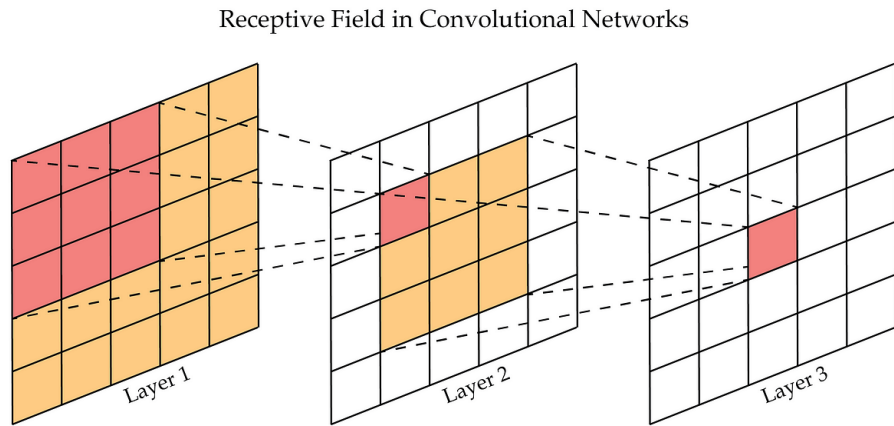


1.4.4 Receptive Field

An important concept for understanding the power of **deep convolutional networks** is the **receptive field**. This concept relates to the power that chained convolutions have.

Consider an input image. Apply a 3×3 convolution to it. Now, apply another 3×3

convolution to the output of the first convolution. Observe this output image. How much information does each pixel contain about its neighbors?



The answer is that each pixel contains information from a region of size 5×5 around it! This is the receptive field of these neurons.

A common misconception is that, since the receptive field of two 3×3 convolutions is 5×5 , two 3×3 convolutions have the same **expressive power** as a 5×5 convolution. This **is not true**. A 5×5 convolution has 25 parameters, while two chained 3×3 convolutions only have 18 parameters.

Chapter 2

GradCAM

One of the most prominent model-specific methods to acquire explanations for classification with CNNs is GradCAM. The intuition for the method is simple. At the last convolutional layers, we have several channels that represent each a different feature. Those features are used by the next part of the network to produce the final output. If we want to know which parts of the image are being more useful to the network, we can look at the feature maps and observe which parts of the image are generating the signal used by the rest of the network.

The problem with this approach is that the features have informations about all the output classes. How we know what features are more important to the decision? The idea behind GradCAM is to **average the feature maps weighted by the gradient** of each channel with respect to a specific class.

However, this will still highlight the regions that have a negative influence to the decision. To filter out those regions, the result is passed through ReLU. The result is a coarse heatmap of the image highlighting important regions.

The formula for the heatmap with regard to the class c is:

$$H = \text{ReLU}\left(\sum_k \alpha_k^c A^k\right)$$

Where α_k^c , the weight of the k -th feature map for the class c , is defined as:

$$\alpha_k^c = \frac{1}{Z} \sum_i \sum_j \frac{\partial y^c}{\partial A_i^k j}$$

Where A^k is the k -th feature map, y^c is the output for the c class, and Z is the number of neurons in each map.

2.1 Guided Backpropagation

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2.2 Guided GradCAM

 The output of GradCAM has the dimensions of the last convolutional layer of the network, and has to be upsampled to be overlayed on top of the input image. This results in a very coarse heatmap, with rough borders and lost details. To solve this, we can multiply (pixel by pixel) the heatmap with other simpler method, as guided-backpropagation.

Chapter 3

Feature Visualization

Feature Visualization is a technique that involves maximizing values of neurons, sets of neurons or even layers of a Neural Network in order to understand concepts learned by the model. By maximizing a neuron's value, we can better understand what set of features each part of our network is learning to capture and verify if the network is aligned with human judgement. In simple terms, Feature Visualization can be described by the following optimization problem:

$$\text{img}^* = \underset{\text{img}}{\operatorname{argmax}} h_{n,x,y,z}(\text{img}) \quad (3.1)$$

where h represents the activation of a neuron, img is the input of the network, x and y represent the spatial position of the neuron, n is the layer of the network and z is the channel index. This expression represents the problem of maximizing a value of a single neuron. For a set of neurons, the problem can be described as the formula:

$$\text{img}^* = \underset{\text{img}}{\operatorname{argmax}} \sum_{(n,x,y,z) \in A} h_{n,x,y,z}(\text{img}) \quad (3.2)$$

where $A \subseteq \mathbb{N}^4$ is a set of combinations of network layer, channel index and spatial position vectors.

