SWAD: Domain Generalization by Seeking Flat Minima

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Abstract

Domain generalization (DG) methods aim to achieve generalizability to an unseen target domain by using only training data from the source domains. Although a variety of DG methods have been proposed, a recent study shows that under a fair evaluation protocol, called DomainBed, the simple empirical risk minimization (ERM) approach works comparable to or even outperforms previous methods. Unfortunately, simply solving ERM on a complex, non-convex loss function can easily lead to sub-optimal generalizability by seeking sharp minima. In this paper, we theoretically show that finding flat minima results in a smaller domain generalization gap. We also propose a simple yet effective method, named Stochastic Weight Averaging Densely (SWAD), to find flat minima. SWAD finds flatter minima and suffers less from overfitting than does the vanilla SWA by a dense and overfit-aware stochastic weight sampling strategy. SWAD shows state-of-the-art performances on five DG benchmarks, namely PACS, VLCS, OfficeHome, TerraIncognita, and DomainNet, with consistent and large margins of +1.6% averagely on outof-domain accuracy. We also compare SWAD with conventional generalization methods, such as data augmentation and consistency regularization methods, to verify that the remarkable performance improvements are originated from by seeking flat minima, not from better in-domain generalizability. Last but not least, SWAD is readily adaptable to existing DG methods without modification; the combination of SWAD and an existing DG method further improves DG performances.

1 Introduction

Independent and identically distributed (i.i.d.) condition is the underlying assumption of machine learning experiments. However, this assumption may not hold in real-world scenarios, *i.e.*, the training and the test data distribution may differ significantly by *distribution shifts*. For example, a self-driving car should adapt to adverse weather or day-to-night shifts [1, 2]. Even in a simple image recognition scenario, systems rely on wrong cues for their prediction, *e.g.*, geographic distribution [3], demographic statistics [4], texture [5], or backgrounds [6]. Consequently, a practical system should require generalizability to distribution shift, which is yet often failed by traditional approaches.

Domain generalization (DG) aims to address *domain shift* simulated by training and evaluating on different domains. DG tasks assume that both task labels and domain labels are accessible. For example, PACS dataset [7] has seven task labels (*e.g.*, "dog", "horse") and four domain labels (*e.g.*, "photo", "sketch"). Previous approaches explicitly reduced domain gaps in the latent space [8–12], obtained well-transferable model parameters by the meta-learning framework [13–16], data

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Table 1: **Comparisons with SOTA.** The proposed SWAD outperforms other state-of-the-art DG methods on five different DG benchmarks with significant gaps (+1.6pp in the average).

	PACS	VLCS	OfficeHome	TerraInc	DomainNet	Avg.
ERM [34]	85.5	77.5	66.5	46.1	40.9	63.3
Best SOTA competitor	86.6 [35]	78.8 [36]	68.7 [36]	48.6 [37]	43.6 [15, 38]	65.3
SWAD (proposed)	88.1	79.1	70.6	50.0	46.5	66.9

augmentation [17–19], or capturing causal relation [20, 21]. Despite numerous previous attempts for a decade, Gulrajani and Lopez-Paz [22] showed that a simple empirical risk minimization (ERM) approach works comparably or even outperforms the previous attempts on diverse DG benchmarks under a fair evaluation protocol, called "DomainBed".

Unfortunately, although ERM showed surprising empirical success on DomainBed, simply minimizing the empirical loss on a complex and non-convex loss landscape is typically not sufficient to arrive at a good generalization [23–26]. In particular, the connection between the generalization gap and the flatness of loss landscapes has been actively discussed under the i.i.d. condition [23–28]. Izmailov et al. [25] argued that seeking flat minima will lead to robustness against the loss landscape shift between training and test datasets, while a simple ERM converges to the boundary of a wide flat minimum and achieves insufficient generalization. In the DG scenario, because training and test loss landscapes differ more drastically due to the domain shift, we conjecture that the generalization gap between flat and sharp minima is larger than expected in the i.i.d. scenario.

To show that flatter minima generalize better to unseen domains, we formulate a robust risk minimization (RRM) problem defined by the worst-case empirical risks within neighborhoods in parameter space [26, 29]. We theoretically show that the generalization gap of DG, i.e., the error on the target domain, is upper bounded by RRM, i.e., a flat optimal solution. In practice, we approximate a flat minimum by a simple yet effective method, named Stochastic Weight Averaging Densely (SWAD). Specifically, SWAD introduces a dense and overfit-aware stochastic weight sampling strategy for SWA [25]. First, we suggest to sample weights *densely*, *i.e.*, for every iteration. Also, we search the start and end iterations for averaging by considering the validation loss to avoid overfitting. We empirically show that SWAD finds flatter minima than the vanilla SWA does, results in better generalization to unseen domains. Similarly, SWAD performs better than other generalization methods, such as data augmentation and consistency regularization methods, for out-of-domain generalization. In our extensive experiments on five DG benchmarks (PACS [7], VLCS [30], OfficeHome [31], TerraIncognita [32], and DomainNet [33]), SWAD achieves state-of-the-art performances with a large margin of 1.6pp on average compared to the previous best performers (See Table 1). Furthermore, since our SWAD does not change the objective function or the model architecture, SWAD can be combined with existing DG methods and improve the performances as shown in Table 4.

2 A Theoretical Analysis on Flat Minima and Domain Generalization Gap

Let $\mathcal{D}:=\left\{\mathcal{D}_i\right\}_i^I$ be a set of training domains, where \mathcal{D}_i is a distribution over input space \mathcal{X} , and I is the total number of domains. From each domain, we observe n training data points which consist of input x and target label y, $(x_j^i, y_j^i)_{j=1}^n \sim \mathcal{D}_i$. We also define a set of target domain $\mathcal{T}:=\left\{\mathcal{T}_i\right\}_i^T$ similarly, where the number of target domains T is usually set to one. Domain generalization (DG) aims to find a model parameter $\theta \in \Theta$ which generalizes well over both multiple training domains \mathcal{D} and unseen target domain \mathcal{T} . More specifically, let us consider a bounded instance loss function $\ell: \mathcal{Y} \times \mathcal{Y} \to [0,c]$, such that $\ell(y_1,y_2)=0$ holds if and only if $y_1=y_2$ where \mathcal{Y} is a set of labels. For simplicity, we set c to one in our proofs, but we note that $\ell(\cdot,\cdot)$ can be generalized for any bounded loss function. Then, we can define a population loss over multiple domains by $\mathcal{E}_{\mathcal{D}}(\theta)=\frac{1}{I}\sum_{i=1}^{I}\mathbb{E}_{x^i\sim\mathcal{D}_i}[\ell(f(x^i;\theta),y^i))]$, where $f(\cdot;\theta)$ is a model parameterized by θ . Formally, the goal of DG is to find a model which minimizes both $\mathcal{E}_{\mathcal{D}}(\theta)$ and $\mathcal{E}_{\mathcal{T}}(\theta)$ by only minimizing an empirical risk $\hat{\mathcal{E}}_{\mathcal{D}}(\theta):=\frac{1}{In}\sum_{i=1}^{I}\sum_{j=1}^{I}\ell(f(x^i;\theta),y^i))$ over training domains \mathcal{D} .

In practice, ERM, *i.e.*, $\arg\min_{\theta} \hat{\mathcal{E}}_{\mathcal{D}}(\theta)$, can have multiple solutions that provide similar values of the training losses but significantly different generalization performances on $\mathcal{E}_{\mathcal{D}}(\theta)$ and $\mathcal{E}_{\mathcal{T}}(\theta)$. Unfortunately, the typical optimization methods, such as SGD and Adam [39], often lead sub-optimal generalizability as finding sharp and narrow minima even under the i.i.d. assumption [23–28]. In

the DG scenario, the generalization gap between empirical loss and target domain loss becomes even worse due to domain shift. Here, we provide a theoretical interpretation of the relationship between finding a flat minimum and minimizing the domain generalization gap, inspired by previous studies [23-28].

We consider a robust empirical loss function defined by the worst-case loss within neighborhoods in the parameter space as $\hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta) := \max_{\|\Delta\| \leq \gamma} \hat{\mathcal{E}}_{\mathcal{D}}(\theta + \Delta)$, where $\|\cdot\|$ denotes the L2 norm and γ is a radius which defines neighborhoods of θ . Intuitively, if γ is sufficiently larger than the "radius" of a sharp optimum θ_s of $\mathcal{E}_{\mathcal{D}}(\theta)$, θ_s is no longer an optimum of $\hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta)$ as well as its neighborhoods within the γ -ball. On the other hand, if an optimum θ_f has larger "radius" than γ , there exists a local optimum within γ -ball – See Figure 1. Hence, solving the robust risk minimization (RRM), *i.e.*, $\arg\min_{\theta} \hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta)$, will find RRM will find flat minima.

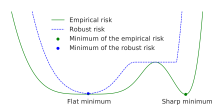


Figure 1: Robust risk minimization (RRM) and flat minima. With proper γ ,

a near solution of a flat optimum showing better generalizability [26, 29]. However, as domain shift worsen the generalization gap by breaking the i.i.d. assumption, it is not trivial that RRM will find an optimum with better DG performance. To answer the question, we first show the generalization bound between $\hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}$ and $\mathcal{E}_{\mathcal{T}}$ as follows:

Theorem 1. Consider a set of N covers $\{\Theta_k\}_{k=1}^N$ such that the parameter space $\Theta \subset \bigcup_k^N \Theta_k$ where $diam(\Theta) := \sup_{\theta, \theta' \in \Theta} \|\theta - \theta'\|_2$, $N := \left\lceil (diam(\Theta)/\gamma)^d \right\rceil$ and d is dimension of Θ . Let v_k be a VC dimension of each Θ_k . Then, for any $\theta \in \Theta$, the following bound holds with probability at least $1 - \delta$,

$$\mathcal{E}_{\mathcal{T}}(\theta) < \hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta) + \frac{1}{2I} \sum_{i=1}^{I} \mathbf{Div}(\mathcal{D}_{i}, \mathcal{T}) + \max_{k \in [1, N]} \sqrt{\frac{v_{k} \ln (m/v_{k}) + \ln(N/\delta)}{m}}, \tag{1}$$

where m = nI is the number of the training samples and $\mathbf{Div}(\mathcal{D}_i, \mathcal{T}) := 2 \sup_A |\mathbb{P}_{\mathcal{D}_i}(A) - \mathbb{P}_{\mathcal{T}}(A)|$ is a divergence between two distributions.

