Out-of-Distribution Generalization via Risk Extrapolation

David Krueger 1 2 Ethan Caballero 1 Jörn-Henrik Jacobsen 3 4 Amy Zhang 1 5 6 Jonathan Binas 1 2 Rémi Le Priol 1 2 Aaron Courville 1 2

Abstract

Generalizing outside of the training distribution is an open challenge for current machine learning systems. A weak form of out-of-distribution (OoD) generalization is the ability to successfully interpolate between multiple observed distributions. One way to achieve this is through robust optimization, which seeks to minimize the worst-case risk over convex combinations of the training distributions. However, a much stronger form of OoD generalization is the ability of models to extrapolate beyond the distributions observed during training. In pursuit of strong OoD generalization, we introduce the principle of Risk Extrapolation (REx). REx can be viewed as encouraging robustness over affine combinations of training risks, by encouraging strict equality between training risks. We show conceptually how this principle enables extrapolation, and demonstrate the effectiveness and scalability of instantiations of REx on various OoD generalization tasks. Our code can be found at https://github.com/capybaralet/REx_code_release.

1. Introduction

Improving the generalization of deep learning models has become a major research topic, with many different threads of research including Bayesian deep learning (Neal, 1996; Gal, 2016), adversarial (Engstrom et al., 2019; Jacobsen et al., 2018) and non-adversarial (Hendrycks & Dietterich, 2019; Yin et al., 2019) robustness, causality (Arjovsky et al., 2019), and other works aimed at distinguishing statistical features from semantic features (Gowal et al., 2019; Geirhos et al., 2018). While neural networks often exhibit superhuman generalization performance on the training distribution, they can be extremely sensitive to minute changes in distribution (Su et al., 2019; Engstrom et al., 2017; Recht et al., 2019). This presents a major bottleneck for their practical application.

In this work, we consider out-of-distribution (OoD) generalization, where a model must generalize to new distributions at test time without seeing any training data from them. We assume a fixed underlying task, and access to labeled data from multiple training environments. We also assume that variation in these environments is somewhat representative of the variations we will see at test time. However, we also seek to make good predictions even when these variations are extreme in magnitude.

As a motivating example, consider the colored MNIST (CMNIST) dataset from Arjovsky et al. (2019): there are two training environments in which digits are colored either red or green, with the color being either 80% or 90% correlated with the (binary) label, respectively, and a test environment where color is 10% correlated (i.e. 90% anti-correlated) with the label. Generalization to the test set can be viewed as extrapolating over possible values of the correlation (0% to 100%) between color and label. Thus seeing minor variation in this correlation in the training environments is a hint that the correlation could vary even more, and even be reversed, at test time. Like the Invariant Risk Minimization (IRM) approach of Arjovsky et al. (2019), our method of Risk Extrapolation (REx) promotes generalization when “spurious” features (such as color, in this example) are predictive during training, but not at test time. For example, in the colored MNIST example, a model that bases its predictions on color will actually do worse than random guessing at test time; the correct way to generalize is to use the “stable” features relating to digit shape.

The classic example of spurious features is the use of background features in object recognition. Most pictures of cows appear in pastures, and so common machine learning approaches struggle to classify pictures of cows on the beach (Beery et al., 2018). In order to build models based on stable features, we can use data collected from different naturally occurring environments (such as geographical locations), with the hope that the natural variations in distribution across environments will highlight spurious features. How spurious features relate to OoD generalization as a whole is currently unclear, although there is evidence that
the lack of robustness of current computer vision models is related to the problem of spurious features (Geirhos et al., 2018; Ilyas et al., 2019).

To provide a more specific explanation for how performing worse on training environments could actually help generalization, we return to the colored MNIST example. Since color is more correlated with the label in the second environment than in the first, a predictor which uses color as a feature to predict the labels will do better on the second environment. By encouraging performance to be equal across training environments, we can prevent the predictor from using color as a feature, and thus generalize to the test environment where it is anti-correlated with the label.

Our contributions are as follows:

1. We introduce the principle of Risk Extrapolation (REx) and explain geometrically and probabilistically how it is able to learn stable features.

2. We derive two practical instantiations of REx. Minimax REx (MM-REx) as a form of robust optimization, and Variance REx (V-REx) minimizing the variance of training risks.

3. We identify crucial considerations for successful training of REx and the related IRM method (Arjovsky et al., 2019).

4. We compare our method to IRM and ERM on a diverse suite of tasks, demonstrating comparable or superior performance in most cases, as well as a limitation of REx in the case of environment-dependent label noise.

2. Background

We consider a multi-environment setting, where our goal is to find parameters $\theta$ that perform well on unseen environments, given a set of training environments, $\mathcal{E} = \{e_1, \ldots, e_m\}$. We assume the loss function, $\ell$, and the input and target spaces are fixed across environments. Each environment, $e$ has an associated dataset $D_e$ and data distribution $\mathcal{D}_e$, and a corresponding risk function, $\mathcal{R}_e : \mathbb{R}^n \rightarrow \mathbb{R}$, which maps $\theta$ to the expected loss on that environment’s data distribution:

$$\mathcal{R}_e(\theta) = \mathbb{E}_{(x,y) \sim \mathcal{D}_e} \ell(\theta(x), y) \quad (1)$$

We refer to members of the set $\{\mathcal{R}_e | e \in \mathcal{E}\}$ as the training risks or simply risks, and we use $\mathcal{R}_i$ to refer to the risk on the $i$-th environment (and similarly for $D_i$ and $\mathcal{D}_i$).

The Empirical Risk Minimization (ERM) approach simply minimizes the average loss across all training examples, regardless of environment:

$$\mathcal{R}_{\text{ERM}}(\theta) = \mathbb{E}_{(x,y) \sim \bigcup_{e \in \mathcal{E}} D_e} \ell(\theta(x), y) \quad (2)$$

Empirical risk minimization has strong theoretical foundations in the case of i.i.d. data (Vapnik, 1992), and often
works well in practice, outperforming more sophisticated methods (Chen et al., 2019). However, it can fail dramatically when test environments differ significantly from training environments (Tzeng et al., 2017).

