### Reinforcement Learning as a Parsimonious Alternative to Prediction Cascades: A Case Study on Image Segmentation

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#### Abstract

Deep learning architectures have achieved state-of-the-art (SOTA) performance on computer vision tasks such as object detection and image segmentation. This may be attributed to the use of over-parameterized, monolithic deep learning architectures executed on large datasets. Although such large architectures lead to increased accuracy, this is usually accompanied by a larger increase in computation and memory requirements during inference. While this is a non-issue in traditional machine learning (ML) pipelines, the recent confluence of machine learning and fields like the Internet of Things (IoT) has rendered such large architectures infeasible for execution in low-resource settings. For some datasets, large monolithic pipelines may be overkill for simpler inputs. To address this problem, previous efforts have proposed decision cascades where inputs are passed through models of increasing complexity until desired performance is achieved. However, we argue that cascaded prediction leads to sub-optimal throughput and increased computational cost due to wasteful intermediate computations. To address this, we propose PaSeR (Parsimonious Segmentation with Reinforcement Learning) a non-cascading, cost-aware learning pipeline as an efficient alternative to cascaded decision architectures. Through experimental evaluation on both real-world and standard datasets, we demonstrate that PaSeR achieves better accuracy while minimizing computational cost relative to cascaded models. Further, we introduce a new metric IoU/GigaFlop to evaluate the balance between cost and performance. On the real-world task of battery material phase segmentation, PaSeR yields a minimum performance improvement of 174% on the IoU/GigaFlop metric with respect to baselines. We also demonstrate PaSeR's adaptability to complementary models trained on a noisy MNIST dataset, where it achieved a minimum performance improvement on IoU/GigaFlop of 13.4% over SOTA models. Code and data are available at github.com/scailab/paser.

#### Introduction

Recent advances in deep learning (DL) and the internetof-things (IoT) have led to the burgeoning application of DL in manufacturing pipelines (Hussain et al. 2020; Meng et al. 2020; Mohammadi et al. 2018; Tang et al. 2017). In many such applications, ML / DL models are often deployed on devices with low memory and computational ca-

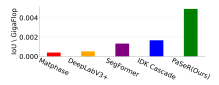


Figure 1: Performance w.r.t IoU/GigaFlop metric (higher is better) of SOTA models and our proposed PaSeR model on the battery material phase segmentation task.

pabilities (edge) in conjunction with DL models that are deployed in less constrained environments (fog, cloud). These edge-fog-cloud (EFC) systems are commonly used in areas such as smart manufacturing (Chen et al. 2018a) and healthcare (Mutlag et al. 2021) where precision machines such as electrocardiograms collect and preprocess high density data while integrating with a local computer as well as cloud based resources to accurately and efficiently provide critical information. Although the fog and cloud environments enable the deployment of larger DL models, querying them is costly (due to communication network and model latency). Hence, such real-world contexts require a high-throughput pipeline to balance task accuracy and computational cost.

A popular solution to deal with this problem is the I Don't Know (IDK) Cascade (Wang et al. 2017) in which models of increasing complexity (starting with the least cost model) are sequentially queried until a model yields a prediction exceeding a preset confidence threshold. Multi-exit models (Kouris et al. 2022) follow a similar cascading architecture but require a potentially costly neural architecture search during training. We argue that such pipelines, although wellmotivated, lead to high computational costs due to excess computations incurred as a function of the sequential cascading constraint. In this paper, we argue that reinforcement learning (RL) can be employed as an effective substitute to circumvent the cascading restriction. We employ RL to directly select which of a set of models to query with a particular input such that the learned policy maximizes task performance while minimizing computational cost. To this end, we propose the PaSeR framework and demonstrate its performance on the challenging task of battery material phase segmentation.

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Application Background. Lithium-ion batteries are extensively used in many industrial applications, (e.g., smartphones, laptops, and electric vehicles) due to their efficient energy storage capability. The electrode coatings of these batteries consist of composite active materials (e.g., Lithium, Nickel, Manganese) and a polymeric binder (Carbon). The microstructure of these composite electrode coatings consists of the spatial distribution of active and binder materials. The physical parameters of a microstructure, (e.g., homogeneity of coating thickness, porosity) influence battery performance. Resolving the locations of the active and binder materials and their phase transitions (i.e., the task of battery material phase segmentation) can help deduce these physical parameters, thereby providing an understanding of phenomena like battery degradation. Existing techniques to address this problem use expensive highresolution X-ray computed tomography images (Lu et al. 2020). Low-resolution (low-res) microtomography images have also been used, but they cannot readily distinguish between spatial distributions of the composite active materials. Recently, DL segmentation models like MatPhase by (Tabassum et al. 2022) have been developed to identify (pixel-wise) these composite materials and their phase transitions from low-res images, however, these approaches are computationally expensive to execute.

