

EVALUATION OF IMAGE RECOGNITION NEURAL NETWORK INTERPRETATION METHODS: AN IN-DEPTH LOOK

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Sworn Declaration

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Linz, August 30, 2023

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Abstract

This paper focuses on the accuracy of interpretability methods for machine learning models. The main research problem is the lack of ground truth for evaluation method in interpretability methods. While there are inherent interpretative models, black-box networks perform better and have developed rapidly. Existing interpretability methods for neural networks such as ROAR, KAR, BAM and Real Time Image Saliency for Black Box Classifiers provide a numerical evaluation method but they do still suffer from ambiguity. This paper summarizes the different evaluation methods, compares them and also calculate ROAR on 2 different saliency methods.

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1 Introduction

Artificial Intelligence (AI) has undergone rapid development in the last years. In today's modern era of mobile phones and computers, algorithm's are used on a daily basis to have quick access to information and improve the efficiency of the daily life.

While various Algorithms (e.g.: Decision Trees, Linear Regression, Support Vector Machines, etc.), which are comprehensible by design, have been developed, the spotlight has turned to Deep Neural Networks(DNN). This shift is attributed to the increase in computational power and the exponential increase in accessible data. Despite their remarkable accuracy, DNN remain opaque black-boxes, which we struggle to understand. Nevertheless, the immense improvement in performance and their ability to handle massive datasets have led to widespread adoption in contemporary devices. It is predicted, that algorithms based on Neural Networks will be becoming increasingly popular in the next years.

However, one of the primary difficulties with Neural Networks is the lack of reliable interpretation techniques. Numerous interpretation methods exist, yet a universally reliable method remains missing. Particularly in the domain of image analysis, encompassing critical applications like automated driving and facial recognition, no solution is present. The decision-making rationale of neural networks remain unclear, attributed to factors like background elements, peripheral objects or lighting conditions. Efforts to address this issue have given rise to gradient methods, aiming to assign significance values to pixels and represent their importance on neural network decisions. Another alternative option to mitigate the black-box nature of algorithms involves employing model-agnostic methods. These methods offer an computational linkage between inputs and outputs, irrespective which model is used. Although highly effective for smaller datasets, they begin to struggle as the data size and their complexity increases. Because of this, they do not offer a reliable way to quickly make Neural Networks interpretable.

In light of these prevalent problems, the object of this thesis is to recapitulate interpretation algorithms Neural networks in computer vision. Emphasis is placed on the evaluation of post-hoc interpretability techniques, forecasting potential future developments and focusing on the strengths and weaknesses of distinct techniques. Concluding the theoretical segment, a practical demonstration showcasing the application of ROAR is shown.

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1.1 Structure of the thesis

- 1. The first part presents a overview of contemporary machine learning algorithms, categorizing them into two main groups: algorithms with inherent interpretability and those without. The goal is to make clear how supervised methods can be applied to image recognition tasks.
- 2. Subsequent sections delve into interpretability, emphasizing global and local model-agnostic techniques. These methods offer insights into overall model behavior, regardless of algorithm specifics.
- 3. Additionally, in the domain of interpretability techniques for Neural Networks, the paper explores ad-hoc methods for Neural Networks. Features visualization and Gradient-focused methods are explained.
- 4. After introducing existing interpretation methods, the paper's focus transitions to evaluating post-hoc interpretation methods. Various approaches to assess the effectiveness and dependability of these methods in offering meaningful insights into intricate models are introduced and discussed. Additionally, the advantages and disadvantages of these approaches are carefully examined to provide a comprehensive understanding of their applicability.
- 5. To exemplify the discussed concepts, the practical application of the ROAR methodology using the food101 data set [5] and MNIST dataset [8] is presented. This real-world instance illustrates the current state of art of interpretability techniques in image recognition.

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2 Machine Learning and their Interpretability

Before going into detail into the different methods of interpretability an overview of current supervised machine learning is given. Unsupervised machine learning is argueable intrinsically understandable, as the goal is to find a structure in the data. [2] With those examples, it should be explained why interpretability is necessary.

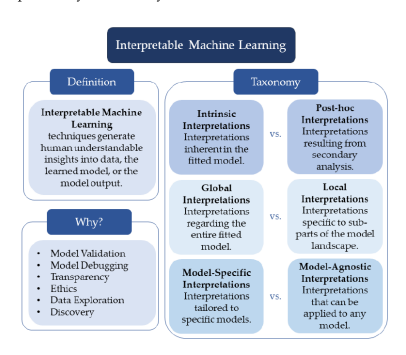


Figure 2.1: Feature Visualization [2]

Interpretability methods can be defined in different categories:

Intrinsic vs Post-hoc: Intrinsic methods) are understandable by nature. Pos-hoc interpretations however are done by an analysis of the model afterwards.

Global Interpretations vs Local Interpretations[22]: Global Interpretations regard the entire fitted model. A global interpretation explains the whole model. Local Interpretations only explain a small part of the model landscape. For example the boundary between two classes.

