Measuring Feature Importance Through Counterfactual Distributions

Abstract

Understanding the importance of individual features in machine learning models is critical for interpretability, especially in a field where no definitive ground-truth exists. This paper provides an alternative perspective by proposing a novel local feature importance method that can be applied to any model. The general idea is to obtain two sets of positive and negative counterfactuals, estimate their underlying distributions with Kernel Density Estimations (KDE), and rank the features where positive and negative counterfactuals differ the most. We anchor our approach within a solid mathematical framework, demonstrating that it satisfies key properties to serve as a measure of dissimilarity and distance.

We demonstrate the effectiveness of our method by illustrating and comparing the results with traditional and well-established local feature importance scores. Furthermore, we incorporate the faithfulness metric as an additional point of comparison, complementing our analysis and providing a more comprehensive evaluation of the differences and similarities between metrics. Our findings highlight the discordance between different metrics for estimating feature importance, underscoring the necessity of employing multiple approaches to capture various mathematical and distributional aspects of the data.

1 INTRODUCTION

Explainable AI (XAI) addresses one of the most pressing challenges in modern machine learning: understanding and interpreting how a model produces its outputs. With the extensive adoption of black-box models in sensitive and high-stakes domains—such as medical diagnosis, credit scoring,

and criminal justice—the need for interpretable methods has become not merely desirable but indispensable. In these contexts, the consequences of opaque decision-making can be severe, ranging from ethical concerns to life-altering outcomes (see [Solon Barocas, 2023], [Ingram, 2020], [Balasubramaniam et al., 2023]). One way to tackle this problem is to detect which features are more involved in generating the output. To do so, the scientific community mostly developed two classes of methods: intrinsic and post-hoc methods. Intrinsic methods leverage the internals of a given classifier to measure the degree to which each feature contributes to the model's predictions, whereas post-hoc methods operate independently of the classifier by analyzing its outputs. The choice of method type is subject of scientific debate, with conflicting opinions and arguments ([Rajbahadur et al., 2022]). Among the various post-hoc methods proposed, counterfactual (CF) explanations, introduced by Wachter et al. ([Wachter et al., 2017]), have emerged as a common apporach for model interpretability. A CF explanation describes a causal situation in the form: "If X had not occurred, Y would not have occurred". Counterfactuals are humanfriendly explanations, because they are contrastive to the current instance and because they are selective, meaning they usually focus on a small number of feature changes. In mathematical terms, let us consider a black-box model $M \colon \mathbb{R}^d \to \{0,1\}$ and a certain input $x \in \mathbb{R}^d$ for which the output y = M(x) is observed. Counterfactuals are solutions

$$x_{cf} = \operatorname*{arg\,min}_{x'} \mathcal{L}(M(x'), y_{target}) + \lambda d(x, x')$$

where \mathcal{L} is a loss function, for instance $(M(x')-y_{target})^2$, λ is a regularization parameter, y_{target} is the desired outcome and $d(\cdot,\cdot)$ is a distance function function ensuring that x and x' remain close. Building on this foundation, [Dandl et al., 2020] proposed an alternative formulation that minimizes a four-objective loss function:

$$\begin{split} &L(x,x',y',X^{obs}) = \\ &= (o_1(\hat{f}(x'),y'),o_2(x,x'),o_3(x,x'),o_4(x',X^{obs})) \end{split}$$

where o_i are proper objective function that encodes desirable properties on the counterfactual: o_1 reflects that the prediction of our counterfactual x' should be close to the desired outcome, o_2 encodes the similarity between x' and x, o_3 sparsity and o_4 that counterfactuals have likely feature values/combinations. From an operational point of view, when generating CFs, we provide a list of x_{cf} based on different notions as proximity, diversity, sparsity, and actionability (see papers [Mothilal et al., 2020], [S., 2023] and [Karimi et al., 2019]). The generation of CFs often relies on techniques such as trial and error or optimization algorithms like NSGA-II ([Deb et al., 2002]). The advantages and limitations of CFs have been widely studied in recent years. Please refer to [Verma et al., 2020] for a wider discussion. In this work, we propose a novel approach to rank dimensions where positive and negative CFs differ the most, enhancing interpretability and enabling better decision-making in high-stakes applications. Section 3 starts with the intuition and definition of formula (2) exploring its properties and geometrical meaning. We conclude the section with two important results, Theorems 3.1 and 3.2, showing that our equation defines a measure of dissimilarity and a distance. Section 4 is devoted to the experiments. Here, we compare formula (2) with DiCE local feature importance scores as well as other metrics on various datasets. In general, the contributions of this work can be summarized as follows:

- We propose an innovative local feature importance method that uses positive and negative counterfactuals to capture the relevance of features in generating a specific output. Our formulation takes into account value support and distributional density, thus capturing nuances in the data
- 2. We effectively prove that our proposed measure satisfies the mathematical properties required to qualify as a measure of dissimilarity, and for k=1, it adheres to the axioms of a metric. This formal grounding ensures that the measure is not only conceptually valid but also mathematically rigorous.
- We perform extensive analysis on a variety of datasets for several variables comparing our metric with other standard feature importance scores, underlying the value of our metrics.
- 4. We further highlight the plurality of feature importance scores as well as the diversity of them. Our comparison experiments in Sections 4.2 and 4.4 underline the lack of a ground-truth in explaining a model with its input.

2 RELATED WORK

Local interpretation methods are used to explain individual predictions in machine learning models where the relationship between features and outputs is complex or opaque. These methods attempt to uncover the contributions of individual features to the model's decision process for a particular prediction, in contrast to global interpretation methods that explain overall model behavior. Broadly speaking, most local methods focus on calculating the importance of each feature for the specific instance under analysis ([Molnar, 2022]). Several methods have been proposed in recent years to tackle this challenge. For instance, Individual Conditional Expectation (ICE), first introduced by Goldstein et al. [2015], is a technique that visualizes how the model's prediction changes when a feature varies for a single instance. ICE plots display one line per instance, illustrating how the prediction responds to changes in that instance's features. This method allows for an instance-level understanding of feature impact, but it can be difficult to interpret when dealing with high-dimensional data. LIME (Local Interpretable Model-agnostic Explanations) ([Ribeiro et al., 2016]) provides another approach, where it approximates the complex model locally with an interpretable surrogate model. The idea is to perturb the input data around the instance of interest and fit a simpler model to these perturbed instances, which can then be interpreted in a straightforward manner. While LIME has become quite popular for providing insights into black-box models, its reliance on local linearity makes it sensitive to the choice of proximity measures and perturbation methods ([Thomas Altmann, 2019]).