In Theorem 1, the test loss $\mathcal{E}_{\mathcal{T}}(\theta)$ is bounded by three terms: (1) the robust empirical loss $\hat{\mathcal{E}}_{\mathcal{T}}^{\gamma}(\theta)$, (2) the discrepancy between training distribution and test distribution, i.e., the quantity of domain shift, and (3) a confidence bound related to the radius γ and the number of the training samples m. More details of Theorem 1, including proof and discussions on the confidence bound, are in Appendix.

From Theorem 1, one can conjure that minimizing the robust empirical loss is directly related to the generalization performances on the target distribution. We show that the domain generalization gap on the target domain \mathcal{T} by the optimal solution of RRM, $\hat{\theta}^{\gamma}$, is upper bounded as follows:

Theorem 2. Let $\hat{\theta}^{\gamma}$ denote the optimal solution of the RRM, i.e., $\hat{\theta}^{\gamma} := \arg\min_{\theta} \hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta)$, and let v be a VC dimension of the parameter space Θ . Then, the gap between the optimal test loss, $\min_{\theta'} \mathcal{E}_{\mathcal{T}}(\theta')$, and the test loss of $\hat{\theta}^{\gamma}$, $\mathcal{E}_{\mathcal{T}}(\hat{\theta}^{\gamma})$, has the following bound with probability at least $1 - \delta$.

$$\mathcal{E}_{\mathcal{T}}(\hat{\theta}^{\gamma}) - \min_{\theta'} \mathcal{E}_{\mathcal{T}}(\theta') \leq \hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\hat{\theta}^{\gamma}) - \min_{\theta''} \hat{\mathcal{E}}_{\mathcal{D}}(\theta'') + \frac{1}{I} \sum_{i=1}^{I} \mathbf{Div}(\mathcal{D}_{i}, \mathcal{T}) + \max_{k \in [1, N]} \sqrt{\frac{v_{k} \ln(m/v_{k}) + \ln(2N/\delta)}{m}} + \sqrt{\frac{v \ln(m/v) + \ln(2/\delta)}{m}}$$
(2)

Proof is in Appendix. It implies that if we find the optimal solution of the RRM (i.e., $\hat{\theta}^{\gamma}$), then the generalization gap in the test domain (i.e., $\mathcal{E}_{\mathcal{T}}(\hat{\theta}^{\gamma}) - \min_{\theta'} \mathcal{E}_{\mathcal{T}}(\theta')$) is upper bounded by the gap between the RRM and ERM (i.e., $\hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\hat{\theta}^{\gamma}) - \min_{\theta''} \hat{\mathcal{E}}_{\mathcal{D}}(\theta'')$). Other terms in Theorem 2 are the discrepancy between the train domains \mathcal{D} and the target domain \mathcal{T} , and the confidence bounds caused by sample means. We remark that if we choose a proper γ , the optimal solution of the RRM will find a point near a flat optimum of ERM as shown in Figure 1. Hence, Theorem 2 and the intuition from Figure 1 imply that seeking a flat minimum of ERM will lead to a better domain generalization gap.

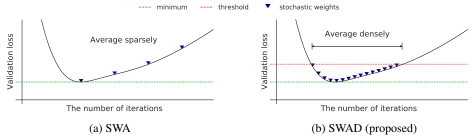


Figure 2: Comparison between SWA and SWAD. (a) SWA collects stochastic weights for every K epochs from the pre-defined K_0 epochs to the final epoch. (b) Our SWAD collects stochastic weights *densely*, *i.e.*, for every iteration, to obtain sufficiently many weights. SWAD collects the weights from the start iteration t_s to the end iteration t_e , where t_s and t_e are obtained by monitoring the validation loss (*overfit-aware* scheduling).

3 SWAD: Domain Generalization by Seeking Flat Minima

We have shown that flat minima will bring a better domain generalization. In this section, we propose Stochastic Weight Averaging Densely (SWAD) algorithm, and provide empirical quantitative and qualitative analyses on SWAD and flatness to understand why SWAD works better than ERM.

3.1 Dense and overfit-aware stochastic weight sampling strategy (SWAD)

SWA [25] is a simple weight ensemble approach to find flat minima. More specifically, SWA updates a pretrained model (namely, a model trained with sufficiently enough training epochs, K_0) with a cyclical [40] or high constant learning rate scheduling. SWA gathers model parameters for every K epochs during the update and averages them for the model ensemble. Intuitively, SWA finds a solution that is an ensemble of different local optima found by a sufficiently large learning rate to escape a local minimum. We illustrate an overview of SWA in Figure 2a. SWA has advantages over other flatness-aware methods [26, 41] in that it does not change the objective function to penalize sharp minima and does not require any additional computation for sharpness related quantities.

Despite its advantages, directly applying SWA to DG task has two problems. First, SWA averages a few weights (usually less than ten) by sampling weights for every K epochs, results in an inaccurate approximation of flat minima on a high-dimensional parameter space (e.g., 23M for ResNet-50 [42]). Furthermore, a common DG benchmark protocol uses relatively small training epochs (e.g., Gulrajani and Lopez-Paz [22] trained with less than two epochs for DomainNet benchmark), resulting in insufficient stochastic weights for SWA. From this motivation, we propose a "dense" sampling strategy for gathering sufficiently enough stochastic weights.

In addition, widely used DG datasets, such as PACS (\approx 10K images, 7 classes) and VLCS (\approx 11K images, 5 classes), are relatively smaller than large-scale datasets, such as ImageNet [43] (\approx 1.2M images, 1K classes). In this case, we observe that a simple ERM approach is rapidly reached to a local optimum only within a few epochs, and easily suffers from the overfitting issue, *i.e.*, the validation loss is increased after a few training epochs. It implies that directly applying the vanilla SWA will suffer from the overfitting issue by averaging sub-optimal solutions (*i.e.*, overfitted parameters). Hence, we need an "overfit-aware" sampling scheduling to omit the sub-optimal solutions for SWA.

The main idea of Stochastic Weight Averaging Densely (SWAD) is a dense and overfit-aware stochastic weight gathering strategy. First, instead of collecting weights for every K epochs, SWAD collects weights for every iteration. This dense sampling strategy easily collects sufficiently many weights than the sparse one. We also employ overfit-aware sampling scheduling by considering traces of the validation loss. Instead of sampling weights from K_0 pretraining epochs to the final epoch, we search the start iteration (when the validation loss achieves a local optimum for the first time) and the end iteration (when the validation loss is no longer decreased, but keep increasing). More specifically, we introduce three parameters: an optimum patient parameter N_s , an overfitting patient parameter N_e , and the tolerance rate r for searching the start iteration t_s and the end iteration t_e . First, we search t_s which satisfies $\mathcal{E}_{\text{val}}^{(t_s)} = \min_{i \in [0, \dots, N_s - 1]} \mathcal{E}_{\text{val}}^{(t_s + i)}$, where $\mathcal{E}_{\text{val}}^{(i)}$ denotes the validation loss at iterations. Simply, t_s is the first iteration where the loss value is no longer decreased during N_s iterations. Then, we find t_e satisfying $\min_{i \in [0, 1, \dots, N_e - 1]} \mathcal{E}_{\text{val}}^{(t_e + i)} > r \mathcal{E}_{\text{val}}^{(t_s)}$. In other words, t_e is the first iteration where the validation loss values exceed the tolerance r during N_e iterations.

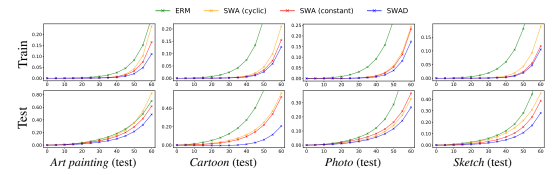


Figure 3: Local flatness comparisons. We plot the local flatness via loss gap, i.e., $\mathcal{F}_{\gamma}(\theta) = \mathbb{E}_{\|\theta'\|=\|\theta\|+\gamma}[\mathcal{E}(\theta')-\mathcal{E}(\theta)]$, of ERM, SWA, and SWAD by varying radius γ on different domains of PACS dataset. For each figure, Y-axis indicates the flatness $\mathcal{F}_{\gamma}(\theta)$ and X-axis indicates the radius scale $\|\gamma\|$. We measure the train flatness on seen domains $\mathcal{F}_{\gamma}^{\mathcal{T}}(\theta)$ in the first row and the test flatness on unseen domain $\mathcal{F}_{\gamma}^{\mathcal{T}}(\theta)$ in the second row. Each point is computed by Monte-Carlo approximation with 100 random samples. This comparisons show SWAD finds flatter minima than not only ERM but also SWA, in both train and test domains.

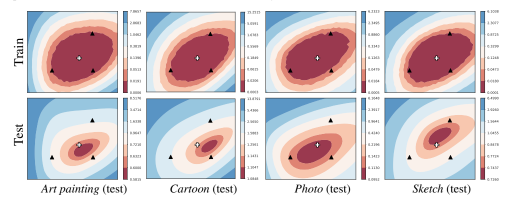


Figure 4: Loss surfaces on model parameters in PACS dataset for each target domain. The three triangles indicate model weights chosen at the end of training phase with equal intervals. Each plane is defined by the three weights and losses upon the plane are visualized with contours. The center cross mark is averaged point of the three weights. The first and second rows show the averaged training loss and the test loss surfaces, respectively.

