ROBUST MODELS ARE MORE INTERPRETABLE BECAUSE ATTRIBUTIONS LOOK NORMAL

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ABSTRACT

Recent work has found that adversarially-robust deep networks used for image classification are more interpretable: their feature attributions tend to be sharper, and are more concentrated on the objects associated with the image’s ground-truth class. We show that smooth decision boundaries play an important role in this enhanced interpretability, as the model’s input gradients around data points will more closely align with boundaries’ normal vectors when they are smooth. Thus, because robust models have smoother boundaries, the results of gradient-based attribution methods, like Integrated Gradients and DeepLift, will capture more accurate information about nearby decision boundaries. This understanding of robust interpretability leads to our second contribution: boundary attributions, which aggregate information about the normal vectors of local decision boundaries to explain a classification outcome. We show that by leveraging the key factors underpinning robust interpretability, boundary attributions produce sharper, more concentrated visual explanations—even on non-robust models. Any example implementation can be found at https://github.com/zifanw/boundary.

1 INTRODUCTION

Feature attribution methods are widely used to explain the predictions of neural networks (Binder et al., 2016; Dhamdhere et al., 2019; Fong & Vedaldi, 2017; Leino et al., 2018; Montavon et al., 2015; Selvaraju et al., 2017; Shrikumar et al., 2017; Simonyan et al., 2013; Smilkov et al., 2017; Springenberg et al., 2014; Sundararajan et al., 2017). By assigning an importance score to each input feature of the model, these techniques help to focus attention on parts of the data most responsible for the model’s observed behavior. Recent work (Croce et al., 2019; Etmann et al., 2019) has observed that feature attributions in adversarially-robust image models, when visualized, tend to be more interpretable—the attributions correspond more clearly to the discriminative portions of the input.

One way to explain the observation relies on the fact that robust models do not make use of non-robust features (Ilyas et al., 2019) whose statistical meaning can change with small, imperceptible changes in the source data. Thus, by using only robust features to predict, these models naturally tend to line up with visibly-relevant portions of the image. Etmann et al. take a different approach, showing that the gradients of robust models’ outputs more closely align with their inputs, which explains why attributions on image models are more visually interpretable.

In this paper, we build on this geometric understanding of robust interpretability. With both analytical (Sec. 3) and empirical (Sec. 5) results, we show that the gradient of the model with respect to its input, which is the basic building block of all gradient-based attribution methods, tends to be more closely aligned with the normal vector of a nearby decision boundary in robust models than in “normal” models. Leveraging this understanding, we propose Boundary-based Saliency Map (BSM) and Boundary-based Integrated Gradient (BIG), two variants of boundary attributions (Sec. 4), which base attributions on information about nearby decision boundaries (see an illustration in Fig. 1a). While BSM provides theoretical guarantees in the closed-form, BIG generates both quantitatively and qualitatively better explanations. We show that these methods satisfy several desirable formal properties, and that even on non-robust models, the resulting attributions are more focused (Fig. 1b) and less sensitive to the “baseline” parameters required by some attribution methods.
To summarize, our main contributions are as follows. (1) We present an analysis that sheds light on the previously-observed phenomenon of robust interpretability, showing that alignment between the normal vectors of decision boundaries and models’ gradients is a key ingredient (Proposition 1, Theorem 1). (2) Motivated by our analysis, we introduce boundary attributions, which leverage the connection between boundary normal vectors and gradients to yield explanations for non-robust models that carry over many of the favorable properties that have been observed of explanations on robust models. (3) We empirically demonstrate that one such type of boundary attribution, called Boundary-based Integrated Gradients (BIG), produces explanations that are more accurate than prior attribution methods (relative to ground-truth bounding box information), while mitigating the problem of baseline sensitivity that is known to impact applications of Integrated Gradients Sundararajan et al. (2017) (Section 6).

2 BACKGROUND

We begin by introducing our notations. Throughout the paper we use italicized symbols $x$ to denote scalar quantities and bold-face $\mathbf{x}$ to denote vectors. We consider neural networks with ReLU as activations prior to the top layer, and a softmax activation at the top. The predicted label for a given input $\mathbf{x}$ is given by $F(\mathbf{x}) = \arg \max_c f_c(\mathbf{x}), \mathbf{x} \in \mathbb{R}^d$, where $F(\mathbf{x})$ is the predicted label and $f_i(\mathbf{x})$ is the output on the class $i$. As the softmax layer does not change the ranking of neurons in the top layer, we will assume that $f_i(\mathbf{x})$ denotes the pre-softmax score. Unless otherwise noted, we use $||\mathbf{x}||$ to denote the $\ell_2$ norm of $\mathbf{x}$, and the $\ell_2$ neighborhood centered at $\mathbf{x}$ with radius $\epsilon$ as $B(\mathbf{x}, \epsilon)$.

**Explainability.** Feature attribution methods are widely-used to explain the predictions made by DNNs, by assigning importance scores for the network’s output to each input feature. Conventionally, scores with greater magnitude indicate that the corresponding feature was more relevant to the predicted outcome. We denote feature attributions by $\mathbf{z} = g(\mathbf{x}, f)$, $\mathbf{z}, \mathbf{x} \in \mathbb{R}^d$. When $f$ is clear from the context, we simply write $g(\mathbf{x})$. While there is an extensive and growing literature on attribution methods, our analysis will focus closely on the popular gradient-based methods, Saliency Map (Simonyan et al., 2013), Integrated Gradient (Sundararajan et al., 2017) and Smooth Gradient (Smilkov et al., 2017), shown in Defs 1-3.

**Definition 1 (Saliency Map (SM))** The Saliency Map $g_S(\mathbf{x})$ is given by $g_S(\mathbf{x}) := \frac{\partial f(\mathbf{x})}{\partial \mathbf{x}}$. 