CFs ([Wachter et al., 2017]) offer an alternative by identifying the minimal change in features required to alter the model's prediction. CFs focus on generating "what if" scenarios, offering users a counterfactual instance that illustrates what would need to happen for a different outcome. The main limitation of counterfactual explanations is the fact that they are instance-specific, i.e. no general information about the model reasoning as a whole is extracted as showed in [Guidotti, 2022] and [Setzu et al., 2021].

Anchors, introduced by [Ribeiro et al., 2018], describe a prediction as being anchored by certain feature values, which lock the model's prediction in place. The idea behind Anchors is to find a local subset of features that sufficiently explain a prediction by ensuring that the prediction would not change when those features remain fixed.

Shapley Values ([Lundberg and Lee, 2017]) come from cooperative game theory and represent the average marginal contribution of a feature across all possible coalitions. Shapley values assign a score to each feature based on how much it contributes to the prediction in comparison to all other possible subsets of features. SHAP has become one of the most widely used methods for feature attribution, as it provides a theoretically grounded explanation that satisfies several desirable properties, such as fairness and consistency, even tough it may suffer from correlation among features ([Marcilio and Eler, 2020].

Although several techniques exist, in the literature LIME, SHAP and CF are among the most commonly referred approaches (Mishra et al. [2021], [Collaris et al., 2022]) and have been widely adopted in various applications.

However, it is important to note that different feature importance methods can yield different rankings of features, even for the same dataset and model. As highlighted by [Rajbahadur et al., 2022], papers in this field rarely provide a clear justification for choosing one particular method over others. This lack of transparency makes it challenging to evaluate or select the best method for any given problem. Additionally, since there is no universally agreed-upon "ground truth" in model interpretability ([Chakraborty et al., 2017], [Haufe et al., 2024], Harel et al. [2023]), the validity of these methods often relies on empirical validation rather than rigorous theoretical proofs.

In this context, our approach, which to the best of our knowledge is the only one with this specific formulation, seeks to provide a mathematically sound alternative by offering a method that is based on distributional discrepancies between local neighborhoods of the data. Our approach directly measures the distance between distributions to assess feature importance, providing an interpretable framework for understanding model behaviour. It is important to emphasise that our method is based on generating the two sets C^+ and C^- with the DiCE library and we use KDE with Gaussian kernel to approximate the distribution of the data.

3 RANKING DIMENSIONS WITH POSITIVE AND NEGATIVE CFS

Let us assume the generation of two sets of CFs: positive CFs C^+ and negative CFs C^- each with cardinality m. Positive CFs C^+ are those that successfully flip the output y, while negative CFs C^- fail to achieve the desired outcome. Our goal is to rank the dimensions in the input space \mathbb{R}^d (or in the activation space of a neural network layer, for neural models) where C^+ and C^- differ the most. This ranking provides insights into which dimensions are most critical in determining the model's output.

A straightforward approach is to compute the variability of each dimension between C^+ and C^- . Specifically, for each dimension i we calculate:

$$Var_i = \frac{1}{m} \sum_{j=1}^{m} (x_{i,j}^+ - x_{i,j}^-)^2.$$

where $x_{i,j}^+$ and $x_{i,j}^-$ represent the *i*-th entry of the *j*-th counterfactual in C^+ and C^- respectively.

While variability provides a simple and effective heuristic, it does not account for the underlying data distribution. To address this, we propose an alternative method based on Kernel Density Estimation (KDE). For continuous or ordinal variables, KDE allows us to estimate the probability distributions of the dimensions in C^+ and C^- . The estimated

distributions are respectively given by

$$P(x)_i = \frac{1}{mh_1} \sum_{j=1}^m K\left(\frac{x - x_{i,j}^+}{h_1}\right),$$

$$Q(x)_{i} = \frac{1}{mh_{2}} \sum_{j=1}^{m} K\left(\frac{x - x_{i,j}^{-}}{h_{2}}\right)$$

where $K(\cdot)$ is the kernel function (e.g., Gaussian or Epanechnikov kernel; see [Weglarczyk, 2018]), h_1 and h_2 are bandwidth parameters for C^+ and C^- . By comparing the above distributions $P(x)_i$, $Q(x)_i$ for each dimension we can capture more nuanced differences in how positive and negative CFs behave. This method is particularly useful in cases where feature variability alone may not fully capture the importance of a dimension.

In order to ground this intuition within a mathematical framework we introduce a notion that captures how the two distribution differs.

3.1 THE NOTION OF OVERLAP OF FUNCTIONS

To formalize the intuition about how two distributions p and q differ, we introduce a measure of overlap o(p,q). Before presenting the formal definition, we outline some desirable properties of such a measure:

- 1. If $p=\mathbf{1}_{[0,1]}$ and $q=\mathbf{1}_{[1,2]}$, the overlap measure should yield a small value, as the support of p and q are disjoint.
- 2. On the other hand, if p = q then d(p, q) should attain its maximum value, indicating high overlap.
- 3. Beyond support overlap, the measure should account for the densities of p and q. For instance, if supp(p) = [0, 1] and supp(q) = [0, 100], but

$$\int_{1}^{100} q(x) \, dx \approx \epsilon$$

where ϵ is very small, the overlap measure o(p,q) should remain high, indicating a limited contribution of q outside supp(p).

Given these considerations, we define the notion of overlap as

Definition 1. Let p, q real positive integrable functions, the notion of overlap is defined as

$$o(p,q) = \frac{\int_{\text{supp}(p)\cap\text{supp}(q)} \min(p(x), q(x)) dx}{\int_{\text{supp}(p)\cup\text{supp}(q)} \max(p(x), q(x)) dx}$$
(1)

and given $k \in \mathbb{R}$ such that $k \geq 1$, we define the measure $d_k(p,q)$ as

$$d_k(p,q) = k - \frac{\int_{\text{supp}(p) \cap \text{supp}(q)} \min(p(x), q(x)) dx}{\int_{\text{supp}(p) \cup \text{supp}(q)} \max(p(x), q(x)) dx}.$$
 (2)

In figure 1 we provide a graphical example of the computation of the above formula for several variables from *Diabetes* dataset. In concrete, to compute integrals we use np.trapz from *Numpy* which applies the composite trapezoidal rule for approximating the value.