We illustrate the overview of SWAD and the comparison of SWAD to SWA in Figure 2. Detailed pseudo code is provided in Appendix. We compare SWAD with other possible SWA strategies in §4.3 and show that our design choice works better for DG tasks.

3.2 Empirical analysis of SWAD and flatness

Here, we analyze solutions found by SWAD in terms of flatness. We first verify that the SWAD solution is flatter than those of ERM and SWA. Our loss surface visualization shows that the SWAD solution is located on the center of the flat minimum, while ERM finds a boundary solution. Finally, we show that the sharp boundary solutions by ERM are not generalized well, resulting in sensitivity to the model selection. All following empirical analyses are conducted on PACS dataset, validating by all four domains (art painting, cartoon, photo, and sketch).

Local flatness anaylsis. To begin with, we quantify the local flatness of a model parameter θ by assuming that flat minima will have smaller changes of loss value within its neighborhoods than sharp minima. For the given model parameter θ , we compute the expected loss value changes between θ and parameters on the sphere surrounding θ with radius γ , i.e., $\mathcal{F}_{\gamma}(\theta) = \mathbb{E}_{\|\theta'\|=\|\theta\|+\gamma}[\mathcal{E}(\theta')-\mathcal{E}(\theta)]$. In practice, $\mathcal{F}_{\gamma}(\theta)$ is approximated by Monte-Carlo sampling with 100 samples. Note that the proposed local flatness $\mathcal{F}_{\gamma}(\theta)$ is computationally efficient than measuring curvature using the Hessian-based quantities. Also, $\mathcal{F}_{\gamma}(\theta)$ has an unbiased finite sample estimator, while the worst-case loss value, i.e., $\max_{\|\theta'\|=\|\theta\|+\gamma}[\mathcal{E}(\theta')-\mathcal{E}(\theta)]$ has no unbiased finite sample estimator.

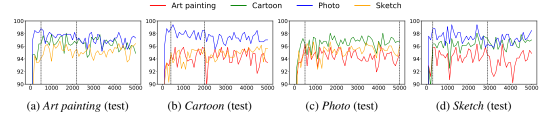


Figure 5: Validation accuracies for in-domains. The X- and Y-axis indicate the training iterations and accuracy, respectively, about the validation domains (legend) and the test domain (caption). The vertical dot lines represent start and end iterations, t_s and t_e , identified by the overfit-aware sampling strategy of SWAD.

In Figure 3, we compare $\mathcal{F}_{\gamma}(\theta)$ of ERM, SWA with cyclic learning rate, SWA with constant learning rate, and SWAD by varying radius γ . The solution found by SWA and SWAD shows lower local flatness quantities in all experiments. We also observe that our dense and overfit-aware sampling strategy, *i.e.*, SWAD, finds flatter optima than SWA does.

Loss surface visualization. We visualize the loss landscapes by choosing three model weights on the optimization trajectory $(\theta_1, \theta_2, \theta_3)^2$, and computing the loss values by linear combinations of $\theta_1, \theta_2, \theta_3^3$ as [25]. More details are in Appendix. In Figure 4, we observe that for all cases, ERM solutions are located at the boundary of a flat minimum of training loss, resulting in poor generalizability in test domains, that is aligned with our theoretical analysis and empirical flatness analysis. Since ERM solutions are located on the boundary of a flat loss surface, we observe that ERM solutions are very sensitive to model selection. In Figure 5, we illustrate the validation accuracies for each train-test domain combination of PACS by ERM, over training iterations (one epoch is equivalent to 83 iterations). We first observe that ERM rapidly reaches the best accuracy within only a few training epochs, namely less than 6 epochs. Furthermore, the ERM validation accuracies fluctuate a lot, and the final performance is very sensitive to the model selection criterion.

On the other hand, we observe that SWA solutions are located on the center of the training loss surfaces as well as of the test loss surfaces (Figure 4). Also, our overfit-aware stochastic weight gathering strategy (denoted as the vertical dot lines in Figure 5) prevents the ensembled weight from overfitting and makes SWAD model selection-free.

4 Experiments

4.1 Evaluation protocols

Dataset and optimization protocol. Following Gulrajani and Lopez-Paz [22], we exhaustively evaluate our method and comparison methods on various benchmarks: PACS [7] (9,991 images, 7 classes, and 4 domains), VLCS [30] (10,729 images, 5 classes, and 4 domains), OfficeHome [31] (15,588 images, 65 classes, and 4 domains), TerraIncognita [32] (24,788 images, 10 classes, and 4 domains), and DomainNet [33] (586,575 images, 345 classes, and 6 domains).

For a fair comparison, we follow training and evaluation protocol by Gulrajani and Lopez-Paz [22], including the dataset splits, hyperparameter (HP) search and model selection (while SWAD does not need it) on the validation set, and optimizer HP, except the HP search space and the number of iterations for DomainNet. We use a reduced HP search space to reduce the computational costs. We also tripled the number of iterations for DomainNet from 5,000 to 15,000 because we observe that 5,000 is not sufficient to convergence. We re-evaluate ERM with 15,000 iterations, and observe 3.1pp average performance improvement $(40.9\% \rightarrow 44.0\%)$ in DomainNet. For training, we choose a domain as the target domain and use the remaining domains as the training domain where 20% samples are used for validation and model selection. ImageNet [43] trained ResNet-50 [42] is employed as the initial weight, and optimized by Adam [39] optimizer with a learning rate of 5e-5. We construct a mini-batch containing all domains where each domain has 32 images. We set SWAD

²We choose weights at iteration 2500, 3500, 4500 during the training.

³Each point is defined by two axes u and v computed by $u = \theta_2 - \theta_1$ and $v = \frac{(\theta_3 - \theta_1) - (\theta_3 - \theta_1, \theta_2 - \theta_1)}{\|\theta_2 - \theta_1\|^2 \cdot (\theta_2 - \theta_1)}$.

Table 2: Comparison with domain generalization methods and SWAD. Out-of-domain accuracies on five domain generalization benchmarks are shown. We highlight the **best results** and the <u>second best results</u>. Note that ERM (reproduced), Mixstyle are reproduced numbers, and other numbers are from the original literature and Gulrajani and Lopez-Paz [22] (denoted with †). Our experiments are repeated three times.

Algorithm	PACS	VLCS	OfficeHome	TerraInc	DomainNet	Avg.
MASF [14]	82.7	-	-	-	-	-
DMG [38]	83.4	-	-	-	43.6	-
MetaReg [15]	83.6	-	-	-	43.6	-
ER [12]	85.3	-	-	-	-	-
pAdaIN [44]	85.4	-	-	-	-	-
EISNet [45]	85.8	-	-	-	-	-
DSON [35]	<u>86.6</u>	-	-	-	-	-
ERM [†] [34]	85.5	77.5	66.5	46.1	40.9	63.3
ERM (reproduced)	84.2	77.3	67.6	47.8	<u>44.0</u>	64.2
IRM^{\dagger} [20]	83.5	78.6	64.3	47.6	33.9	61.6
GroupDRO [†] [46]	84.4	76.7	66.0	43.2	33.3	60.7
I-Mixup [†] [47–49]	84.6	77.4	68.1	47.9	39.2	63.4
MLDG [†] [13]	84.9	77.2	66.8	47.8	41.2	63.6
CORAL [†] [36]	86.2	<u>78.8</u>	<u>68.7</u>	47.7	41.5	64.5
MMD^{\dagger} [50]	84.7	77.5	66.4	42.2	23.4	58.8
DANN [†] [9]	83.7	78.6	65.9	46.7	38.3	62.6
$CDANN^{\dagger}$ [10]	82.6	77.5	65.7	45.8	38.3	62.0
MTL [†] [51]	84.6	77.2	66.4	45.6	40.6	62.9
SagNet [†] [37]	86.3	77.8	68.1	<u>48.6</u>	40.3	64.2
ARM [†] [16]	85.1	77.6	64.8	45.5	35.5	61.7
VREx [†] [21]	84.9	78.3	66.4	46.4	33.6	61.9
RSC [†] [52]	85.2	77.1	65.5	46.6	38.9	62.7
Mixstyle [17]	85.2	77.9	60.4	44.0	34.0	60.3
SWAD (ours)	88.1 (±0.4)	79.1 (±0.4)	70.6 (±0.3)	50.0 (±0.4)	46.5 (±0.2)	66.9

HPs N_s to 3, N_e to 6, and r to 1.2 for VLCS and 1.3 for the others by HP search on the validation sets. Additional dataset details and implementation details, such as other HPs, are given in Appendix.

Evaluation metrics. We report out-of-domain accuracies for each domain and their average, *i.e.*, a model is trained and validated on training domains and evaluated on the unseen target domain. Each out-of-domain performance is an average of three different runs with different train-validation splits.

4.2 Main results

Comparison with domain generalization methods. We report the full out-of-domain performances on five DG benchmarks in Table 2. The full tables including outof-domain accuracies for each domain are in Appendix. In all experiments, our SWAD achieves significant performance gain against ERM as well as the previous best results: +2.6pp in PACS, +0.3pp in VLCS, +1.4pp in TerraIncognita, +1.9pp in OfficeHome, and +2.9pp in DomainNet comparing to the previous best results. We observe that SWAD provides two practical advantages comparing to previous methods. First, SWAD does not need any modification on training objectives or model architecture, i.e., it is universally applicable to any other methods. As an example, we show that SWAD actually improves the performances of other DG methods, such as CORAL [36] in Table 4. Moreover, as we discussed before, SWAD is free to the model selection, resulting in

Table 3: Comparison between generalization methods on PACS. The scores are averaged over all settings using different target domains. (†) and (\downarrow) indicate statistically significant improvement and degradation from ERM.