Model-Specific Interpretations vs Model-Agnostic Interpretations [22]: Model Specific interpretations can only be used by a single class. For example gradient-methods for Neural networks. Model-agnostic interpretations can be applied to any method. For example LIME or SHAP.

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A broad definition is given by [2]: "Interpretable machine learning (IML) is the use of machine learning techniques to generate human-understandable insights into data, the learned model, or the model output."

Interpretability is needed for various reasons. Many have proposed a number of reasons and uses for IML. [23] [27] [22] [12] [19] [10] [6] [9]:

Model Validation: Does the learned model makes sense and behave in consistent behavior matching with my prior expectations and knowledge about the system? Validation by humans require interpretable machine learning models.

Model Debugging: When unexpected behavior occurs, how can one diagnose the problem in the machine learning system? Interpreting and understanding machine learning systems is critical for diagnosing, debugging and fixing systems.[16]

Transparency, Accountability & Trust: IML makes black-box and other machine learning systems easier for human to understand and more transparent. The utilization in high-stakes societal applications requires accountability and trust of machine learning systems [28] [29] [32].

Ethics: Machine learning algorithms can possibly be trained on biased data lead to unfair predictions that are discriminatory. To improve the fairness of machine learning algorithms understandable techniques need to be deployed. [12]

Data Exploration: Insights into major patterns, trends, groups, or artifacts of the data can be gained by applying human-interpretable techniques. This data exploration insights are used to adjust the data preprocessing and model influence decisions. [23] [3] [4]

Discovery: IML can help making data-driven discoveries in the realm of data analytic.

2.1 Supervised Machine Learning

In supervised machine learning, there are several base methods of classification. They are shortly introduced and analyzed for their interpretability. This should give readers a simply structure on why this research is needed. This section analyzes the base functionality of each singular model. Furthermore, an analysis of the interpretability from a human perspective is made. Methods are developing rapidly, therefore only the most common methods are included.

There are several methods on how algorithms can be classified in their interpretability: Algorithm Transparency: How does the algorithm learn the model from the data? What relationships can it learn? [22] A linear regression model using least squares method is easier to understand than a deep learning model. Pushing a gradient through a network with millions of weights is less well understood and considered less transparent.

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2.1.1 Linear Models

When it comes to predicting outcomes, a simple method is to use a linear regression model. This model predicts by adding up different features, each multiplied by a weight. The linear nature of this model makes it easy to understand. Mathematically, the predictive output, denoted as \hat{y} , is captured in the equation:

$$\hat{y} = \alpha_0 + \alpha_1 x_1 + \alpha_2 x_2 + \dots + \alpha_n x_n + \epsilon$$

The alphas α_i indicate the significance of each feature. The initial coefficient α_0 is known as the intercept, signifying the baseline. The noise ϵ encapsulates the inevitable errors stemming from inherent non-linearity in real-world dynamics or measurement inaccuracies.

To train model, the MSE-Loss or the absolute loss can be applied. When using regularization methods, the absolute loss is taken to be more resilient to outliers.

$$MSE = \frac{1}{n} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

$$ABS = \frac{1}{n} \sum_{i=1}^{n} y_i - \hat{y}_i$$

The interpretability of the model is very simple. The factors are given through the coefficient matrix. Each feature has a distinctive importance to the model and it can be seen easily how important each factor is, when the data is normalized.

$$\alpha = \begin{bmatrix} \alpha_0 \\ \alpha_1 \\ \dots \\ \alpha_n \end{bmatrix}$$

Although linear models possess comprehensibility and provide a straightforward method for prediction and are inherent understandable, their application is limited to linear relationships and small datasets. One can use advanced techniques like L1 and L2 regularization [11] to achieve regularization against outliers and in case of correlation between factors. In the domain of image recognition it is not applicable.

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2.1.2 Distance-Based methods

K-Nearest Neighbors is used for classification and uses the nearest neighbars as classification. KNN is not interpretable by default, as there are no parameters to learn and analyze. However KNN is inherently easy to understand, as it simply looks for the most similar samples. One is able to visualize fewer features or use clustering algorithms to reduce the dimensionality, but the relationship between input and output is unclear. KNN also struggles with many features, therefore it is not recommended.

Support Vector Machines (SVM) aim to find a hyperplane that maximizes the margin between different classes of data points.

$$Minimize \frac{1}{2}||w||^2 + C\sum_{i=1}^n \xi_i$$

Subject to
$$y_i(w \cdot x_i + b) \ge 1 - \xi_i$$
, $\xi_i \ge 0$ for $i = 1, ..., n$

By using the kernel trick, SVM can be used for non-linear data. In higher dimensionality, SVM becomes non-interpretable, as displaying the weight matrix is not understandable. SVM also struggles with big datasets. Therefore also SVM is not a good choice for image classification.

2.1.3 Decision Tree-Based Methods

Decision Trees based on minimizing the gini-index are inherently interpretable. As the depth increases, the models become less understandable. An example for the titanic dataset is visible in 2.2 Their bad generalization method gave rise to ensemble methods.