3.2 MATHEMATICAL AND GEOMETRICAL PROPERTIES OF $d_1(p,q)$

Remark 3.1. We now explore a set of properties and results that will help clarify the meaning of the overlap measure and provide the foundation for the main results of this paper, Theorems 3.2. and 3.1. The discussion will be done for k=1 as it is the most relevant, all the results can be immediately adapted to $d_k(p,q)$.

First of all we note that since $p(x), q(x) \ge 0$ and

$$supp(p) \cap supp(q) \subset supp(p) \cup supp(q)$$

we can conclude that

$$0 \le o(p,q) \le 1$$

which implies that

$$0 < d_1(p,q) < 1.$$

If p=q then $d_1(p,q)=0$ and if the two density estimations share a 0 measure support, as $\mu(\operatorname{supp}(p)\cap\operatorname{supp}(q))=0$, we obtain that $d_1(p,q)=1$. Furthermore, we have a monotonicity property that takes into account densities: if p(x), g(x), q(x) are three density estimations sharing the same support with the property that, almost everywhere,

$$p(x) \le g(x) \le g(x) \implies d_1(p,q) \le d_1(q,q).$$

It is important to underline since we are working with integrals, we have to consider statements as p < q or $p \neq q$ almost everywhere for a given measure μ . More properly, (2) is defined on

$$L^1(\mathbb{R})\times L^1(\mathbb{R}) = \frac{\mathcal{L}^1(\mathbb{R})\times \mathcal{L}^1(\mathbb{R})}{\sim_{\mu}}$$

which is the quotient space of $\mathcal{L}^1(\mathbb{R}) \times \mathcal{L}^1(\mathbb{R})$ with the equivalence relation induced by μ .

Finally, we are left to show that the notion (2) introduced, satisfies property 3 in Section 3.1.

Proposition 3.1. Let p, q be two real probability distributions, supp(p) = [a, b], supp(q) = [a, c] such that b < c. Moreover, assume that

$$|p(x) - q(x)| < \delta$$
 for $x \in [a, b]$, $\int_b^c q(x) = \epsilon$.

Then o(p,q) attains values close to 1, and so $d_1(p,q)$ to 0.

Proof. See Appendix.

There is an important corollary of previous Proposition. Several density function are defined on \mathbb{R} , even though they are concentrated on a small region as in the case of $N(\sigma,\mu^2)$ and $\Gamma(\alpha,\beta)$ (a list of possible Kernel functions can be found in paper [Weglarczyk, 2018]). Assume now that p(x),q(x) are two density estimation defined on \mathbb{R} , then we can concentrate our overlap measure on a finite proper interval D without any relevant information loss. This is a result of the continuity of $d_1(p,q)$, Proposition 3.1 and the following trivial remark

Remark 3.2. Let $f \in L^1(\mathbb{R})$ then $\exists a \in \mathbb{R}_{\geq 0}$ such that $\forall \epsilon > 0$

$$\int_{a}^{\infty} |f(x)| < \epsilon, \quad \int_{-\infty}^{-a} |f(x)| < \epsilon.$$

Example 3.1. Assume the following density functions

$$p(x) = \begin{cases} \frac{1}{2} & \text{if } x \in [0,2] \\ 0 & \text{otherwise} \end{cases}, \quad q(x) = \begin{cases} \frac{19}{40} & \text{if } x \in [0,2] \\ \frac{1}{60} & \text{if } x \in (2,5] \\ 0 & \text{otherwise}. \end{cases}$$

In this case, the overlap value is

$$d_1(p,q) = 1 - \frac{19/20}{1 + 1/20} = \frac{2}{21} \approx 0.095,$$

indicating that the two densities share highly the same domain.

In the appendix we prove the following important Proposition.

Proposition 3.2. Given p, q real positive integrable functions such that $p \neq q$, then $d_k(p, q) \geq k - 1$.

For $k \ge 2$ in the additional material we show how to use Proposition 3.2 along with some computation to derive the following

Theorem 3.1. The notion of overlap $d_k(p,q)$ is a metric dissimilarity measure on Ω for $k \geq 2$, where

$$\Omega = \{ f \in L^1(\mathbb{R}) \mid supp(f) \neq \emptyset \text{ and } f(x) \geq 0 \}$$

Our focus, as we stated at the beginning of the Section, is k=1, for which we can obtain a stronger result for $d_1(p,q)$. In short, invoking previous results, using the fact that Jaccard distance satisfy triangle inequality and by dominated convergence, we can obtain the following central Theorem

Theorem 3.2. $d_1(p,q)$ is a metric on Ω . Moreover, $d_k(p,q)$ is a dissimilarly measure on Ω for $k \in [1,2)$.

As a consequence, Theorems 3.1 and 3.2 convey that formula defined in (2) actually represents a way of calculating in mathematical terms the dissimilarity between two probability distributions. There is more, Theorem 3.2 establishes that $d_1(p,q)$ satisfies the axioms of a distance: non-negativity, symmetry, the triangle inequality, and that $d_1(p,q)=0$ if and only if p=q. This result is significant because it implies that d_1 induces a metric topology on the space Ω . This topology provides a rigorous framework for comparing probability distributions. From a topological perspective, the metric d_1 ensures that the space Ω is metrizable. For instance, d_1 can be used to define neighborhoods of probability distributions, facilitating the study of their stability, convergence, or variation under perturbations.

4 EXPERIMENTAL RESULTS

To demonstrate the applicability of the proposed framework, we conduct experiments on numerical features on classical datasets: *Diabetes*, *Heart Disease*, *Adult Income* and *Student Depression*. To compute CFs we use the Diverse Counterfactual Explanations (DiCE) for ML library which is based on [Mothilal et al., 2020] that generates CF explanations for any ML model. This library also allow us to generate positive and negative CFs. We take inspiration from Paper [Wiegreffe and Pinter, 2019] and compare the results using $d_k(p,q)$ with local feature importance scores using DiCE. In order to provide a complete picture, for the analyzed variables we also compute local SHAP values ([Lundberg and Lee, 2017]), local LIME values ([Ribeiro et al., 2016]) and global scores as pearson correlation and feature permutation importance ([Altmann et al., 2010]).