	Out-of-domain	In-domain
ERM	85.3±0.4	96.6±0.0
EMA	85.5±0.4(-)	97.0±0.1(↑)
SAM	$85.5 \pm 0.1(-)$	97.4±0.1(†)
Mixup	84.8±0.3(-)	97.3±0.1(↑)
CutMix	$83.8 \pm 0.4 (\downarrow)$	97.6±0.1(↑)
VAT	85.4 ± 0.6 (-)	$96.9 \pm 0.2 (\uparrow)$
Π -model	$83.5 \pm 0.5 (\downarrow)$	96.8±0.2(†)
SWA	85.9±0.1(†)	97.1±0.1(†)
SWAD	87.1 ±0.2(↑)	97.7 ±0.1(†)

stable performances (i.e., small standard errors) on various benchmarks. Note that we only compare

Table 4: **Application of SWAD.** The scores are averaged over every target domain case. The performances of ERM, CORAL, and SAM are optimized by HP searches of DomainBed. In contrast, for the combinations, SWAD is simply applied without any HP search. Note that ERM + SWAD is same as "SWAD" in Table 2.

	PACS	VLCS	OfficeHome	TerraInc	DomainNet	Avg. (Δ)
ERM ERM + SWAD	85.5 ± 0.2 88.1 ± 0.1	77.5 ± 0.4 79.1 ± 0.1	66.5 ± 0.3 70.6 ± 0.2	$46.1 \pm 1.8 \\ 50.0 \pm 0.3$	$40.9 \pm 0.1 \\ 46.5 \pm 0.1$	63.3 66.9 (+3.6)
CORAL CORAL + SWAD	$86.2 \pm 0.3 \\ 88.3 \pm 0.1$	$78.8 \pm 0.6 \\ 78.9 \pm 0.1$	$68.7 \pm 0.3 \\ 71.3 \pm 0.1$	$47.6 \pm 1.0 \\ 51.0 \pm 0.1$	$41.5 \pm 0.1 \\ 46.8 \pm 0.0$	64.5 67.3 (+2.8)
SAM SAM + SWAD	85.8 ± 0.2 87.1 ± 0.2	79.4 ± 0.1 78.5 ± 0.2	$69.6 \pm 0.1 \\ 69.9 \pm 0.1$	43.3 ± 0.7 45.3 ± 0.9	44.3 ± 0.0 46.5 ± 0.1	64.5 65.5 (+1.0)

results with ResNet-50 backbone for a fair comparison. We describe the implementation details of each comparison method and the hyperparameter search protocol in Appendix.

Comparison with conventional generalization methods. We also compare SWAD with other conventional generalization methods to show that the remarkable domain generalization gaps by SWAD is not achieved by better generalization, but by seeking flat minima. The comparison methods include flatness-aware optimization methods, such as SAM [26], ensemble methods, such as EMA [53], data augmentation methods, such as Mixup [54] and CutMix [55], and consistency regularization methods, such as VAT [56] and Π -model [57]. We also split in-domain datasets into training (60%), validation (20%), and test (20%) splits, while no in-domain test set used for Table 2. Every experiment is repeated three times.

The results are shown in Table 3. We observe that all conventional methods helps in-domain generalization, *i.e.*, performing better than ERM on in-domain test set. However, their out-of-domain performances are similar to or even worse than ERM. For example, CutMix and II-model improve in-domain performances by 1.0pp and 0.2pp but degrade out-of-domain performances by 1.5pp and 1.8pp. SAM, another method for seeking flat minima, slightly increases both in-domain and out-of-domain performances but the out-of-domain performance is not statistically significant. We will discuss performances of SAM in other benchmarks later. In contrast, the vanilla SWA and our SWAD significantly improve both in-domain and out-of-domain performances. SWAD improves the performances by SWA with statistically significantly gaps: 1.2pp on the out-of-domain and 0.6pp on the in-domain. Further comparison between SWA and SWAD is provided in §4.3.

Combinations with other methods. Since SWAD does not require any modification on training procedures and model architectures, SWAD is universally applicable to any other methods. Here, we combine SWAD with ERM, CORAL [36], and SAM [26]. Results are shown in Table 4. Both CORAL and SAM solely show better performances than ERM with +1.2pp average out-of-domain accuracy gap. Note that SAM is not a DG method but a sharpness-aware optimization method to find flat minima. It supports our theoretical motivation: DG can be achieved by seeking flat minima.

By applying SWAD on the baselines, the performances are consistently improved by 3.6pp on ERM, 2.8pp on CORAL, and 1.0pp on SAM. Interestingly, CORAL + SWAD show the best performances with both incorporating different advantages of utilizing domain labels and seeking flat minima. We also observe that SAM + SWAD shows worse performance than ERM + SWAD, while SAM performs better than ERM. We conjecture that it is because the objective control by SAM restricts the model parameter diversity durinig training, reducing the diversity for SWA ensemble. However, applying SWAD on SAM still leads to better performances than the sole SAM. The results demonstrate that the application of SWAD on other baselines is a simple yet effective method for DG.

4.3 Ablation study

Table 5 provides ablative studies on the starting and ending iterations for averaging, the learning rate schedule, and the sampling interval. $SWA_{w/\,cyclic}$ (SWA in Table 3) and $SWA_{w/\,constant}$ are vanilla SWAs with fixed sampling positions. We also report SWAD by eliminating three factors: the dense sampling strategy, and searching the start iteration, searching the end iteration. The dense sampling strategy lets SWAD estimate a more accurate approximation of flat minima: showing 0.8pp

Table 5: Ablation studies of the stochastic weights selection strategies on PACS and VLCS. In the configuration, " t_s ", " t_e ", "lr", and "interval" indicate start and end iterations of sampling, a learning rate schedule, and a stochastic weight sampling interval, respectively. "Opt" and "Overfit" indicate the start and end iterations identified by our overfit-aware sampling strategy, and "Val" means the start and end iterations whose averaging shows the best accuracy on the validation set. "Cyclic" and "Const" represent cyclic and constant learning rate schedules. All experiments are repeated three times.

	Configuration			Out-of-domain			In-domain			
	t_s	t_e	lr	interval	PACS	VLCS	Avg.	PACS	VLCS	Avg.
SWA _{w/ cyclic}	4000	5000	Cyclic	100	85.9 ±0.1	76.6 ± 0.1	81.2	97.1 ±0.1	85.0 ± 0.2	91.0
SWA _{w/const}	4000	5000	Const	100	86.5 ± 0.3	76.7 ± 0.2	81.6	97.3 ± 0.1	85.0 ± 0.2	91.1
$SWAD_{w/o\ Dense}$	Opt	Overfit	Const	100	86.5 ± 0.4	78.0 ± 0.7	82.2	97.6 ± 0.1	85.8 ± 0.4	91.7
SWAD _{w/o Opt-Overfit}	4000	5000	Const	1	86.6 ± 0.6	76.9 ± 0.3	81.7	97.5 ± 0.1	85.2 ± 0.1	91.3
SWAD _{w/o Overfit}	Opt	5000	Const	1	87.1 ±0.3	77.6 ± 0.1	82.4	97.7 ±0.1	$85.8 \pm \hspace{-0.05cm}\pm \hspace{-0.05cm}0.3$	91.8
SWAD _{fit-on-val}	Val	Val	Const	1	86.2 ± 0.2	78.6 ± 0.1	82.4	97.5 ± 0.2	$85.8 \pm \hspace{-0.05cm} \pm \hspace{-0.05cm} 0.3$	91.7
SWAD (proposed)	Opt	Overfit	Const	1	87.1 ±0.2	$\textbf{78.9} \pm 0.2$	83.0	97.7 ±0.1	$\textbf{86.1} \pm 0.5$	91.9

degeneration in the average out-of-domain accuracy (SWAD_{w/o Dense}). When we take an average from t_s to the final iteration, the out-of-domain performance degrades by 0.6pp (SWAD_{w/o Overfit}). Similarly, a fixed scheduling without the overfit-aware scheduling only shows very marginal improvements from the vanilla SWA (SWAD_{w/o Opt-Overfit}). We also evaluate SWAD_{fit-on-val} that uses the range achieving the best performances on the validation set, but it becomes overfitted to the validation, results in lower performances than SWAD. The results demonstrate the benefits of combining "dense" and "overfit-aware" sampling strategies of SWAD.

5 Discussion and Limitations

Despite many benefits from SWAD, such as the significant performance improvements, model selection-free property, working plug-and-play manner for various methods, there are some potential limitations. Here, we discuss the limitations of SWAD for further improvements.

Confidence error in Theorem 1. While the confidence error in Theorem 1 tells the effect of γ on generalization error bound, there exists a limitation in that the confidence error term shows improper behavior with respect to γ if γ is close to zero. The behavior we expect is that the confidence error of RRM converges to the confidence error of ERM as γ decreases to zero, however, the current theorem does not show such tendency since the confidence bound diverges to infinity when γ goes to zero. However, we would like to note that this limitation is not a drawback of RRM, but it is caused by the looseness of the union bound which is a mathematical technique used to derive the confidence error of RRM. Our RRM formulation has a similarity to previous works [26, 29] and we note that the counter-intuitive behavior of the confidence bound and γ also appears in Foret et al. [26].

SWAD is not a perfect flatness-aware optimization method. Note that SWAD is not a perfect and theoretically guaranteed solver for flat minima, but a heuristic approximation with empirical benefits. However, even if a better flatness-aware optimization method is proposed, our theoretical contribution still holds: showing the relationship between flat minima and DG.