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Figure 1: (a) Visualizations of geometrical interpretations of Saliency Map (SM), Boundary-based Saliency Map (BSM), Integrated Gradient (IG) and Boundary-based Integrated Gradient (BIG). Gradient computations can be viewed as projecting the input onto a particular decision boundary. While SM projects to a nearby boundary ($H_1$), BSM projects to the nearest one ($H_2$). IG (the red dashed path) from a global baseline $\mathbf{x}_g$, i.e. zeros, aggregates boundaries in colorful shaded areas; BIG (the green dashed path) integrates from the point $\mathbf{x}_{bd}$ on the nearest boundary $H_2$ to $\mathbf{x}$ and therefore aggregates nearby boundaries, $H_1$ and $H_2$ in gray shaded areas. (b) Visualizations of Integrated Gradient and the proposed improvement of it, Boundary-based Integrated Gradient, which is sharper, more concentrated and less noisy.
**Definition 2 (Integrated Gradient (IG))** Given a baseline input \( x_b \), the Integrated Gradient \( g_{IG}(x; x_b) \) is given by \( g_{IG}(x; x_b) := (x - x_b) \int_0^1 \frac{\partial f((x-x_b) + t(x-x_b))}{\partial x} \, dt \).

**Definition 3 (Smooth Gradient (SG))** Given a zero-centered Gaussian distribution \( N \) with a standard deviation \( \sigma \), the Smooth Gradient \( g_{SG}(x; \sigma) \) is given by \( g_{SG}(x; \sigma) := \mathbb{E}_{\varepsilon \sim N(0, \sigma^2)} \frac{\partial f(x + \varepsilon)}{\partial x} \).

Besides, we will also include results from DeepLIFT (Shrikumar et al., 2017) and \( \text{grad} \times \text{input} \) (element-wise multiplication between Saliency Map and the input) (Simonyan et al., 2013) in our empirical evaluation. As we show in Section 3.2, Dets [13] satisfy axioms that relate to the local linearity of ReLU networks, and in the case of randomized smoothing (Cohen et al., 2019), their robustness to input perturbations. We further discuss these methods relative to others in Sec. 7.

**Robustness.** Two relevant concepts about adversarial robustness will be used in this paper: prediction robustness that the model’s output label remains unchanged within a particular \( \ell_p \) norm ball and attribution robustness that the feature attributions are similar within the same ball. Recent work has identified the model’s Lipschitz continuity as a bridge between these two concepts (Wang et al., 2020c) and some loss functions in achieving prediction robustness also bring attribution robustness (Chalasani et al., 2020). We refer to robustness as prediction robustness if not otherwise noted.

### 3 EXPLAINABILITY, DECISION BOUNDARIES, AND ROBUSTNESS

In this section, we begin by discussing the role of decision boundaries in constructing explanations of model behavior via feature attributions. We first illustrate the key relationships in the simpler case of linear models, which contain exactly one boundary, and then generalize to piecewise-linear classifiers as they are embodied by deep ReLU networks. We then show how local robustness causes attribution methods to align more closely with nearby decision boundaries, leading to explanations that better reflect these relationships.

#### 3.1 ATTRIBUTIONS FOR LINEAR MODELS

Consider a binary classifier \( C(x) = \text{sign}(w^T x + b) \) that predicts a label \( \{-1, 1\} \) (ignoring “tie” cases where \( C(x) = 0 \), which can be broken arbitrarily). In its feature space, \( C(x) \) is a hyperplane \( H \) that separates the input space into two open half-spaces \( S_1 \) and \( S_2 \) (see Fig. 2a). Accordingly, the normal vector \( \hat{n} \) of the decision boundary is the only vector that faithfully explains the model’s classification while other vectors, while they may describe directions that lead to positive changes in the model’s output score, are not faithful in this sense (see \( v \) in Fig. 2a for an example). In practice, to assign attributions for predictions made by \( C \), SM, SG, and the integral part of IG (see Sec. 2) return a vector characterized by \( x = k_1 \hat{n} + k_2 \) (Ancona et al., 2018), where \( k_1 \neq 0 \) and \( k_2 \in \mathbb{R} \), regardless of the input \( x \) that is being explained. In other words, these methods all measure the importance of features by characterizing the model’s decision boundary, and are equivalent up to the scale and position of \( \hat{n} \).

#### 3.2 GENERALIZING TO PIECEWISE-LINEAR BOUNDARIES

In the case of a piecewise-linear model, such as a ReLU network, the decision boundaries comprise a collection of hyperplane segments that partition the feature space, as in \( H_1, H_2 \) and \( H_3 \) in the example shown in Figure 2b. Because the boundary no longer has a single well-defined normal, one intuitive way to extend the relationship between boundaries and attributions developed in the previous section is to capture the normal vector of the closest decision boundary to the input being explained. However, as we show in this section, the methods that succeeded in the case of linear models (SM, SG, and the integral part of IG) may in fact fail to return such attributions in the more general case of piecewise-linear models, but local robustness often remedies this problem. We begin by reviewing key elements of the geometry of ReLU networks (Jordan et al., 2019).

**ReLU activation polytopes.** For a neuron \( u \) in a ReLU network \( f(x) \), we say that its status is ON if its pre-activation \( u(x) \geq 0 \), otherwise it is OFF. We can associate an activation pattern denoting the status of each neuron for any point \( x \) in the feature space, and a half-space \( A_u \) to the activation constraint \( u(x) \geq 0 \). Thus, for any point \( x \) the intersection of the half-spaces corresponding to its activation pattern defines a polytope \( P \) (see Fig. 2b), and within \( P \) the network is a linear function.
such that \( \forall \mathbf{x} \in P, f(\mathbf{x}) = \mathbf{w}_p^\top \mathbf{x} + b_p \), where the parameters \(\mathbf{w}_p\) and \(b_p\) can be computed by differentiation (Fromherz et al. 2021). Each facet of \(P\) (dashed lines in Fig. 2b) corresponds to a boundary that “flips” the status of its corresponding neuron. Similar to activation constraints, decision boundaries are piecewise-linear because each decision boundary corresponds to a constraint \(f_i(\mathbf{x}) \geq f_j(\mathbf{x})\) for two classes \(i, j\) (Fromherz et al. 2021; Jordan et al. 2019).

