## **MACHINE LEARNING IN BIOINFORMATICS**

Explainability - XAI

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#### INTRODUCTION

- Machine learning studies the relationship between
  - independent or predictor variables X
  - dependent or response variables Y
- Machine learning and statistics may have multiple goals [Zhao and Hastie, 2021]:
  - Prediction: Predict the response variables Y as accurate at possible from X
  - Science: If X → Y is a causal relationship, we may want to understand the *laws of nature* that determine this relationship

#### INTRODUCTION

Two opposing cultures of statistical analysis [Breiman, 2001]:

- ► Data modeling culture: Assume a parametric function f such that  $Y = f(X) + \epsilon$ , where  $\epsilon$  models the aleatoric uncertainty. The parameters of f are often easy to interpret and the model is used to understand the laws of nature
- Algorithmic modeling culture: Use of black-box models that are very complex and optimized to maximize predictive accuracy. Black-box models are notoriously difficult to interpret and do barely allow to draw any conclusions about the laws of nature
- If we have a black-box model, how can we still gain some interpretation?

## OUTLINE I

- Assume we have a black-box machine learning model *f*
- Can we gain some *limited* understanding of the predictions of *f*?
- Understanding the predictions increases our *trust* in *f*

## OUTLINE II

- Given a fixed input x, what is the *contribution* of each feature to the prediction y = f(x)?
   (Attribution Map / Saliency Maps)
  - Occlusion
  - Layer-wise relevance propagation (LRP) / DeepLIFT
  - Integrated gradients
  - Shapley values
  - SHAP
- Given a fixed input x, is there an interpretable model that approximates f locally?

## OUTLINE III

- Local interpretable model-agnostic explanations (LIME)
- Taylor approximations
- What would *f* predict if we vary one or more features?
  - Partial dependence plots (PDP)
  - Individual conditional expectation (ICE)
- What is the most likely input x for a given prediction y = f(x)?
  - Input optimization

# ATTRIBUTION MAPS

#### ATTRIBUTION MAPS

- Attribution maps are very popular with images, where the attribution of each pixel can be easily visualized
- Each input feature is assigned an attribution score (feature attribution)



[Kim et al., 2019]

## OCCLUSION I

- Occlusion is a perturbation method that masks part of the input and measures the effect on the output of the network [Ancona et al., 2017]
- This method requires to evaluate the model for many perturbations
- The size of the mask is of particular importance



## OCCLUSION II

- Assume we have a network that detects whether an image contains a cat
- If there are multiple cats in the image, occluding parts of the image with image patches (i.e. occluding at most one cat at a time) will not change the output of the network
- In this case, we would require masking multiple regions at the same time
- This leads to a combinatorial explosion

## **GRADIENT BASED EXPLANATIONS**

#### LOCAL EXPLANATIONS



- Given a fixed input x and the corresponding output y = f(x), what input features contribute most to the output value y?
- Note that for many applications (e.g. images) it is not very valuable to know which features (e.g. pixels) contribute most to the output of a neural network unless a specific input is considered
- The provided level of interpretability is hence limited to individual input data points

### **GRADIENT** I

- Let f be a neural network or any other differentiable machine learning model
- Using the first-order Taylor expansion of f at an input  $\tilde{x}$  we approximate f as a linear function

$$f(x) \approx f(\tilde{x}) + \nabla_x^\top f(\tilde{x})(x - \tilde{x})$$

[Simonyan et al., 2013]

• With 
$$w = \nabla_x f(\tilde{x})$$
 and  $x' = x - \tilde{x}$  we obtain

$$f(\mathbf{x}') \approx f(\tilde{\mathbf{x}}) + \mathbf{w}^{ op} \mathbf{x}'$$

where the gradient *w* can be easily interpreted as *feature importances* 

#### $\mathsf{Gradient} \times \mathsf{input} \, \mathsf{I}$

- Using the gradient alone is problematic
- Let the network be defined as

$$f(x) = \max\{0, x - 10\}$$

i.e. a single linear unit with ReLU activation

The gradient is given by

$$abla_x f(x) = egin{cases} 1 & ext{if } x > 10 \ 0 & ext{otherwise} \end{cases}$$