Robust optimization (RO) (Ben-Tal et al., 2009) is another approach to solving the above problem, designed to handle distributional shift. Robust optimization minimizes the worst performance over a set of possible environments \( F \), that is, \( \max_{x \in F} R_e(\theta) \). Given a single training environment, a set of possible environments is often generated using some set of allowed perturbations, e.g., \( P_e(x) \) varying within a KL-divergence \( \epsilon \)-ball of the empirical distribution (Bagnell, 2005). When multiple training environments are available, a straightforward approach is to use the empirical distribution of environments, setting \( F \) to \( E \):

\[
\mathcal{R}_{\text{RO}}(\theta) = \max_{e \in E} R_e(\theta) - r_e
\]

where \( r_e \) is an optional “baseline” risk for each environment, which could represent the difficulty or irreducible risk of that environment. We discuss baselines in more detail in the Appendix.

Invariant Risk Minimization (IRM) (Arjovsky et al., 2019) seeks for an invariant representation of inputs from different environments. The IRM principle states: “An invariant representation \( \Phi(X) \) is one such that the optimal linear predictor, \( w \), is the same across all environments, \( e \).” Robust optimization may fail to be invariant in this sense, since it may make use of features which are differentially useful across environments (such as color in CMNIST). This leads Arjovsky et al. (2019) to reject this minimax approach in favor of a bilevel optimization problem formulation of the IRM principle:

\[
\min_{\Phi, w} \sum_e R_e(w^\top \Phi(X^e)) \\
\text{s.t. } w \in \arg \min \, R_e(w^\top \Phi(X^e)), \quad \forall e \in E
\]

Arjovsky et al. (2019) also propose IRMv1 as a practical algorithm for solving the IRM problem:

\[
\min_{\Phi} \sum_e R_e(\Phi(X^e)) + \lambda \|\nabla_w R_e(w^\top \Phi(X^e))\|^2
\]

in which \( w \) is a fixed vector of ones, and \( \lambda \) controls the strength of the penalty term on gradients on \( w \). Large gradients on \( w \) indicate that \( w \) is not optimal for some environment. We frequently refer to “IRMv1” as simply “IRM”.

Despite being a form of, REx is more similar to IRM in spirit than Eqn. 3, as we share their motivation of discovering stable features as a means of generalizing OoD. At the same time, although IRM and REx behave quite similarly in most of our experiments, it is easy to show that satisfying the IRM objective does not necessarily lead to the risk across environments being equal. For instance, the IRM solution to the simplest structural equation model in Arjovsky et al. (2019) is an invariant prediction rule for which the risks do differ across environments, due to the environment-dependent noise on the targets.

3. Methods

We now explain and motivate the principle of risk extrapolation, which has two goals:

1. Reducing training risks
2. Increasing similarity of training risks

In general, these goals can be at odds with each other; decreasing the risk in the environment with the lowest risk also decreases the overall similarity of training risks. In this section, we explain how sacrificing performance on individual environments in order to perform more similarly across environments can help with OoD generalization, and present two methods, Minimax REx and Variance REx, for achieving, and balancing, goals (1) and (2).

3.1. Minimax REx

Here we present a robust optimization method that achieves the aims of risk extrapolation. We begin by observing that we can frame the robust optimization objective from eqn. 3 as a minimax objective over convex combinations of training risks:

\[
\mathcal{R}_{\text{RI}}(\theta) = \max_{\sum \lambda_e = 1} \sum_e \lambda_e R_e(\theta)
\]

Naturally, we refer to this objective as Risk Interpolation (RI). To arrive at Minimax REx (MM-REx), we relax this objective to allow a more general set of affine combinations of training risks:

\[
\mathcal{R}_{\text{MM-REx}}(\theta) = \max_{\sum \lambda_e = 1} \sum_e \lambda_e R_e(\theta)
\]

where \( m \) is the number of environments, and the hyperparameter \( \beta \) controls how much we seek to extrapolate in the space of risk functions. As \( \beta \to \infty \), this criterion enforces strict equality between training risks. If \( \beta \) is negative, it is simply an interpolation coefficient between RI \( (\beta = 0) \) and ERM \( (\beta = -1/m) \).

However, for positive values of \( \beta \), this criterion places negative weights on the risk of all but the worst-case environment.
The resulting $\mathcal{R}_{\text{MM-REx}}$ is an extrapolation of the training risks, exaggerating variations in the training distributions beyond what has been observed. Larger values of $\beta$ correspond to extrapolating farther from the convex hull of the training risks, and thus encourage a flatter “risk-plane” (see Figure 1). Note that the constraint that $\sum \lambda_i = 1$ means that there is always some pressure to decrease the average risk (but only by aggressively decreasing the highest risk). In practice, we found that this objective would typically enter a regime in which risks are close enough that which risk is highest rapidly changes between iterations, and a gradual decrease of average risk can take place.

### 3.2. REx via regularized optimization

A simpler approach that we find works well in practice is to use the average mean squared error (MSE) between risks, or equivalently, their variance. This yields Variance REx (V-REx), which is the algorithm we focus on in our experiments:

$$\mathcal{R}_{\text{V-REx}} \doteq \beta \text{Var}\{\mathcal{R}_1, ..., \mathcal{R}_m\} + \sum_{e=1}^{m} \mathcal{R}_e$$

(9)

Here $\beta \in [0, \infty)$ controls the balance between reducing average risk and enforcing equality of risks, with $\beta = 0$ recovering ERM, and $\beta \to \infty$ leading V-REx to focus entirely on making the risks equal. We plot the gradient vector field for the total risk as a function of two training risks $\mathcal{R}$ in Figure 3.2, and note that the gradients change smoothly at the diagonal for $\mathcal{R}_{\text{V-REx}}$, where risks are equal, but not for $\mathcal{R}_{\text{MM-REx}}$. Empirically, we also found that $\mathcal{R}_{\text{V-REx}}$ can lead to slightly more stable learning, see Appendix.