Random forests are an ensemble of multiple decision trees. Their advantage is a smoother predictive power. But it suffers from being less interpretable. Using special techniques like SHAP values [20] or partial dependence trees can make them more understandable.

Gradient boosting like XGBoost, LightGBM and CatBoost are similar to random forest and decision trees, but with differential learned weights for each decision. They suffer from the same interpretability issues as random forests.

While decision tree-based methods can be applied for image classification, their accuracy is poor.

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Survival of passengers on the Titanic

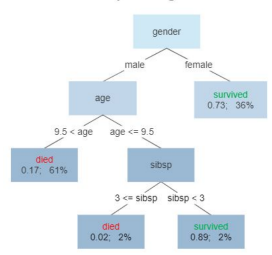


Figure 2.2: Decision Tree Example: By Gilgoldm - Own work, CC BY-SA 4.0, https://commons.wikimedia.org/w/index.php?curid=90405437

2.1.4 Neural Networks

The rise of neural networks and their strong predictive power makes them a common choice for classification task. But as they increase in size, understandability by taking a look at the weights becomes impossible. With the rise of CNN in 2012 [17] Neural Networks have become state of art for image prediction. Special interpretability methods considering the learned weights are looked in detail in 3.3.

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3 Interpretation of Image Recognition Neural Networks

In this chapter, evaluation method are described to analyze the behavior of not inherently interpretable models. Only evaluation methods which are commonly used for neural networks are introduced.

3.1 Global Model-Agnostic Methods

Global model-agnostic methods describe expected outcomes based on the distribution of the data. They can show a correlation between singular or multiple features and an outcome.

- 1. **Partial Dependency Plots:** Partial Dependency Plots (PDP) are easy to interpret but using it with several features becomes increasingly difficult. Therefore, using a PDP is not possible in Neural Networks.
- 2. **Accumulated Local Effects:** Accumulated Local Effects (ALE) are an an advancement of PDP. While it solves some problems PDP suffers from, it can't be applied to a Neural Network because of the complexity of the datasets.
- 3. **Feature Interaction:** Feature Interaction also analyzes how features correlate. Doesn't work due to the same problem of the complexity of the dataset.
- 4. **Functional Decomposition:** Functional Decomposition is commonly used in Neural Networks. See Chapter 3.3.1.
- 5. **Permutation Feature Importance:** Permutation Feature Importance is regularly used in visual machine learning tasks.
- 6. **Prototype and Criticism:** Prototype and Criticism is used as adversarial attacks in Neural Networks.

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3.2 Local Model-Agnostic Methods

To explain individual predictions, Local model-agnostic methods are used. A single outcome is correlated to some features and explain the model.

- 1. **LIME: Local Interpretablle Model-agnostic Explanations:** Lime generates locally faithful explanations by training interpretable models on perterurbed instances of the original data.
- 2. **Local surrogate models:** Local surrogate models can be programmed to select a singular instance to explain.
- 3. **Scoped Rules (Anchors):** Find so called anchors to explain the predictions.
- 4. **Individual conditional expectation curves:** Practically not useful in image recognition, as the computation is too high.
- 5. **Counterfactual explanations:** Try to find a change while still resembling the original image.
- 6. **SHAPly:** Calculate SHAP values for an image prediction to determine how each pixel contributes to the prediction's deviation from the average prediction across all images. Positive SHAP values indicate pixels that push the prediction up, while negative values indicate pixels that pull it down.

3.3 Neural Network Interpretation: Model specific Interpretations

In the domain of Natural Language Processing (NLP) and Computer Vision, Deep Learning has proven very successful. By passing the input data through a sequence of layers, characterized by matrix-multiplications with kernel weights and nonlinear transformations functions, a prediction is computed. Depending on the specific task, additional elements like Long Short-Time Memory(LSTM) layers and Convolutional layers (CNN) are utilized. Given the immense amount of mathematical operations underlying a single prediction, humans are not fit to apprehend the mapping. To interpret predictions, we would have to decipher the intricate learned knowledge of numerous different kernels and weights. Recognizing that it's impossible for humans to grasp millions of weights, the demand for evaluation methods is high. To assess the behavior and predictions of Deep Neural networks, specific interpretation methods were developed. These methods calculate the likelihood of an input entry being responsible for the result.

While model-agnostic methods offer an approach to understand Neural Networks, the sheer size of the data used to train and test Neural Networks make this task extremely hard. For instance, an image with the dimensions of 3x224x224, as commonly encountered in Food-101, the data entries

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exceed 150.000. In NLP tasks, where vocabularies often encompass around 20.000 words, the computational complexity renders most model-agnostic techniques as too expensive.

In the pursuit of comprehending the complexity of Deep Neural Networks it makes sense to utilize the weights in the model. The information saved in the hidden layers as learned weights can be used to evaluate the network. Moreover, the gradients can be taken into consideration as well. In the following subsections several concepts for understanding Deep Neural Networks are introduced.

