DISTRIBUTIONALLY ROBUST NEURAL NETWORKS FOR GROUP SHIFTS: ON THE IMPORTANCE OF REGULARIZATION FOR WORST-CASE GENERALIZATION

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ABSTRACT

Overparameterized neural networks can be highly accurate on average on an i.i.d. test set yet consistently fail on atypical groups of the data (e.g., by learning spurious correlations that hold on average but not in such groups). Distributionally robust optimization (DRO) allows us to learn models that instead minimize the worst-case training loss over a set of pre-defined groups. However, we find that naively applying group DRO to overparameterized neural networks fails: these models can perfectly fit the training data, and any model with vanishing average training loss also already has vanishing worst-case training loss. Instead, their poor worst-case performance arises from poor generalization on some groups. By coupling group DRO models with increased regularization—stronger-than-typical \( \ell_2 \) regularization or early stopping—we achieve substantially higher worst-group accuracies, with 10-40 percentage point improvements on a natural language inference task and two image tasks, while maintaining high average accuracies. Our results suggest that regularization is critical for worst-group generalization in the overparameterized regime, even if it is not needed for average generalization. Finally, we introduce and give convergence guarantees for a stochastic optimizer for the group DRO setting, underpinning the empirical study above.

1 INTRODUCTION

Machine learning models are typically trained to maximize average accuracy on a test set that is independent and identically distributed (i.i.d.) to the training set. However, even models that perform well on average can consistently fail on rare and atypical examples (Hovy & Sgaard, 2015; Blodgett et al., 2016; Tatman, 2017; Hashimoto et al., 2018; Duchi et al., 2019). Such models are problematic when they violate equity considerations (Jurgens et al., 2017; Buolamwini & Gebru, 2018) or rely on spurious correlations: misleading heuristics that work for most training examples but do not hold on different test distributions. For example, in natural language inference (NLI)—determining if two sentences agree or contradict—negation words like ‘never’ are strongly correlated with contradiction due to artifacts in crowdsourced training data (Gururangan et al., 2018). A model could achieve high average accuracy on an i.i.d. test set by learning this spurious correlation, but perform poorly on test sets where the correlation does not hold (McCoy et al., 2019).

In this paper, we show that in the modern regime of training overparameterized models—i.e., large neural networks that can perfectly fit the training data—this problem is one of generalization: these models can generalize well on average, as in Zhang et al. (2017), but do not generalize well on some (sub)groups of data (e.g., the group of contradictions with no negations in the NLI example above).

To avoid the pitfalls of optimizing for average loss, we instead optimize for the worst-case loss over various groups in the data, using prior knowledge of spurious associations (e.g., between negation

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and contradiction) to choose these groups. This is an instance of distributionally robust optimization (DRO), which studies worst-case performance over potential test distributions (Ben-Tal et al., 2013; Duchi et al., 2016). In this group DRO setting, the data is distributed as a mixture of groups with mixture weights that can vary at test time (Hu et al., 2018; Oren et al., 2019). Optimizing for the worst-case group shift—i.e., shift in group mixture weights at test time—is equivalent to minimizing the worst-case loss over each group.

Prior work has applied DRO as a data-dependent regularizer (Maurer & Pontil, 2009; Shafieezadeh-Abadeh et al., 2015; Namkoong & Duchi, 2017) and to learn models that are robust over different data subsets (Duchi & Namkoong, 2018; Hashimoto et al., 2018). However, a major challenge with DRO in the overparameterized regime is that models can achieve vanishing training loss (Zhang et al., 2017); any such model would be (near) optimal for both the average and worst-case training losses, so there is little incentive for their minimizers to differ (Wen et al., 2014). Indeed, existing work on DRO has focused on either generative models or convex predictive models with limited capacities, both of which cannot reach vanishing training loss.

In this paper, we study group DRO in the context of large, high-performing neural networks in three applications (Figure 1)—natural language inference with the MultiNLI dataset (Williams et al., 2018), facial attribute recognition with CelebA (Liu et al., 2015), and bird photograph recognition with our modified version of the CUB dataset (Wah et al., 2011). In the vanishing-training-loss regime, we find that group DRO models do no better than ERM models: both models achieve nearly perfect training accuracies across groups and continue to perform well on average at test time, but suffer low test accuracies on the worst-case group (Section 3.1). In other words, the generalization gap is small on average, but the worst-case generalization gap over groups is large.

However, group DRO models can significantly outperform ERM models when we account for these generalization gaps—through appropriate regularization, such as strong $l_2$ regularization or early stopping (Section 3.2), and through group adjustments that explicitly handle the differences in generalization gaps between groups (Section 3.3). Across the three applications, group DRO improves worst-case test accuracies by 10 to 40 percentage points while maintaining high average test accuracies. These results give a new perspective on generalization in neural networks: regularization might not be important for good average performance (e.g., models can “train longer and generalize better” on average (Hoffer et al., 2017)) but it appears critical for good worst-case performance.

Finally, we introduce a stochastic optimizer for group DRO that underpins the experiments above and scales to large models and datasets. We derive convergence guarantees for our algorithm in the convex case and empirically show that it behaves well in our non-convex models (Section 5).

2 Setup

Consider predicting labels $y \in \mathcal{Y}$ from input features $x \in \mathcal{X}$. Given a model family $\Theta$, loss $\ell : \Theta \times (\mathcal{X} \times \mathcal{Y}) \rightarrow \mathbb{R}_+$, and training data drawn from some distribution $P$, the standard goal is to find a model $\theta \in \Theta$ that minimizes the expected loss $\mathbb{E}_P[\ell(\theta; (x, y))]$ under the same distribution $P$. 
The uncertainty set \(Q\) optimizes for implausible worst-case scenarios \((\text{Duchi et al., 2019})\). In other words, without any \(\hat{P}\) where \(\hat{P}\) is the empirical distribution over the training data.

In distributionally robust optimization (DRO) \((\text{Ben-Tal et al., 2013} \text{ Duch et al., 2016})\), we aim instead to minimize the worst-case loss over an uncertainty set of distributions \(Q\). \[\min_{\theta \in \Theta} \left\{ \mathcal{R}(\theta) := \sup_{Q \in Q} \mathbb{E}_{(x,y) \sim P} [\ell(\theta; (x,y))] \right\}. \] (2)

The uncertainty set \(Q\) encodes the possible test distributions that we want our model to perform well on. Choosing a general family \(Q\), such as a divergence ball around the training distribution \(P\), confers robustness to a wide set of distributional shifts but can be too pessimistic because it optimizes for implausible worst-case scenarios \((\text{Duchi et al., 2019})\). In other words, without any prior knowledge or structural assumptions on the distributional shifts, it is difficult to specify an uncertainty set \(Q\) that is not too conservative but can still capture realistic shifts.