SWAD does not strongly utilize domain-specific information. In Theorem 2, the domain generalization gap is bounded by three factors: flat minima, domain discrepancy, and confidence bound. Most of the existing approaches focus on domain discrepancy, reducing the difference between the source domains and the target domain by domain invariant learning [8–12]. SWAD focuses on the first factor, the flat minima. While the domain labels are used to construct a mini-batch, SWAD does not strongly utilize domain-specific information. It implies that if one can consider both flatness and domain discrepancy, better domain generalization can be achievable. Table 4 gives us a clue: the combination of CORAL (utilizing domain-specific information) and SWAD (seeking flat minima) shows the best performance among all comparison methods. As a future research direction, we encourage studying a method that can achieve both flat optima and small domain discrepancy.

6 Concluding Remarks

In this paper, we theoretically and empirically demonstrate that DG is achievable by seeking flat minima. We propose SWAD that collects a number of valid weights and captures flatter minima than the vanilla SWA does. The extensive experiments on five benchmarks show superior performances of SWAD comparing to existing DG methods. Comparison with conventional generalization methods supports that the significant improvement by SWAD is from seeking flat minima, not from better generalization performances. In addition, we show that combinations of SWAD and other DG methods can achieve state-of-the-art performances on the DG benchmarks.

Acknowledgement

NAVER Smart Machine Learning (NSML) [58] platform has been used in experiments.

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A Potential Societal Impacts

In this study, we theoretically and empirically demonstrate that domain generalization (DG) is achievable by seeking flat minima, and propose SWAD to find flat minima. With SWAD, researchers and developers can develop a model robust to domain shift, a realistic deployment environment, without relying on a task-dependent prior, a modified objective function, or a specific model architecture. Accordingly, SWAD has potential positive impacts by developing machines less biased towards ethical aspects, as well as potential negative impacts, *e.g.*, improving weapon or surveillance systems under unexpected environment changes.

B Implementation Details

B.1 Hyperparameters of SWAD

The evaluation protocol by Gulrajani and Lopez-Paz [22] is computationally too expensive; it requires about 4,142 models for every DG algorithm. Hence, we reduce the search space of SWAD for computational efficiency; batch size, learning rate are fixed to 32 for each domain and 5e-5, respectively. We set dropout probability and weight decay to zero. We only search N_s , N_e and r. Here, N_s and N_e are searched in PACS dataset, and the searched values are used for all experiments, while r is searched in [1.2, 1.3] depending on dataset. As a result, we use $N_s=3$, $N_e=6$, and r=1.2 for VLCS and r=1.3 for the others. We initialize our model by ImageNet-pretrained ResNet-50 and batch normalization statistics are frozen during training. The number of total iterations is 15,000 for DomainNet and 5,000 for others, which are sufficient numbers to be converged. Finally, we slightly modify the evaluation frequency because it should be set to small enough to detect the moments that the model is optimized and overfitted. However, too small frequency brings large evaluation overhead, thus we compromise between exactness and efficiency: 50 for VLCS, 500 for DomainNet, and 100 for others.

B.2 Hyperparameter search protocol for reproduced results

Table 6: **Hyperparameter search space comparison.** U and list indicate Uniform distribution and random choice, respectively.

Parameter	Default value	DomainBed	Ours
batch size	32	2 ^{U(3,5.5)}	32
learning rate	5e-5	$10^{\text{U}(-5,-3.5)}$	[1e-5, 3e-5, 5e-5]
ResNet dropout	0	[0.0, 0.1, 0.5]	[0.0, 0.1, 0.5]
weight decay	0	$10^{\text{U}(-6,-2)}$	[1e-4, 1e-6]

We evaluate recently proposed methods, SAM [26] and Mixstyle [17], and compare them with previous results. For a fair comparison, we follow the hyperparameter (HP) search protocol proposed by Gulrajani and Lopez-Paz [22], with a modification to reduce computational resources. They searched HP by training a total of 58,000 models, corresponding to about 4,142 runs for each algorithm. It is too much computational burden to train 4,142 models whenever evaluate a new algorithm. Therefore, we re-design the HP search protocol efficiently and effectively. In the HP search protocol of DomainBed [22], training domains and algorithm-specific parameters are included in the HP search space, and HP is found for every data split independently by random search. Instead, we do not sample training domains, use HP found in the first data split to the other splits, search algorithm-specific HP independently, and conduct grid search on the more effectively designed HP space as shown in Table 6. Through the proposed protocol, we find HP for an algorithm under only 396 runs. Although the number of total runs is reduced to about 10% (4, $142 \rightarrow 396$), the results of reproduced ERM is improved 0.9pp in average (63.3% \rightarrow 64.2%). It demonstrates both the effectiveness and the efficiency of our search protocol.

B.3 Algorithm-specific hyperparameters

We search the algorithm-specific hyperparameters independently in PACS dataset, based on the values suggested from each paper. For Mixstyle [17], we insert Mixstyle block with domain label after the 1st, 2nd, and 3rd residual blocks with $\alpha=0.1$ and p=0.5. We train SAM [26] with $\rho=0.05$, and VAT [56] with $\epsilon=1.0$ and $\alpha=1.0$. In Π -model [57], $w_{max}=1$ is chosen among various w_{max} values such as 1, 10, 100, and 300. We use EMA [53] with decay=0.99, Mixup [54] with $\alpha=0.2$, and CutMix [55] with $\alpha=1.0$ and p=0.5.

B.4 Pseudo code

Algorithm 1: Stochastic Weight Averaging Densely

B.5 Loss surface visualization

Following Garipov et al. [24], we choose three model weights $\theta_1, \theta_2, \theta_3$ and define two dimensional weight plane from the weights:

$$u = \theta_2 - \theta_1, \qquad v = \frac{(\theta_3 - \theta_1) - \langle \theta_3 - \theta_1, \theta_2 - \theta_1 \rangle}{\|\theta_2 - \theta_1\|^2 \cdot (\theta_2 - \theta_1)},$$
 (3)

where $\hat{u} = u/\|u\|$ and $\hat{v} = v/\|v\|$ are orthonormal bases of the weight plane. Then, we build Cartesian grid near the weights on the plane. For each grid point, we calculate the weight corresponding to the point and compute loss from the weight. The results are visualized as a contour plot, as shown in Figure 4 in the main text.

C Proof of Theorems

C.1 Technical Lemmas

Consider an instance loss function $\ell(y_1,y_2)$ such that $\ell:\mathcal{Y}\times\mathcal{Y}\to[0,1]$ and $\ell(y_1,y_2)=0$ if and only if $y_1=y_2$. Then, we can define a functional error as $\mathcal{E}_{\mathcal{P}}(f(\cdot;\theta),h):=\mathbb{E}_{\mathcal{P}}[\ell(f(x;\theta),h(x))]$. Note that if we set h as a true label function which generates the label of inputs, y=h(x), then, it becomes a population loss $\mathcal{E}_{\mathcal{P}}(\theta)=\mathcal{E}_{\mathcal{P}}(f(\cdot;\theta),h)$. Given two distributions, \mathcal{P} and \mathcal{Q} , the following lemma shows that the difference between the error with \mathcal{P} and the error with \mathcal{Q} is bounded by the divergence between \mathcal{P} and \mathcal{Q} .

Lemma 1. $|\mathcal{E}_{\mathcal{P}}(h_1, h_2) - \mathcal{E}_{\mathcal{Q}}(h_1, h_2)| \leq \frac{1}{2} \mathbf{Div}(\mathcal{P}, \mathcal{Q})$

Proof. We employ the same technique in Zhao et al. [59] for our loss function ℓ . From the Fubini's theorem, we have,

$$\mathbb{E}_{x \sim \mathcal{P}}[\ell(h_1(x), h_2(x))] = \int_0^\infty \mathbb{P}_{\mathcal{P}}\left(\ell(h_1(x), h_2(x)) > t\right) dt \tag{4}$$

By using this fact,

$$\left| \mathbb{E}_{x \sim \mathcal{P}} \left[\ell(h_1(x), h_2(x)) \right] - \mathbb{E}_{x \sim \mathcal{Q}} \left[\ell(h_1(x), h_2(x)) \right] \right| \tag{5}$$

$$= \left| \int_0^\infty \mathbb{P}_{\mathcal{P}} \left(\ell(h_1(x), h_2(x)) > t \right) dt - \int_0^\infty \mathbb{P}_{\mathcal{Q}} \left(\ell(h_1(x), h_2(x)) > t \right) dt \right| \tag{6}$$

$$\leq \int_0^\infty |\mathbb{P}_{\mathcal{P}}\left(\ell(h_1(x), h_2(x)) > t\right) - \mathbb{P}_{\mathcal{Q}}\left(\ell(h_1(x), h_2(x)) > t\right)| dt \tag{7}$$

$$\leq M \sup_{t \in [0,M]} |\mathbb{P}_{\mathcal{P}} \left(\ell(h_1(x), h_2(x)) > t \right) - \mathbb{P}_{\mathcal{Q}} \left(\ell(h_1(x), h_2(x)) > t \right) | \tag{8}$$

$$\leq M \sup_{h_1, h_2} \sup_{t \in [0, M]} |\mathbb{P}_{\mathcal{P}} \left(\ell(h_1(x), h_2(x)) > t \right) - \mathbb{P}_{\mathcal{Q}} \left(\ell(h_1(x), h_2(x)) > t \right)| \tag{9}$$

$$\leq M \sup_{\bar{h} \in \bar{\mathcal{H}}} \left| \mathbb{P}_{\mathcal{P}} \left(\bar{h}(x) = 1 \right) - \mathbb{P}_{\mathcal{Q}} \left(\bar{h}(x) = 1 \right) \right| \tag{10}$$

$$\leq M \sup_{A} |\mathbb{P}_{\mathcal{P}}(A) - \mathbb{P}_{\mathcal{Q}}(A)| \tag{11}$$

where
$$\bar{\mathcal{H}} := \{ \mathbb{I}[\ell(h(x), h'(x)) > t] | h, h' \in \mathcal{H}, t \in [0, M] \}.$$

Lemma 2. Consider a distribution S on input space and global label function $f: \mathcal{X} \to \mathcal{Y}$. Let $\{\Theta_k \subset \mathbb{R}^d, k = 1, \cdots, N\}$ be a finite cover of a parameter space Θ which consists of closed balls with radius $\gamma/2$ where $N := \left\lceil (diam(\Theta)/\gamma)^d \right\rceil$. Let $\theta_k \in \arg\max_{\Theta_k \cap \Theta} \mathcal{E}_S(\theta)$ be a local maximum in the k-th ball. Let a VC dimension of Θ_k be v_k . Then, for any $\theta \in \Theta$, the following bound holds with probability at least $1 - \delta$.

$$\mathcal{E}_{S}(\theta) - \hat{\mathcal{E}}_{S}^{\gamma}(\theta) \le \max_{k} \sqrt{\frac{\left(v_{k} \left[\ln\left(n/v_{k}\right) + 1\right] + \ln\left(N/\delta\right)\right)}{2n}}$$
(12)

where $\hat{\mathcal{E}}_{S}^{\gamma}(\theta_{k})$ is an empirical robust risk with n samples.