3.3.1 Feature visualization and Network Dissection [25]

Modern Neural Networks like ResNet50 or Bard consist of several million layers. Network dissection attempts to overcome this challenge by breaking down separate layers and connecting them with ideas.

The higher-level features in these networks relate to clear concepts, shown in Figure 3.1. As the input image moves through layers, it changes at each layer. In each convolutional layer, the network gains new and more complex features. The smooth joining of fully connected layers then changes image-based data into predictions.

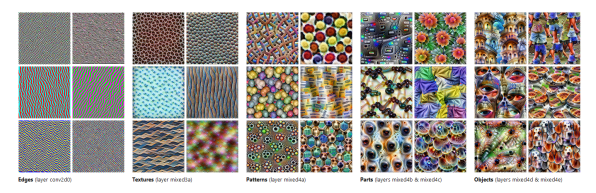


Figure 3.1: Feature Visualization [25]

The image explains this process. The first convolutional layers find simple features like edges and basic textures. Later, they recognize more detailed patterns. The deepest layers learn about parts and objects. This object information passes to the other hidden layers, which then finally make a prediction.

Feature visualization is based on activating one kernel in the network. This involves maximizing the activation of a specific neuron (Visible in 3.2). There are two methods for achieving this. First, we can make use of the training image that triggers the highest activation. Yet, this approach faces a significant problem. When an image contains multiple objects, it's hard to pinpoint which object causes the activation. Because of this an alternative route is adopted: generating new images

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from random noise. This is accomplished through methods like Generative Adversarial Networks (GANs) or other diffusion-based techniques.



Figure 3.2: Activation Maximization [25]

Advantages of Feature Visualization:

- 1. **Initial Model Insights:** Feature visualization offer an initial view into a model's behavior, improving the understanding of its inner layers.
- 2. **Enhanced Domain Understanding:** It has the potential to enrich domain understanding by aligning learned features with domain-specific knowledge. An example can be seen in the medical industry
- 3. **Debugging and Improvement:** Feature visualization assists in debugging and refining models, contributing to their overall performance enhancement.

Disadvantages of Feature Visualization:

- 1. **Unclear Decision-Making:** While activations are evident, understanding the meaning behind them and how they contribute to decision-making remains challenging.
- 2. **Subjective Interpretation:** The interpretation of visualized features can be subjective, potentially leading to differing conclusions among observers.
- 3. **Limited Applicability to Visual Data:** Feature visualization's applicability is limited to visual data types.

3.4 Saliency Maps

Saliency maps are visualizations that highlight the regions of an input image that have the most significant impact on a model's output. By revealing the areas that strongly influence a prediction, saliency maps bridge the gap between the model's "black-box" nature and human understanding.

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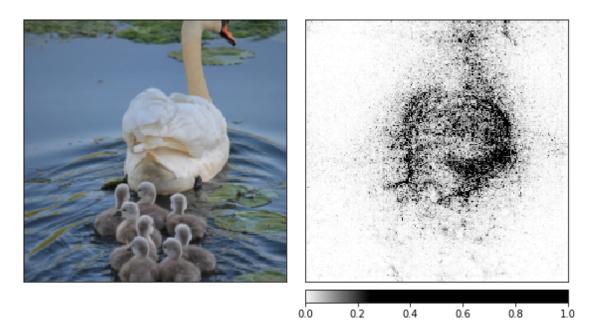


Figure 3.3: Saliency Map - Source: https://captum.ai/tutorials/Resnet_TorchVision_Interpret

Saliency maps are commonly calculated using SHAP [20] or gradient methods. Saliency maps provide a direct and intuitive way to understand which parts of an input data are influencing a model's decision. The main difficulty is the generation of reliable saliency maps.

3.4.1 Vanilla Gradient & Deconv Net

Vanilla Gradient[31] focuses on computing the gradients of the network. A forward pass of an image is generated. The gradients of the class score is computed. Then the gradients are visualized. It is the same process as neural network backpropogation. This method has 2 problems: When ReLU is used and the gradients are negative, then information is lost. Furthermore, in pooling layers, there are no gradients and the information is lost.

Deconv Net [34] takes care of the gradient problem of Pooling and Convolutional Layers.

A standard convolutional network with convolutional layers and fully connected layers and a activation funtion is chosen. An image input image x_i maps to a probability layer y_i . A deconvnet [35] is attached to each of it's layers proving a path back to the image pixels. Unpooling, Rectifying and filtering is done to reconstruct the activity in the layers.

Unpooling: The pooling operation is non-invertible. By recording the original locations of the maxima using a set of switch variables, the reconstructions can be created.

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Rectification: Each layer passes through a relu non-linearity. Each layer backwards also passes through a relu non-linearity to keep all signals positive.

Filtering: Filtering is done by using inverted original filters. The devoncnet uses transposed versions of the same filters. This means flipping each filter vertically and horizontally.