**Gradients might fail.** Saliency maps, which we take to be simply the gradient of the model with respect to its input, can thus be seen as a way to project an input onto a decision boundary. That is, a saliency map is a vector that is normal to a nearby decision boundary segment. However, as others have noted, a saliency map is not always normal to any real boundary segment in the model’s geometry (see the left plot of Fig. 2c), because when the closest boundary segment is not within the activation polytope containing \(\mathbf{x}\), the saliency map will instead be normal to the linear extension of some other hyperplane segment (Fromherz et al. 2021). In fact, the fact that iterative gradient descent typically outperforms the Fast Gradient Sign Method (Goodfellow et al. 2015) as an attack demonstrates that this is often the case.

**When gradients succeed.** While saliency maps may not be the best approach in general for capturing information about nearby segments of the model’s decision boundary, there are cases in which it serves as a good approximation. Recent work has proposed using the Lipschitz continuity of an attribution method to characterize the difference between the attributions of an input serves as a good approximation. Recent work has proposed using the Lipschitz continuity of an attribution method to characterize the difference between the attributions of an input.

**Definition 4 (Attribution Robustness)** An attribution method \(g(\mathbf{x})\) is \((\lambda, \delta)\)-locally robust at the evaluated point \(\mathbf{x}\) if \(\forall \mathbf{x}' \in B(\mathbf{x}, \delta), \|g(\mathbf{x}') - g(\mathbf{x})\| \leq \lambda \|\mathbf{x}' - \mathbf{x}\|\).

**Proposition 1** Suppose that \(f\) has a \((\lambda, \delta)\)-robust saliency map \(g_\delta\) at \(\mathbf{x}\). \(\mathbf{x}'\) is the closest point on the closest decision boundary segment to \(\mathbf{x}\) and \(\|\mathbf{x}' - \mathbf{x}\| \leq \delta\), and that \(\mathbf{n}\) is the normal vector of that boundary segment. Then \(\|\mathbf{n} - g_\delta(\mathbf{x})\| \leq \lambda \|\mathbf{x} - \mathbf{x}'\|\).

Proposition 1 therefore provides the following insight: for networks that admit robust attributions (Chen et al. 2019; Wang et al. 2020c), the saliency map is a good approximation to the boundary vector. As prior work has demonstrated the close correspondence between robust prediction and robust attributions (Wang et al. 2020c; Chalasani et al. 2020), this in turn suggests that explanations on robust models will more closely resemble boundary normals.

As training robust models can be expensive, and may not come with guarantees of robustness, post-processing techniques like randomized smoothing (Cohen et al. 2019), have been proposed as an alternative. Dombrowski et al. (2019) noted that models with softplus activations \(y = 1/\beta \log(1 + \exp(\beta x))\) approximate smoothing, and in fact give an exact correspondence for single-layer networks. Combining these insights, we arrive at Theorem 1 which suggests that the saliency map on a smoothed model approximates the closest boundary normal vector well; the similarity is inversely proportional to the standard deviation of the noise used to smooth the model.
Theorem 1 Let \( m(x) \) be a one-layer ReLU network, and denote its smoothed counterpart under the Gaussian with standard deviation \( \sigma \) as \( m_s(x) \), and \( g(x) \) be the saliency map for \( m_s(x) \). If \( x' \) is the closest adversarial example to \( x \), and \( \forall x'' \in B(x, ||x - x'||) \cdot ||g(x'')|| \geq c \), then the following statement holds: \[ ||g(x) - g(x')|| \lesssim \lambda \] where \( \lambda \approx O\left(\frac{1}{\sigma}\right) \).

Theorem 1 suggests that when randomized smoothing is used, the normal vector of the closest decision boundary segment and the saliency map are similar, and this similarity increases with the smoothness of the model’s boundaries. Because the saliency map for a smoothed model is equivalent to smooth gradient of its non-smooth counterpart (Wang et al., 2020c), the smooth gradient is a better choice whenever computing the exact boundary is too expensive in a standard model. We provide empirical validation of this in Figure 10.

4 Boundary-Based Attribution

Without the properties introduced by robust learning or randomized smoothing, the local gradient, i.e. saliency map, may not be a good approximation of decision boundaries. In this section, we build on the insights of our analysis to present a set of novel attribution methods that explicitly incorporate the normal vectors of nearby boundary segments. Importantly, these attribution methods can be applied to models that are not necessarily robust, to derive explanations that capture many of the beneficial properties of explanations for robust models.

Using the normal vector of the closest decision boundary to explain a classifier naturally leads to Definition 5 which defines attributions directly from the normal of the closest decision boundary.

Definition 5 (Boundary-based Saliency Map (BSM)) Given \( f \) and an input \( x \), we define Boundary-based Saliency Map \( B_S(x) \) as follows: \( B_S(x) := \partial f_c(x')/\partial x' \), where \( x' \) is the closest adversarial example to \( x \), i.e. \( c = F(x) \neq F(x') \) and \( \forall x_m, ||x_m - x|| < ||x' - x|| \rightarrow F(x) = F(x_m) \).

Incorporating More Boundaries. The main limitation of using Definition 5 as a local explanation is obvious: the closest decision boundary only captures one segment of the entire decision surface. Even in a small network, there will be numerous boundary segments in the vicinity of a relevant point. Taking inspiration from Integrated Gradients, Definition 6 proposes the Boundary-based Integrated Gradient (BIG) by aggregating the attributions along a line between the input and its closest boundary segment.

Definition 6 (Boundary-based Integrated Gradient (BIG)) Given \( f \) and an input \( x \), we define Boundary-based Integrated Gradient \( B_I(x) \) as follows: \( B_I(x) := g_{IG}(x; x') \), where \( x' \) is the nearest adversarial example to \( x \), i.e. \( c = F(x) \neq F(x') \) and \( \forall x_m, ||x_m - x|| < ||x' - x|| \rightarrow F(x) = F(x_m) \).