Proof. We first show that the following inequality holds for the local maximum of N covers,

$$\mathbb{P}\left(\max_{k} \left[\mathcal{E}_{S}(\theta_{k}) - \hat{\mathcal{E}}_{S}(\theta_{k})\right] > \epsilon\right) \leq \sum_{k=1}^{N} \mathbb{P}\left(\mathcal{E}_{S}(\theta_{k}) - \hat{\mathcal{E}}_{S}(\theta_{k}) > \epsilon\right)$$
(13)

$$\leq \sum_{k=1}^{N} \mathbb{P} \left(\sup_{\theta \in \Theta_k} \left[\mathcal{E}_S(\theta) - \hat{\mathcal{E}}_S(\theta) \right] > \epsilon \right)$$
 (14)

$$\leq \sum_{k=1}^{N} \left(\frac{en}{v_k}\right)^{v_k} e^{-2n\epsilon^2}.$$
 (15)

Now, we introduce a confidence error bound $\epsilon_k := \sqrt{\frac{(v_k[\ln(n/v_k)+1]+\ln(N/\delta))}{2n}}$. Then, we set $\epsilon := \max_k \epsilon_k$. Then, we get,

$$\mathbb{P}\left(\max_{k} \left[\mathcal{E}_{S}(\theta_{k}) - \hat{\mathcal{L}}_{S}(\theta_{k})\right] > \epsilon\right) \leq \sum_{k=1}^{N} \left(\frac{en}{v_{k}}\right)^{v_{k}} e^{-2n\epsilon^{2}}$$
(16)

$$\leq \sum_{k=1}^{N} \left(\frac{en}{v_k}\right)^{v_k} e^{-2n\epsilon_k^2} \tag{17}$$

$$=\sum_{k=1}^{N} \frac{\delta}{N} = \delta,\tag{18}$$

since $\epsilon > \sqrt{\frac{(v_k[\ln(n/v_k)+1] + \ln(N/\delta))}{2n}}$ for all k. Hence, the inequality holds with probability at least $1-\delta$.

Based on this fact, let us consider the set of events such that $\max_k \left[\mathcal{E}_S(\theta_k) - \hat{\mathcal{E}}_S(\theta_k) \right] \leq \epsilon$. Then, for any θ , there exists k' such that $\theta \in \Theta_{k'}$. Then, we get

$$\mathcal{E}_S(\theta) - \hat{\mathcal{E}}_S^{\gamma}(\theta) \le \mathcal{E}_S(\theta) - \hat{\mathcal{E}}_S(\theta_{k'}) \tag{19}$$

$$\leq \mathcal{E}_S(\theta) - \mathcal{E}_S(\theta_{k'}) + \epsilon \tag{20}$$

$$\leq \mathcal{E}_S(\theta_{k'}) - \mathcal{E}_S(\theta_{k'}) + \epsilon = \epsilon,$$
 (21)

where the second inequality holds since $\mathcal{E}_S(\theta_{k'}) - \hat{\mathcal{E}}_S(\theta_{k'}) \leq \max_k \left[\mathcal{E}_S(\theta_k) - \hat{\mathcal{E}}_S(\theta_k)\right] \leq \epsilon$ and the final inequality holds since $\theta_{k'}$ is the local maximum in $\Theta_{k'}$. In this regards, we know that $\max_k \left[\mathcal{E}_S(\theta_k) - \hat{\mathcal{E}}_S(\theta_k)\right] \leq \epsilon$ implies $\mathcal{E}_S(\theta) - \hat{\mathcal{E}}_S^{\gamma}(\theta) \leq \epsilon$. Consequently, $\mathcal{E}_S(\theta) - \hat{\mathcal{E}}_S^{\gamma}(\theta) \leq \epsilon$ holds with probability at least $1 - \delta$.

C.2 Proof of Theorem 1

Proof. The proof consists of two parts. First, we show that the following inequality holds with high probability.

$$\mathcal{E}_{\mathcal{T}}(\theta) \leq \hat{\mathcal{E}}_{\mathcal{S}}^{\gamma}(\theta) + \frac{1}{2} \mathbf{Div}\left(\mathcal{S}, \mathcal{T}\right) + \max_{k} \sqrt{\frac{\left(v_{k} \left[\ln\left(n/v_{k}\right) + 1\right] + \ln\left(N/\delta\right)\right)}{2n}}.$$

Then, secondly, we apply the inequality for multiple source domains.

The first part can be proven by simply combining Lemma 1 and Lemma 2. Then, we get,

$$\mathcal{E}_{\mathcal{T}}(\theta) \le \mathcal{E}_{\mathcal{S}}(\theta) + \frac{1}{2} \mathbf{Div}(\mathcal{S}, \mathcal{T})$$
 (22)

$$\leq \hat{\mathcal{E}}_{S}^{\gamma}(\theta) + \frac{1}{2} \mathbf{Div}\left(\mathcal{S}, \mathcal{T}\right) + \max_{k} \sqrt{\frac{\left(v_{k}\left[\ln\left(n/v_{k}\right) + 1\right] + \ln\left(N/\delta\right)\right)}{2n}}$$
(23)

where $\mathbf{Div}\left(\mathcal{S},\mathcal{T}\right)$ is a divergence between \mathcal{S} and \mathcal{T} .

For the second part, we set $\mathcal{D}:=\sum_{i=1}^I \mathcal{D}_i/I$ which is a mixture of source distributions. Then, by applying \mathcal{D} to the first part, we obtain the following inequality,

$$\mathcal{E}_{\mathcal{T}}(\theta) \leq \hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta) + \frac{1}{2} \mathbf{Div}\left(\mathcal{D}, \mathcal{T}\right) + \max_{k} \sqrt{\frac{\left(v_{k} \left[\ln\left(In/v_{k}\right) + 1\right] + \ln\left(N/\delta\right)\right)}{2In}}$$
(24)

$$\leq \hat{\mathcal{E}}_{\mathcal{D}}^{\gamma}(\theta) + \frac{1}{2I} \sum_{i=1}^{I} \mathbf{Div} \left(\mathcal{D}_{i}, \mathcal{T} \right) + \max_{k} \sqrt{\frac{\left(v_{k} \left[\ln \left(In / v_{k} \right) + 1 \right] + \ln \left(N / \delta \right) \right)}{2In}}$$
(25)

where the total number of training data set is In and, for the second inequality, we use the fact that $\frac{1}{2}\mathbf{Div}\left(\mathcal{D},\mathcal{T}\right)\leq\frac{1}{2I}\sum_{i=1}^{I}\mathbf{Div}\left(\mathcal{D}_{i},\mathcal{T}\right)$, which has been proven in [59].

C.3 Proof of Theorem 2

Proof. First, let $\bar{\theta} \in \arg \max_{\theta \in \Theta} \mathcal{E}_{\mathcal{T}}(\theta)$. Then, from generalization error bound of $\mathcal{E}_{\mathcal{D}}(\bar{\theta})$, the following inequality holds with probability at most $\frac{\delta}{2}$,

$$\hat{\mathcal{E}}_{\mathcal{D}}(\bar{\theta}) - \mathcal{E}_{\mathcal{D}}(\bar{\theta}) > \sqrt{\frac{v \ln(In/v) + \ln(2/\delta)}{In}},\tag{26}$$

where v is a VC dimension of Θ . Furthermore, from Theorem 1, we have the following inequality with probability at most $\frac{\delta}{2}$,

$$\mathcal{E}_{\mathcal{T}}(\hat{\theta}^{\gamma}) > \mathcal{E}_{\mathcal{D}}^{\gamma}(\hat{\theta}^{\gamma}) + \frac{1}{2}\mathbf{Div}(\mathcal{D}, \mathcal{T}) + \max_{k \in [1, N]} \sqrt{\frac{v_k \ln(In/v_k) + \ln(2N/\delta)}{In}}.$$
 (27)

Finally, let us consider the set of event such that $\hat{\mathcal{E}}_{\mathcal{D}}(\bar{\theta}) - \mathcal{E}_{\mathcal{D}}(\bar{\theta}) \leq \sqrt{\frac{v \ln(In/v) + \ln(2/\delta)}{In}}$ and $\mathcal{E}_{\mathcal{T}}(\hat{\theta}^{\gamma}) \leq \mathcal{E}_{\mathcal{D}}^{\gamma}(\hat{\theta}^{\gamma}) + \frac{1}{2} \mathbf{Div}(\mathcal{D}, \mathcal{T}) + \max_{k \in [1,N]} \sqrt{\frac{v_k \ln(In/v_k) + \ln(2N/\delta)}{In}}$ whose probability is at least greater than $1 - \delta$. Then, under this set of event, we have,

$$\min_{\theta'} \hat{\mathcal{E}}_{\mathcal{D}}(\theta') \le \hat{\mathcal{E}}_{\mathcal{D}}(\bar{\theta}) \le \mathcal{E}_{\mathcal{D}}(\bar{\theta}) + \sqrt{\frac{v \ln (In/v) + \ln (2/\delta)}{In}}$$
(28)

$$\leq \mathcal{E}_{\mathcal{T}}(\bar{\theta}) + \frac{1}{2}\mathbf{Div}(\mathcal{D}, \mathcal{T}) + \sqrt{\frac{v \ln(In/v) + \ln(2/\delta)}{In}}$$
(29)

$$\leq \min_{\theta'} \mathcal{E}_{\mathcal{T}}(\theta') + \frac{1}{2} \mathbf{Div}(\mathcal{D}, \mathcal{T}) + \sqrt{\frac{v \ln (In/v) + \ln (2/\delta)}{In}}$$
(30)