We thus consider the group DRO setting \((\text{Hu et al., 2018} \text{ Oren et al., 2019})\) where the training distribution \(P\) is assumed to be a mixture of \(m\) groups \(P_g\) indexed by \(G = \{1, 2, \ldots, m\}\). We use the term group to refer to a sub-distribution of either the population or empirical training distribution \((P\) or \(\hat{P})\); in the latter, each group \(\hat{P}_g\) corresponds to a subset of training examples drawn from \(P_g\).\footnote{In our main experiments, \(m = 4\) or \(6\); we also use \(m = 64\) in our supplemental experiments.}

Further, we assume the mixture weights can differ arbitrarily at train and test time, i.e., \(Q := \{\sum_{g=1}^{m} \omega_g \hat{P}_g : \omega \in \Delta_m\}\), where \(\Delta_m\) is the \((m-1)\)-dimensional probability simplex;\footnote{In our main experiments, \(m = 4\) or \(6\); we also use \(m = 64\) in our supplemental experiments.} this choice of \(Q\) allows us to learn models that are robust to group shifts. Because the optimum of a linear program is attained at a vertex, the worst-case risk over this choice of \(Q\) is equivalent to a maximum over the expected loss of each group,

\[\mathcal{R}(\theta) = \max_{g \in G} \mathbb{E}_{(x,y) \sim P_g} [\ell(\theta; (x,y))]. \] (3)

We assume that we know which group each training point comes from—i.e., the training data comprises \((x, y, g)\) triplets—though we do not assume we observe \(g\) at test time, so the model cannot use \(g\) directly. Instead, we use the training data to learn the group DRO model that minimizes the empirical robust risk \(\hat{R}(\theta)\):

\[\hat{\theta}_{\text{DRO}} := \arg \min_{\theta \in \Theta} \left\{ \hat{R}(\theta) := \max_{g \in G} \mathbb{E}_{(x,y) \sim \hat{P}_g} [\ell(\theta; (x,y))] \right\}. \] (4)

where \(\hat{P}_g\) is the empirical distribution over all training points \((x, y, g')\) with \(g' = g\). Group DRO learns models with good robust training loss across groups. This need not imply good robust test loss because of the generalization gap \(\delta := \mathcal{R}(\hat{\theta}_{\text{DRO}}) - \mathcal{R}(\hat{\theta})\). We will show that for overparameterized neural networks, \(\delta\) is large unless we apply sufficient regularization.

### 2.1 Applications

In the rest of this paper, we study three applications that share a similar structure (Figure 1): each data point \((x, y)\) has some input attribute \(a(x) \in A\) that is spuriously correlated with the label \(y\), and we use this prior knowledge to form \(m = |A| \times |\mathcal{Y}|\) groups, one for each value of \((a, y)\). We expect that models that learn the correlation between \(a\) and \(y\) in the training data would do poorly on groups for which \(a \neq y\), and hence do worse on the robust loss \(\mathcal{R}(\theta)\).

**Object recognition with correlated backgrounds (Waterbirds dataset).** Object recognition models can learn to make predictions from image backgrounds instead of the actual object \((\text{Ribeiro et al., 2016})\). We investigate this by constructing a new dataset, Waterbirds, which combines bird photographs from the Caltech-UCSD Birds-200-2011 (CUB) dataset \((\text{Wah et al., 2011})\) with image backgrounds from the Places dataset \((\text{Zhou et al., 2017})\). We label each bird as one of \(\mathcal{Y} = \{\text{waterbird, landbird}\}\) and place it on one of \(A = \{\text{water background, land background}\}\), with waterbirds (landbirds) more frequently appearing against a water (land) background (Appendix C.1). There are \(n = 4795\) training examples and 56 in the smallest group (waterbirds against land).
Object recognition with correlated demographics (CelebA dataset). Object recognition models (and other ML models more generally) have also been shown to learn spurious associations between the target label and demographic information like gender and ethnicity (Buolamwini & Gebru, 2018). We examine this on the CelebA celebrity face dataset (Liu et al., 2015), using hair color ($Y = \{\text{blond, dark}\}$) as the target and gender ($A = \{\text{male, female}\}$) as the spurious attribute. There are $n = 162770$ training examples, with 1387 in the smallest group (blond-haired males).

Natural language inference (MultiNLI dataset). In natural language inference, the task is to determine if a given hypothesis is entailed by, neutral with, or contradicts a given premise. Prior work has shown that crowdsourced training datasets for this task have significant annotation artifacts, such as the spurious correlation between contradictions and the presence of the negation words nobody, no, never, and nothing (Gururangan et al., 2018). We divide the MultiNLI dataset (Williams et al., 2018) into $m = 6$ groups, one for each pair of labels $Y = \{\text{entailed, neutral, contradictory}\}$ and spurious attributes $A = \{\text{no negation, negation}\}$. There are $n = 206175$ examples in the training set, with 1521 examples in the smallest group (entailment with negations); we modify the standard MultiNLI split to better estimate accuracy on small groups (Appendix C.1).

3 COMPARISON BETWEEN GROUP DRO AND ERM

To study the behavior of group DRO vs. ERM in a modern setting, we fine-tuned ResNet50 models (He et al., 2016) on Waterbirds and CelebA, and a BERT model (Devlin et al., 2019) on MultiNLI; these are popular and high-performing models for image and natural language tasks, respectively.

We train the ERM models using standard (minibatch) stochastic gradient descent. For the DRO models, we introduce a stochastic minibatch algorithm based on online mirror descent that optimizes for an adaptively-weighted training distribution at each iteration; we defer discussion of this to Section 5. We tune the learning rate for ERM and use the same setting for DRO (Appendix C.2).

For each model, we measure its average (in-distribution) accuracy over training and test sets drawn from the same distribution, as well as its robust accuracy on the worst-performing group.

3.1 ERM AND DRO HAVE POOR ROBUST ACCURACY IN THE OVERPARAMETERIZED REGIME

Overparameterized neural networks can perfectly fit the training data and still generalize well on average (Zhang et al., 2017). We start by investigating if these overparameterized models also generalize well on the worst-case group—i.e., if they attain high robust test accuracy on top of high average test accuracy—after they are trained to convergence using standard hyperparameter settings (He et al., 2016; Devlin et al., 2019).

ERM. As expected, ERM models attain near-perfect robust training accuracies of at least 99.9% on all three datasets and also obtain high average test accuracies (97.3%, 94.8%, and 82.5% on Waterbirds, CelebA, and MultiNLI). However, they perform poorly on the worst-case group at test time with robust accuracies of 60.0%, 41.1%, and 65.7% respectively (Table 1, Figure 2). Their low robust accuracies imply that these models are brittle under group shifts.

DRO. In this vanishing-training-loss regime, there exist models that do almost perfectly on both the ERM and DRO objectives. We train group DRO models in the same way as above, and indeed find that they perform similarly to ERM models, attaining near-perfect training accuracies and high average test accuracies, but poor robust test accuracies (Table 1, Figure 2).