3.4.2 Grad-CAM, Guided Grad-CAM and Smooth Gradient

Gradient-weighted Class Activation Map (Grad-CAM) provides visual explanations for CNN decisions. The goal is to understand at which parts of an image a convolutional layer looks for a certain classification.

$$\alpha_k^c = \frac{1}{Z} \sum_i \sum_j \frac{\delta y^c}{\delta A_{ij}^k}$$

 $c = class of interest A^k = feature map i, j = width and height dimension Z = average term$

GRAD-CAM sets all classes which are not interesting to 0. Then it backpropagates the gradients of the main class. Then the gradients are scaled between [0,1] and displayed. GRAD-CAM does take all convolutional feature maps but the last. This is done, because the last features are responsible for all classes and are therefore not informative. A main problem of GRAD-Cam is that it is very coarse.

Guided Grad-CAM computes Grad-CAM with another method to have a better localization. It's basically multiplied with another method to achieve stable results. GRAD-Cam works as a lenses to focus on specific parts of the pixel-wise attribution map.

SmoothGrad [smilkov2017smoothgrad] is a smoothing method for other gradient calculation methods. By adding noise to the input image and calculating several input images and taking the gradient, noise should cancel out and be removed.

3.4.3 Top-Down Neural Attention by Excitation Backprop [36]

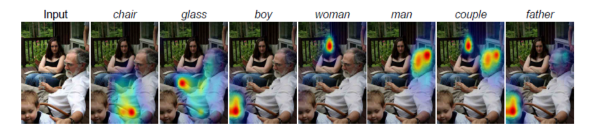


Figure 3.4: CNN classifier's top-down attention map [36]

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By calculating the importance of a neuron set and saving it during the computational step, the gradient step can be combined with the importance and localize exact objects. The probabilistic WTA formulation produce well-normalized attention maps that enable direct subtraction.

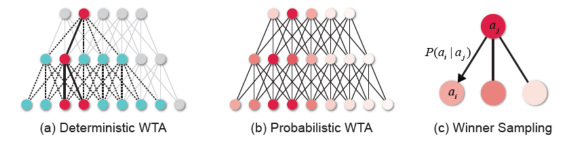


Figure 3.5: Identifying task-relevant neurons in the network. The red shading of a dot indicates its relative likelihood of winning against the other ones in the same layer. [36]

3.4.4 Advantages and Disadvantages of Saliency Maps

The advantage of saliency maps is that they are fast to compute and easy to interpret. The explanations are visual and can immediately be recognized. Furthermore, there are many methods to choose from.

The main disadvantage is that it is unclear whether an explanation is correct or not. In the next chapter possible evaluation methods are explained.

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4 Evaluation of post-hoc interpretability methods

Gradient-methods which generate saliency maps are hard to measure. While humans can evaluate the maps by giving a general statement, this can not be applied to thousands of images. Despite many significant recent contributions to saliency maps, the valuable effort of explaning machine learning models face this methodological challenge: the difficulty of assessing the scope and quality of model explanations [1].

4.1 Evaluation metrics, ground truths [13]

Different evaluation methods exist to evaluate saliency methods. Evaluations can be extrinsic [13], involving human evaluations and comparing the results to certain ground truth explanations [36]. Intrinsic methods use computations involving the net itself and the saliency map, without human evaluation or retraining a net. These methods are based on creating a new composite input using the heat map and the original input. Then they are evaluated using the original trained model. E.g.[7] These methods suffer from violating one key assumption in machine learning: the training and evaluation data must come from the same distribution. [15] Without re-training it is not clear if the degrade in performance stems from the distribution shirt or because informative features were removed.

[24] states that comparing the rank correlation between attention weights and feature-additive methods is not appropriate. This means, using agreement as evaluation (using a method as standard to compare to) should not be used. In the paper a low empirical evidence was found, that models are similar. This means that just evaluating similar methods and taking the common results is not applicable. More rigorous defined metrics need be considered.

4.2 Completeness and Soundness

[13] proposes Soundness and Completeness, two concepts which are required for evaluation metrics which involve using a composition of a heat map and the original input.

Soundness is needed: The masked input method proves that a certain part of the image "caused" the networks output.

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Completeness means, for any composition of an input image of label a with a well functioning saliency mask the model must still be able to identify the correct label.

$$\forall x : f(x \to a)$$

$$x = input(a) \odot mask$$

Soundness means, for any composition of an input image of label b with a well functioning saliency mask them model must not return a wrong label.

$$\nexists x: f(x \to a)$$

$$x = input(b) \odot mask$$

The AUC metric [26] of the insertion game:

For s = [1,dim(x)] take the top s pixels as per saliency map m and plot the probability f(x,a) given by the model. The top s pixels of x are retained and the remaining pixels are assigned a default value). Return the area under the curve.