4.1 DISSIMILARITY ANALYSIS ON DATASETS

In the datasets we performed iteratively the same analysis. In particular, we extracted the variables for which KDE can be computed, then we generate with DiCE 50 elements for each set C^+ and C^- . It is true that with one-hot encoding could have been applied to categorical variables and then we could have performed KDE on these features, but KDE is highly sensitive on the range of values, so we decided not to perform our analysis on categorical features. Moreover, in order to reduce randomness caused by the algorithm of DiCE, to obtain our dissimilarity measure $d_1(p,q)$ and the DiCE local feature importance we performed the analysis 100 times. As a consequence in the tables below, we report the mean value, while we show the variability in the violin plots in Figure 2. For the model we choose randomly among RandomForestClassifier, KNeighborsClassifier, DecisionTreeClassifier and LogisticRegression. The local scores we report refer to the first entry of the test dataset.

First of all, let us visualize an instance of $d_1(p, q)$. We recall that $d_1(p, q)$ conveys how different are the two distributions:

the more dissimilar are the two distributions, the higher the value $d_1(p,q)$ would be reflecting the fact that different values must be used for that variable to alter the output result.

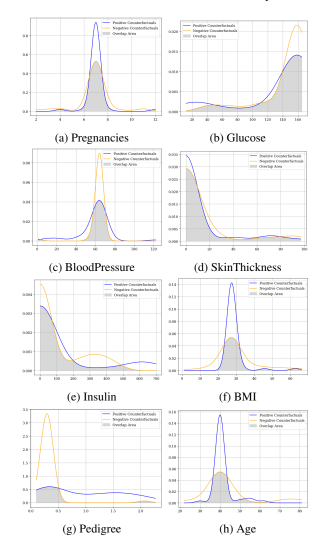
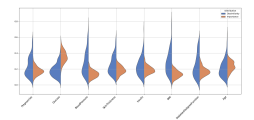


Figure 1: An instance of one iteration of the dissimilarity analysis for the *Diabetes* dataset based on the various entries of C^+ and C^- for the first element of the test dataset. In grey it is highlighted the overlap area.

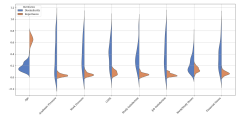
We are now ready to show the comparison between the dissimilarity measure and DiCE local scores. The results are summarized in Figure 2.

4.2 COMPARISON WITH OTHER MEASURES

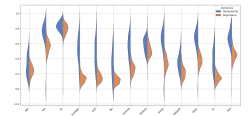
In order to validate our framework and formulation, we computed for all the dataset for the same model and instance SHAP values, LIME values, Pearson correlations and permutation importance scores. The complete results are summarized in the Tables 6, 7, 8 and 9, one for each dataset. For reasons of visibility we call the features v_i and put the names in the caption.



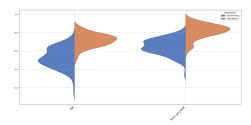
(a) Analysis on the Diabetes dataset.



(b) Analysis on the Student Depression dataset.



(c) Analysis on the Heart Disease dataset.



(d) Analysis on the Adult Income dataset.

Figure 2: The distribution of the feature importance scores for a specific instance over the iterations for $d_1(p,q)$ and DiCE scores.

Even tough, as we already mentioned, there is no ground-truth in feature importance scores and approaches ([Rajba-hadur et al., 2022]), we use *FeatureAgreement* introduced in [Krishna et al., 2024] with k=4, to evaluate the agreement/disagreement on the feature importance scores across the various metrics we used. In short, given two explanations E_a and E_b , feature agreement is formulated as

$$\frac{|TF(E_a,k) \cap TF(E_b,k)|}{k} \tag{3}$$

where TF(E,k) returns the set of top-k features of explanation E based on the magnitude of the feature importance values. Formula (3) it computes the fraction of common

features between the sets of top-k features of two explanations. Clearly for dataset *Adult Income* we used k=2 as we only analyzed 2 features. In Table 3 d_1 and Dice show

	d_1	DiCE	SHAP	LIME
d_1	1.00	0.50	0.50	0.25
DiCE	0.50	1.00	0.50	0.25
SHAP	0.50	0.50	1.00	0.50
LIME	0.25	0.25	0.50	1.00

Table 1: Feature Agreement Matrix for the Diabetes dataset.

	d_1	DiCE	SHAP	LIME
d_1	1.00	0.50	0.25	0.50
DiCE	0.50	1.00	0.50	0.25
SHAP	0.25	0.50	1.00	0.75
LIME	0.50	0.25	0.75	1.00

Table 2: Feature Agreement Matrix for the *Student Depression* dataset.

	d_1	DiCE	SHAP	LIME
d_1	1.00	0.75	0.00	0.50
DiCE	0.75	1.00	0.25	0.50
SHAP	0.00	0.25	1.00	0.00
LIME	0.50	0.50	0.00	1.00

Table 3: Feature Agreement Matrix for the *Heart Disease* dataset.

	d_1	DiCE	SHAP	LIME
d_1	1.00	1.00	0.00	1.00
DiCE	1.00	1.00	0.00	1.00
SHAP	0.00	0.00	1.00	0.00
LIME	1.00	1.00	0.00	1.00

Table 4: Feature Agreement Matrix for the *Adult Income* dataset.

high agreement (0.75), while d_1 and SHAP no agreement. Conversely, in Table 2, d_1 and DiCE show low agreement (0.25). Here, SHAP and LIME achieve high agreement (0.75), contrary to Table 3 where there is no agreement. Table 1 displays more moderate and consistent agreement levels between most metric pairs. These findings align with prior literature emphasizing the lack of a universal ground truth in feature importance scores. They also reinforce the necessity of analyzing multiple metrics to obtain a more comprehensive understanding of feature importance.