Discussion. The ERM and DRO models attain near-perfect training accuracy and vanishing training loss even in the presence of default regularization (batch normalization and $\ell_2$ regularization for ResNet50, and dropout for BERT). However, they do not generalize well on the worst-case group,
|                | Average Accuracy | Robust Accuracy |
|----------------|------------------|-----------------|
|                | ERM | DRO | ERM | DRO |
| **Waterbirds** |     |     |     |     |
| Train          | 100.0 | 100.0 | 100.0 | 100.0 |
| Test           | 97.3  | 97.4  | 60.0  | 76.9  |
| **CelebA**     |     |     |     |     |
| Train          | 100.0 | 100.0 | 99.9  | 100.0 |
| Test           | 94.8  | 94.7  | 41.1  | 41.1  |
| **MultiNLI**   |     |     |     |     |
| Train          | 99.9  | 99.3  | 99.9  | 99.0  |
| Test           | 82.5  | 82.0  | 65.7  | 66.4  |

Table 1: Average and robust accuracies for each training method. Both ERM and DRO models perform poorly on the worst-case group in the absence of regularization (top). With regularization (middle, bottom), DRO achieves high worst-group performance, significantly improving from ERM. Cells are colored by accuracy, from low (red) to medium (white) to high (blue) accuracy.

and consequently suffer from low robust accuracies. This gap between average and robust test accuracies arises not from poor robust training performance—the models are near-perfect at training time, even on the worst-case groups—but from variations in the generalization gaps across groups.

### 3.2 Group DRO Improves Robust Accuracy Under Appropriate Regularization

Classically, we can control the generalization gap with regularization techniques that constrain the model family’s capacity to fit the training data. In the modern overparameterized regime, explicit regularization is not critical for average performance: models can do well on average even when all regularization is removed [Zhang et al., 2017], and default regularization settings (like in the models trained above) still allow models to perfectly fit the training data. Here, we study if increasing regularization strength—until the models no longer perfectly fit the training data—can rescue worst-case performance. We find that departing from the vanishing-training-loss regime allows group DRO models to significantly outperform ERM models on robust test accuracy while maintaining high average accuracy. We investigate two types of regularization:
\( \ell_2 \) regularization. The default \( \ell_2 \) regularization strength (i.e., the coefficient of the \( \ell_2 \)-norm penalty \( \lambda ||\theta||_2^2 \)) in ResNet50 is \( \lambda = 0.0001 \) (He et al. [2016]). We find that increasing \( \lambda \) by several orders of magnitude—to \( \lambda = 1.0 \) for Waterbirds and \( \lambda = 0.1 \) for CelebA—does two things: 1) it prevents both ERM and DRO models from achieving perfect training accuracy, and 2) substantially reduces the generalization gap for each group.

Both ERM and DRO models still achieve high average test accuracies that are comparable to each other and to the standard models with weaker regularization. However, because no model can achieve perfect training accuracy in this regime, ERM models sacrifice training accuracy on the worst-case group (robust accuracies of 35.7% and 40.4% on Waterbirds and CelebA; Table 1, Figure 2) and consequently obtain poor robust test accuracies (21.3% and 37.8%, respectively).

In contrast, DRO models optimize for high robust training accuracy (97.5% and 93.4% on Waterbirds and CelebA). The small generalization gap in the strong \( \ell_2 \) regularization regime means that high robust training accuracy translates to high robust test accuracy: with group DRO, robust test accuracy improves from 21.3% to 84.6% on Waterbirds and from 37.8% to 86.7% on CelebA.

While these results show that strong \( \ell_2 \) regularization has a striking impact on ResNet50 models for Waterbirds and CelebA, our initial experiments did not show a similar effect on BERT model for MultiNLI. \( \ell_2 \) regularization is not typically tuned for this task (Devlin et al. [2019])—our default hyperparameters set \( \lambda = 0 \) (Appendix C.2)—so we turn to other forms of regularization.

Early stopping. A different, implicit form of regularization is early stopping (Hardt et al. [2016b]). We use the same settings in Section 3.1 but only train each model for a fixed (small) number of epochs (Section C.2). As with strong \( \ell_2 \) regularization, curtailing training reduces the generalization gap and prevents models from fitting the data perfectly. In this setting, DRO also does substantially better than ERM on robust test accuracy, improving from 6.7% to 86.0% on Waterbirds, 25.0% to 88.3% on CelebA, and 66.0% to 77.7% on MultiNLI. Average test accuracies are comparably high in both ERM and DRO models, though there is a small drop of 1-3% for DRO (Table 1, Figure 2).

Discussion. We conclude that regularization—preventing the model from perfectly fitting the training data—does matter for robust accuracy. Specifically, regularization can control the generalization gap across each group, even on the worst-case group; good robust test accuracy then becomes a question of good robust training accuracy. ERM and DRO models make different training trade-offs in the regularized regime, since no model can perfectly fit the training data; ERM models sacrifice robust for average training accuracy and therefore do poorly at test time, while DRO models maintain high robust training accuracy and therefore do well at test time. Our findings raise additional questions about the nature of generalization in neural networks, which has been predominantly studied in the context of average accuracy (Zhang et al. [2017]; Hoffer et al. [2017]).

3.3 ACCOUNTING FOR GENERALIZATION THROUGH GROUP ADJUSTMENTS IMPROVES DRO

We have optimized thus far for the robust training loss via DRO (4), relying on regularization to translate good training loss to good test loss by controlling the generalization gap \( \delta \). Here, we show that we can improve performance by directly optimizing for an estimated upper bound on the robust test loss, using ideas from structural risk minimization (Vapnik [1992]).

The key consideration is that each group corresponding to \( g \) has its own generalization gap \( \delta_g = \mathbb{E}_{(X,Y) \sim P_g} [\ell(\theta; (X,Y))] - \mathbb{E}_{(X,Y) \sim P_g} [\ell(\theta; (X,Y))] \). To approximate optimizing for the robust test loss \( \mathcal{R}(\theta) = \max_{g \in \mathcal{G}} \mathbb{E}_{(X,Y) \sim P_g} [\ell(\theta; (X,Y))] + \delta_g \), we propose using the simple, parameter-independent heuristic \( \hat{\delta}_g = C/\sqrt{n_g} \), where \( n_g \) is the group size for \( g \) and \( C \) is a model capacity constant which we treat as a hyperparameter. This gives the group-adjusted DRO estimator

\[
\hat{\theta}_{\text{adj}} := \arg\min_{\theta \in \Theta} \max_{g \in \mathcal{G}} \left\{ \mathbb{E}_{(X,Y) \sim P_g} [\ell(\theta; (X,Y))] + \frac{C}{\sqrt{n_g}} \right\}.
\]

The scaling with \( 1/\sqrt{n_g} \) reflects how smaller groups are more prone to overfitting than larger groups, and is inspired by the general size dependence of model-complexity-based generalization bounds (see, e.g., Cao et al. [2019]). Indeed, even in the regularized setting of Section 3.2 different groups \( g \) can have significant differences in their generalization gaps \( \delta_g \); for example, in the strong-\( \ell_2 \)-regularization DRO model, the smallest group in Waterbirds has a train-test robust accuracy gap of
|                | Average Accuracy | Robust Accuracy |
|----------------|------------------|-----------------|
|                | Naïve  | Adjusted | Naïve | Adjusted |
| Waterbirds     | 96.6   | 93.7     | 84.6  | 90.5     |
| CelebA         | 93.5   | 93.4     | 86.7  | 87.8     |

Table 2: Average and robust test accuracies with and without group adjustments. Group adjustments improve robust accuracy, though average accuracy drops for Waterbirds.