### 3.3. A probabilistic view on risk extrapolation

Since the operation of integration is a linear functional, and since we assume a fixed loss function, we have $\mathcal{R} = \int f_y(x, y) \ell(f_\theta(x), y)$ is linear in $P(x, y)$. This means we can understand risk extrapolation “term-wise” by looking at its effects at every point $(x, y)$. In particular, linear combinations of risk functions can be understood in terms of the corresponding linear combinations of probability density functions (PDFs). Figure 3.3 shows a graphical example of how REx allows us to effectively synthesize environments with distributions very different from those of the training environments, in a way that risk interpolation does not, by using negative weights on the training risks.

While convex combinations of PDFs yield mixture distributions, a non-convex combination will not generally be a distribution, itself. Affine combinations do maintain the constraint that their integral is equal to 1, but may also assign “negative probabilities” to some points. We reinterpret these negative probabilities as a positive probability of the same example occurring, but with the loss function negated. The equivalence is exact, and results from simultaneously negating $P(x, y)$ and $\ell(f_\theta(x), y)$ for the corresponding terms in the integral.

Since typical loss functions, such as mean-squared error and negative log-likelihood, can take on unboundedly high values, negating the loss function could hypothetically lead to unboundedly low loss. However, this is not a concern for the REx methods we propose, since this would also correspond to a larger gap in the training risks, which would be penalized. In particular, $\mathcal{R}_{\text{MM-REx}} \geq \mathcal{R}_{\text{ERM}}$, since the max in $\mathcal{R}_{\text{MM-REx}}$ could always set the $\lambda_i = 1/m$ to produce the ERM risk. Meanwhile, $\mathcal{R}_{\text{V-REx}}$ penalizes the difference of an individual training risk from the mean risk quadratically, outweighing the linear reduction in mean risk that assigning a negative weight to some examples allows.

Overall, the net effect of such negative probabilities is to prevent the model from making predictions that are overconfident (in classification) or extreme (in regression) on the basis of features that vary across environments, since doing so would lead to differences in the training risks that would be heavily penalized.

### 4. Experiments

We evaluate REx and compare with IRM on a range of tasks that require OoD generalization, and investigate the challenges of training these methods. REx performs comparably or better than IRM on a wide range of tasks, including 1) the PACS domain generalization image dataset, 2) continuous control tasks corrupted with spurious features, 3) predicting financial indicators, and 4) the tasks proposed in Arjovsky et al. (2019). The one exception is for “heteroskedastic”
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Figure 3. The probabilistic view of risk extrapolation, illustrated. **Top:** Input distributions $P(x)$ for environments $e_1$ and $e_2$. **Middle:** Interpolating between risk functions $R_1(x)$ and $R_2(x)$ corresponds to point-wise interpolations of $P_{e_1}(x)$ and $P_{e_2}(x)$. **Bottom:** Extrapolation can yield a distribution with zero, or even negative probability for examples from the left mode. These negative probabilities can be interpreted as flipping the sign of the loss function at those points.

tasks, in which the amount of intrinsic noise in the targets varies across environments. Equalizing risks across these environments does not make sense because the loss in environments with noisier labels will remain higher even for the optimal stable predictor, see Appendix for details.

4.1. Methodology for OoD

Out-of-distribution generalization (OoD) presents unique challenges for experimental design. In contrast to many commonly-used machine learning benchmarks, in the case of OoD, we cannot assume access to the test distribution. This makes it hard to tune hyperparameters in a principled way and care must be taken to not accidentally leak information about the test distribution into this process.

The main challenge is to identify a suitable validation distribution. By definition one should not have access to any distribution that is very close to the test distribution. On the other hand, it is also important to have access to a validation distribution that is sufficiently different from the training distribution, as it is otherwise impossible to properly tune for good OoD generalization.

To resolve the issue in this work, we clearly explain why and when we chose to tune on the test set (e.g. illustrative experiments). In all other cases we strive for good validation practices (e.g. tune on VLCS and apply to PACS).

| Method                  | train envs acc | test env acc |
|-------------------------|----------------|--------------|
| REx (V-REx) (ours)      | 71.5 ± 1.0     | 68.7 ± 0.9   |
| IRM                     | 70.8 ± 0.9     | 66.9 ± 2.5   |
| REx (MM-REx) (ours)     | 72.4 ± 1.8     | 66.1 ± 1.5   |
| RI                      | 88.9 ± 0.3     | 22.3 ± 4.6   |
| ERM                     | 87.4 ± 0.2     | 17.1 ± 0.6   |
| ERM, grayscale (oracle) | 73.5 ± 0.2     | 73.0 ± 0.4   |
| Theoretical Optimum     | 75             | 75           |
| Random Guessing         | 50             | 50           |

**Table 1.** Accuracy (percent) of different method on the Colored MNIST task. REx and IRM learn to discard spurious color feature and rely on stable shape feature. We use *strikethrough* to denote results achieved via tuning on the test set.

4.2. CMNIST

Arjovsky et al. (2019) construct a binary classification problem (with 0-4 and 5-9 each collapsed into a single class) based on the MNIST dataset, using color as a spurious feature. Specifically, digits are either colored red or green, and there is a strong correlation between color and label, which is reversed at test time. The goal is to learn the stable “digit shape” feature and ignore the unstable “digit color” feature. The learner has access to three environments:

1. A training environment where green digits have a 80% chance of belonging to class 1 (digits 5-9).
2. A training environment where green digits have a 90% chance of belonging to class 1.
3. A test environment where green digits have a 10% chance of belonging to class 1.

We use the exact same hyperparameters as Arjovsky et al. (2019), only replacing the IRMv1 penalty with MM-REx or V-REx penalty. These methods achieve similar performance, see Table 1.

We emphasize, however, that this task is methodologically unsuited for benchmarking, since, following Arjovsky et al. (2019) we assume access to the test distribution for hyperparameter tuning. Ultimately, these experiments should be interpreted only as a demonstration that, unlike ERM or RO, REX and IRM are both capable of OoD generalization in the face of spurious features, when properly tuned.