The method is α complete on f,x,a if:

$$g_{AUC}(x,a,m) >= \alpha f(x,a)$$

The method is β sound on f,x,a if:

$$g_{AUC}(x,a,m) <= \frac{1}{\beta} \alpha f(x,a)$$

$$\alpha(x,a) = min(\frac{max(g_{AUC}(x,a,n),\epsilon_1)}{f(x,a)},1)$$

$$\alpha(x,a) = min(\frac{max(f(x,a),\epsilon_2)}{g_{AUC}(x,a,n)},1)$$

Then
$$\alpha(m) = E_x[min_a(\alpha(x,a))]$$

Then
$$\beta(m) = E_x[min_a(\beta(x,a))]$$

If alpha is close to 1: The blocked input predicts correctly.

If beta is close to 1: The blocked input mimics the behavior and is not overconfident.

x: input to the model

a: true label

f: model

m: saliency method.

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 ϵ = term if x is very small

Older evaluation methods only try to maximize the g_{AUC} curve. This completely ignores if the method is overconfident.

Critics to the method: No human interpretability is included. Furthermore, the masked images do suffer from not having the same distribution as in the original trained maps. This could cause a unwanted drop in performance.

4.3 Perturbation based [30]

A heatmap is an array of pixel-wise scores that indicate which pixels are relevant.

This paper describes a method, where the most important pixels are replaced by randomly sampled values. IT then is measured how the values f(x) the label value goes down. Depending on the method used, the method works superior or worse. If the edges are the most important in a saliency map, then deleting them results in a steep decrease.

This method has high computational costs.

4.4 A benchmark for interpretability methods in deep neural networks [15]

In this paper two methods to estimate the effectiveness of saliency maps are presented. "RemOve And Retrain (ROAR)" and "Keep And Retrain (KAR)".

ROAR proposes a numerical solution to evaluate attribution maps. It is a algorithm which estimates the effectiveness of saliency maps. This is done, by removing supposedly informative features from the input and observing the reaction of the neural network. ROAR is applicable to any visual domain. In Section 5, an example for MNIST and Food-101 is done.

RoaR works as follows:

1. Selection of the most important pixels from an attribution map

An attribution map provides a mapping to the most important pixels. Those pixels are ranked based on their importance. Depending on the algorithm, different values differ on the importance. In the case of Integrated Gradient, the pixels with the highest absolute values are considered the most important.

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2. Replacement of x% of the pixels with the mean value

In the Roar algorithm, 0.1, 0.3, 0.5, 0.7 or 0.9 % of the most important pixels are identified and replaced by the mean value. Both the training and the testing data undergo this process.

3. Retraining the model using the modified dataset

To ensure the consistency of the model, retraining is necessary. Afterwards, the model is evaluated using several seed runs on the test set. The same split should be used for all runs. This is crucial, because the training data and the test data must be drawn from the same distribution. Without retraining the model this property is ignored.

4. Comparison of the attribution map with a random baseline

For each training setup, respectively with 0.1, 0.3, 0.5, 0.7 or 0.9 % pixels replaced, a random baseline is considered. The random baseline also replaces the same percentage of pixels. The random baseline is expected to perform worse than a sophisticated method.

5. Evaluation

An attribution method is deemed effective, if it consistently outperforms the random baseline across various setups.

KAR works similar: Instead of removing the most important pixels, the least important pixels are identified. Because KAR performs worse than ROAR, it is not described further.

The conclusion of [15] is that commonly used based estimator, Gradients, Integrated Gradient and Guided BackProp are worse than a random assignment. The effectiveness of Smooth-Grad-Squared and VarGrad was proofed.

4.5 Benchmarking Attribution Methods (BAM) [33]

BAM with relative feature importance explores wether features are important or not. In reality, we do not know how important a feature is. However, we can calculate how important a feature is to model relative to another model. Given the relative feature importance, the metrics compare attributions between pairs of models and pairs of input.

The main idea of BAM is: If you paste a grey square to every training image for all classes, it is expected that this square matters less than the original image region being covered. The same expectation should hold if instead an object (which is not represented by the dataset) is pasted in every image. Any explanations that assign higher attribution to the inserted object than to the original pixels are false positives.

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BAM solves the problem of an distribution shift by inserting objects by only inserting objects which don't strongly change the distribution of the image. E.g. an object with mean similar to the mean of the dataset is chosen.

The BAM algorithm works as follows:

1. BAM dataset construction

The BAM dataset is constructed by pasting object pixels from MSCOCO [18] into scene images from MiniPlaces [37]. An object is re-scaled to between 1/3 to 1/2 of a scene image at a randomly chosen location. Resulting images have an object label and a scene label. Either can be used to train a classifier. Every object class appears in every scene class and vice versa. Scenes which contain original BAM objects are not used.

2. Common features and commonality

Common features are defined as a set of pixels with semantic meaning (e.g. looks like a dog) which commonly appear in all examples of one or more classes. For example, a dog which appears in all bamboo forests is less common than a dog which appears in all images of bamboo forests, bedroom and corn field.

3. 2 Classifiers are trained and attribution maps are created

An object detection classifier and a scene detection classifier are trained. Objects should be significantly more important than the scene to the object detector than to the scene detector. To verify this intuition, the objects are removed. With this knowledge, attribution maps can be tested by checking if the pixels are assigned noticeably higher attributions.