■ In this simple example, the larger x the larger the output y = f(x) (assuming x > 10)

- However, for f(20) we obtain the same attribution value as for f(1000), i.e. 1 in both cases
- Multiplying the gradient with the input x seems to improve results [Shrikumar et al., 2016]
- For f(20) we would obtain 20 as attribution value, whereas for f(1000) the attribution is 1000

#### INTEGRATED GRADIENTS |

 Integrated gradients (IG): Consider the gradient along an entire path from a baseline x<sub>o</sub> to an input x̃
 [Sundararajan et al., 2017]

$$\mathrm{IG}_{j}(\tilde{x}) = (\tilde{x}^{(j)} - x_{\mathsf{o}}^{(j)}) \int_{[\mathsf{o},\mathsf{1}]} \frac{\partial f(\alpha \tilde{x} + (\mathsf{1} - \alpha) x_{\mathsf{o}})}{\partial \tilde{x}^{(j)}} \mathsf{d}\alpha$$

IG satisfies several convincing axioms that other methods violate

#### Axiom 1: Sensitivity

Assume that  $x_0$  and  $\tilde{x}$  differ in at least one feature and that  $f(x_0) \neq f(\tilde{x})$ . Clearly the features that differ between  $x_0$  and  $\tilde{x}$  have some influence on the prediction. Hence, non-zero attribution should be given to these features

- $\blacksquare$  Gradient and Gradient  $\times$  input fail this axiom
- Consider the following example with just one feature

$$f(x) = 1 - \operatorname{ReLU}(1 - x) \xrightarrow[-10]{0.5}{0.50} - \frac{0.5}{0.50} - \frac{0$$

#### INTEGRATED GRADIENTS III

- For  $x_0 = 0$  and  $\tilde{x} = 2$  we obtain  $f(x_0) = 0$  and  $f(\tilde{x}) = 1$
- However, the gradient at  $\tilde{x} = 2$  is zero
- The sensitivity axiom is the complement of the dummy property of Shapley values

#### Axiom 2: Implementation invariance

Let f and f' be two machine learning models. The two models are functionally equivalent if f(x) = f'(x) for all x. Attributions should be identical for functionally equivalent models

- Gradients are invariant to implementations
- Several attribution methods fail this axiom, e.g. LRP and DeepLIFT

#### Axiom 3: Linearity

Let f be a machine learning model such that

$$f(x) = af_1(x) + bf_2(x)$$

where *a* and *b* are weights. The attribution for *f* is the sum of attributions of  $f_1$  and  $f_2$  weighted by *a* and *b* 

 Attribution methods should preserve any linearity in the machine learning models

#### **Axiom 4: Completeness**

Let f be a machine learning model. The attribution of all features at a point  $\tilde{x}$  should sum up to

$$f(\tilde{x}) - f(x_0)$$

where  $x_0$  is a baseline

The completeness axiom corresponds to the efficiency property of Shapley values for  $f(x_0) = \mathbb{E}f(X)$ 

# LAYER-WISE RELEVANCE PROPAGATION (LRP)

## LAYER-WISE RELEVANCE PROPAGATION (LRP) I

- Layer-wise relevance propagation (LRP) [Bach et al., 2015] exploints the layered structure of neural networks
- Let *f* be a neural network with *L* layers
- $R^{(l)} \in \mathbb{R}^{p_l}$  denotes a vector of relevance scores, one for each neuron in the *l*-th layer
- LRP satisfies the following law of conservation

$$f(x) = \dots = \sum_{j=1}^{p_{l+1}} R_j^{(l+1)} = \sum_{j=1}^{p_l} R_j^{(l)} = \sum_{j=1}^{p_1} R_j^{(1)}$$

where  $R_j^{(1)}$  are the relevances of the input features

## LAYER-WISE RELEVANCE PROPAGATION (LRP) II

- The output of the neural network f for a given input x is the total relevance, which is distributed among neurons in previous layers
- More specifically, we call any relevance attribution method LRP if it satisfies

$$R_{i}^{(l)} = \sum_{\substack{k:i \to k}} R_{i \leftarrow k}^{(l,l+1)}$$
$$R_{k}^{(l+1)} = \sum_{\substack{i:i \to k}} R_{i \leftarrow k}^{(l,l+1)}$$

where  $R_{i \leftarrow k}^{(l,l+1)}$  is the relevance sent from neuron k to i between layers l and l + 1