4.2.1. HOW TO LEARN STABLE FEATURES

In order to begin addressing this issue, we run a series of experiments aimed at understanding the sensitivity of IRM and REx to the choice of hyperparameters. In particular, we note that increasing the relative weight of the penalty term
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after 100 epochs of training (using a so-called “waterfall” schedule (Desjardins et al., 2015)) is critically important to performance on the colored MNIST task, see Figure 4(b). In light of this finding, we hypothesize that successful learning of stable features using REx or IRM should proceed in two stages. In the first stage, predictive features are learned. In the second stage, stable features are selected and/or predictive features are fine-tuned for stability. This viewpoint suggests that we could use overfitting on the training tasks as an indicator for when to apply (or increase) the IRM or REx penalty. This insight could provide a methodologically sound way of tuning the penalty term of REx or IRMv1 in the absence of a representative validation set.

In this section, we provide additional support for our hypothesis. In Figure 4, we demonstrate that the optimal point to apply the waterfall is after predictive features have been learned, but before the model starts to memorize training examples. Before predictive features are available, the penalty terms push the model to learn a constant predictor, impeding further learning. And after the model starts to memorize, it become difficult to distinguish spurious and stable features. This second effect is because neural networks often have the capacity to memorize all training examples given sufficient training time, achieving and near-0 loss (Zhang et al., 2016). In the limits of this memorization regime, the differences between losses become small, and gradients of the loss typically do as well, and so the REx and IRMv1 penalties no longer provide a strong or meaningful training signal, see Figure 5.

Figure 4. Stability penalties should be applied around when traditional overfitting begins, to ensure that the model has learned predictive features, and that penalties still give meaningful training signals. **Top:** Test accuracy as a function of epoch at which penalty term weight is increased (learning rate is simultaneously decreased proportionally). Choosing this hyperparameter correctly is essential for good performance. **Middle:** Generalization gap on a validation set with 85% correlation between color and label (the same as the average training correlation). The best test accuracy is achieved by increasing the penalty when the generalization gap begins to increase. The increase clearly indicates memorization because color and shape are only 85%/75% correlated with the label, and so cannot be used to make predictions with higher than 85% accuracy. **Bottom:** Accuracy on training/test sets, as well as an auxiliary grayscale set. Training/test performance reach 85%/15% after a few epochs of training, but grayscale performance improves, showing that meaningful features are still being learned.

Figure 5. Given sufficient training time, empirical risk minimization (ERM) minimizes both REx and IRMv1 penalty terms on Colored MNIST (without including either term in the loss function). This is because the model (a deep network) has sufficient capacity to fit the training sets almost perfectly. This prevents these penalties from having the intended effect, once the model has started to overfit. The y-axis is in log-scale.

4.3. Domain Generalization: VLCS and PACS

In this section we compare REx, IRM and ERM performance on the VLCS (Torralba & Efros, 2011) and PACS (Li et al., 2017) image datasets. Both datasets are commonly-used for multi-source domain generalization. The task is to train on three domains and generalize to a fourth one at test time. For VLCS, the domains are To avoid tuning our hyperparameters on the test distribution, we resort to tuning hyperparameters on VLCS and apply the best setting to PACS without any further tuning.

We use the exact same architecture, training procedure and data augmentation strategy as the state-of-the-art Jigsaw Puzzle approach (Carlucci et al., 2019) (except with IRM or V-REx instead of JigSaw as auxiliary loss) for all three
methods, experimental details as well as hyperparameter selection results for VLCS can be found in the Appendix. As runs are very noisy, we ran each experiment 10 times, and report average test accuracies extracted at the time of the highest validation accuracy on each run.

On PACS we find that REx outperforms IRM and IRM outperforms ERM on average, while all are worse than the state-of-the-art Jigsaw method.

4.4. Reinforcement Learning with Spurious Correlations

We take tasks from the Deepmind Control Suite (Tassa et al., 2018) and modify the original state, s, to include noise and spurious features. Specifically, a scaled copy of 1 or 2 of the dimensions of s is concatenated with s, and then \( N(0, 0.01) \) noise is added to the corresponding dimensions of the original state vector, to form a new representation of the state, \( \hat{s} \). The magnitude of the scaling factor differs across environments, with \( n = 1 \) and \( n = 2 \) for the two training environments. The agent takes \( \hat{s} \) as input and learns a single representation that is shared across training environments. The representation is used by the Soft Actor-Critic (Haarnoja et al., 2018) policy learning algorithm, as well as an auxiliary reward predictor, which is trained to predict the next 3 rewards conditioned on the next 3 actions. The representation is trained using gradients from the reward predictor, as well as the critic. The entire model is evaluated on the unseen test environment, which uses a larger scaling factor \( n = 3 \). Like for CMNIST, since the spurious features are copied from the state before the noise is added, they are more informative for the reward prediction task. In Figure 6, we see that REx outperforms both IRM and ERM in unseen evaluation environments. We use cartpole\_swingup as a development set for tuning hyperparameters, and perform a single evaluation with 10 random seeds on each of finger\_spin and walker\_walk. We tune hyperparameters on cartpole\_swingup and apply to finger\_spin and walker\_walk. Details and results with other scaling factors in Appendix.

Figure 6. Performance on walker\_walk (left) and finger\_spin (right), with 10 seeds and 1 standard error shaded.

Figure 7. Financial indicators tasks. The left panel indicates the set of training environments; the middle and right panels show the test accuracy on the respective environments relative to ERM (a black dot corresponds to a training environment; a colored patch indicates the test accuracy on the respective environment.)

4.5. Financial indicators

To compare the types of solutions found by IRM and REx, we apply our method to a prediction task in the finance domain, where countless confounding and unknown factors make prediction based on a small number of features hard-to-impossible. The dataset is split into five years, 2014–18, containing 37 publicly reported financial indicators of several thousand publicly listed companies each. The task is to predict if a company’s value will increase or decrease in the following year (see Appendix for dataset details.) We consider each year a different environment, and create 20 different tasks by selecting all possible combinations of environments where three environments represent the training sets, one environment the validation set, and another one the test set. We train an MLP using the validation set to determine an early stopping point, with \( \beta = 10^4 \). The per-task results summarized in fig. 7 indicate substantial differences between ERM and IRM, and ERM and REx. The predictions produced by IRM and REx, however, only differ insignificantly, highlighting the similarity of IRM and REx. While performance on specific tasks differs significantly between ERM and IRM/REx, performance averaged over tasks is not significantly different, see Appendix.