Figure 3: Training (light) and validation (dark) accuracies for each group over training progress, for different values of the adjustment constant $C$. When $C = 0$, the generalization gap for waterbirds on land is large, dragging down robust accuracy; at $C = 2$ (which has the best robust validation accuracy), the accuracies are balanced; and at $C = 4$, we overcompensate for group sizes, so smaller groups (e.g., waterbirds on land) do better at the expense of larger groups (e.g., landbirds on land).

Empirically, we find that group adjustments also help in the early stopping setting of Section 3.2 (in the benchmark in the next section, we consider models with group adjustments and early stopping across a grid of $\ell_2$ regularization strengths, and report on the one with highest validation accuracy). However, it is difficult to rigorously study the effects of early stopping (e.g., because the group losses have not converged to a stable value), so we leave a more thorough investigation of the interaction between early stopping and group adjustments to future work.

4 COMPARISON BETWEEN GROUP DRO AND IMPORTANCE WEIGHTING

Our results above show that appropriately-regularized group DRO models can be significantly more robust than ERM models. Here, we show theoretically and empirically that DRO also outperforms a strong importance weighting baseline that is commonly used in machine learning tasks where the train and test distributions differ (Shimodaira, 2000; Byrd & Lipton, 2019). Recall that in our setting, the test distribution can be any mixture of the group distributions. For some assignment of weights $w \in \Delta_m$ to groups, an importance-weighted estimator would learn

$$\hat{\theta}_w := \arg \min_{\theta \in \Theta} \mathbb{E}_{(x, y, g) \sim \hat{P}} [w_g \ell(\theta; (x, y))].$$

Empirical benchmark. We consider an importance-weighted baseline with weights set to the inverse training frequency of each group, $w_g = 1/\mathbb{E}_{g' \sim \hat{P}}[\mathbb{1}(g' = g)]$. This optimizes for a test distribution with uniform group frequencies and is analogous to the common reweighting technique for label shifts (Cui et al., 2019; Cao et al., 2019). Concretely, we train our weighted model by sampling with equal probability from each group for each minibatch (Shen et al., 2016), since a recent study found this to be more effective than similar methods (Buda et al., 2018).

Unlike group DRO, resampling for uniform group size does not necessarily yield uniformly low training losses across groups in practice, as some groups are easier to fit than others. To compare
resampling (RS) with ERM and group DRO, we benchmark each objective, training models across the same grid of $\ell_2$ regularization strengths and early stopping at the epoch with best robust validation accuracy (Table 3). In CelebA and Waterbirds, resampling performs much better than ERM but is slightly outperformed by group DRO. However, resampling fails on MultiNLI, achieving lower clean and robust accuracies than even ERM. With resampling, it appears that the rare group is overemphasized and extremely low training loss is achieved for that group at the cost of others.

| Objective       | Average Accuracy | Robust Accuracy |
|-----------------|------------------|-----------------|
|                 | ERM (0.2)        | RS (0.3)        | DRO (0.2) |
|                 | 97.0 (0.2)       | 95.1 (0.3)      | 93.5 (0.3) |
|                 | 81.2 (0.1)       | 81.4 (0.1)      | 81.2 (0.1) |
|                 | 94.9 (0.2)       | 92.9 (0.2)      | 92.9 (0.2) |
|                 | 82.8 (0.1)       | 81.4 (0.1)      | 81.4 (0.1) |
|                 | 66.4 (1.6)       | 64.8 (1.6)      | 77.7 (1.4) |

Table 3: Average and robust test accuracies of the models with the best robust validation accuracies for ERM, resampling (RS), and group DRO, with binomial standard deviation in parenthesis. For each objective, we conduct a grid search over regularization, number of epochs, and group adjustments and report the performance of the model with highest validation accuracy.

Figure 4: Toy example illustrating that DRO and importance weighting are not equivalent. The robust solution is $\theta^*$, while any importance weighting would result in solutions at $\theta_1$ or $\theta_2$.

**Theoretical comparison.** Should we expect importance weighting to learn models with good worst-case loss? We show that importance weighting and DRO can learn equivalent models in the convex setting under some importance weights, but not necessarily when the models are non-convex.

For this analysis, we shift to a more general framework of importance weighting, in which we minimize the expected loss $\mathbb{E}_{z \sim P}[w(z)\ell(\theta; z)]$ over some source distribution $P$. Here, we assign weights to each data point $z$, abstracting away from our particular choice of assigning the weights according to its group. Minimizing the weighted objective is equivalent to minimizing the expected unweighted loss $\mathbb{E}_{z \sim Q}[\ell(\theta; z)]$ over a target distribution $Q$ such that $Q(z) \propto w(z)P(z)$.

However, we want good worst-case performance over a family of $Q \in \mathcal{Q}$ instead of optimizing for a single $Q$. Are there weights $w$ such that the resulting model $\hat{\theta}_w$ achieves optimal robust risk? In the convex regime, standard duality arguments show that this is true (see Appendix [A.1] for the proof):

**Proposition 1.** Suppose that the loss $\ell(\cdot; z)$ is continuous and convex for all $z$ in $\mathcal{Z}$, and let the uncertainty set $\mathcal{Q}$ be a set of distributions supported on $\mathcal{Z}$. Assume that $\mathcal{Q}$ and the model family $\Theta \subseteq \mathbb{R}^d$ are convex and compact, and let $\theta^* \in \Theta$ be a minimizer of the robust objective $R(\theta)$. Then there exists a distribution $Q^* \in \mathcal{Q}$ such that $\theta^* \in \arg \min_{Q \in \mathcal{Q}} \mathbb{E}_{z \sim Q}[\ell(\theta; z)]$.

However, this equivalence breaks down when the loss $\ell$ is non-convex (as in neural networks).

**Counterexample 1.** Consider a uniform data distribution $P$ supported on two points $\mathcal{Z} = \{z_1, z_2\}$, and let $\ell(\theta; z)$ be as in Figure 2 with $\Theta = [0, 1]$. The robust solution $\theta^*$ achieves a worst-case loss of $R(\theta^*) = 0.6$. Now consider any weights $(w_1, w_2) \in \Delta_2$ and w.l.o.g. let $w_1 \geq w_2$. The minimizer of the weighted loss $w_1\ell(\theta; z_1) + w_2\ell(\theta; z_2)$ is $\theta_1$, which only attains a worst-case loss of $R(\theta^*) = 1.0$.

This negative result implies that in the non-convex setting, there may not be any choice of weights $w$ such that the resulting minimizer $\hat{\theta}_w$ is robust. Even if such weights did exist, their values are
unknown a priori. In particular, the weights depend on $\theta^*$, highlighting the benefit of algorithms like group DRO that simultaneously learn $\theta$ and $Q$. Common choices of weights, such as inverse group size, are heuristics that may not yield robust solutions (as observed for MultiNLI in Table 3).