4.3 ANALYSIS OF RESULTS

The variability observed Figure 2 reflects both the use of a random counterfactual generator and the inherent nature of

feature importance, which can be volatile due to the complexity of the relationships between inputs and outputs. To address this, we mitigated randomness by running multiple experiments and reporting the average values.

Interestingly, the variability appears to be influenced by the dataset's complexity. For instance, in the simpler Adult Income dataset in Figure 2d (with only two features analyzed), the variability is relatively low. However, for the Heart Disease dataset in Figure 2c, which involves 13 features, the variability increases. This suggests that the complexity of the dataset—and the underlying relationships we aim to infer—plays a crucial role in explainability.

The metric also leverages Kernel Density Estimation (KDE) to identify the distribution of the values analyzed, providing an alternative insight into the interplay between features and their impact. This aspect becomes particularly important in more complex datasets, where a single value might not fully capture the nuances of feature importance. These findings demonstrate that the metric is sensitive to the data's complexity. When compared to existing methods, our metric provides a distinct perspective. For example, in Table 7, summarizing the results for the *Student Depression* dataset, our metric produces more uniform scores across the variables, in contrast to DiCE. However, it is notable that even when considering SHAP importance and LIME importance, there is no consistent agreement in ranking the most significant features. Conversely, for the Diabetes dataset, as shown in Table 6, our metric aligns closely with DiCE, SHAP, and LIME importance. These insights, as we discussed in Section 4.2 conform to the literature on feature importance ([Harel et al., 2023], [Rajbahadur et al., 2022], [Jacovi and Goldberg, 2020]).

This diversity in outcomes highlights the complementary nature of our approach. Just as in other domains where different metrics offer unique insights ([Mishra and Sadia, 2023], [Rajbahadur et al., 2022]), our metric enriches the analysis by capturing a novel aspect of explainability. Specifically, while LIME relies on a local linear approximation, DiCE uses perturbations and gradients, and SHAP employs a cooperative game-theoretic approach to attribute importance based on Shapley values, our metric focuses on analyzing the distributional discrepancies between two sets of points near the instance in question. This effectively quantifies the "distance" between them, offering a new perspective to understand the importance of features. Integrating our metric with existing ones offers a more comprehensive way to evaluate feature importance, especially when understanding distributions is crucial.

4.4 EVALUATION OF FAITHFULNESS METRICS

The results obtained using the Feature Agreement metric (3) are not conclusive, highlighting the inherent plurality of approaches to estimating feature importance. To complement the qualitative analysis, we additionally estimate

the *correctness* of an explanation. In particular, we follow the frameworks proposed in [Zhou and Shah, 2023] and [Chan et al., 2022], which assess the faithfulness of feature attribution evaluations. The key intuition behind these metrics is that altering an important feature should significantly impact the model's prediction and the magnitude of this impact reflects the quality of the explanation.

Among the various methods to estimate faithfulness, we adopt **comprehensiveness** introduced in [DeYoung et al., 2020]. Given an input $x=(x_1,\ldots,x_L)$ and an explanation $e=(e_1,\ldots,e_L)$, if we denote $\tilde{x}_e^{(l)}$ the input with l most important features removed according to e, comprehensiveness is defined as

$$k(x,e) = \frac{1}{L+1} \sum_{l=0}^{L} f(x) - f(\tilde{x}_e^{(l)})$$
 (4)

where f is the function we want to explain. In simple terms, comprehensiveness measures how much the model prediction deviates from its original value when important features are removed sequentially (larger values indicate better explanations). In our experiments, we compute (4) using the predict-proba function as f for the numerical features, setting k=4. The explanations are generated using d_1 , DiCE, LIME, and SHAP. Since we are working with local feature importance estimations, in order to obtain $\tilde{x}_e^{(l)}$ we mask the top relevant features, according to the explanation, using the mean ([Covert et al., 2021]). In addition, we plot the trend of predict-proba when gradually removing relevant features.

	d_1	DiCE	SHAP	LIME
Diabetes	-0.1750	0.6985	-0.0044	0.5848
Student	0.0490	0.1328	0.0600	-0.0093
Heart	0.3950	0.4900	0.2300	0.3500
Adult	0.00	0.00	0.00	0.00

Table 5: The comprehensiveness measure across the various datasets.

The remaining plots can be found in the supplementary materials. As shown in Table 5, the DiCE library achieves the best results in terms of comprehensiveness and, consequently, in visualizing the predict-probability values. The values obtained with d_1 are generally close to those of SHAP and LIME. It is worth noting that the k(x,e) values obtained for the Adult dataset are null, as we only modified the two numerical features, which are not the most influential in determining the final prediction. Additionally, the test instance already had a predict-probability of 1 for the income = 0 class, making it more challenging to observe variations. Nevertheless, with the exception of the Diabetes dataset, the method we introduced aligns with the results of other metrics.

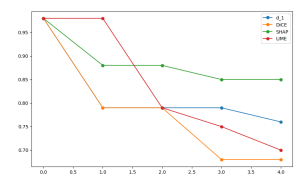


Figure 3: The trend of k(x,e) when gradually removing the relevant features according the different explanations in *Heart Disease* dataset.

5 DISCUSSION

The formula proposed in Definition 2 provides a method to quantify the dissimilarity between two data distributions, offering a novel perspective on how features contribute to a given output. Importantly, this approach can naturally extend beyond binary classifiers. For instance, in the case of a continuous output space \mathbb{R} , it is sufficient to define a subset $A \in \mathbb{R}$ where C^+ represents the data points for which $f(x) \in A$, and C^- corresponds to those with $f(x) \in A^c$. In our analysis of feature importance using this framework, we treated feature importance independently, a common simplification in mathematical modeling (see [Mothilal et al., 2020], [Molnar, 2022], [Schölkopf et al., 2021]). Nevertheless, following the results of Section 4, the formula (2) offers an alternative perspective to understand how the various inputs of a specific data point participate in producing a certain output for model M. As a consequence, it may be included to complement existing methods for feature importance analysis.