Geometric View of BIG. BIG explores a linear path from the boundary point to the target point. Because points on this path are likely to traverse different activation polytopes, the gradient of intermediate points used to compute \( g_{IG} \) are normals of linear extensions of their local boundaries. As the input gradient is identical within a polytope \( P_i \), the aggregate computed by BIG sums each gradient \( w_i \) along the path and weights it by the length of the path segment intersecting with \( P_i \). In other words, one may view IG as an exploration of the model’s global geometry that aggregates all boundaries from a fixed reference point, whereas BIG explores the local geometry around \( x \). In the former case, the global exploration may reflect boundaries that are not particularly relevant to model’s observed behavior at a point, whereas the locality of BIG may aggregate boundaries that are more closely related (a visualization is shown in Fig. 1a).

Finding nearby boundaries. Finding the exact closest boundary segment is identical to the problem of certifying local robustness (Kolter & Wong, 2018; Lee et al., 2020; Leino et al., 2021b; Tjeng et al., 2019; Weng et al., 2018), which is NP-hard for piecewise-linear models (Sinha et al., 2020). To efficiently find an approximation of the closest boundary segment, we leverage and ensemble techniques for generating adversarial examples, i.e. PGD (Madry et al., 2018), AutoPGD (Croce & Hein, 2020) and CW (Carlini & Wagner, 2017), and use the closest one found given a time budget. The details of our implementation are discussed in Section 5 where we show that this yields good results in practice.
Table 1: Results of several attribution methods over 1500 images of ImageNet using a standard and robust ResNet50 (training $\epsilon$ is reported in the first column). BIG: Boundary-based Integrated Gradient. BSM: Boundary-based Saliency Map. AGI: Adversarial Gradient Integration. SM: Saliency Map. GTI: $\text{grad} \times \text{input}$. SG: Smoothed Gradient. IG: Integrated Gradient. See Appendix E for the corresponding boxplot.

5 EVALUATION

In this section, we first validate that the attribution vectors are more aligned to normal vectors of nearby boundaries in robust models (Fig. 3a). We secondly show that boundary-based attributions provide more “accurate” explanations – attributions highlight features that are actually relevant to the label – both visually (Fig. 4 and 5) and quantitatively (Table 1). Finally, we show that in a standard model, whenever attributions more align with the boundary attributions, they are more “accurate”.

General Setup. We conduct experiments over two data distributions, ImageNet (Russakovsky et al., 2015) and CIFAR-10 (Krizhevsky et al.). For ImageNet, we choose 1500 correctly-classified images from ImageNette (Howard), a subset of ImageNet, with bounding box area less than 80% of the original source image. For CIFAR-10, we use 5000 correctly-classified images. All standard and robust deep classifiers are ResNet50. All weights are pretrained and publicly available (Engstrom et al., 2019). Implementation details of the boundary search (by ensembling the results of PGD, CW and AutoPGD) and the hyperparameters used in our experiments, are included in Appendix B.2.

5.1 ROBUSTNESS → BOUNDARY ALIGNMENT

In this subsection, we show that SM and IG better align with the normal vectors of the decision boundaries in robust models. For SM, we use BSM as the normal vectors of the nearest decision boundaries and measure the alignment by the $\ell_2$ distance between SM and BSM following Proposition 1. For IG, we use BIG as the aggregated normal vectors of all nearby boundaries because...
IG also incorporates more boundary vectors. Recently, Pan et al. (2021) also provides Adversarial Gradient Integral (AGI) as an alternative way of incorporating the boundary normal vectors into IG. We first use both BIG and AGI to measure how well IG aligns with boundary normals and later compare them in Sec. 5.2 followed by a formal discussion in Sec. 7.

Aggregated results for standard models and robust models are shown in Fig. 3. It shows that adversarial training with bigger $\epsilon$ encourages a smaller difference between the difference between attributions and their boundary variants. Particularly, using $\ell_{2}$ norm and setting $\epsilon = 3.0$ are most effective for ImageNet compared to $\ell_{\infty}$ norm bound. One possible explanation is that the $\ell_{2}$ space is special because training with $\ell_{\infty}$ bound may encourage the gradient to be more Lipschitz in $\ell_{1}$ because of the duality between the Lipschitzness and the gradient norm, whereas $\ell_{2}$ is its own dual.

### 5.2 Boundary Attribution → Better Localization

In this subsection, we show boundary attributions (BSM, BIG and AGI) better localize relevant features. Besides SM, IG and SG, we also focus on other baseline methods including Grad $\times$ Input (GTI) (Simonyan et al., 2013) and DeepLIFT (rescale rule only) (Shrikumar et al., 2017) that are reported to be more faithful than other related methods (Adebayo et al., 2018; 2020).

In an image classification task where ground-truth bounding boxes are given, we consider features within a bounding box as more relevant to the label assigned to the image. Our evaluation is performed over ImageNet only because no bounding box is provided for CIFAR-10 data. The metrics used for our evaluation are: 1) Localization (Loc.) (Chattopadhyay et al., 2017) evaluates the intersection of areas with the bounding box and pixels with positive attributions; 2) Energy Game (EG) (Wang et al., 2020a) instead evaluates computes the portion of attribute scores within the bounding box. While these two metrics are common in the literature, we propose the following additional metrics: 3) Positive Percentage (PP) evaluates the portion of positive attributions in the bounding box because a naive assumption is all features within bounding boxes are relevant to the label (we will revisit this assumption in Sec. 6); and 4) Concentration (Con.) sums the absolute value of attribution scores over the distance between the “mass” center of attributions and each pixel within the bounding box. Higher Loc., EG, PP and Con. are better results. We provide formal details for the above metrics in Appendix B.1.

We show the average scores for ResNet50 models in Table 1 where the corresponding boxplots can be found in Appendix B.4. BIG is noticeably better than other methods on Loc. EG, PP and Con. scores for both robust and standard models and matches the performance of SG on EG for a standard model. Notice that BSM is not significantly better than others in a standard model, which confirms our motivation of BIG – that we need to incorporate more nearby boundaries because a single boundary may not be sufficient to capture the relevant features.