5 ALGORITHM

To train group DRO models efficiently, we introduce an online optimization algorithm with convergence guarantees. Prior work on group DRO has either used batch optimization algorithms, which do not scale to large datasets, or stochastic optimization algorithms without convergence guarantees.

In the convex and batch case, there is a rich literature on distributionally robust optimization which treats the problem as a standard convex conic program (Ben-Tal et al. 2013; Duchi et al. 2016; Bertsimas et al. 2018; Lam & Zhou 2015). For general non-convex DRO problems, two types of stochastic optimization methods have been proposed: (i) stochastic gradient descent (SGD) on the Lagrangian dual of the objective (Duchi & Namkoong 2018; Hashimoto et al. 2018), and (ii) direct minimax optimization (Namkoong & Duchi, 2016). The first approach fails for group DRO because the gradient of the dual objective is difficult to estimate in a stochastic and unbiased manner. A second type of algorithm has been proposed for group DRO (Oren et al. 2019), but prior work does not provide convergence guarantees and we observed instability in practice under some settings.

Recall that we aim to solve the optimization problem

$$
\min_{\theta} \sup_{Q \in \Delta_m} \sum_{q=1}^m q_g E_{(X,Y) \sim P_g}[ \ell(\theta; (X,Y)) ].
$$

Extending existing minimax algorithms for DRO (Namkoong & Duchi 2016; Oren et al. 2019), we interleave gradient-based updates on $\theta$ and $q$. Intuitively, we maintain a distribution $q$ over groups, with high masses on high-loss groups, and update on each example proportionally to the mass on its group. Concretely, we interleave SGD on $\theta$ and exponentiated gradient ascent on $q$ (Algorithm 1). The key improvement from the existing group DRO algorithm is that $q$ is updated using gradients instead of picking the group with worst average loss at each iteration, which is important for algorithmic stability and obtaining convergence guarantees. In practice, we implement the algorithm with minibatching and evaluate model performance with the run time of the algorithm is similar to that of SGD for a given number of epochs (less than 5% difference), as run time is dominated by the computation of the loss and its gradient.

Algorithm 1: Online optimization algorithm for group DRO

Input: Step sizes $\eta_\theta, \eta_q$; $P_g$ for each $g \in G$

Initialize $\theta^{(0)}$ and $q^{(0)}$

For $t = 1, \ldots, T$ do

- $q \sim \text{Uniform}(1, \ldots, m)$
- $x, y \sim P_g$
- $q' \leftarrow q^{(t-1)}$
- $q'_g \leftarrow q'_g \exp(\eta_q \ell(\theta^{(t-1)}; (x, y)))$
- $q^{(t)} \leftarrow q'/\sum_g q'_g$
- $\theta^{(t)} \leftarrow \theta^{(t-1)} - \eta_\theta \nabla \ell(\theta^{(t-1)}; (x, y))$

We analyze the convergence rate by studying the error $\varepsilon_T$ of the average iterate $\bar{\theta}^{(1:T)}$:

$$
\varepsilon_T = \max_{q \in \Delta_m} L(\bar{\theta}^{(1:T)}, q) - \min_{\theta} \max_{q \in \Delta_m} L(\theta, q),
$$

where $L(\theta, q) := \sum_{g=1}^m q_g E_{(x,y) \sim P_g} [\ell(\theta; (x,y))]$ is the expected worst-case loss. Applying results from Nemirovski et al. (2009), we can show that Algorithm 1 has a standard convergence rate of $O(1/\sqrt{T})$ in the convex setting.

---

4 The dual optimization problem for group DRO is $\min_{\theta, Q} \frac{1}{\alpha} \mathbb{E}_P [\max(0, \mathbb{E}_{x,y} \sim P_g [\ell(\theta; (x,y)) | g] - \beta)] + \beta$ for constant $\alpha$. The max over expected loss makes it difficult to obtain an unbiased, stochastic gradient estimate.
Proposition 2. Suppose that the loss $\ell((x,y))$ is non-negative, convex, $B_2$-Lipschitz continuous, and bounded by $B$ for all $(x,y)$ in $\mathcal{X} \times \mathcal{Y}$, and $\|\theta\|_2 \leq B_0$ for all $\theta \in \Theta$ with convex $\Theta \subseteq \mathbb{R}^d$. Then, the average iterate of Algorithm 1 achieves an expected error at the rate

$$E[\varepsilon_T] \leq 2m \sqrt{\frac{10B^2B_0^2 + B_0^2 \log m}{T}}.$$  \hfill (9)

where the expectation is taken over the randomness of the algorithm.

6 RELATED WORK

The problem of non-uniform accuracy. Other approaches to addressing non-uniform accuracy over the data distribution include domain adaptation techniques for known target distributions (Ben-David et al., 2006; Ganin & Lempitsky, 2015) and work in ML fairness (Dwork et al., 2012; Hardt et al., 2016a; Kleinberg et al., 2017). As we discuss in Section 4, importance weighting is a classic example of the former (Shimodaira, 2000). Byrd & Lipton (2019) empirically study importance weighting in neural networks and demonstrate that it has little effect unless regularization is applied. This is consistent with the theoretical analysis in Wen et al. (2014), which points out that weighting has little impact in the zero-loss regime, and with our own observations in the context of DRO.

Distributionally robust optimization. Prior work in DRO typically defines the uncertainty set $Q$ as a divergence ball around the training distribution over $(x,y)$ (Ben-Tal et al., 2013; Lam & Zhou, 2015; Duchi et al., 2016; Miyato et al., 2018; Esfahani & Kuhn, 2018; Bertsimas et al., 2018; Blachet & Murthy, 2019). With small divergence balls of radii $O(1/n)$, DRO acts as a regularizer (Shafieezadeh-Abadeh et al., 2015; Namkoong & Duchi, 2017). However, when the radius is larger, the resulting $Q$ can be too pessimistic. To better control $Q$, prior work proposed group DRO in the context of label shifts (Hu et al., 2018) and shifts in data sources (Oren et al., 2019). Our work studies group DRO in the overparameterized regime with vanishing training loss and poor worst-case generalization. In contrast, prior work on DRO has operated in settings with training loss trade-offs and reasonable generalization. Most DRO work has focused on the classic (underparameterized) model setting (Namkoong & Duchi, 2017; Hu et al., 2018; Duchi et al., 2019); Sinha et al. (2018) study neural networks but with a more conservative Wasserstein uncertainty set that leads to non-vanishing training loss; and Oren et al. (2019) study neural networks but for generative modeling.