Furthermore, the results from both the numerical and distributional analysis and those based on feature agreement and faithfulness reveal important insights into the variability and sensitivity of interpretability methods. Despite employing three well-established and widely used metrics (alongside the novel approach introduced in this work), substantial differences are observed in the results. These discrepancies can often be traced back to the specific characteristics of individual datasets or even to variations within the same dataset. This sensitivity underscores a broader challenge in interpretability: the lack of a universally "correct" method or metric. Traditional measures of interpretability in data science, while widely adopted, are not necessarily definitive or universally applicable.

This variability calls for caution when interpreting results and highlights the need for a more nuanced understanding of the strengths and limitations of different interpretability methods. It also suggests that employing multiple complementary metrics can provide a more robust view of feature importance.

This work represents an initial exploration of this dissimilarity framework, with promising results across diverse datasets. However, several open questions remain. For instance, future studies could investigate the impact of dependencies among features; the possibility of extending this framework to multivariate setting using multivariate kernel density estimation or dealing with the limitation of continuous or ordinal variables.

We believe this approach has potential applications in improving model interpretability, especially in scenarios where traditional feature importance methods might struggle to account for distributional differences. While the current implementation demonstrates the formula's theoretical foundation and practical applicability, further testing could expand its utility in real-world scenarios.

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Measuring Feature Importance Through Counterfactual Distributions (Supplementary Material)

A APPENDIX

Proof of Proposition 3.1

Proof. To show this, suppose the following scenario: supp(p) = [a, b], supp(q) = [a, c] such that b < c. Moreover, assume that

$$|p(x) - q(x)| < \delta$$
 for $x \in [a, b]$, $\int_b^c q(x) = \epsilon$.

Under this assumptions,

$$\begin{split} d_k(p,q) &= k - \frac{\int_a^b \min(p(x),q(x))dx}{\int_a^c \max(p(x),q(x))dx} = k - \frac{\int_a^b \min(p(x),q(x))dx}{\int_a^b \max(p(x),q(x))dx + \int_b^c q(x)dx} \\ &= k - \frac{\int_a^b \min(p(x),q(x))dx}{\int_a^b \max(p(x),q(x))dx + \epsilon} \\ &= k - \frac{\int_a^b \min(p(x),q(x))dx}{\int_a^b \max(p(x),q(x)) + \frac{\epsilon}{b-a}dx} \\ &= k - \frac{\int_a^b \min(p(x),q(x))dx}{\int_a^b \max(p(x),q(x))dx} \\ &= k - \frac{\int_a^b \min(p(x),q(x))dx}{\int_a^b \max(p(x),q(x))dx} \end{split}$$

Now, from the last equation, we can obtain

$$k - \frac{1 - \epsilon}{1 - \delta(b - a)} \le d(p, q) \le k - \frac{1 - \epsilon - \delta(b - a)}{1 + \delta(b - a)}$$

since

$$\int_{a}^{b} q(x) = 1 - \epsilon.$$

The inequality can be obtained by observing that $|p(x) - q(x)| < \delta$ and so $q(x) - \delta < p(x) < q(x) + \delta$ which implies

$$\min(p(x), q(x)) \le q(x), \quad \max(p(x), q(x)) \ge q(x) - \delta$$

and

$$\min(p(x), q(x)) \ge q(x) - \delta, \quad \max(p(x), q(x)) \le q(x) + \delta.$$

Now, as desired, if ϵ and δ are relatively small, d(p,q) takes on a value close to k-1, i.e., indicating an high overlap. \Box

Derivation of Theorem 3.1

Proposition 5.1. Given p, q real positive integrable functions such that $p \neq q$, then $d_k(p,q) > k-1$.

Proof. Consider two functions p, q such that $p \neq q$. It is straightforward that if $supp(p) \neq supp(q)$ then d(p,q) > k-1 by how it is defined the measure of overlap. Indeed, in the worst case scenario p = q in $supp(p) \cap supp(q)$, but as $supp(p) \neq supp(q)$, then

$$\operatorname{supp}(p) \cap \operatorname{supp}(q) \subsetneq \operatorname{supp}(p) \cup \operatorname{supp}(q)$$

which implies

$$\frac{\int_{\operatorname{supp}(p)\cap\operatorname{supp}(q)}\min(p(x),q(x))dx}{\int_{\operatorname{supp}(p)\cup\operatorname{supp}(q)}\max(p(x),q(x))dx}<1.$$

As a consequence, let us assume supp(p) = supp(q) = D. Since $p \neq q$, at least one among

$$A = \{x \in D \mid p(x) < q(x)\}, \quad B = \{x \in D \mid q(x) < p(x)\}\$$

has a positive measure. Assume $\mu(A) > 0$, therefore

$$\frac{\int_{D} \min(p(x), q(x)) dx}{\int_{D} \max(p(x), q(x)) dx} = \frac{\int_{A} p(x) + \int_{D \setminus A} q(x)}{\int_{A} q(x) + \int_{D \setminus A} \max(p(x), q(x))} < 1$$

because

$$\int_{D \setminus A} q(x) \le \int_{D \setminus A} \max(p(x), q(x)),$$
$$\int_{A} p(x) < \int_{A} q(x).$$

The case $\mu(B) > 0$ is analogous.

Now let us recall the following definition.

Definition 2. A dissimilarity measure (DM) d on X is a function $d: X \times X \to \mathbb{R}$ such that

$$\exists d_0 \text{ s.t. } -\infty < d_0 \leq d(x,y) < \infty \quad \forall x,y \in X,$$

$$d(x,x) = d_0 \quad \forall x \in X,$$

$$d(x,y) = d(y,x) \quad \forall x,y \in X.$$

If in addition

$$d(x,y) = d_0 \iff x = y,$$

$$d(x,z) \le d(x,y) + d(y,z) \quad \forall x, y, z \in X,$$

d is called a metric DM on X.