We also measure the correlation between the alignment of SM and BS with boundary normals and the localization abilities, respectively. For SM, we use BSM to represent the normal vectors of the boundary. For IG, we use AGI and BIG. For each pair $X-Y$ in $\{\text{SM-BSM, IG-AGI, IG-BIG}\}$, we measure the empirical correlation coefficient between $-\|X - Y\|_{2}$ and the localization scores of $X$ in a standard ResNet50 and the result is shown in Fig. 3b. Our results suggest that when the attribution methods better align with their boundary variants, they can better localize the relevant features in terms of the Loc. and EG. However, PP and Con. have weak and even negative correlations. One possible explanation is that the high PP and Con. of BIG and AGI compared to IG (as shown in Table 1) may also come from the choice of the reference points. Namely, compared to a zero vector, a reference point on the decision boundary may better filter out noisy features.

We end our evaluations by visually comparing the proposed method, BIG, against all other attribution methods for the standard ResNet50 in Fig. 4 and for the robust ResNet50 in Fig. 5 which demonstrates that BIG can easily and efficiently localize features that are relevant to the prediction. More visualizations can be found in the Appendix E.

**Summary.** Taken together, we close the loop and empirical show that standard attributions in robust models are visually more interpretable because they better capture the nearby decision boundaries. Therefore, the final take-away from our analytical and empirical results is if more resources are devoted to training robust models, effectively identical explanations can be obtained using much less costly standard gradient-based methods, i.e. IG.
6 DISCUSSION

Baseline Sensitivity. It is naturally to treat BIG frees users from the baseline selection in explaining non-linear classifiers. Empirical evidence has shown that IG is sensitive to the baseline inputs (Sturmfels et al., 2020). We compare BIG with IG when using different baseline inputs, white or black images. We show an example in Fig 6b. For the first two images, when using the baseline input as the opposite color of the dog, more pixels on dogs receive non-zero attribution scores. Whereas backgrounds always receive more attribution scores when the baseline input has the same color as the dog. This is because $g_{IG}(x)_{i} \propto (x - x_b)_i$ (see Def. 2) that greater differences in the input feature and the baseline feature can lead to high attribution scores. The third example further questions the readers using different baselines in IG whether the network is using the white dog to predict Labrador retriever. We demonstrate that conflicts in IG caused by the sensitivity to the baseline selection can be resolved by BIG. BIG shows that black dog in the last row is more important for predicting Labrador retriever and this conclusion is further validated by our counterfactual experiment in Appendix D. Overall, the above discussion highlights that BIG is significantly better than IG in reducing the non-necessary sensitivity in the baseline selection.

Limitations. We identify two limitations of the work. 1) Bounding boxes are not perfect ground-truth knowledge for attributions. In fact, we find a lot of examples where the bounding boxes either fail to capture all relevant objects or are too big to capture relevant features only. Fixing mislabeled bounding boxes still remain an open question and should benefit more expandability research in general. 2) Our analysis only targets on attributions that are based on end-to-end gradient computations. That is, we are not able to directly characterize the behavior of perturbation-based approaches, i.e. Mask (Fong & Vedaldi, 2017), and activation-based approaches, i.e. GradCAM (Selvaraju et al., 2017) and Feature Visualization (Olah et al., 2017).

7 RELATED WORK

Ilyas et al. (2019) shows an alternative way of explaining why robust models are more interpretable by showing robust models usually learn robust and relevant features, whereas our work serves as a geometrical explanation to the same empirical findings in using attributions to explain deep models. Our analysis suggests we need to capture decision boundaries in order to better explain classifiers,
Figure 6: (a): Qualitative comparisons between IG with black baseline, BIG and AGI. BIG can use any boundary search approaches or an ensemble of them while AGI uses PGD only. AGI fails to meet the symmetry axiom \cite{Sundararajan2017}, where BIG satisfies all axioms that IG satisfies, i.e. completeness. (b): Comparisons of IG with black and white baselines with BIG. Predictions are shown in the first column.

8 CONCLUSION

In summary, we rethink the target question an explanation should answer for a classification task, the important features that the classifier uses to place the input into a specific side of the decision boundary. We find the answer to our question relates to the normal vectors of decision boundaries in the neighborhood and propose BSM and BIG as boundary attribution approaches. Empirical evaluations on STOA classifiers validate that our approaches provide more concentrated, sharper and more accurate explanations than existing approaches. Our idea of leveraging boundaries to explain classifiers connects explanations with the adversarial robustness and help to encourage the community to improve model quality for explanation quality.
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A Theorems and Proofs

A.1 Proof of Proposition

Proposition 1 Suppose that $f$ has a $(\lambda, \delta)$-robust saliency map $g_{\delta}$ at $x$, $x'$ is the closest point on the closest decision boundary segment to $x$ and $||x' - x|| \leq \delta$, and that $n$ is the normal vector of that boundary segment. Then $||n - g_{\delta}(x)|| \leq \lambda||x - x'||$.

To compute $n$ can be efficiently computed by taking the derivative of the model’s output w.r.t to the point that is on the decision boundary such that $n = \frac{\partial f(x)}{\partial x}$ and $\forall x, \in \mathbb{R}^{d}, F(x) = F(x)$ if $||x_m - x|| \leq ||x' - x||$.

Because we assume $||x - x'|| \leq \delta$, and the model has $(\lambda, \delta)$-Saliency Map, then by Def. 4 we have

$$||n - g_{\delta}(x)|| \leq \lambda||x - x'||$$

A.2 Proof of Theorem

Theorem 1 Let $m(x) = \text{ReLU}(w^{T}x)$ be a one-layer network and when using randomized smoothing, we write $m_{\sigma}(x)$. Let $g(x)$ be the SM for $m_{\sigma}(x)$ and suppose $\forall x'' \in B(x, ||x - x'||), ||g(x'')|| \geq c$ where $x'$ is the closest adversarial example, we have the following statement holds: $||g(x) - g(x')|| \propto O(\frac{1}{\sigma^{2}})$.

Before we start the proof, we firstly introduce Randomized Smoothing and its theorem that certify the robustness.