In this context, we propose PaSeR as a low-cost but effective and robust solution to address the task of battery material segmentation from low-res microtomography images. Our contributions are as follows: (C1) We develop a novel computationally parsimonious DL framework (employing reinforcement learning with cost-aware rewards) to balance cost with task performance. (C2) Through qualitative and quantitative experiments, we demonstrate that PaSeR yields competitive performance with SOTA models on the battery material phase segmentation task while also being the most computationally efficient. (C3) We demonstrate the effectiveness of the learned RL policy in unseen (noisy) contexts as well as with task models having complementary strengths. (C4) Finally, we introduce a novel metric called IoU per GigaFlop (IoU/GigaFlop) which measures the segmentation performance obtained per GigaFlop of computation expended, an effective metric for evaluating such lowcost learning pipelines (see Fig. 1).

#### **Related Work**

We review two areas of research related to our work, lowcost ML and image segmentation models.

Low-Cost & Tiny ML. There have been many past efforts to develop low-cost DL pipelines for use in low memory, low storage, high-throughput IoT contexts. *Knowledge distillation* (KD) and employing *decision cascades* are two popular approaches in this context. While the primary goal of KD (Hinton, Vinyals, and Dean 2015; Gou et al. 2021; Phuong and Lampert 2019) is to learn smaller models to *mimic* larger models, this goal isn't fully aligned with the scope of the current work, which is to learn optimal decision pipelines to create a low cost, high performance ML by incorporating multi-models. However, the other research thread of employing decision cascades is directly relevant to

our work. Decision Cascades, originally introduced in (Cai, Saberian, and Vasconcelos 2015; Angelova et al. 2015) were recently re-popularized by the work of IDK Cascades (Wang et al. 2017). The IDK cascade framework imposes a sequential model architecture, where each model is queried in order of increasing complexity until prediction confidence exceeds a threshold. Yet another paradigm of *Tiny-ML* (Rajapakse, Karunanayake, and Ahmed 2023; Ren, Anicic, and Runkler 2022) also aims to develop ML models but with the goal of deploying them on extremely low-cost hardware devices. Our goal is aligned with but complementary to this as our proposed decision pipeline can be employed with such lowcost models along with higher-cost models (on the cloud) to maximize performance and minimize computational cost.

**Image Segmentation**. The field of image segmentation has also seen many successes in multiple domains (Chen et al. 2017; Li et al. 2018; Chen et al. 2019) with popular architectures like the U-Net (Ronneberger, Fischer, and Brox 2015) and the recent Segment Anything (Kirillov et al. 2023) *foundation model*. Our PaSeR framework is flexible enough to incorporate any of these SOTA segmentation models as we have developed a decision pipeline that can leverage multiple models to maximize performance on a target task while minimizing computational cost. Finally, efforts in intelligent data sampling (Uzkent, Yeh, and Ermon 2020; Uzkent and Ermon 2020) which may possess a motivation in terms of employing RL for maximal task performance at minimal cost, differ in the actual application of the RL pipeline and learning task.

#### **Problem Formulation**

In this work, our goal is to develop a learnable decision pipeline that is computationally parsimonious (i.e., minimizes wasteful computations) and also yields competitive performance (compared to SOTA models) on the target task. To develop such a decision pipeline, we leverage reinforcement learning (RL). Specifically, we propose the PaSeR framework (see Fig. 2 for architecture details) composed of an RL policy model  $f_{RL}$ , a small/efficient task model  $f_0$ , and m large task models  $\{f_1, \ldots, f_m\}$ . In this paper, we demonstrate the performance of PaSeR in the context of image segmentation. Algorithm 1 outlines the training procedure of PaSeR in the context of our target task (i.e., image segmentation), but we note that PaSeR is task independent and can be applied to other learning contexts with a few appropriate modifications. Code and data are available at github.com/scailab/paser.