Generalization of robust models. There is extensive work investigating generalization of neural networks in terms of average loss, theoretically and empirically (Hardt et al., 2016b; Szegedy et al., 2016; Hoffer et al., 2017). However, analysis on robust losses is limited. For label shifts, prior work has observed overfitting on rare labels and proposed algorithms to mitigate it (Buda et al., 2018; Cui et al., 2019; Cao et al., 2019). In the DRO literature, generalization bounds on the DRO objective exist for particular uncertainty sets (Duchi & Namkoong, 2018), but these bounds do not apply directly to group DRO. Invariant prediction models from the causal inference literature similarly aim to achieve high performance on a range of test distributions (Peters et al., 2016; Bühlmann & Meinshausen, 2016; Heinze-Deml & Meinshausen, 2017; Rothenhäusler et al., 2018; Yang et al., 2019). The maximin regression framework (Meinshausen & Bühlmann, 2015) also assumes group-based shifts, but focuses on settings without the generalization problems identified in our work.

7 DISCUSSION

In this paper, we analyzed group DRO in the context of overparameterized neural networks and highlighted the importance of regularization for worst-case group generalization. When appropriately regularized, group DRO significantly improves robust accuracy at a small cost in average accuracy.

As an application of group DRO, we showed that it can prevent models from learning pre-specified spurious correlations. While handling shifts beyond pre-specified group shifts is important future work, existing work has identified many distributional shifts that can be expressed with pre-specified groups: e.g., batch effects in biology (Leek et al., 2010), or image artifacts (Oakden-Rayner et al., 2019) and patient demographics (Badgerley et al., 2019) in medicine. Our supplemental experiments also suggest that group DRO models can maintain high robust accuracy even when groups are imperfectly specified (Appendix E).
More generally, our observations call for a deeper analysis of average vs. worst-case generalization in the overparameterized regime. Such analysis may shed light on the failure modes of deep neural networks as well as provide more precise tools (beyond strong $\ell_2$ regularization or early stopping) to counter poor worst-case generalization while maintaining high average accuracy.

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REPRODUCIBILITY

Code for training group DRO models is available at [https://github.com/kohpangwei/group_DRO](https://github.com/kohpangwei/group_DRO). We are working on releasing scripts that will replicate the experiments above. The constructed Waterbirds dataset will also be publicly available for download, with a script to adjust its generation (e.g., to choose different object backgrounds or proportions).

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## A Proofs

### A.1 Equivalence of DRO and Importance Weighting in the Convex Setting

**Proposition** Suppose that the loss $\ell(\cdot; z)$ is continuous and convex for all $z$ in $\mathcal{Z}$, and let the uncertainty set $\mathcal{Q}$ be a set of distributions supported on $\mathcal{Z}$. Assume that $\mathcal{Q}$ and the model family $\mathcal{X} \subseteq \mathbb{R}^d$ are convex and compact, and let $\theta^* \in \Theta$ be a minimizer of the robust objective $R(\theta)$. Then there exists a distribution $Q^* \in \mathcal{Q}$ such that $\theta^* \in \arg\min_{\theta \in \Theta} \mathbb{E}_{z \sim Q}[\ell(\theta; z)]$.

**Proof.** Let $h(\theta, Q) := \mathbb{E}_{Z \sim Q}[\ell(\theta; Z)]$. Since the loss $\ell(\theta; Z)$ is continuous and convex in $\theta$ for all $Z$ in $\mathcal{Z}$, we have that $h(\theta, Q)$ is continuous, convex in $\theta$, and concave (linear) in $Q$. Moreover, since convexity and lower semi-continuity are preserved under arbitrary pointwise suprema, $\sup_{Q \in \mathcal{Q}} h(\theta, Q)$ is also convex and lower semi-continuous (therefore proper).

Together with the compactness of $\Theta$ and $\mathcal{Q}$, the above conditions imply (by Weierstrass’ theorem, proposition 3.2.1, Bertsekas [2009]), that the optimal value of the DRO objective

$$\inf_{\theta \in \Theta} R(\theta) = \inf_{\theta \in \Theta} \sup_{Q \in \mathcal{Q}} h(\theta, Q).$$

is attained at some $\theta^* \in \Theta$. 

A similar argument implies that the sup-inf objective
\[
\sup_{Q \in \mathcal{Q}} \inf_{\theta \in \Theta} h(\theta, Q)
\]
attains its optimum at some \( Q^* \in \mathcal{Q} \).
Moreover, because \( \Theta \) and \( \mathcal{Q} \) are compact and \( h \) is continuous, we have the max-min equality (see, e.g., Ex 5.25 in Boyd & Vandenberghe (2004))
\[
\sup_{Q \in \mathcal{Q}} \inf_{\theta \in \Theta} h(\theta, Q) = \inf_{\theta \in \Theta} \sup_{Q \in \mathcal{Q}} h(\theta, Q).
\]
(12)
Together, the above results imply that \((\theta^*, Q^*)\) form a saddle point (proposition 3.4.1, Bertsekas (2009)), that is,
\[
\sup_{Q \in \mathcal{Q}} h(\theta^*, Q) = h(\theta^*, Q^*) = \inf_{\theta \in \Theta} \sup_{Q \in \mathcal{Q}} h(\theta, Q^*).
\]
(13)
In particular, the second equality indicates that the optimal DRO model \( \theta^* \) also minimizes the weighted risk \( h(\theta, Q^*) = \mathbb{E}_{Z \sim Q} [\ell(\theta; Z)] \), as desired.

A.2 CONVERGENCE RATE OF ALGORITHM

Proposition 2. Suppose that the loss \( \ell(\cdot; (x, y)) \) is non-negative, convex, \( B_\nabla \)-Lipschitz continuous, and bounded by \( B_\ell \) for all \( (x,y) \) in \( \mathcal{X} \times \mathcal{Y} \), and \( \|\theta\|_2 \leq B_\theta \) for all \( \theta \in \Theta \) with convex \( \Theta \subseteq \mathbb{R}^d \).

Then, the average iterate of Algorithm 1 achieves an expected error at the rate
\[
\mathbb{E}[\varepsilon_T] \leq 2m \sqrt{\frac{10[B_\nabla^2 B_\ell^2 + B_\ell^2 \log m]}{T}}.
\]
(14)
where the expectation is taken over the randomness of the algorithm.

Proof. Our proof is an application of the regret bound for online mirror descent on saddle point optimization from Nemirovski et al. (2009).

We first introduce the existing theorem. Consider the saddle-point optimization problem
\[
\min_{\theta \in \Theta} \max_{q \in \Delta_m} \sum_{g=1}^m q_g f_g(\theta)
\]
under the following assumptions:

Assumption 1. \( f_g \) is convex on \( \Theta \).

Assumption 2. \( f_g(\theta) = \mathbb{E}_{\xi \sim q}[F_g(\theta; \xi)] \) for some function \( F_g \).

Assumption 3. We generate i.i.d. examples \( \xi \sim q \). For a given \( \theta \in \Theta \) and \( \xi \in \Xi \), we can compute \( F_g(\theta, \xi) \) and unbiased stochastic subgradient \( \nabla F_g(\theta; \xi) \), that is, \( \mathbb{E}_{\xi \sim q}[\nabla F_g(\theta; \xi)] = \nabla f_g(\theta) \).