Definition 7 (Randomized Smoothing (Cohen et al., 2019)) Suppose $F(x) = \arg \max_c f_c(x)$, the smoothed classifier $G(x)$ is defined as

$$G(x) := \arg \max_c \Pr[F(x + \epsilon) = c]$$ (1)

where $\epsilon \sim \mathcal{N}(0, \sigma^2 I)$

Theorem 2 (Theorem 1 from Cohen et al. (Cohen et al., 2019)) Suppose $F$ and $G$ are defined in Def. 7. For a target instance $x$, suppose $\Pr[F(x) = c_A]$ is lower-bounded by $p_A$ and $\max_{c \neq c_A} \Pr[F(x) = c]$ is upper-bounded by $p_B$, then

$$G(x') = c_A \ \forall x' \in B(x, R_{\sigma})$$ (2)

where

$$R_{\sigma} = \frac{\sigma}{2}[\Phi^{-1}(p_A) - \Phi^{-1}(p_B)]$$ (3)

and $\Phi$ is the c.d.f of Gaussian.

We secondly introduce a theorem that connects Randomized Smoothing and Smoothed Gradient.

Theorem 3 (Proposition 1 from Want et al. (Wang et al., 2020c)) Suppose a model $f(x)$ satisfies $\max |f(x)| < \infty$. For a Smoothed Gradient $g_{SG}(x)$, we have

$$g_{SG}(x) = \frac{\partial (f \circ q)(x)}{\partial x}$$ (4)

where $q(x) = \mathcal{N}(0, \sigma^2 I)$ and $\circ$ denotes the convolution operation.

Finally, we introduce two theorems that connects Smoothed Gradient with Softplus network.

Theorem 4 (Theorem 1 from Dombrowski et al. (Dombrowski et al., 2019)) Suppose $m(x)$ is a feed-forward network with softplus-$\beta$ activation and $\|\frac{\partial m(x)}{\partial x}\| \geq c$ if $x \in B(x, \epsilon)$, then

$$\|\frac{\partial m(x)}{\partial x} - \frac{\partial m(x')}{\partial x'}\| \leq \beta C \delta_m(x, x')$$ (5)
Proof:

We now begin our proof for Theorem 1.

Denote a one-layer ReLU network as \( f(x) = \text{ReLU}(w^\top x) \) and a one-layer softplus network as \( m_{\beta}(x) = \text{softplus}_{\beta}(w^\top x) \), then the following statement holds:

\[
\mathbb{E}_{\epsilon_i \sim p_{\beta}} \left[ \frac{\partial m(x + \epsilon_i)}{\partial x} \right] = \frac{\partial m_{\beta}(x)}{\partial x}.
\]

(6)

where \( p_{\beta}(\epsilon_i) = \frac{\text{exp}(\beta \epsilon_i / 2) + \text{exp}(-\beta \epsilon_i / 2))}{\text{exp}(\beta \epsilon_i / 2) + \text{exp}(-\beta \epsilon_i / 2)} \).

We now begin our proof for Theorem 2.

Proof:

Given a one-layer ReLU network \( m(x) = \text{ReLU}(w^\top x) \) that takes an input \( x \in \mathbb{R}^d \) and outputs the logit score for the class of interest, WLOG we assume \( w \in \mathbb{R}^d \) is the \( j \)-th column of the complete weight matrix \( W \in \mathbb{R}^{d \times n} \), where \( j \) is the class of interest. With Theorem 5 we know that

\[
\mathbb{E}_{\epsilon_i \sim p_{\beta}} \left[ \frac{\partial m(x + \epsilon_i)}{\partial x} \right] = \frac{\partial m_{\beta}(x)}{\partial x}.
\]

(7)

Dombrowski et al. (Dombrowski et al., 2019) points out that the random distribution \( p_{\beta}(\epsilon_i) = \frac{\text{exp}(\beta \epsilon_i / 2) + \text{exp}(-\beta \epsilon_i / 2))}{\text{exp}(\beta \epsilon_i / 2) + \text{exp}(-\beta \epsilon_i / 2)} \) closely resembles a normal distribution with a standard deviation

\[
\sigma = \sqrt{\frac{\log(2) \sqrt{2\pi}}{\beta}}
\]

(8)

Therefore, we have the following relation

\[
\mathbb{E}_{\epsilon_i \sim p_{\beta}} \left[ \frac{\partial m(x + \epsilon_i)}{\partial x} \right] \approx \mathbb{E}_{\epsilon_i \sim N(0, \sigma^2 I)} \left[ \frac{\partial m(x + \epsilon_i)}{\partial x} \right]
\]

(9)

The LHS of the above equation is Smoothed Gradient, or equivalently, it is the Saliency Map of a smoothed classifier \( m_{\beta} \) due to Theorem 2. Eq 7 and 9 show that we can analyze randomized smoothing with the tool of an intermediate softplus network \( m_{\beta}/||w|| \).

\[
\frac{\partial m_{\sigma}(x)}{\partial x} = \mathbb{E}_{\epsilon_i \sim N(0, \sigma^2 I)} \left[ \frac{\partial m(x + \epsilon_i)}{\partial x} \right] \approx \frac{\partial m_{\beta}(x)}{\partial x}
\]

(10)

and we denote \( g(x) := \frac{\partial m_{\sigma}(x)}{\partial x} \). Now consider Theorem 4 we have

\[
||g(x) - g(x')|| \leq \frac{\beta}{c||w||}||W||_F^2 ||x - x'||
\]

(11)

(12)

where \( x' \) is the closest adversarial example. Because \( m_{\sigma} \) is certified to be robust within the neighborhood \( B(x, R_{\sigma}) \), therefore, the closest decision boundary is at least \( R_{\sigma} \) distance away from the evaluated point \( x \). We then have

\[
||g(x) - g(x')|| \leq \frac{\beta}{c||w||}||W||_F^2 R_{\sigma}
\]

(13)

Now let's substitute \( \beta \) and \( R_{\sigma} \) with \( \sigma \) using Eq. 8 and 3 we arrive at

\[
||g(x) - g(x')|| \leq \lambda
\]

(14)
where
\[
\lambda = \frac{||W||^2_2}{c||W||} \log(2) \frac{\sqrt{2\pi}}{\sigma^2} \left[ \Phi^{-1}(p_A) - \Phi^{-1}(p_B) \right]
\] (15)
\[
= \frac{1}{\sigma} \frac{\sqrt{2\pi} ||W||^2_2}{2c||W||} \log(2) [\Phi^{-1}(p_A) - \Phi^{-1}(p_B)]
\] (16)
\[
\propto O \left( \frac{1}{\sigma} \right)
\] (17)

We use $\preceq$ instead of $\leq$ because of the fact we approximate the randomized smoothing with a similar distribution $p_{\beta}(\epsilon_i) = \frac{\beta}{\exp(\beta \epsilon_i/2) + \exp(-\beta \epsilon_i/2)}$. In fact, a more rigorous proof exists when considering $p_{\beta}$ other than Gaussian. We refer the readers who are interested in the content to Yang et al. (Yang et al., 2020).