5. Related Work

Out-of-distribution generalization is typically tackled by pre-specified learning invariances that aid generalization far outside of the training distribution. There are many approaches to OoD generalization; one is robust learning in the face of worst-case perturbations, another one is causal discovery, which aims to determine an underlying causal graph, and therefore achieve extrapolation. Other approaches are broadly termed domain generalization and often incorporate prior knowledge about the overall problem structure into the
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| PACS   | Art Painting | Cartoon | Sketch | Photo | Average |
|--------|--------------|---------|--------|-------|---------|
| REx (ours) | 67.04       | 67.97   | 59.81  | 89.74 | 71.14   |
| IRM    | 67.05       | 68.49   | 57.81  | 89.39 | 70.69   |
| ERM    | 66.22       | 67.59   | 57.90  | 89.69 | 70.35   |
| Jigsaw (SOTA) | 67.43   | 69.49   | 62.74  | 89.64 | 72.32   |

Table 2. Accuracy (percent) of different methods on the PACS task. Results are test accuracy at the time of the highest validation accuracy, averaged over 10 runs. REx outperforms ERM and IRM on average, but performs worse than the state-of-the-art.

training procedure. The assumptions made and approaches proposed can be quite different in these various settings.

One direction in domain generalization is to explicitly project out superficial domain-specific statistics (Wang et al., 2019) to reduce sensitivity to the domain. Another approach increases sensitivity to shape features of natural images like global spatial coherence of objects (Carlucci et al., 2019) and therefore implicitly reduces reliance on local and superficial cues. Other approaches focus on adversarially learning representations that are invariant with respect to domain-specific features (Tzeng et al., 2017; Long et al., 2018; Li et al., 2018; Albuquerque et al., 2019). Rothenhausler et al. (2018) note that OoD generalization can be viewed as the solution to a minimax of the worst case risk under different distribution shifts.

Recently, invariance as a proxy for causal inference has been proposed as a way of learning a causal graph. Peters et al. (2016) propose a method called Invariant Causal Prediction that utilizes the invariance property of causal models (Schölkopf et al., 2012; Pearl, 2009) to obtain a feature set that is a causal predictor of a target. Arjovsky et al. (2019) propose an extension of that work, Invariant Risk Minimization (IRM), with the goal of learning a data representation that does not rely on spurious correlations.

Others in causal inference propose disentanglement as a way to uncover and separate causal variables from spurious correlations. Heinze-Deml & Meinshausen (2017) assume “grouping” knowledge, that one has access to the same object under varying conditions, in order to disentangle what features are invariant conditional on a target variable, and what are simply “style”. Similarly, Gowal et al. (2019) attempt to find a representation in which “style” (i.e. spurious) and “semantic” (i.e. stable) features are disentangled. This allows them to manipulate the style of training images as a form of adversarial training (Madry et al., 2017) with semantic perturbations.

6. Discussion

While the high level goals of IRM and REx are similar, IRM has a much more developed mathematical foundation. On the other hand, REx has simple geometric and probabilistic interpretations, which may be more intuitive.

Empirically, REx and IRM perform on par in many settings. However, on the PACS domain generalization benchmark REx slightly outperforms IRM and in our reinforcement learning experiments we observe a clear advantage of REx over IRM. Despite the encouraging empirical performance, REx has a shortcoming when compared to IRM. It may not succeed across environments with different levels of intrinsic label noise, whereas IRM can deal with such fluctuations (see Appendix).

Thus, a key unresolved challenge for REx is how to fairly compare risk functions across environments that contain different levels of innate noise, since it may not make sense to enforce exact equality of risks in that case. The use of “baselines” designed to account for these differences exists in the literature (Meinshausen et al., 2015), but the baseline proposed by Meinshausen et al. (2015), $Var(Y)$, only provides an upper bound on the true noise-level.

7. Conclusion

We have introduced risk extrapolation as a new method to generalize outside of the training distribution. Our method is a simple generalization of the idea underlying robust optimization, inspired by Invariant Risk Minimization, and their goal of learning stable features. We have shown that maximizing the overall risk over a specified set of affine combinations of the training risks means that all training risks should be made as similar as possible. This can be implemented efficiently with a variance penalty on the environment risks. We show that our method performs competitively on a range of out-of-distribution and domain generalization tasks.

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A. Overview

Our appendix contains details and further results from the experiments reported in the paper in Section B. In Section C, we provide results on the synthetic structural equation models from Arjovsky et al. (2019). We also discuss an observed limitation of REx and prospects for addressing this issue.

We include some simple mathematical derivations of results in the paper in Section D. Finally, in Section E, we define interpolation and extrapolation, and argue for the importance of extrapolation in machine learning.

B. Experiments from the paper

B.1. Domain Generalization

Here we provide details and further results on the experiments in Section 4.3. For our domain generalization experiments we closely follow the experimental setup in the Jigsaw paper (Carlucci et al., 2019) as it is listed as the method with highest average accuracy on the PACS leaderboard. Since every domain in PACS is used as a test set when training on the other three domains, it is not possible to perform a methodologically sound evaluation on PACS after examining results on any of the data. Thus to avoid performing any tuning on test distributions, we use VLCS to tune hyperparameters and then apply these exact same settings to PACS and report the final average over 10 runs on each domain.

We use all hyperparameters from the original Jigsaw codebase. We use Imagenet pre-trained AlexNet features and chose batch-size, learning rate, as well as penalty weights based on performance on the VLCS dataset where test performance on the holdout domain was used for the set of parameters producing the highest validation accuracy. The best performing parameters on VLCS were then applied to the PACS dataset without further changes. We searched over batch-sizes in {128, 384}, over penalty strengths in {0.0001, 0.001, 0.01, 0.1, 1, 10}, learning rates in {0.001, 0.01} and used average performance over all 4 VLCS domains to pick the best performing hyperparameters. Table 3 shows results on VLCS with the best performing hyperparameters.