Online mirror descent with some \( c \)-strongly convex norm \( \|\cdot\|_{\theta} \), yielding iterates \( \theta^{(1)}, \ldots, \theta^{(T)} \) and \( q^{(1)}, \ldots, q^{(T)} \), has the following guarantee.

Theorem 1 (Nemirovski et al. (2009), Eq 3.23). Suppose that Assumptions 1-3 hold. Then the pseudo-regret of the average iterates \( \bar{q}_g^{(1:T)} \) and \( \bar{q}_g^{(1:T)} \) can be bounded as
\[
\mathbb{E} \left[ \max_{q \in \Delta_m} \sum_{g=1}^m q_g f_g(\bar{q}_g^{(1:T)}) - \min_{\theta \in \Theta} \sum_{g=1}^m q_g f_g(\theta) \right] \leq 2 \sqrt{\frac{10[B_\nabla^2 M_\nabla^2 + M_\nabla^2 \log m]}{T}},
\]
(16)
where
\[
\mathbb{E} \left[ \left\| \nabla_{\theta} \sum_{g=1}^{m} q F_g(\theta; \xi) \right\|_{*}^{2} \right] \leq M_{\star, \theta}^{\star}
\]
(17)
\[
\mathbb{E} \left[ \left\| \nabla_{q} \sum_{g=1}^{m} q F_g(\theta; \xi) \right\|_{*}^{2} \right] \leq M_{\star, q}^{\star}
\]
(18)
\[
R_{\theta}^{2} = \frac{1}{c} \left( \max_{\theta} \|\theta\|_{\theta}^{2} - \min_{\theta} \|\theta\|_{\theta}^{2} \right)
\]
(19)

for online mirror descent with c-strongly convex norm \(\| \cdot \|_{\theta}\).

It remains to formulate our algorithm as an instance of online mirror descent applied to the saddle-point problem above. We start by defining the following:

**Definition 1.** Let \( q \) be a distribution over \( \xi = (x, y, g) \) that is a uniform mixture of individual group distributions \( P_{g} \):\n
\[
(x, y, g) \sim q := \frac{1}{m} \sum_{g'=1}^{m} P_{g'}.
\]
(20)

**Definition 2.** Let \( F_{g'}(\theta; (x, y, g)) := m \mathbb{I}[g = g'] \ell(\theta; (x, y)) \). Correspondingly, let \( f_{g'} := \mathbb{E}_{P_{g'}}[\ell(\theta; (x, y))] \).

We now check that Assumptions 1-3 hold under the original assumptions in the statement of Theorem 2:

1. We assume that the loss \( \ell(\cdot; (x, y)) \) is non-negative, continuous, and convex for all \((x, y)\) in \(\mathcal{X} \times \mathcal{Y}\). As a result, \( f_{g}(\theta) \) is non-negative, continuous, and convex on \(\Theta\).

2. The expected value of \( F_{g}(\theta) \) over distribution \( q \) is \( f_{g}(\theta) \):

\[
\mathbb{E}_{x,y,g \sim q}[F_{g'}(\theta; (x, y, g))] = \frac{1}{m} \sum_{i=1}^{m} \mathbb{E}_{P_i}[F_{g'}(\theta; (x, y, g)) | g = i]
\]
\[
= \frac{1}{m} \mathbb{E}_{P_{g'}}[F_{g'}(\theta; (x, y, g)) | g = g']
\]
\[
= \frac{1}{m} \mathbb{E}_{P_{g'}}[m \ell(\theta; x, y) | g = g']
\]
\[
= \mathbb{E}_{P_{g'}}[\ell(\theta; x, y) | g = g']
\]
\[
= f_{g'}(\theta).
\]

3. We can compute an unbiased stochastic subgradient \( \nabla F_{g'}(\theta; (x, y, g)) \)

\[
\mathbb{E}_{x,y,g \sim q}[\nabla F_{g'}(\theta; (x, y, g))] = \mathbb{E}_{x,y,g \sim q}[\nabla m \mathbb{I}[g = g'] \ell(\theta; (x, y))]
\]
\[
= \frac{1}{m} \sum_{i=1}^{m} \mathbb{E}_{P_i}[\nabla m \mathbb{I}[g = g'] \ell(\theta; (x, y))]
\]
\[
= \mathbb{E}_{Q_{g'}}[\nabla \ell(\theta; (x, y))]
\]
\[
= \nabla f_{g'}(\theta).
\]

Finally, we compute the constants required for the regret bound in Theorem 2. Recalling the original assumptions of Theorem 2:

1. Bounded losses: \( \ell(\theta; (x, y)) \leq B_{\ell} \) for all \( x, y, \theta \)

2. Bounded gradients: \( \| \nabla \ell(\theta; (x, y)) \|_{2} \leq B_{\psi} \) for all \( \theta, x, y \)
3. Bounded parameter norm: \( \|\theta\|_2 \leq B_\theta \) for all \( \theta \in \Theta \),

we obtain:

\[
\mathbb{E} \left[ \left\| \nabla_\theta \sum_{g'=1}^m q_{g'} F_{g'}(\theta; (x, y, g)) \right\|_{\ast, \theta}^2 \right] \leq m^2 B_C^2 = M_{\ast, \theta} \tag{21}
\]

\[
\mathbb{E} \left[ \left\| \nabla_q \sum_{g'=1}^m q_{g'} F_{g'}(\theta; (x, y, g)) \right\|_{\ast, q}^2 \right] \leq m^2 B_{\ell}^2 = M_{\ast, q} \tag{22}
\]

\[
R_\theta^2 = \max_{\theta} \|\theta\|_2^2 - \min_{\theta} \|\theta\|_2^2 = B_\theta^2. \tag{23}
\]

Plugging in these constants into the regret bound from Theorem\[1\] we obtain

\[
\mathbb{E} \left[ \max_{q \in \Delta_m} \sum_{g=1}^m q_g f_g(\tilde{\theta}^{(1:T)}) - \min_{\theta \in \Theta} \sum_{g=1}^m \tilde{q}_g^{(1:T)} f_g(\theta) \right] \leq 2m \sqrt{\frac{10|B_{\ell}^2 B_C^2 + B_{\ell}^2 \log m|}{T}} \tag{24}
\]

This implies Theorem\[2\] because the minimax game is convex-concave.

\[\square\]

### B Supplementary Experiments

Group DRO can maintain high robust accuracy even when spurious attributes are not perfectly specified. We repeat the CelebA experiment on strong \( \ell_2 \) regularization (Section\[3.2\]) but with inexact group specifications:

1. Instead of the ground-truth spurious attribute \textit{Male}, we provide a related attribute \textit{Wearing Lipstick}, and
2. We also specify four distractor/non-spurious attributes (\textit{Eyeglasses, Smiling, Double Chin,} and \textit{Oval Face}).

Optimizing for worst-case performance over all \( 2^5 = 64 \) groups (for all combinations of 5 attributes and 1 label), the DRO model attains 78.9% robust accuracy across the 4 original groups (dark-haired males and females, and blond males and females). These robust accuracies are not far off from the original DRO model with just the ground-truth spurious attribute (86.7%) and significantly outperform the ERM model (37.8%).