## LAYER-WISE RELEVANCE PROPAGATION (LRP) III

- The relevances are in both the forward and backward direction sums of the relevances from connecting neurons
- Multiple solutions satisfy these constraints [Montavon et al., 2019], e.g.:

► Basic rule (LRP-0)

$$R_{i\leftarrow k}^{(l,l+1)} = R_k^{(l+1)} \frac{a_i w_{ik}}{\sum_j a_j w_{jk}}$$

Epsilon rule (LRP- $\epsilon$ )

$$R_{i\leftarrow k}^{(l,l+1)} = R_k^{(l+1)} \frac{a_i w_{ik}}{\epsilon + \sum_j a_j w_{jk}}$$

 a<sub>i</sub> denotes the activation of neuron *i* (i.e. the output of a neuron before the non-linear activation is applied)  DeepLIFT [Shrikumar et al., 2017] is an extension of LRP that backpropagates relevance values of

$$f(\tilde{x}) - f(x_0)$$

where  $x_0$  is a user defined point that provides a baseline prediction (note that LRP uses  $f(x_0) = 0$ )

# LOCAL INTERPRETABLE MODEL-AGNOSTIC EXPLANATIONS (LIME)

## LIME - BASIC IDEA

- Local interpretable model-agnostic explanations (LIME)
- Model-agnostic: We can evaluate the model f but do not make any further assumptions about the model
- In particular, we do not require the model to be differentiable
- LIME locally approximates a machine learning model f using a simple interpretable model g at a specific point x such that

$$f(\tilde{x}) = g(\tilde{x})$$

and  $f(x) \approx g(x)$  whenever x is close to  $\tilde{x}$ 

■ *g* is typically a linear regression model

## LIME - BASIC IDEA

Given a model class G, we determine a local interpretable model ĝ by solving

$$\hat{g} = rgmin_{g\in \mathsf{G}} \mathcal{L}(f, g, \pi_{\widetilde{X}}) + \Omega(g)$$

- G could be the class of linear models
- $\blacksquare \ \mathcal{L}$  is the main loss function we want to minimize
- Ω(g) is a penalty on the complexity of g, which for instance gives preference to models with fewer parameters
- $\pi_{\tilde{x}}(x)$  is a weight function that measures the proximity of x to  $\tilde{x}$ , whereby a local approximation of f is enforced

#### LIME - BASIC IDEA

- Let  $x_1, \ldots, x_n$  be a set of *n* training points
- $\blacksquare$  For regression problems, the  $\mathcal L$  function can be

$$\mathcal{L}(f, g, \pi_{\tilde{X}}) = \sum_{i} \left( f(x_i) - g(x_i) \right)^2 \pi_{\tilde{X}}(x_i)$$

The weight function can be given by an exponential kernel

$$\pi_{\tilde{x}}(x_i) = \exp\left\{-\frac{d(\tilde{x}, x_i)^2}{\sigma^2}\right\}$$

where *d* is a distance function

 $\blacksquare \ \sigma^{\rm 2}$  controls how local the approximation should be

- The data used for training f might not be appropriate for estimating g
- We require samples (*x<sub>i</sub>*)<sub>*i*</sub> close to *x̃*, where LIME uses the following scheme:
  - Draw a binary vector b<sub>i</sub> of length p at random
  - Compute  $x_i = h_{\tilde{x}}(b_i)$
- The function  $h_{\tilde{x}}$  creates a sample  $x_i$  from  $b_i$  by masking parts of the reference  $\tilde{x}$  (occlusion)

## LIME - IN PRACTICE II

For instance,  $h_{\tilde{x}}$  could mask features by replacing them with feature means  $\bar{x}^{(j)}$ , i.e. the *j*th feature of  $x_i = h_{\tilde{x}}(b_i)$  is given by

$$x_i^{(j)} = \begin{cases} \tilde{x}^{(j)} & \text{if } b_i^{(j)} = 1\\ \bar{x}^{(j)} & \text{if } b_i^{(j)} = 0 \end{cases}$$