Consequently, we have,

$$\mathcal{E}_{\mathcal{T}}(\hat{\theta}^{\gamma}) - \min_{\theta'} \mathcal{E}_{\mathcal{T}}(\theta')$$

$$\leq \mathcal{E}_{\mathcal{D}}^{\gamma}(\hat{\theta}^{\gamma}) - \min_{\theta'} \hat{\mathcal{E}}_{\mathcal{D}}(\theta') + \mathbf{Div}(\mathcal{D}, \mathcal{T}) + \max_{k \in [1, N]} \sqrt{\frac{v_{k} \ln{(In/v_{k})} + \ln{(2N/\delta)}}{In}}$$

$$+ \sqrt{\frac{v \ln{(In/v)} + \ln{(2/\delta)}}{In}}$$

$$\leq \mathcal{E}_{\mathcal{D}}^{\gamma}(\hat{\theta}^{\gamma}) - \min_{\theta'} \hat{\mathcal{E}}_{\mathcal{D}}(\theta') + \frac{1}{I} \sum_{i=1}^{I} \mathbf{Div}(\mathcal{D}_{i}, \mathcal{T}) + \max_{k \in [1, N]} \sqrt{\frac{v_{k} \ln{(In/v_{k})} + \ln{(2N/\delta)}}{In}}$$

$$+ \sqrt{\frac{v \ln{(In/v)} + \ln{(2/\delta)}}{In}}$$

$$(32)$$

D Full Results

In this section, we show detailed results of Table 2 in the main text. † and ‡ indicate results from DomainBed's and our HP search protocols, respectively. Standard errors are reported from three trials, if available.

D.1 PACS

Table 7: Out-of-domain accuracies (%) on PACS.

Algorithm	A	С	P	S	Avg
$CDANN^{\dagger}$	84.6 ± 1.8	75.5 ± 0.9	96.8 ± 0.3	73.5 ± 0.6	82.6
MASF	82.9	80.5	95.0	72.3	82.7
DMG	82.6	78.1	94.5	78.3	83.4
IRM^{\dagger}	84.8 ± 1.3	76.4 ± 1.1	96.7 ± 0.6	76.1 ± 1.0	83.5
MetaReg	87.2	79.2	97.6	70.3	83.6
DANN^\dagger	86.4 ± 0.8	77.4 ± 0.8	97.3 ± 0.4	73.5 ± 2.3	83.7
ERM^{\ddagger}	85.7 ± 0.6	77.1 ± 0.8	97.4 ± 0.4	76.6 ± 0.7	84.2
GroupDRO†	83.5 ± 0.9	79.1 ± 0.6	96.7 ± 0.3	78.3 ± 2.0	84.4
MTL^\dagger	87.5 ± 0.8	77.1 ± 0.5	96.4 ± 0.8	77.3 ± 1.8	84.6
I-Mixup	86.1 ± 0.5	78.9 ± 0.8	97.6 ± 0.1	75.8 ± 1.8	84.6
MMD^\dagger	86.1 ± 1.4	79.4 ± 0.9	96.6 ± 0.2	76.5 ± 0.5	84.7
$VREx^{\dagger}$	86.0 ± 1.6	79.1 ± 0.6	96.9 ± 0.5	77.7 ± 1.7	84.9
MLDG^\dagger	85.5 ± 1.4	80.1 ± 1.7	97.4 ± 0.3	76.6 ± 1.1	84.9
ARM^\dagger	86.8 ± 0.6	76.8 ± 0.5	97.4 ± 0.3	79.3 ± 1.2	85.1
RSC^{\dagger}	85.4 ± 0.8	79.7 ± 1.8	97.6 ± 0.3	78.2 ± 1.2	85.2
Mixstyle [‡]	86.8 ± 0.5	79.0 ± 1.4	96.6 ± 0.1	$78.5 \pm \! 2.3$	85.2
ER	87.5	79.3	98.3	76.3	85.3
pAdaIN	85.8	81.1	97.2	77.4	85.4
ERM^\dagger	84.7 ± 0.4	80.8 ± 0.6	97.2 ± 0.3	79.3 ± 1.0	85.5
EISNet	86.6	81.5	97.1	78.1	85.8
$CORAL^{\dagger}$	88.3 ± 0.2	80.0 ± 0.5	97.5 ± 0.3	78.8 ± 1.3	86.2
SagNet [†]	87.4 ± 1.0	80.7 ± 0.6	97.1 ± 0.1	80.0 ± 0.4	86.3
DSON	87.0	80.6	96.0	82.9	86.6
Ours	89.3 ±0.2	83.4 ±0.6	97.3 ± 0.3	82.5 ± 0.5	88.1

D.2 VLCS

Table 8: Out-of-domain accuracies (%) on VLCS.

Algorithm	С	L	S	V	Avg
GroupDRO [†]	97.3 ± 0.3	63.4 ± 0.9	69.5 ± 0.8	76.7 ± 0.7	76.7
RSC^{\dagger}	97.9 ± 0.1	62.5 ± 0.7	72.3 ± 1.2	75.6 ± 0.8	77.1
MLDG^\dagger	97.4 ± 0.2	65.2 ± 0.7	71.0 ± 1.4	75.3 ± 1.0	77.2
MTL^\dagger	97.8 ± 0.4	64.3 ± 0.3	71.5 ± 0.7	75.3 ± 1.7	77.2
ERM^{\ddagger}	98.0 ± 0.3	64.7 ± 1.2	71.4 ± 1.2	75.2 ± 1.6	77.3
I-Mixup	98.3 ± 0.6	64.8 ± 1.0	72.1 ± 0.5	74.3 ± 0.8	77.4
ERM^\dagger	97.7 ± 0.4	64.3 ± 0.9	73.4 ± 0.5	74.6 ± 1.3	77.5
MMD^\dagger	97.7 ± 0.1	64.0 ± 1.1	72.8 ± 0.2	75.3 ± 3.3	77.5
$CDANN^{\dagger}$	97.1 ± 0.3	65.1 ± 1.2	70.7 ± 0.8	77.1 ± 1.5	77.5
ARM^\dagger	98.7 ± 0.2	63.6 ± 0.7	71.3 ± 1.2	76.7 ± 0.6	77.6
SagNet [†]	97.9 ± 0.4	64.5 ± 0.5	71.4 ± 1.3	77.5 ± 0.5	77.8
Mixstyle [‡]	98.6 ± 0.3	64.5 ± 1.1	72.6 ± 0.5	75.7 ± 1.7	77.9
$VREx^{\dagger}$	98.4 ± 0.3	64.4 ± 1.4	74.1 ± 0.4	76.2 ± 1.3	78.3
IRM^\dagger	98.6 ± 0.1	64.9 ± 0.9	73.4 ± 0.6	77.3 ± 0.9	78.6
DANN^\dagger	99.0 ±0.3	65.1 ± 1.4	73.1 ± 0.3	77.2 ± 0.6	78.6
$CORAL^{\dagger}$	98.3 ± 0.1	66.1 ± 1.2	73.4 ± 0.3	77.5 ± 1.2	78.8
Ours	98.8 ± 0.1	63.3 ±0.3	75.3 ±0.5	79.2 ±0.6	79.1

D.3 OfficeHome

Table 9: Out-of-domain accuracies (%) on OfficeHome.

Table 7. Out-of-domain accuracies (70) on differentime.								
Algorithm	A	C	P	R	Avg			
Mixstyle [‡]	51.1 ± 0.3	53.2 ± 0.4	68.2 ± 0.7	69.2 ± 0.6	60.4			
IRM^\dagger	58.9 ± 2.3	52.2 ± 1.6	72.1 ± 2.9	74.0 ± 2.5	64.3			
ARM^\dagger	58.9 ± 0.8	51.0 ± 0.5	74.1 ± 0.1	75.2 ± 0.3	64.8			
RSC^{\dagger}	60.7 ± 1.4	51.4 ± 0.3	74.8 ± 1.1	75.1 ± 1.3	65.5			
$CDANN^{\dagger}$	61.5 ± 1.4	50.4 ± 2.4	74.4 ± 0.9	76.6 ± 0.8	65.7			
$DANN^\dagger$	59.9 ± 1.3	53.0 ± 0.3	73.6 ± 0.7	76.9 ± 0.5	65.9			
GroupDRO†	60.4 ± 0.7	52.7 ± 1.0	75.0 ± 0.7	76.0 ± 0.7	66.0			
MMD^\dagger	60.4 ± 0.2	53.3 ± 0.3	74.3 ± 0.1	77.4 ± 0.6	66.4			
MTL^\dagger	61.5 ± 0.7	52.4 ± 0.6	74.9 ± 0.4	76.8 ± 0.4	66.4			
$VREx^{\dagger}$	60.7 ± 0.9	53.0 ± 0.9	75.3 ± 0.1	76.6 ± 0.5	66.4			
ERM^\dagger	61.3 ± 0.7	52.4 ± 0.3	75.8 ± 0.1	76.6 ± 0.3	66.5			
MLDG^\dagger	61.5 ± 0.9	53.2 ± 0.6	75.0 ± 1.2	77.5 ± 0.4	66.8			
ERM^{\ddagger}	63.1 ± 0.3	51.9 ± 0.4	77.2 ± 0.5	78.1 ± 0.2	67.6			
I-Mixup	62.4 ± 0.8	54.8 ± 0.6	76.9 ± 0.3	78.3 ± 0.2	68.1			
SagNet [†]	63.4 ± 0.2	54.8 ± 0.4	75.8 ± 0.4	78.3 ± 0.3	68.1			
CORAL [†]	65.3 ± 0.4	54.4 ± 0.5	76.5 ± 0.1	78.4 ± 0.5	68.7			
Ours	66.1 ±0.4	57.7 ±0.4	78.4 ±0.1	80.2 ±0.2	70.6			

D.4 TerraIncognita

Table 10: Out-of-domain accuracies (%) on TerraIncognita.