The final parameters for all methods on PACS were a batch size of 384 with 30 epochs of training with Adam, using a learning rate of 0.001, and multiplying it by 0.1 after 24 epochs (this step schedule was taken from the Jigsaw repo). The penalty weight chosen for Jigsaw was 0.9; for IRM and REx it was 0.1. We used the same data-augmentation pipeline as the original Jigsaw code for ERM, IRM, Jigsaw and REx to allow for a fair comparison. In Table 4, we include results with standard errors, which we’d previously omitted in the main paper; these experiments were performed independently from the results in the main paper.

| VLCS    | CALTECH | SUN   | PASCAL | LABELME | Average |
|---------|---------|-------|--------|---------|---------|
| REx (ours) | 96.72   | 63.68 | 72.41  | 60.40  | 73.30   |
| IRM     | 95.99   | 62.85 | 71.71  | 59.61  | 72.84   |
| ERM     | 94.76   | 61.92 | 69.03  | 60.55  | 71.56   |
| Jigsaw (SOTA) | 96.46 | 63.84 | 70.49  | 60.06  | 72.74   |

Table 3. Accuracy (percent) of different methods on the VLCS task. Results are test accuracy at the time of the highest validation accuracy, averaged over 10 runs. On VLCS REx outperforms all other methods. Numbers are shown in strike-through because we selected our hyperparameters based on highest test set performance; the goal of this experiment was to find suitable hyperparameters for the PACS experiment.

B.2. Reinforcement Learning Experiments

Here we provide details and further results on the experiments in Section 4.4. The hyperparameters used for training Soft Actor-Critic can be found in Table 5. We used cartpole_swingup as a development task to tune the hyperparameters of penalty weight (chosen from [0.01, 0.1, 1, 10]) and number of iterations before the penalty is turned up (chosen from [5000, 10000, 20000]), both for REx and IRM. The plots with the hyperparameter sweep are in Figure 8.

1https://domaingeneralization.github.io/
2https://github.com/fmcarlucci/JigenDG
Table 4. Accuracy (percent) of different methods on the PACS task. Results are test accuracy at the time of the highest validation accuracy, averaged over 10 runs. REx outperforms ERM on average, and performs similar to IRM and Jigsaw (the state-of-the-art).

B.3. Financial Indicators

Here we provide details and further results on the experiments in Section 4.5. We use v1 of the dataset published on \(^3\) and prepare the data as described in.\(^4\) We further remove all the variables that are not shared across all 5 years, leaving us with 37 features, and whiten the data through centering and normalizing by the standard deviation.

On each subtask, we train an MLP with two hidden layers of size 128 with tanh activations and dropout (p=0.5) after each layer. We optimize the binary cross-entropy loss using Adam (learning rate 0.001, \(\beta_1 = 0.9, \beta_2 = 0.999, \epsilon = 10^{-8}\)), and an L2 penalty (weight 0.001). In the IRM/REx experiments, the respective penalty is added to the loss (\(\beta = 1\)) and the original loss is scaled by a factor \(10^{-4}\) after 1000 iterations. Experiments are run for a maximum of 9000 training iterations with early stopping based on the validation performance. All results are averaged over 3 trials. The overall performance of the different models, averaged over all tasks, is summarized in Tab. 6. The difference in average performance between ERM, IRM, and REx is not statistically significant, as the error bars are very large.

C. Structural equation model (SEM) experiments and the issue of baselines

Here we present experiments on the structural equation model tasks introduced by Arjovsky et al. (2019). Structural equation models (SEMs) are a way of specifying causal relationships between variables. The left hand side of an SEM is expressed as a deterministic function of the right hand side which includes the values of other variables, including i.i.d. random noise variables. Arjovsky et al. (2019) construct several varieties of SEM where the task is to predict targets \(Y\) from inputs
Table 5. A complete overview of hyperparameters used for reinforcement learning experiments.

| Parameter name                        | Value    |
|---------------------------------------|----------|
| Replay buffer capacity                | 1000000  |
| Batch size                            | 1024     |
| Discount $\gamma$                     | 0.99     |
| Optimizer                             | Adam     |
| Critic learning rate                  | $10^{-5}$|
| Critic target update frequency        | 2        |
| Critic Q-function soft-update rate $\tau_Q$ | 0.005   |
| Critic encoder soft-update rate $\tau_{enc}$ | 0.005   |
| Actor learning rate                   | $10^{-5}$|
| Actor update frequency                | 2        |
| Actor log stddev bounds               | $[-5, 2]$|
| Encoder learning rate                 | $10^{-5}$|
| Decoder learning rate                 | $10^{-5}$|
| Decoder weight decay                  | $10^{-7}$|
| L1 regularization weight              | $10^{-5}$|
| Temperature learning rate             | $10^{-4}$|
| Temperature Adam’s $\beta_1$         | 0.9      |
| Init temperature                      | 0.1      |

Table 6. Test accuracy of models trained on the financial domain dataset, averaged over all 20 tasks, as well as min./max. accuracy across the tasks.

| Overall accuracy | Min acc. | Max acc. |
|------------------|----------|----------|
| ERM              | 54.6 ± 4.6 | 47.6     | 66.2     |
| IRM              | 55.3 ± 5.9 | 45.9     | 67.5     |
| REx              | **55.5 ± 6.0** | 47.2     | **68.0** |

$X_1, X_2$, where $X_1$ are (non-spurious) causes of $Y$, and $X_2$ are (spurious) effects of $Y$. We refer the reader to Section 5.1 and Figure 3 of Arjovsky et al. (2019) for more details. We use the same experimental settings as Arjovsky et al. (2019) (except we only run 7 trials), and report results in Table 7.

These experiments include several variants of a simple SEM, given by:

$$
X_1 = N_1 \\
Y = W_{1 \rightarrow Y} X_1 + N_Y \\
X_2 = W_{Y \rightarrow 2} Y + N_2
$$

Where $N_1, N_Y, N_2$ are all sampled i.i.d. from normal distributions. The variance of these distributions may vary across environments.