## LIME - IN PRACTICE III

- Example topic models:
  - A data point x is a word count vector, where each entry x<sup>(j)</sup> denotes the number of occurrences of word j in a text document
  - $h_{\tilde{x}}(b_i)$  could mask features by replacing them with zeros, i.e.

$$h_{\tilde{x}}(b_i) = \begin{cases} \tilde{x}^{(j)} & \text{if } b_i^{(j)} = 1 \\ 0 & \text{if } b_i^{(j)} = 0 \end{cases}$$

Hence, samples x<sub>i</sub> are created from the reference x̃ by replacing some of the counts with zeros

## LIME - IN PRACTICE IV

- Example image classification:
  - x is a an image, where each entry x<sup>(j)</sup> denotes a pixel or super-pixel
  - A sample x<sub>i</sub> = h<sub>x̃</sub>(b<sub>i</sub>) consists of the reference image x̃ where a some pixels or super-pixels have been masked, as defined by the binary vector b<sub>i</sub>



**Original Image** 



Interpretable Components

- The interpretable model g is typically defined on the binarized points b<sub>i</sub>
- The loss for regression problems then becomes

$$\mathcal{L}(f, g, \pi_{\tilde{X}}) = \sum_{i} \left( f(x_i) - g(b_i) \right)^2 \pi_{\tilde{X}}(x_i)$$
# **SHAPLEY VALUES**

#### **SHAPLEY VALUES - MOTIVATION**

Assume a linear model

$$f(x) = \theta_1 x^{(1)} + \theta_2 x^{(2)} + \dots + \theta_p x^{(p)}$$

- If features are standardized we can interpret the coefficient θ<sub>i</sub> as the global importance of the jth feature
- Given a specific input *x*, the contribution of feature *j* to the prediction *f*(*x*) is given by

$$egin{aligned} \phi_j(f, \mathbf{x}) &= heta_j \mathbf{x}^{(j)} - \mathbb{E}\left[ heta_j \mathbf{X}^{(j)}
ight] \ &= heta_j\left(\mathbf{x}^{(j)} - \mathbb{E}\left[\mathbf{X}^{(j)}
ight]
ight) \end{aligned}$$

assuming features are independent

# SHAPLEY VALUES - MOTIVATION I

- For non-linear models we need a more advanced definition
- Let *F* denote the set of *p* features and  $S \subseteq F$  a subset
- Furthermore, let f(x<sup>(S)</sup>) be the prediction of a machine learning model where only a subset of features S is used
- Let *S* = *F*\{*j*}, then the contribution of the *j*th feature can be measured as

$$f(x^{(S \cup \{j\})}) - f(x^{(S)})$$

- $f(x^{(S \cup \{j\})})$  is the prediction with feature *j*
- $f(x^{(S)})$  the prediction without feature j

- In practice, features are rarely independent, i.e. feature j might only be informative in combination with other features
- In this case we have to attribute some of feature *j*th contribution to those features
- We have to test for all subsets  $S \subseteq F \setminus \{j\}$

#### SHAPLEY VALUES - EXAMPLE I

- Let the feature set *F* consist of p = 3 elements, i.e.  $F = \{1, 2, 3\}$
- Assume we observe the following predictions

$$\begin{array}{l} f(x^{\{1\}}) = 100, \quad f(x^{\{1,2\}}) = 500, \\ f(x^{\{2\}}) = 100, \quad f(x^{\{1,3\}}) = 300, \quad f(x^{\{1,2,3\}}) = 1100 \\ f(x^{\{3\}}) = 100, \quad f(x^{\{\{2,3\}\}}) = 300, \end{array}$$

- Clearly, features are not contributing independently to the predictions
- For independent features we would expect

$$f(x^{(\{1,2\})}) = f(x^{(\{1\})}) + f(x^{(\{2\})})$$

#### SHAPLEY VALUES - EXAMPLE II

- How much should we attribute to each feature?
- We fix a particular feature j and evaluate its contribution to all subsets  $S \subseteq F \setminus \{j\}$
- To simplify notation, let

$$\xi_j(S) = f(x^{(S \cup \{j\})}) - f(x^{(S)})$$

• For j = 3 and  $S = \{1, 2\}$  we have

$$\xi_j(S) = 1100 - 500 = 600$$

For j = 2 and  $S = \{3\}$  we have

$$\xi_j(S) = 300 - 100 = 200$$

#### SHAPLEY VALUES - EXAMPLE III

- $\xi_j(S)$  denotes the contribution of feature *j* to the prediction based on features *S*
- The Shapley value for feature j is the average over all contributions
- We evaluate all *p*! permutations of *p* features, i.e.