Algorithm	L100	L38	L43	L46	Avg
MMD^\dagger	41.9 ± 3.0	34.8 ± 1.0	57.0 ± 1.9	35.2 ± 1.8	42.2
GroupDRO†	41.2 ± 0.7	38.6 ± 2.1	56.7 ± 0.9	36.4 ± 2.1	43.2
Mixstyle [‡]	54.3 ± 1.1	34.1 ± 1.1	55.9 ± 1.1	31.7 ± 2.1	44.0
ARM^\dagger	49.3 ± 0.7	38.3 ± 2.4	55.8 ± 0.8	38.7 ± 1.3	45.5
MTL^\dagger	49.3 ± 1.2	$39.6 \pm \! 6.3$	55.6 ± 1.1	37.8 ± 0.8	45.6
$CDANN^{\dagger}$	47.0 ± 1.9	41.3 ± 4.8	54.9 ± 1.7	39.8 ± 2.3	45.8
ERM^\dagger	49.8 ± 4.4	42.1 ± 1.4	56.9 ± 1.8	35.7 ± 3.9	46.1
$VREx^{\dagger}$	48.2 ± 4.3	41.7 ± 1.3	56.8 ± 0.8	38.7 ± 3.1	46.4
RSC^{\dagger}	50.2 ± 2.2	39.2 ± 1.4	56.3 ± 1.4	40.8 ± 0.6	46.6
DANN^\dagger	51.1 ± 3.5	40.6 ± 0.6	57.4 ± 0.5	37.7 ± 1.8	46.7
IRM^\dagger	54.6 ± 1.3	39.8 ± 1.9	56.2 ± 1.8	39.6 ± 0.8	47.6
$CORAL^{\dagger}$	51.6 ± 2.4	42.2 ± 1.0	57.0 ± 1.0	39.8 ± 2.9	47.7
MLDG^\dagger	54.2 ± 3.0	44.3 ± 1.1	55.6 ± 0.3	36.9 ± 2.2	47.8
I-Mixup	59.6 ± 2.0	42.2 ± 1.4	55.9 ± 0.8	33.9 ± 1.4	47.9
SagNet [†]	53.0 ± 2.9	43.0 ± 2.5	57.9 ± 0.6	40.4 ± 1.3	48.6
ERM [‡]	54.3 ± 0.4	$42.5 \pm \hspace*{-0.5em} \pm \hspace*{-0.5em} 0.7$	55.6 ± 0.3	38.8 ± 2.5	47.8
Ours	55.4 ±0.0	44.9 ±1.1	59.7 ±0.4	39.9 ± 0.2	50.0

D.5 DomainNet

Table 11: Out-of-domain accuracies (%) on DomainNet.

Algorithm	clip	info	paint	quick	real	sketch	Avg
MMD^\dagger	32.1 ± 13.3	11.0 ± 4.6	26.8 ± 11.3	8.7 ± 2.1	32.7 ± 13.8	28.9 ± 11.9	23.4
GroupDRO†	47.2 ± 0.5	17.5 ± 0.4	33.8 ± 0.5	$9.3 \pm \! 0.3$	51.6 ± 0.4	40.1 ± 0.6	33.3
$VREx^{\dagger}$	47.3 ± 3.5	16.0 ± 1.5	$35.8 \pm \! 4.6$	10.9 ± 0.3	49.6 ± 4.9	42.0 ± 3.0	33.6
IRM^\dagger	48.5 ± 2.8	15.0 ± 1.5	$38.3 \pm \! 4.3$	10.9 ± 0.5	$48.2 \pm \! 5.2$	42.3 ± 3.1	33.9
Mixstyle [‡]	51.9 ± 0.4	13.3 ± 0.2	37.0 ± 0.5	12.3 ± 0.1	46.1 ± 0.3	43.4 ± 0.4	34.0
ARM^\dagger	49.7 ± 0.3	16.3 ± 0.5	40.9 ± 1.1	9.4 ± 0.1	53.4 ± 0.4	43.5 ± 0.4	35.5
$CDANN^{\dagger}$	54.6 ± 0.4	17.3 ± 0.1	43.7 ± 0.9	12.1 ± 0.7	56.2 ± 0.4	45.9 ± 0.5	38.3
DANN^\dagger	53.1 ± 0.2	18.3 ± 0.1	44.2 ± 0.7	11.8 ± 0.1	55.5 ± 0.4	46.8 ± 0.6	38.3
RSC^\dagger	55.0 ± 1.2	18.3 ± 0.5	44.4 ± 0.6	12.2 ± 0.2	55.7 ± 0.7	47.8 ± 0.9	38.9
I-Mixup	55.7 ± 0.3	18.5 ± 0.5	44.3 ± 0.5	12.5 ± 0.4	55.8 ± 0.3	48.2 ± 0.5	39.2
SagNet [†]	57.7 ± 0.3	19.0 ± 0.2	45.3 ± 0.3	12.7 ± 0.5	58.1 ± 0.5	48.8 ± 0.2	40.3
MTL^\dagger	57.9 ± 0.5	18.5 ± 0.4	46.0 ± 0.1	12.5 ± 0.1	59.5 ± 0.3	49.2 ± 0.1	40.6
ERM^\dagger	58.1 ± 0.3	18.8 ± 0.3	46.7 ± 0.3	12.2 ± 0.4	59.6 ± 0.1	49.8 ± 0.4	40.9
MLDG^\dagger	59.1 ± 0.2	19.1 ± 0.3	45.8 ± 0.7	13.4 ± 0.3	59.6 ± 0.2	50.2 ± 0.4	41.2
$CORAL^{\dagger}$	59.2 ± 0.1	19.7 ± 0.2	46.6 ± 0.3	13.4 ± 0.4	59.8 ± 0.2	50.1 ± 0.6	41.5
MetaReg	59.8	25.6	50.2	11.5	64.6	50.1	43.6
DMG	65.2	22.2	50.0	15.7	59.6	49.0	43.6
ERM [‡]	63.0 ± 0.2	21.2 ± 0.2	50.1 ± 0.4	13.9 ± 0.5	63.7 ± 0.2	52.0 ± 0.5	44.0
Ours	66.0 ±0.1	22.4 ±0.3	53.5 ±0.1	16.1 ±0.2	65.8 ±0.4	55.5 ±0.3	46.5

E Assets

In this section, we discuss about licenses, copyrights, and ethical issues of our assets, such as code and datasets.

E.1 Code

Our work is built upon DomainBed [22]⁴, which is released under the MIT license.

E.2 Datasets

While we use public datasets only, we track how the datasets were built to discuss licenses, copyrights, and potential ethical issues. For DomainNet [33] and OfficeHome [31], we use the datasets for non-profit academic research only following their fair use notice. TerraIncognita [32] is a subset of Caltech Camera Traps (CCT) dataset, distributed under the Community Data License Agreement (CDLA) license. PACS [7] and VLCS [30] datasets have images collected from the web and we could not find any statements about licenses, copyrights, or whether consent was obtained. Considering that both datasets contain person class and images of people, there may be potential ethical issues.

F Reproducibility

To provide details of our algorithm and guarantee reproducibility, our source code is given in the supplementary material. The code also specifies detailed environments, dependencies, how to download datasets, and instructions to reproduce the main results (Table 1 and 2 in the main text).

F.1 Infrastructures

Every experiment is conducted on a single NVIDIA Tesla P40, Python 3.8.6, PyTorch 1.7.0, Torchvision 0.8.1, and CUDA 9.2.

⁴https://github.com/facebookresearch/DomainBed

F.2 Runtime Analysis

The total runtime varies depending on datasets and the moment detected to overfit. It takes about 4 hours for PACS and VLCS, 8 hours for OfficeHome, 8.5 hours for TerraIncognita, and 56 hours for DomainNet on average. Each experiment includes the leave-one-out cross-validations for all domains in each dataset.

F.3 Complexity Analysis

The only additional time overhead incurs from stochastic weights selection, which requires further evaluations. To analyze the overhead, let the forward time t_f , backward time t_b , training and validation split ratio $r = |X^{train}|/|X^{valid}|$, total in-domain samples n, and evaluation frequency v that indicates how many evaluations are conducted for each epoch. For conciseness, we assume $t = t_f = t_b$ and do not consider early stopping.

For one epoch, training time is 2tnr/(r+1), and evaluation time is vtn/(r+1). The total runtime for one epoch is tn(2r+v)/(r+1). Final overhead ratio is $(2r+v)/(2r+v_b)$ where v_b is the evaluation frequency of a baseline. In our main experiments, we use r=4. Compared to the default parameters of DomainBed [22], we use $v=2v_b$ for DomainNet, $v=6v_b$ for VLCS, and $v=3v_b$ for the others. Then, the total runtime of our algorithm takes from 1.07 (PACS) to 1.27 (DomainNet) times more than the ERM baseline. In practice, it can be improved by conducting approximated evaluations using sub-sampled validation set.

In terms of memory complexity, our method does not require additional GPU memory. Instead, we leverage CPU memory to minimize training time overhead, which takes up to $\max(N, M)$ times more than the baseline.