Let us investigate the triangular inequality

$$d_k(p,q) \le d_k(p,q) + d_k(q,q)$$

which in our case becomes

$$k - \frac{\int_{\operatorname{supp}(p) \cap \operatorname{supp}(r)} \min(p,r)}{\int_{\operatorname{supp}(p) \cup \operatorname{supp}(r)} \max(p,r)} \leq k - \frac{\int_{\operatorname{supp}(p) \cap \operatorname{supp}(q)} \min(p,q)}{\int_{\operatorname{supp}(p) \cup \operatorname{supp}(q)} \max(p,q)} + k - \frac{\int_{\operatorname{supp}(q) \cap \operatorname{supp}(r)} \min(q,r)}{\int_{\operatorname{supp}(q) \cup \operatorname{supp}(r)} \max(q,r)}.$$

and simplified it becomes

$$\frac{\int_{\operatorname{supp}(p) \cap \operatorname{supp}(q)} \min(p,q)}{\int_{\operatorname{supp}(p) \cup \operatorname{supp}(q)} \max(p,q)} + \frac{\int_{\operatorname{supp}(q) \cap \operatorname{supp}(r)} \min(q,r)}{\int_{\operatorname{supp}(q) \cup \operatorname{supp}(r)} \max(q,r)} - \frac{\int_{\operatorname{supp}(p) \cap \operatorname{supp}(r)} \min(p,r)}{\int_{\operatorname{supp}(p) \cup \operatorname{supp}(r)} \max(p,r)} \leq k$$

which is always satisfied if $k \ge 2$ because every term on the left is positive and bounded by 1. As a consequence of this result combined with Proposition 3.2, noticing that the symmetry is guaranteed and that d_0 in Definition 2 in our case is k-1, we have the following

Theorem 5.1. The notion of overlap $d_k(p,q)$ is a metric dissimilarity measure on $\Omega \subset L^1(\mathbb{R})$ s.t. $supp(f) \neq \emptyset$ and $f(x) \geq 0$ for all $f \in \Omega$, for $k \geq 2$.

Derivation of Theorem 3.2 According to our definition, we are left to analyze the case for $k \in [1, 2)$ in (2). First of all we note that the Jaccard distance defined fro two sets A, B

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

satisfies the triangle inequality (cfr. Paper Kosub [2019]). As a consequence, if we consider the functions $p(x) = \mathbf{1}_A$, $q(x) = \mathbf{1}_B$, $r(x) = \mathbf{1}_C$

$$d_1(p,q) \le d_1(p,q) + d_1(q,q)$$

and of course if we add on the lhs k-1 and on the rhs 2(k-1) the inequality still holds, hence

$$d_k(p,q) \le d_k(p,g) + d_k(g,q).$$

Now, the idea is to prove if the inequality is valid for general characteristic functions, therefore if

$$d_k(\alpha \mathbf{1}_A, \gamma \mathbf{1}_C) \le d_k(\alpha \mathbf{1}_A, \beta \mathbf{1}_B) + d_k(\beta \mathbf{1}_B, \gamma \mathbf{1}_C) \quad \alpha, \beta, \gamma \in \mathbb{R}_{\ge 0}.$$

Without loss of generality we can assume that $\alpha \leq \beta \leq \gamma$, so we have to prove that

$$1 - \frac{\alpha |A \cap C|}{\gamma |A \cup C|} \le 1 - \frac{\alpha |A \cap B|}{\beta |A \cup B|} + 1 - \frac{\beta |B \cap C|}{\gamma |B \cup C|}$$

which can be rewritten as

$$\frac{\beta|B\cap C|}{\gamma|B\cup C|} + \frac{\alpha|A\cap B|}{\beta|A\cup B|} - \frac{\alpha|A\cap C|}{\gamma|A\cup C|} \leq 1$$

but this is true since we know that

$$\begin{split} \frac{|B\cap C|}{|B\cup C|} + \frac{|A\cap B|}{|A\cup B|} - \frac{|A\cap C|}{|A\cup C|} \leq 1, \\ 0 \leq \frac{\alpha}{\beta}, \frac{\alpha}{\gamma}, \frac{\beta}{\gamma} \leq 1. \end{split}$$

We observe that it does not affect the order of α , β , γ since we have a minimum over a maximum and so the ratio is always less or equal than 1. Now, let us assume that p(x), q(x), r(x) are simple functions, i.e.

$$p(x) = \sum_{i=1}^{k} a_i \mathbf{1}_{A_i}, \quad q(x) = \sum_{i=1}^{k} b_i \mathbf{1}_{B_i}, \quad r(x) = \sum_{i=1}^{k} c_i \mathbf{1}_{C_i}.$$

where $k \in \mathbb{N}$, $a_i, b_i, c_i \ge 0$ and A_i, B_i, C_i disjoint. It is trivial now, since the inequality holds for general characteristic functions, that

$$d_k(p,q) \le d_k(p,q) + d_k(q,q).$$

Any function in $L^1(\mathbb{R})$ can be approximated with a sequence of simple functions (cfr. L. and M. [2017]). By dominated convergence theorem we can exchange the limit with the integral and so, combining it with the results for simple functions, we conclude that

$$d_k(p,q) \le d_k(p,g) + d_k(g,q) \quad p,q,r \in L^1(\mathbb{R}).$$

As a conclusion, we can extend Theorem 3.1 to the case $k \ge 1$, and so we obtain Theorem 3.2.

ADDITIONAL TABLES AND FIGURES

Here we show the tables for the various experiments for *Diabates*, *Student Depression*, *Heart Disease* and *Adult Income* datasets. There are also barplots for easy visualisation.

Feature	Dissimilarity	DiCE Importance	SHAP Importance	Lime Importance	PCC	PIMP
$\overline{v_1}$	0.229566	0.174800	-0.006384	0.133056	0.221898	-0.025974
v_2	0.185589	0.357800	0.002810	0.318405	0.466581	0.039610
v_3	0.269329	0.135800	-0.004329	0.014217	0.065068	0.007792
v_4	0.247616	0.183600	-0.004720	0.016571	0.074752	-0.033766
v_5	0.236607	0.190600	-0.035018	-0.017215	0.130548	-0.003896
v_6	0.283659	0.134200	0.014626	-0.140965	0.292695	0.018831
v_7	0.236086	0.174000	-0.000889	-0.052864	0.173844	-0.010390
v_8	0.212042	0.234800	-0.005184	0.011030	0.238356	-0.019481

Table 6: Results for *Diabetes*. The variables involved are: Pregnancies, Glucose, BloodPressure, SkinThickness, Insulin, BMI, DiabetesPedigreeFunction and Age.