1, 2, 3 1, 3, 2 2, 1, 3 2, 3, 1 3, 1, 2 3, 2, 1

### SHAPLEY VALUES - EXAMPLE IV

- A permutation is interpreted as a sequence of features entering the set of features S
- For instance, for 2, 1, 3 we first have feature 2 entering S and afterwards feature 1. Feature 3 is the last to join S
- We then evaluate the contribution of each feature, i.e. for 2, 1, 3 we evaluate  $\xi_2(\{\}), \xi_1(\{2\})$ , and  $\xi_3(\{1,2\})$

#### Shapley values - Example V

	j = 1		j = 2		j = 3	
1,2,3	$\xi_1(\{\})$	= 100	$\xi_2(\{1\})$	= 400	$\xi_3(\{1,2\})$	= 600
1,3,2	$\xi_1(\{\})$	= 100	$\xi_2(\{1,3\})$	= 800	$\xi_3(\{1\})$	= 200
2,1,3	$\xi_1(\{2\})$	= 400	$\xi_2(\{\})$	= 100	$\xi_3(\{1,2\})$	= 600
2,3,1	$\xi_1(\{2,3\})$	= 800	$\xi_2(\{\})$	= 100	$\xi_3(\{2\})$	= 200
3,1,2	<i>ξ</i> <sub>1</sub> ({3})	= 200	$\xi_2(\{1,3\})$	= 800	$\xi_3(\{\})$	= 100
3,2,1	$\xi_1(\{2,3\})$	= 800	$\xi_2(\{3\})$	= 200	$\xi_3(\{\})$	= 100

- The rows are the permutations, the columns represent features to enter the set S
- The Shapley value  $\phi_j(f, x)$  for feature *j* is the average over all p! = |F|! rows in column *j*
- Hence, permutations are assumed to be uniformly distributed

- How often do we observe a particular entry  $\xi_j(S)$  in column *j*?
- We can permute all features before j enters and all features after j enters
- Hence, an entry  $\xi_j(S)$  occurs

|S|!(|F| - |S| - 1)!

times in column j

# Shapley value [Shapley, 1951]

The shapley value for the *j*th feature is defined as

$$\begin{split} \phi_j(f,x) &= \sum_{S \subseteq F \setminus \{j\}} \frac{|S|!(|F| - |S| - 1)!}{|F|!} \xi_j(S) \\ &= \sum_{S \subseteq F \setminus \{j\}} \frac{|S|!(|F| - |S| - 1)!}{|F|!} \left( f(x^{(S \cup \{j\})}) - f(x^{(S)}) \right) \end{split}$$

- The sum is over  $2^{p-1}$  permutations
- For large feature sets the Shapley value is computationally very expensive or even impossible to compute

■ Assume *f* is a linear model of the from

$$f(x) = \theta_1 x^{(1)} + \theta_2 x^{(2)} + \dots + \theta_p x^{(p)}$$

 Given independent features, the Shapley values for this model reduce to

$$\phi_j(f, \mathbf{X}) = \theta_j\left(\mathbf{X}^{(j)} - \mathbb{E}\left[\mathbf{X}^{(j)}\right]\right)$$

[Štrumbelj and Kononenko, 2014]