### C Experimental Details

#### C.1 Datasets

**MultiNLI.** The standard MultiNLI train-test split allocates most examples (approximately 90%) to the training set, with another 5% as a publicly-available development set and the last 5% as a held-out test set that is only accessible through online competition leaderboards (Williams et al., 2018). To accurately estimate performance on rare groups in the validation and test sets, we combine the training set and development set and then randomly re-split it to a 50 – 20 – 30 train-val-test split that allocates more examples to the validation and test sets than the standard split.

We use the provided gold labels as the target, removing examples with no consensus gold label (as is standard procedure). We annotate an example as having a negation word if any of the words \textit{nobody, no, never,} and \textit{nothing} appear in the hypothesis (Gururangan et al., 2018).
**Waterbirds.** The CUB dataset (Wah et al., 2011) contains photographs of birds annotated by species as well as pixel-level segmentation masks of each bird. To construct the Waterbirds dataset, we label each bird as a *waterbird* if it is a seabird (albatross, auklet, cormorant, frigatebird, fulmar, gull, jaeger, kittiwake, pelican, puffin, or tern) or waterfowl (gadwall, grebe, mallard, merganser, guillemot, or Pacific loon). Otherwise, we label it as a *landbird*.

To control the image background, we use the provided pixel-level segmentation masks to crop each bird out from its original background and onto a water background (categories: ocean or natural lake) or land background (categories: bamboo forest or broadleaf forest) obtained from the Places dataset (Zhou et al., 2017). In the training set, we place 95% of all waterbirds against a water background and the remaining 15% against a land background. Similarly, 95% of all landbirds are placed against a land background with the remaining 15% against water.

We refer to this combined CUB-Places dataset as the Waterbirds dataset to avoid confusion with the original fine-grained species classification task in the CUB dataset.

We use the official train-test split of the CUB dataset, randomly choosing 20% of the training data to serve as a validation set. For the validation and test sets, we allocate equal numbers of birds to each of the four groups (i.e., landbirds and waterbirds are uniformly distributed on land and water backgrounds) so as to be able to more accurately measure the performance of the rare groups. This is particularly important for the Waterbirds dataset because of its relatively small size; otherwise, the smaller groups (waterbirds on land and landbirds on water) would have too few samples to accurately estimate performance on. We note that we can only do this for the Waterbirds dataset because we control the generation process; for the other datasets, we cannot generate more samples from the rare groups.

Due to the above procedure, when reporting average test accuracy in our experiments, we calculate the average test accuracy over each group and then report a weighted average, with weights corresponding to the relative proportion of each group in the (skewed) training dataset.

**CelebA.** We use the official train-val-test split that accompanies the CelebA celebrity face dataset (Liu et al., 2015). We use the *Blond Hair* attribute as the target label and the *Male* attribute as the spuriously-associated variable.

### C.2 Models

**ResNet50.** We use the Pytorch torchvision implementation of the ResNet50 model, starting from pretrained weights.

We train the ResNet50 models using stochastic gradient descent with a momentum term of 0.9 and a batch size of 128; the original paper used batch sizes of 128 or 256 depending on the dataset (He et al., 2016). As in the original paper, we used batch normalization (Ioffe & Szegedy, 2015) and no dropout (Srivastava et al., 2014). For simplicity, we train all models without data augmentation.

We use a fixed learning rate instead of the standard adaptive learning rate schedule to make our different model types easier to directly compare, since we expected the scheduler to interact differently with different model types (e.g., due to the different definition of loss). The interaction between batch norm and $\ell_2$ regularization means that we had to adjust learning rates for each different setting of $\ell_2$ regularization (and each dataset). The learning rates below were chosen to be the highest learning rates that still resulted in stable optimization.

For the standard training experiments in Section 3.1 we use a $\ell_2$ regularization strength of $\lambda = 0.0001$ (as in He et al., 2016) for both Waterbirds and CelebA, with a learning rate of 0.001 for Waterbirds and 0.0001 for CelebA. We train the CelebA models for 50 epochs and the Waterbirds models for 300 epochs.

For the early stopping experiments in Section 3.2 we train each ResNet50 model for 1 epoch. For the strong $\ell_2$ regularization experiments in that section, we use $\lambda = 1.0$ for Waterbirds and $\lambda = 0.1$ for CelebA, with both datasets using a learning rate of 0.00001. These settings of $\lambda$ differ because we found that the lower value was sufficient for controlling overfitting on CelebA but not on Waterbirds.

For the group adjustment experiments in Section 3.3 we use the same settings of $\lambda = 1.0$ for Waterbirds and $\lambda = 0.1$ for CelebA, with both datasets using a learning rate of 0.00001. For both
datasets, we use the value of $C \in \{0, 1, 2, 3, 4, 5\}$ found in the benchmark grid search described below.

For the benchmark in Section 4 (Table 3), we grid search over $\ell_2$ regularization strengths of $\lambda \in \{0.0001, 0.1, 1.0\}$ for Waterbirds and $\lambda \in \{0.0001, 0.01, 0.1\}$ for CelebA, using the corresponding learning rates for each $\lambda$ and dataset listed above. (Waterbirds and CelebA at $\lambda = 0.1$, which is not listed above, both use a learning rate of 0.0001.) To avoid advantaging DRO by allowing it to try many more hyperparameters, we only test group adjustments (searching over $C \in \{0, 1, 2, 3, 4, 5\}$) on the $\ell_2$ regularization strengths used in Section 3.3, i.e., $\lambda = 1.0$ for Waterbirds and $\lambda = 0.1$ for CelebA. All benchmark models were evaluated at the best early stopping epoch (as measured by robust validation accuracy).

**BERT.** We use the Hugging Face `pytorch-transformers` implementation of the BERT `bert-base-uncased` model, starting from pretrained weights (Devlin et al., 2019). We use the default tokenizer and model settings from that implementation, including a fixed linearly-decaying learning rate starting at 0.00002, AdamW optimizer, dropout, and no $\ell_2$ regularization ($\lambda = 0$), except that we use a batch size of 32 (as in Devlin et al. (2019)) instead of 8. We found that this slightly improved robust accuracy across all models and made the optimization less noisy, especially on the ERM model.

For the standard training experiments in Section 3.1, we train for 20 epochs.

For the $\ell_2$ regularization experiments in Section 3.2, we tried regularization strengths of $\lambda \in \{0.01, 0.03, 0.1, 0.3, 1.0, 3.0, 10.0\}$. However, these models had similar or worse robust accuracies compared to the default model with no $\ell_2$ regularization.

For the early stopping experiments in Section 3.2, we train for 3 epochs, which is the suggested early-stopping time in Devlin et al. (2019).

For the benchmark in Section 4 (Table 3), we similarly trained for 3 epochs. All benchmark models were evaluated at the best early stopping epoch (as measured by robust validation accuracy).