Feature	Dissimilarity	DiCE Importance	SHAP Importance	Lime Importance	PCC	PIMP
$\overline{v_1}$	0.179028	0.642600	0.342085	-0.005727	-0.226511	0.023423
v_2	0.346901	0.035000	-0.023132	-0.267086	0.474793	0.067545
v_3	0.378021	0.049600	0.000000	0.000000	-0.003351	0.000000
v_4	0.386789	0.078000	0.013947	0.012580	0.022268	-0.002706
v_5	0.332646	0.048000	0.059307	-0.054926	-0.168010	0.002706
v_6	0.389102	0.044200	0.000000	0.048434	-0.003483	0.000000
v_7	0.222857	0.135000	0.000006	0.003201	0.208627	0.008065
v_8	0.358206	0.069800	0.047165	0.136878	0.363591	0.033495

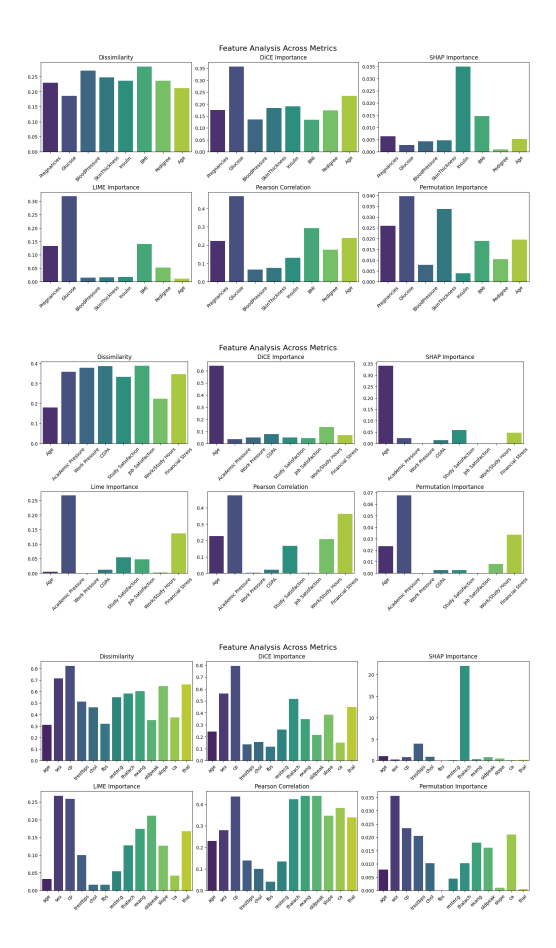
Table 7: Results for *Student Depression*. The variables involved are: Age, Academic Pressure, Work Pressure, CGPA, Study Satisfaction, Job Satisfaction, Work/Study Hours and Financial Stress.

Feature	Dissimilarity	DiCE Importance	SHAP Importance	Lime Importance	PCC	PIMP
$\overline{v_1}$	0.310265	0.242600	-1.074923	-0.032082	-0.229324	-0.007805
v_2	0.714655	0.561200	-0.242341	-0.267094	-0.279501	0.035610
v_3	0.822264	0.793400	-0.772315	-0.258588	0.434854	0.023415
v_4	0.512483	0.135800	3.917459	0.099770	-0.138772	-0.020488
v_5	0.462853	0.156600	-0.907916	-0.016388	-0.099966	-0.010244
v_6	0.318435	0.117000	0.001464	0.016551	-0.041164	0.000000
v_7	0.551675	0.261200	-0.089746	-0.054682	0.134468	-0.004390
v_8	0.582921	0.516600	-22.026003	-0.126994	0.422895	-0.010244
v_9	0.604392	0.346400	-0.336086	-0.173920	-0.438029	-0.018049
v_{10}	0.350887	0.215600	-0.794898	-0.210574	-0.438441	0.016098
v_{11}	0.645632	0.384600	-0.508091	-0.125980	0.345512	0.000976
v_{12}	0.376092	0.149200	-0.089461	-0.041630	-0.382085	0.020976
v_{13}	0.661424	0.449800	0.194678	0.167151	-0.337838	0.000488

Table 8: Results for *Heart Disease*. The variables involved are: age, sex, cp, trestbps, chol, fbs, restecg, thalach, exang, oldpeak, slope, ca and thal.

Feature	Dissimilarity	DiCE Importance	SHAP Importance	Lime Importance	PCC	PIMP
v_1 v_2	0.525798	0.718000	-0.545029	-0.006652	0.233879	0.000537
	0.647884	0.815400	0.020406	-0.110052	0.225602	0.000403

Table 9: Results for Adult Income. The variables involved are: Age and hours-per-week.



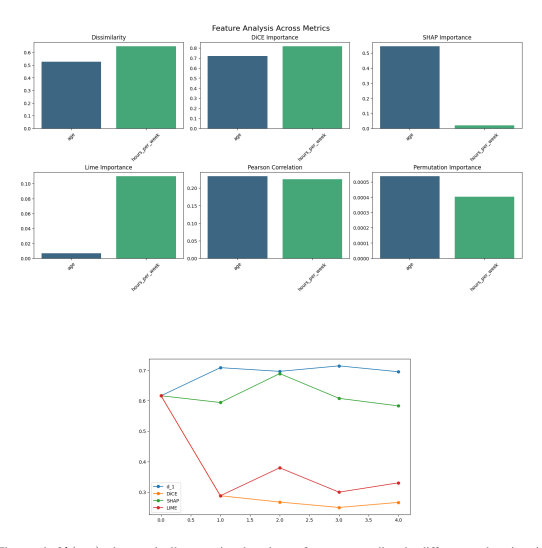


Figure 4: The trend of k(x,e) when gradually removing the relevant features according the different explanations in Diabetes dataset.

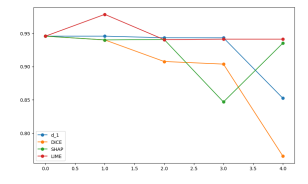


Figure 5: The trend of k(x, e) when gradually removing the relevant features according the different explanations in *Student Depression* dataset.

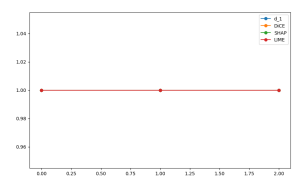


Figure 6: The trend of k(x,e) when gradually removing the relevant features according the different explanations in Adult Income dataset.