This is what we expected from our previous discussion

$$\sum_{j} \phi_{j}(f, x) = f(x) - \mathbb{E}_{X} f(X)$$

Symmetry: If two features *j* and *k* contribute equally to all subsets, then

$$\phi_j(f, \mathbf{x}) = \phi_k(f, \mathbf{x})$$

for all x

Dummy: If feature *j* does not influence the prediction  $f(x^{(S)})$  for all *S*, then

$$\phi_j(f, \mathsf{X}) = \mathsf{O}$$

• Additivity: If  $f(x) = \sum_m f_m(x)$  then

$$\phi_j(f,x) = \sum_m \phi_j(f_m,x)$$

i.e. f could be a random forest or any other bagging method

# SHAPLEY VALUES - IN PRACTICE I

- How do we remove features from the prediction of our machine learning model f?
- The optimal but impractical way would be to train a model fs for each subset S
- Instead, we often use

$$f(\mathbf{X}^{(\mathsf{S})}) = \mathbb{E}\left[f(\mathbf{X}) \,|\, \mathbf{X}^{(\mathsf{S})} = \mathbf{x}^{(\mathsf{S})}\right]$$

where all elements of X that are not given by  $\{X^{(S)} = x^{(S)}\}$  are considered random

The expectation can be estimated from our training data, which however requires many evaluations of the model f

#### SHAPLEY VALUES - IN PRACTICE II

Assuming that our model f is linear, we obtain

$$f(\mathbf{x}^{(5)}) = \mathbb{E}\left[f(X) \,|\, \mathbf{X}^{(5)} = \mathbf{x}^{(5)}\right] = f\left(\mathbb{E}[X \,|\, \mathbf{X}^{(5)} = \mathbf{x}^{(5)}]\right)$$

Furthermore, assuming independent features we obtain

$$f(x^{(S)}) = f(\bar{x}^{(S)})$$

where

$$ar{\mathbf{x}}^{(\mathsf{S})} = egin{cases} \mathbf{x}^{(j)} & ext{if } j \in \mathsf{S} \ \mathbb{E} \, \mathbf{X}^{(j)} & ext{if } j \notin \mathsf{S} \end{cases}$$

i.e. all features not in S have been replaced by their expectation

# SHAPLEY VALUES - MONTE CARLO I

- Summing over 2<sup>*p*-1</sup> contributions is often too expensive
- We may utilize Monte Carlo approximations (law of large numbers) to estimate the Shapley value [Štrumbelj and Kononenko, 2014]
  - ▶ Draw *k* permutation  $\pi_i = (r_1, ..., r_p)$  with  $r_m \in \{1, ..., p\}$  from a uniform distribution
  - For each permutation π<sub>i</sub>, compute the set of features S<sub>ij</sub> from π<sub>i</sub>, i.e. all features until feature j occurs in π<sub>i</sub>
  - The Monte Carlo approximation of the Shapley value is given by

$$\phi_j(f, \mathbf{x}) \approx \frac{1}{k} \sum_{i=1}^k \left( f(\mathbf{x}^{(\mathsf{S}_{ij} \cup \{j\})}) - f(\mathbf{x}^{(\mathsf{S}_{ij})}) \right)$$

### SHAPLEY VALUES - KERNEL SHAP I

- SHapley Additive exPlanations (SHAP) [Lundberg and Lee, 2017]
- Kernel SHAP reformulates the computation of Shapley values as a linear regression problem using the LIME framework
- The interpretable model g is assumed to be a linear regression model

$$g(b_i) = \phi_{\mathsf{o}} + \sum_{j=1}^{p} \phi_j b_i^{(j)}$$

i.e. the contributions of the linear model depend on the weights  $\phi_j$  and the binary values  $b_i^{(j)}$ 

# SHAPLEY VALUES - KERNEL SHAP II

**The weights**  $\phi_i$  are the Shapley values

Notice that LIME with loss function

$$\mathcal{L}(f, g, \pi_{\tilde{x}}) = \sum_{i} \left( f(x_i) - g(b_i) \right)^2 \pi_{\tilde{x}}(x_i)$$

and  $\Omega(g) = 0$  corresponds to weighted ordinary least squares

$$\hat{\theta} = \arg\min_{\theta} \left\| W^{1/2} (y - X\theta) \right\|_{2}^{2}$$
$$= (X^{\top} WX)^{-1} X^{\top} Wy$$

where  $X \in \{0, 1\}^{2^{p} \times p}$  denotes a matrix containing all possible binary vectors  $b_i$  of length p as rows,  $W = (w_{ij})$  is a weight