B EXPERIMENT DETAILS AND ADDITIONAL RESULTS

B.1 METRICS WITH BOUNDBING BOXES

We will use the following extra notations in this section. Let $X$, $Z$ and $U$ be a set of indices of all pixels, a set of indices of pixels with positive attributions, and a set of indices of pixels inside the bounding box for a target attribution map $g(x)$. We denote the cardinality of a set $S$ as $|S|$.

Localization (Loc.) (Chattopadhyay et al., 2017) evaluates the intersection of areas with the bounding box and pixels with positive attributions.

**Definition 8 (Localization)** For a given attribution map $g(x)$, the localization score (Loc.) is defined as
\[
\text{Loc} := \frac{|Z \cap U|}{|U| + |Z \cap (X \setminus U)|}
\] (18)

Energy Game (EG) (Wang et al., 2020a) instead evaluates computes the portion of attribute scores within the bounding box.

**Definition 9 (Energy Game)** For a given attribution map $g(x)$, the energy game EG is defined as
\[
\text{EG} := \frac{\sum_{i \in Z \cap U} g(x)_i}{\sum_{i \in X} \max(g(x)_i, 0)}
\] (19)

Positive Percentage (PP) evaluates the sum of positive attribute scores over the total (absolute value of) attribute scores within the bounding box.

**Definition 10 (Positive Percentage)** Let $V$ be a set of indices of all pixels with negative attribution scores, for a given attribution map $g(x)$, the positive percentage PP is defined as
\[
\text{PP} := \frac{\sum_{i \in Z \cap U} g(x)_i}{\sum_{i \in Z \cap U} g(x)_i - \sum_{i \in V \cap U} g(x)_i}
\] (20)

Concentration (Con.) evaluates the sum of weighted distances by the “mass” between the “mass” center of attributions and each pixel within the bounding box. Notice that the computation of $c_x$ and $c_y$ can be computed with `scipy.ndimage.center_of_mass`. This definition encourages that pixels with high absolute value of attribution scores to be closer to the mass center.

**Definition 11 (Concentration)** For a given attribution map $g(x)$, the concentration Con. is defined as follows
\[
\text{Con.} := \sum_{i \in U} \frac{g(x)_i}{\sqrt{(i_x - c_x)^2 + (i_y - c_y)^2}}
\] (21)
where $\hat{g}$ is the normalized attribution map so that $\hat{g}_i = g_i / \sum_{i \in U} |g_i|$. $i_x, i_y$ are the coordinates of the pixel and

$$c_x = \frac{\sum_{i \in U} i_x \hat{g}(x)_i}{\sum_{i \in U} \hat{g}(x)_i}, \quad c_y = \frac{\sum_{i \in U} i_y \hat{g}(x)_i}{\sum_{i \in U} \hat{g}(x)_i} \quad (22)$$

Besides metrics related to bounding boxes, there are other metrics in the literature used to evaluate attribution methods (Adebayo et al., 2018; Ancona et al., 2017; Samek et al., 2016; Wang et al., 2020b; Yeh et al., 2019). We focus on metrics that use provided bounding boxes, as we believe that they offer a clear distinction between likely relevant features and irrelevant ones.

### B.2 Implementing Boundary Search

Our boundary search uses a pipeline of PGDs, CW and AutoPGD. Adversarial examples returned by each method are compared with others and closer ones are returned. If an adversarial example is not found, the pipeline will return the point from the last iteration of the first method (PGDs in our case). Hyper-parameters for each attack can be found in Table 2. The implementation of PGDs and CW are based on Foolbox (Rauber et al., 2020, 2017) and the implementation of AutoPGD is based on the authors’ public repository1 (we only use apgd-ce and apgd-dlr losses for efficiency reasons). All computations are done using a GPU accelerator Titan RTX with a memory size of 24 GB. Comparisons on the results of the ensemble of these three approaches are shown in Fig. 9a.

1https://github.com/fra31/auto-attack
Figure 9: (a): Pipeline: the methods used for boundary search. Avg Distance: the average $\ell_2$ distance between the input to the boundary. Success Rate: the percentage when the pipeline returns an adversarial example. Time: per-instance time with a batch size of 64. We are using much bigger $\epsilon$s for robust models, so the success rates are higher than a standard model. (b): Hyper-parameters used for AGI. We use the default parameters from the authors’ implementation for ImageNet and make minimal changes for CIFAR-10.

### B.3 Hyper-parameters for Attribution Methods

All attributions are implemented with Captum [Kokhlikyan et al., 2020] and visualized with Trulens [Leino et al., 2021a]. For BIG and IG, we use 20 intermediate points between the baseline and the input and the interpolation method is set to `riemann`, `trapezoid`. For AGI, we base on the authors’ public repository\footnote{https://github.com/pd90506/AGI}. The choice of hyper-paramters follow the default choice from the authors for ImageNet and we make minimal changes to adapt them to CIFAR-10 (see Fig. 9b).

To visualize the attribution map, we use the HeatmapVisualizer with `blur=10`, `normalization_type="signed_max"` and default values for other keyword arguments from Trulens.

### B.4 Detailed Results on Localization Metrics

We show the average scores for each localization metrics in Sec. 5. We also show the boxplots of the scores for each localization metrics in Fig. 7 for the standard ResNet50 model and Fig. 8 for the robust ResNet50 ($\ell_2[3.0]$). All higher scores are better results.