In order to find a solution to the optimization problem, we can use the Gradient Descent technique presented in [Chapter 1](#). However, instead of minimizing an optimization problem, we are looking for a solution that maximizes [Equation 3.1](#) or [Equation 3.2](#). Therefore, instead of subtracting the partial derivative term of [Equation 1.2](#) we just need to add the partial derivative multiplied by the *learning rate* (We will call the technique *Gradient Ascent*). Thus, the following formula is derived for Feature Visualization:

$$\text{img}_{t+1} = \text{img}_t + \eta \sum_{(n,x,y,z) \in A} \frac{\partial h_{n,x,y,z}(\text{img}_t)}{\partial \text{img}_t} \quad (3.3)$$

where t is the iteration t of the algorithm.

The initial image img_0 can be defined in two ways:

- A completely random initialization, with $\text{img}_0(c, x, y) = U[0, 1]$ for each channel c and image spatial positions x and y .
- A user defined image, normally a real world image.

The choice between these two approaches depends on the researcher's ultimate objective when using the algorithm. Providing an initial non-random image allows researchers to focus on specific features within the image and examine their effects on a particular set of neurons. In contrast, using a random image may be better suited for discovering unknown features.

Like most optimization problems, the Feature Visualization problem doesn't strictly have a single optimal solution, but rather multiple viable solutions that maximize the given set of neurons. For example, a set of seemingly random images may activate a neuron fully while at the same time an image of a cute dog may also activate this same neuron to its maximum value. Directly applying *Gradient Ascent* to random images without additional techniques often produces images that poorly align with human perception. This outcome occurs not because human-aligned images fail to maximize the neuron's value, but because solutions comprising seemingly random pixels are closer to the image's initial state.

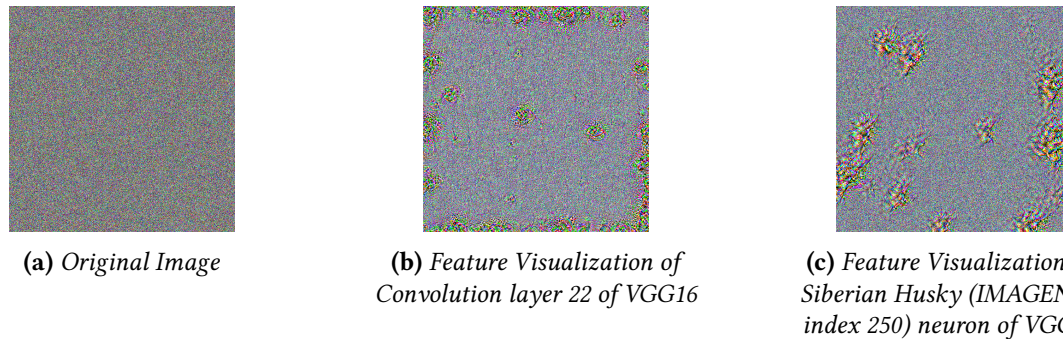
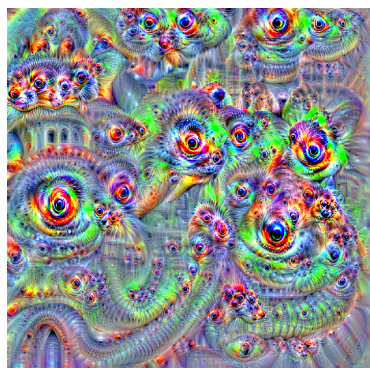


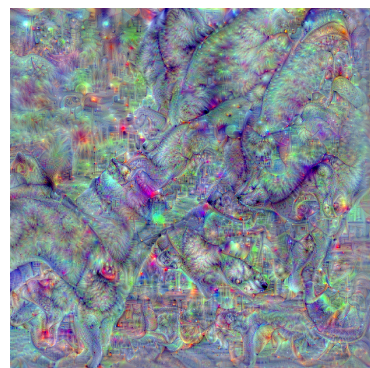
Figure 3.1: Feature Visualization images using solely Gradient Ascent. Little human-recognizable features are present in the resulting images

To identify human-aligned features in our models, we need effective techniques for generating images that closely correspond to those features. A closer examination reveals that Feature Visualization using solely Gradient Ascent typically produces features that occupy only small portions of the final image.

To expand these features across a larger area and create more intricate forms, we can downscale the image and apply the Gradient Ascent step. Once the downscaled image has been refined, it can be upscaled by a specific factor, and Gradient Ascent can be reapplied. Repeating this process until the image reaches its original size enables the generation of more detailed and compelling results, as demonstrated below:



(a) Feature Visualization of Convolution layer 22
of VGG16



(b) Feature Visualization of Siberian Husky
(IMAGENET index 250) neuron of VGG16

References

- [SELVARAJU *et al.* 2019] Ramprasaath R. SELVARAJU *et al.* “Grad-cam: visual explanations from deep networks via gradient-based localization”. *International Journal of Computer Vision* 128.2 (Oct. 2019), pp. 336–359. ISSN: 1573-1405. DOI: [10.1007/s11263-019-01228-7](http://dx.doi.org/10.1007/s11263-019-01228-7). URL: <http://dx.doi.org/10.1007/s11263-019-01228-7> (cit. on p. 4).