While REx achieves good performance in the “homoskedastic” case, it performs poorly in the so called “heteroskedastic” case, where the amount of intrinsic noise, $\sigma^2_Y$ in the target changes across environments.\(^5\) Intuitively, this is because the irreducible error varies across environments in these tasks, meaning that the risk will be larger on some environments than others, even when the model learns the optimal stable predictor. We tried using a “baseline” (see Eqn. 3) of $r_e = Var(Y_e)$ (Meinshausen et al., 2015) to account for the different noise levels in $Y$, but this also did not work.

\(^5\)At the time of writing the current version of Arjovsky et al. (2019) has the names “heteroskedastic” and “homoskedastic” backwards; their code uses the correct names, however.
Average mean-squared error between true and estimated weights on causal ($\hat{Y}$) and non-causal ($X_2$) variables. **Top:** When the level of noise in the spurious features varies across environments, REx performs well (FOU, FOS, POU, POS). **Bottom:** When the level of noise in the targets varies instead, REx performs poorly (FEU, FES, PEU, PES). Using the baselines $r_e = \mathbb{V}(Y)$ does not solve the problem, and indeed, hurts performance on the homoskedastic environments.

Table 7.

|                     | FOU(c) | FOU(nc) | FOS(c) | FOS(nc) | POU(c) | POU(nc) | POS(c) | POS(nc) |
|---------------------|--------|---------|--------|---------|--------|---------|--------|---------|
| IRM                 | 0.001±0.000 | 0.001±0.000 | 0.001±0.000 | 0.001±0.000 | 0.004±0.001 | 0.006±0.003 | 0.002±0.000 | 0.000±0.000 |
| REx, $r_e = 0$      | 0.001±0.000 | 0.008±0.002 | 0.007±0.002 | 0.000±0.000 | 0.004±0.001 | 0.004±0.001 | 0.002±0.000 | 0.000±0.000 |
| REx, $r_e = \mathbb{V}(Y)$ | 0.816±0.149 | 1.417±0.442 | 0.919±0.091 | 0.000±0.000 | 0.915±0.055 | 1.113±0.085 | 0.937±0.090 | 0.000±0.000 |

We include a mathematical analysis of the simple SEM given above in order to better understand why REx succeeds in the homoskedastic, but not the heteroskedastic case. Assuming that $Y, X_1, X_2$ are scalars, this SEM becomes

$$
X_1 = N_1 \\
Y = w_{1 \rightarrow Y} N_1 + N_Y \\
X_2 = w_{y \rightarrow 2} w_{1 \rightarrow y} N_1 + w_{y \rightarrow 2} N_Y + N_2
$$

We consider learning a model $\hat{Y} = \alpha X_1 + \beta X_2$. Then the residual is:

$$
\hat{Y} - Y = (\alpha + w_{1 \rightarrow y} (\beta w_{y \rightarrow 2} - 1)) N_1 + (\beta w_{y \rightarrow 2} - 1) N_Y + \beta N_2
$$

Since all random variables have zero mean, the MSE loss is the variance of the residual. Using the fact that the noise $N_1, N_Y, N_2$ are independent, this equals:

$$
\mathbb{E}[(\hat{Y} - Y)^2] = (\alpha + w_{1 \rightarrow y} (\beta w_{y \rightarrow 2} - 1))^2 \sigma_1^2 + (\beta w_{y \rightarrow 2} - 1)^2 \sigma_Y^2 + \beta^2 \sigma_2^2
$$

Thus when (only) $\sigma_2$ changes, the only way to keep the loss unchanged is to set the coefficient in front of $\sigma_2$ to 0, meaning $\beta = 0$. By minimizing the loss, we then recover $\alpha = w_{1 \rightarrow y}$, i.e. in the homoskedastic setting, the loss equality constraint of REx yields the causal model. On the other hand, if (only) $\sigma_Y$ changes, then REx enforces $\beta = 1/w_{y \rightarrow 2}$, which then induces $\alpha = 0$, recovering the anticausal model.

The relationship between causal features and targets is expected to be more stable, since changing the noise in causal features will also lead to a change in the targets. While REx is able to account for this type of stability, there could also be changes in the environment that change the distribution of $Y$ without directly influencing $X$, causing issues for REx. In contrast, IRM can handle such changes. But IRM also is not designed match the true distribution $P(Y|X)$ on test environments, only its expectation. It is not clear whether IRM can be generalized to cases where one cares about having an accurate predictive distribution at test time, and not just an accurate prediction. On the other hand, it is plausible that an appropriately designed variant of REx may be suited to such a task, but we leave further investigation for future work.

D. Mathematical derivations

D.1. Closed form solutions to risk interpolation and minimax-REx

Here, we show that risk interpolation (Eqn. 6) is equivalent to the robust optimization objective of Eqn. 3. Without loss of generality, let $R_1$ be the largest risk, so $R_c \leq R_1$, for all $c$. Thus we can express $R_c = R_1 - d_c$ for some non-negative $d_c$, with $d_1 = 0 \geq d_c$ for all $c$. And thus we can write the weighted sum of Eqn. 6 as:

$$
R_{R_1}(\theta) = \max_{\lambda_1, \lambda_c \geq 0} \sum_{\lambda_c = 1} \lambda_c R_c(\theta) \\
= \max_{\lambda_1, \lambda_c \geq 0} \sum_{\lambda_c = 1} \lambda_c (R_1(\theta) - d_c) \\
= R_1(\theta) + \max_{\lambda_1, \lambda_c \geq 0} \sum_{\lambda_c = 1} -\lambda_c (d_c)
$$

(10)

(11)

(12)

(13)
We wish to show that this is equal to \( R \). And this holds whenever \( \beta \).