4. Relative importance is calculated.

With this knowledge, the object pixels should be more important than any other pixels for the detection classifier. Additionally for the scene attribution, the attribution maps should be higher if the object pixels are replaced with the original pixels.

The advantage of BAM over ROAR [15] and other methods is the lower computational cost. No retraining or perturbation is required.

4.6 Sanity Checks for Saliency Maps [1]

In this paper an actionable methodology is proposed, which kinds of explanation a given method can and cannot provide. It shows that visual inspection by humans does not provide good information if the explanation is sensitive to the underlying model and data.

Two instances of the framework are tested:

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1. Model Parameter randomization test

The model parameter randomization test compares the output of a salinecy method on a trained model with the output of a randomly initalized untrrained network of the same architecture. The output should differ, otherwise the saliency map is deemed as not helpful.

2. Data randomization test

The data randomization test randomly shuffles the labels of the data and the model is trained on this method. If the saliency maps does not differ to a normally trained model, then the method does not depend on the relationship of the images and labels.

If any of the 2 hypothesis is failed, then the method can safely be rejected. This is a so called sanity check.

Extensive experiments on several explanation methods are done across data sets and model architectures.

On the tested methods, Gradients & GradCam pass the Sanity checks, while Guided Back-Prop & Guided GradCAM fail.

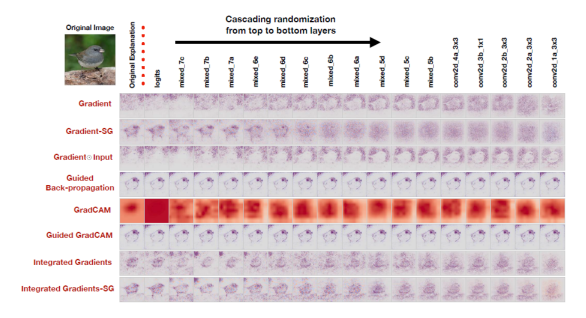


Figure 4.1: Cascading Randomization on Image Net. The figure shows the original explanations. Progression from left to right indicate complete randomization of network weights up to that block inclusive. The last block corresponds to a network with completely reinitialized weights.[1]

4.7 Validating the research

After understanding common machine learning algorithms, one common problem is identified: validating the research [2].

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- 1. Replicability, Reliability and Trust. Evaluation methods need to be replicable to some degree. Predictions and finding should be robust to sensitivity tests like small changes in the data or the model, out-of-sample prediction tests and consistency with domain knowledge [21].
- 2. Why do machine learning interpretations fail?
- a. The machine learning model could be a poor fit for the model and any resulting interpretation approach would poorly reflect the signal in the data.
- b. The interpretation approach could be a poor fit for the model. Especially when fitting a second model to generate the interpretation, like LIME.
- c. There can be a mismatch to the employed interpretation technique and the desired discovery task. For example trying to find out why a model predicts wrongly can't be understood using a poor saliency map.

For prediction tasks well established techniques for validation were developed: Using a split for train, test and validation data. Using this the predictive model should generalize well to new, unseen data. Knowing this, can't one simply employ a training and test set to validate interpretations? [2] This prospect is way more complicated for interpretations. There exists no fair metric that determines which interpretation is the best. Using human evaluation or domain experts could also be an option, but this also struggles from having no good evaluation method. And even experts do not have a common consensus for each dataset.

Validating discoveries from interpretable machine learning is still a challenge and more research is needed to promote trustworthy and reliable data science.

4.8 Research findings: Which evaluation method to use, how to control the result?

Several saliency maps and possible evaluation methods were introduced. Now which one should a data scientist use?

The unsatisfactory answer is: There is no clear right and wrong. Depending on the model and dataset used, different evaluation methods give different results. In [1] and [15] various methods were seen as unsuitable.

Using a SG - GRAD or a VAR-GRAD approach results in a good results for ROAR and SG-Grad also has good results for Sanity Check for saliency maps. Excitation Backprop has not been analyzed yet with this methods, but could also be trustworthy. After all, there is not enough reliable research to say one method is clearly better than another. This can seen as recommendation, but not as clear instruction.

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It remains difficult to evaluate visual explanations and there is no clear truth yet, on which evaluation method is the best. More research is needed to be able to give clear guidelines.

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5 Reconstructing ROAR

5.1 Scientific Motivation and Goal

In RoaR[15] the paper does not list the standard deviation of the trained nets. We expect to validate the results by achieving similar results.

As training 25 image nets requires high computational power we do not have right now, we limited our research to evaluating food-101 [5] using only 2 interpretation methods and comparing it to the baseline.

Additionally, I also add a proof-of-concept evaluation using the MNIST dataset.

5.1.1 Project Setup MNIST

The MNIST dataset [8] was chosen as proof-of-concept. Because calculating the gradients and judging if a mask is meaningful is easier here, it should give a reliable baseline for the results. Only integrated gradients is tested, as is it only an example.