### B.5 Additional Experiment with Smoothed Gradient

We empirically validate Theorem 1 in deeper networks, i.e. ResNet50, which suggests that SM on BSM are more similar on models with randomized smoothing, which are known to be robust [Cohen et al., 2019]. To obtain meaningful explanations on smoothed models, which are implemented by evaluating the model on a large set of noised inputs, we assume that the random seed is known by
Table 2: Hyper-parameters used for adversarial attacks. adaptive means the actual step size is determined by $2 \times \epsilon/\text{max steps}$.  

|                | CIFAR10 | std          | robust        |
|----------------|---------|--------------|---------------|
| $\epsilon$     | [0.2, 0.4, 0.6, 0.8, 1.0] | [0.25, 0.5, 1.0, 1.5, 2.0] |
| max steps      | 100     | 100          |
| step size      | 5e-3    | 5e-3         |
| **PGDs**       |         |              |               |
| ImageNet       |         |              |               |
| $\epsilon$     | [36/255, 64/255, 0.3, 0.5, 0.7, 0.9, 1.1] | [1.0, 2.0, 3.0, 4.0, 5.0, 6.0] |
| max steps      | 100     | 100          |
| step size      | adaptive| adaptive     |
| **CW**         |         |              |               |
| ImageNet       |         |              |               |
| $\epsilon$     | 1.0     | 2.0          |
| max steps      | 100     | 100          |
| step size      | 1e-3    | 1e-3         |
| **CIFAR10**    |         |              |               |
| $\epsilon$     | 1.0     | 2.0          |
| max steps      | 100     | 100          |
| step size      | 6e-3    | 1.6e-2       |
| **AutoPGD**    |         |              |               |
| ImageNet       |         |              |               |
| $\epsilon$     | 1.1     | 6.0          |
| max steps      | 100     | 100          |
| step size      | 2.3e-2  | 1.2e-1       |

Figure 10: $\ell_2$ distances in logarithm between SG and BSG against different standard deviations $\sigma$ of the Gaussian noise. Results are computed on ResNet50. Notice the first column corresponds to $\sigma = 0$.  

the adversarial example generator, and search for perturbations that point towards a boundary on as many of the noised inputs simultaneously as possible. We do the boundary search for a subset of 500 images, as this computation is significantly more expensive than previous experiments. Instead
of directly computing SM and BSM on the smoothed classifier, we utilize the connection between randomized smoothing and SG (see Theorem 3 in the Appendix A); therefore, we compare the difference between SG on the clean inputs and SG on their adversarial examples (referred as BSG). The setup is shown as follows.

To generate the adversarial examples for the smoothed classifier of ResNet50 with randomized smoothing, we need to compute back-propagation through the noises. The noise sampler is usually not accessible to the attacker who wants to fool a model with randomized smoothing. However, our goal in this section is not to reproduce the attack with similar setup in practice, instead, what we are after is the point on the boundary. We therefore do the noise sampling prior to run PGD attack, and we use the same noise across all the instances. The steps are listed as follows:

1. We use `numpy.random.randn` as the sampler for Gaussian noise with its random seed set to 2020. We use 50 random noises per instance.
2. In PGD attack, we aggregate the gradients of all 50 random inputs before we take a regular step to update the input.
3. We set $\epsilon = 3.0$ and we run at most 40 iterations with a step size of $2 \times \epsilon/40$.
4. The early stop criteria for the loop of PGD is that when less than 10% of all randomized points have the original prediction.
5. When computing Smooth Gradient for the original points or for the adversarial points, we use the same random noise that we generated to approximate the smoothed classifier.

We plot the results in Fig. 10. Notably, the trend of the log difference against the standard deviation $\sigma$ used for the Gaussian noise validates that the qualitative meaning of Theorem 1 holds even for large networks.
C  SYMMETRY OF ATTRIBUTION METHODS

Sundararajan et al. (2017) prove that a linear path is the only path integral that satisfies symmetry; that is, when two features’ orders are changed for a network that is not using any order information from the input, their attribution scores should not change. One simple way to show the importance of symmetry by the following example and we refer Sundararajan et al. (2017) to readers for more analysis.

Example 1  Consider a function \( f(x, y) = \min(x, y) \) and to attribute the output of \( f \) to the inputs at \( x = 1, y = 1 \) we consider a baseline \( x = 0, y = 0 \). An example non-linear path from the baseline to the input can be \( (x = 0, y = 0) \rightarrow (x = 1, y = 0) \rightarrow (x = 1, y = 1) \). On this path, \( f(x, y) = \min(x, y) = y \) after the point \( (x = 1, y = 0) \); therefore, gradient integral will return 0 for the attribution score of \( x \) and 1 for \( y \) (we ignore the infinitesimal part of \( (x = 0, y = 0) \rightarrow (x = 1, y = 0) \)). Similarly, when choosing a path \( (x = 0, y = 0) \rightarrow (x = 0, y = 1) \rightarrow (x = 1, y = 1) \), we find \( x \) is more important. Only the linear path will return 1 for both variables in this case.

D  COUNTERFACTUAL ANALYSIS IN THE BASELINE SELECTION

The discussion in Sec. 6 shows an example where there are two dogs in the image. IG with black baseline shows that the body of the white dog is also useful to the model to predict its label and the black dog is a mix: part of the black dog has positive attributions and the rest is negatively contribute to the prediction. However, our proposed method BIG clearly shows that the most important part is the black dog and then comes to the white dog. To validate where the model is actually using the white dog, we manually remove the black dog or the white dog from the image and see if the model retain its prediction. The result is shown in Fig. 11. Clearly, when removing the black dog, the model changes its prediction from Labrador retriever to English foxhound while removing the white dog does not change the prediction. This result helps to convince the reader that BIG is more reliable than IG with black baseline in this case as a more faithful explanation to the classification result for this instance.

E  ADDITIONAL VISUALIZATIONS FOR BIG

More visualizations comparing BIG with other attributions can be found in Fig. 12 and 13.
Figure 12: Visualizations of different attributions for a standard ResNet50
Figure 13: Visualizations of different attributions for a robust ($\ell_2 | 3.0$) ResNet50