The V-REx penalty (Eqn. 9) is equivalent to the average pairwise mean squared error between all training risks (up to a constant factor of 2). Recalling that \( R \) denotes the risk on environment \( i \). We have:

\[
\frac{1}{2n^2} \sum_{i=1}^{m} \sum_{j=1}^{m} (R_i - R_j)^2 = \frac{1}{2n^2} \sum_{i} \sum_{j} (R_i^2 + R_j^2 - 2R_iR_j)
\]

\[
= \frac{1}{2n} \sum_{i} R_i^2 + \frac{1}{2n} \sum_{j} R_j^2 - \frac{1}{n^2} \sum_{i} \sum_{j} R_iR_j
\]

\[
= \frac{1}{n} \sum_{i} R_i^2 - \left( \frac{1}{n} \sum_{i} R_i \right)^2
\]

\[
= \text{Var}(\{R_1, ..., R_m\})
\]

D.2. Minimax-REx and Mean absolute error REx

In the case of only two training risks, MM-REx is equivalent to using a penalty on the mean absolute error (MAE) between training risks. However, penalizing the pairwise absolute errors is not equivalent when there are \( m > 2 \) training risks, as we show below. Without loss of generality, assume that \( R_1 < R_2 < ... < R_m \). Then the \( R_{\text{MAE}} \) penalty term is:

\[
2 \sum_i \sum_{j \leq i} (R_i - R_j) = mR_m - \sum_{j \leq m} R_j + (m - 1)R_{m-1} - \sum_{j \leq m-1} R_j \ldots
\]

\[
= 2 \sum_i iR_i - \sum_i \sum_{j \leq i} R_j
\]

\[
= 2 \sum_i iR_i - (m - i + 1)R_i
\]

\[
= 2 \sum_i (2i - m - 1)R_i
\]

Now, for \( m = 2 \), we have \( R_{\text{MAE}} = 2(1 + 2 \beta_{\text{MAE}})R_2 + (1 - 2 \beta_{\text{MAE}})R_1 \). Now, adding this penalty term with some coefficient \( \beta_{\text{MAE}} \) to the ERM term yields:

\[
R_{\text{MAE}} = R_1 + R_2 + 2\beta_{\text{MAE}}(R_2 - R_1) = (1 + 2\beta_{\text{MAE}})R_2 + (1 - 2\beta_{\text{MAE}})R_1
\]

We wish to show that this is equal to \( R_{\text{MM}} \) for an appropriate choice of learning rate \( \gamma_{\text{MAE}} \) and hyperparameter \( \beta_{\text{MAE}} \). Still assuming that \( R_1 < R_2 \), we have that:

\[
R_{\text{MM}} = (1 + \beta_{\text{MM}})R_2 - \beta_{\text{MM}}R_1
\]

Choosing \( \gamma_{\text{MAE}} = 1/2\gamma_{\text{MM}} \) is equivalent to multiplying \( R_{\text{MM}} \) by 2, yielding:

\[
2R_{\text{MM}} = 2(1 + \beta_{\text{MM}})R_2 - 2\beta_{\text{MM}}R_1
\]

Now, in order for \( R_{\text{MAE}} = 2R_{\text{MM}} \), we need that:

\[
2 + 2\beta_{\text{MM}} = 1 + 2\beta_{\text{MAE}}
\]

\[
-2\beta_{\text{MM}} = 1 - 2\beta_{\text{MAE}}
\]

And this holds whenever \( \beta_{\text{MAE}} = \beta_{\text{MM}} + 1/2 \). When \( m > 2 \), however, these are not equivalent; this can be determined by noting that \( R_{\text{MM}} \) puts equal weight on all but the highest training risk, whereas \( R_{\text{MAE}} \) assigns a different weight to each risk.

D.3. Penalizing pairwise mean squared error (MSE) yields V-REx

The V-REx penalty (Eqn. 9) is equivalent to the average pairwise mean squared error between all training risks (up to a constant factor of 2). Recall that \( R_i \) denotes the risk on environment \( i \). We have:

\[
\frac{1}{2n^2} \sum_{i=1}^{m} \sum_{j=1}^{m} (R_i - R_j)^2 = \frac{1}{2n^2} \sum_{i} \sum_{j} (R_i^2 + R_j^2 - 2R_iR_j)
\]

\[
= \frac{1}{2n} \sum_{i} R_i^2 + \frac{1}{2n} \sum_{j} R_j^2 - \frac{1}{n^2} \sum_{i} \sum_{j} R_iR_j
\]

\[
= \frac{1}{n} \sum_{i} R_i^2 - \left( \frac{1}{n} \sum_{i} R_i \right)^2
\]

\[
= \text{Var}(\{R_1, ..., R_m\})
\]
E. Extrapolation vs. Interpolation

We define interpolation and extrapolation as follows: **interpolation** refers to making decisions about points within the convex hull of the training examples and **extrapolation** refers to making decisions about points outside their convex hull.\(^6\) This generalizes the familiar sense of these terms for one-dimensional functions. An interesting consequence of this definition is: for data of high intrinsic dimension, generalization requires extrapolation (Hastie et al., 2009), even in the i.i.d. setting. This is because the volume of high-dimensional manifolds concentrates near their boundary; see Figure 9.

E.1. Extrapolation in the space of risk functions

The same geometric considerations apply to extrapolating to new environments, or even tasks. Environments and tasks can be highly diverse, varying according to high dimensional attributes, and thus requiring extrapolation to generalize across. For instance, in medical applications, we could view individual patients as different environments. This suggests that risk extrapolation might provide benefits in many types of out-of-distribution problems, regardless of the presence of spurious features, and maybe even multi-task learning as well.

![Illustration of the importance of extrapolation for generalizing in high dimensional space. In high dimensional spaces, mass concentrates near the boundary of objects. For instance, the uniform distribution over a ball in N + 1-dimensional space can be approximated by the uniform distribution over the N-dimensional hypersphere. We illustrate this in 2 dimensions, using the 1-sphere (i.e. the unit circle). Dots represent a finite training sample, and the shaded region represents the convex hull of all but one member of the sample. Even in 2 dimensions, we can see why any point from a finite sample from such a distribution remains outside the convex hull of the other samples, with probability 1. The only exception would be if two points in the sample coincide exactly.](image)

\(^6\) Surprisingly, we were not able to find any existing definition of these terms in the machine learning literature. They have been used in this sense (Hastie et al., 2009; Haffner, 2002), but also to refer to strong generalization capabilities more generally (Sahoo et al., 2018).