### SHAPLEY VALUES - KERNEL SHAP III

matrix with  $w_{ii} = \pi_{\tilde{x}}(x_i)$  and  $y = (y_i)$  is the vector of targets  $y_i = f(x_i)$ 

The coefficients  $\hat{\theta}$  are the Shapley values  $\phi = (\phi_1, \dots, \phi_p)$  for

$$\pi_{\tilde{x}}(x_i) = \frac{p-1}{\binom{p}{k_i}k_i(p-k_i)}$$

where  $k_i = |b_i|$  is the number of ones in the binary representation of the *i*th sample

• Notice that  $|b_i|$  measures the similarity between  $x_i$  and  $x_i$ , therefore this particular choice of  $\pi_{\tilde{x}}$  is indeed a weight based on a distance measured

# SHAPLEY VALUES - KERNEL SHAP IV

- Recall that the *i*th sample x<sub>i</sub> is generated from x by randomly generating a binary representation b<sub>i</sub> and afterwards masking all features j in x̃ where b<sub>i</sub><sup>(j)</sup> = 0
- The linear regression coefficients θ correspond to the Shapley values φ only when we consider all possible binary vectors b<sub>i</sub>
- In practice, Kernel SHAP uses a sampled subset of binary vectors
- An improved method has been proposed [Kwon and Zou, 2022]



# **PARTIAL DEPENDENCE PLOT**

- Let *f* be a black-box model such as a neural network
- What is the effect of individual predictors X<sup>(j)</sup> on the response variable Y as captured by our model f?

Partial dependence plots (PDP) [Friedman, 2001]:

$$\mathrm{PDP}_{j}(x) = \int f(x, x^{(-j)}) \mathrm{pr}(x^{(-j)}) \mathrm{d}x^{(-j)}$$

where  $x^{(-j)} = (x^{(1)}, \dots, x^{(j-1)}, x^{(j+1)}, \dots x^{(p)})$ 

■ In practice we use the training data  $(x_i, y_i)_{i=1}^n$  to estimate the PDP, i.e.

$$\widehat{\text{PDP}}_j(x) = \frac{1}{n} \sum_{i=1}^n f(x, x_i^{(-j)})$$

- Boston housing data: Housing data for 506 census tracts of Boston from the 1970 census
- X: capita crime rate, proportion of non-retail business acres per town, nitric oxides concentration, average number of rooms per dwelling, proportion of owner-occupied units built prior to 1940, ...
- Y: median value of owner-occupied homes in USD 1000's

#### PARTIAL DEPENDENCE PLOT - EXAMPLE

Partial Dependence Plot (PDP)



Housing prices drop when nitric oxides concentration reaches  $\sim$  0.68

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#### PARTIAL DEPENDENCE PLOT - EXAMPLE

Partial Dependence Plot (PDP)



Housing prices drop quickly with crime rate

# PARTIAL DEPENDENCE PLOT - ICE

- The individual conditional expectation (ICE) is an extension of the PDP
- It plots each component of the PDP sum individually, i.e.

$$\widehat{\mathrm{ICE}}_{ij}(x) = f(x, x_i^{(-j)})$$

Hence, we have

$$\widehat{\mathrm{PDP}}_{j}(x) = \frac{1}{n} \sum_{i=1}^{n} \widehat{\mathrm{ICE}}_{ij}(x)$$

# **INPUT OPTIMIZATION**

- Assume that *f* is a classifier for images
- We want to find inputs x not contained in the training set that correspond to predictions of a given classification
- This analysis might help to understand if f is sensitive to the correct features
- For a given output *y* we solve the optimization problem

$$\hat{x} = \underset{x}{\operatorname{arg\,min}} \mathcal{L}(f(x), y)$$

■ The loss function *L* typically corresponds to the loss function used for training *f* 

- As for training f we may use gradient descent to compute  $\hat{x}$
- The result x̂ depends strongly on the initial value for solving the optimization problem
- Using multiple initial conditions allows to generate multiple inputs (x<sub>i</sub>)<sub>i</sub> corresponding to the same prediction y

#### SHAP:

https://shap.readthedocs.io

- iNNvestigate (Keras/Tensorflow):
   https://github.com/albermax/innvestigate
- Captum (PyTorch): https://github.com/pytorch/captum

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