A. Training the model

The model used in this report is a simple convolution neural network architecture consisting of two convolution layers, two pooling layers, and three linear layers. The model was trained for ten epochs with a learning rate of 1e-3 and a SGD optimizer with a momentum term of 0.9. The batch size was 64. Across five separate runs, the model achieved an average accuracy of 97.4% with a standard deviation of 0.4% on the test data.

B. Dataset preparation and Splitting

The MNIST data set consists of 10 classes and 70.000 different images. It was chosen, because it is easy to evaluate the effectiveness of integrated gradients and ROAR: a) The importance of the pixels is clear from a human perspective. b) The model required to achieve a solid performance is very modest and speeds up the retraining. c) The information loss in retraining is clearly visible in the pictures. Furthermore, it is a small, usable data set. The images are 28x28 pixels big and one pixel has a value ranging from [0,1] in gray-scale. The data set used in this report was normalized using the mean and standard deviation, and was split using the PyTorch library's default

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splitting method (60.000 images for the training data, 10.000 images for the test data). The training data was shuffled randomly using the inbuilt functions. The Cross entropy loss was selected for training runs. The same data set split was used for all training runs to ensure consistency in the results.

C. Example of the new dataset

In 5.1 the difference between Integrated Gradients and the Random Baseline is visible. In the first row the random baseline is visible, in the second row Integrated Gradients is shown. [0.1,0.3,0.5,0.7,0.9%] of the pixels are blacked out (left to right).

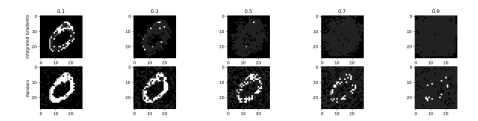


Figure 5.1: Integrated Gradients and Random Baseline Comparison

Judging by human intuition, Integrated Gradients is appearing to perform better over all thresholds. The model is expected to perform worse following retraining.

D. Retraining upon the new datasets

To ensure that the model learns consistently, the same split and learning parameters were utilized as in the standard case to retrain the model on the new data set. An untrained network was created and then retrained for each training run. In order to validate the results and minimize the impact of random seed initialization, the training was performed with 5 different seeds and 5 models were created. In the Appendix, the exact results can be found.

E. Advantages and Disadvantages of the setup

Because of the small size of the data set and the limited amount of classes, training a model is fast and effective. One disadvantage is the limited generalization of this setup for other datasets and models. The majority of use cases primarily involve working with images in the RGB color space, utilizing deep learning models that are often more complex, and dealing with a larger number of distinct classes. Furthermore, the pixels itself are blurred on their importance. In RGB images 3 channels are combined, which make attribution maps more difficult to compute and understand.

F. Results

Integrated Gradient is performing better than the random baseline.

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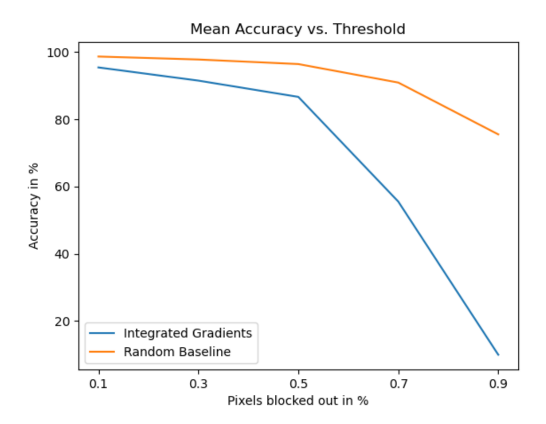


Figure 5.2: Accuracy: Random Baseline vs Integrated Gradient

5.1.2 Project Setup Food-101

The MNIST dataset [8] was chosen as proof-of-concept. Because calculating the gradients and judging if a mask is meaningful is easier here, it should give a reliable baseline for the results. Only integrated gradients is tested, as is it only an example.

A. Training the model

The model used in this report is ResNet50 [14]. The Food-101 [5] dataset was chosen to replicate. Learning rate was adjusted to 0.175, batchsize of 64. Using an SGD optimizer with momentum =0.9 and weight decay = 0.0001 and a scheduler at epoch 30 with gamma=0.1, an accuracy of averagely 70.5% with standard deviation of 0.01 was achieved over 5 separate training runs on the test score. In total 31 epochs were done.

In the original paper an accuracy of 84.54% was achieved on the Food-101 dataset. A smaller accuracy was achieved, as the training was noticeably shorter because only a NVIDIA GTX 1080 was available.

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The data was resized with center-crop to make all images 3x224x224. Afterwards they were normalized. No data augmentation like flips and mirroring was applied. The standard split 75:25 was applied. The altered datasets were calculated with all using the same model.

B. Results

5.1.3 Interpretation of the results

The results from the original paper were confirmed. Integrated Gradients preforms as good as a random baseline for the Food-101 setup. While the MNIST result shows that Integrated Gradient indeed seems to find important pixels, in the Food-101 setup Integrated Gradient fails. Therefore we can say that Integrated Gradient seems to fail for bigger networks.

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