Segmentation Model Pretraining. At the outset of our training procedure, we split the training data into three equal subsets:  $\mathcal{D}_{PT}$ ,  $\mathcal{D}_{RL}$ ,  $\mathcal{D}_{FT}$ . Each subset is comprised of image instances and pixel labels  $(\mathbf{x}, \mathbf{y})$  where  $\mathbf{x} \in \mathbb{R}^{C \times H \times W}$  and  $\mathbf{y} \in \mathbb{R}^{1 \times H \times W}$ . Using the pretraining subset  $\mathcal{D}_{PT}$ , we train the *m* large segmentation models  $f_1, \ldots, f_m$  first by splitting each image  $\mathbf{x}$  into *P* equal size patches (in our case P = 16) with the help of a *patchification* function  $\mathscr{P}(\cdot)$  where  $\mathbf{x}^{(p)}$  denotes the  $p^{th}$  patch. These patches are passed as inputs to each model while optimizing cross entropy loss  $\mathcal{L}(\hat{\mathbf{z}}^{(p)}, \mathbf{y}^{(p)})$  between the prediction logits  $\hat{\mathbf{z}}^{(p)}$ 

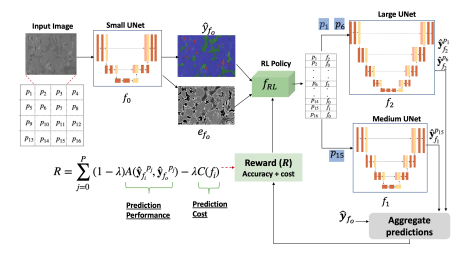


Figure 2: Overview of PaSeR. The small UNet  $(f_0)$  yields the segmentation  $(\hat{\mathbf{y}}_{f_0})$  and corresponding entropy map  $\mathbf{e}_{f_0}$  conditioned on the whole input image (**x**). Then, **x** is divided into 'P' equal sized patches. The RL policy directs each patch  $\mathbf{x}^{(p)}$  of **x** to one of  $f_0, f_1, f_2$  to maximize reward. Based on the RL actions, models  $f_1$  and  $f_2$  yield predictions for the corresponding image patch. All the predicted patches are then aggregated to yield the final segmentation.

and ground truth  $\mathbf{y}^{(p)}$ . Once models  $f_1, \ldots, f_m$ , are pretrained, the smallest model,  $f_0$  is pre-trained using  $\mathcal{D}_{PT}$  on the full image (i.e., no patchification). In addition to using the cross entropy loss  $\mathcal{L}(\hat{\mathbf{z}}, \mathbf{y})$  we also use a knowledge distillation (KD) loss (Hinton, Vinyals, and Dean 2015; Kim et al. 2021) between the outputs of the largest model  $f_m$  and the small model  $f_0$ . We define the KD loss function in Eq. 1.

$$\mathcal{L}_{KD} = \frac{1}{|\mathcal{D}_{PT}|} \sum_{j=1}^{|\mathcal{D}_{PT}|} \left( \hat{\mathbf{z}}_{f_{0},j}^{(p)} - \hat{\mathbf{z}}_{f_{m},j}^{(p)} \right)^{2}$$
(1)

The term  $\hat{\mathbf{y}}_{f_0,j}^{(p)}$  indicates the segmentation predictions for patch p of instance j yielded by model  $f_0$ .  $\hat{\mathbf{y}}_{f_m,j}^{(p)}$  is the corresponding prediction yielded by model  $f_m$ . This loss encourages outputs of  $f_0$  to be closer to the largest model  $f_m$ , thereby transferring information from the representations learned by  $f_m$  to  $f_0$  improving its performance without increasing its size.

**RL Training.** We incorporate reinforcement learning as the decision paradigm to develop a compute-efficient segmentation pipeline. Specifically, our RL policy is conditioned upon states s, constituted by the image segmentation  $\hat{\mathbf{y}}_{f_0}$  and entropy maps  $\mathbf{e}_{f_0}$  of the smallest model  $f_0$  to output an action which specifies a set of patch and model pairs for each image to be passed *upstream* to more sophisticated models in the pipeline. States are of the form  $(\hat{\mathbf{y}}_{f_0}, \mathbf{e}_{f_0})$  and actions are defined as  $\mathbf{a} \in$  $\{0, \ldots, m\}^P$ . We define the patch-model selection policy as  $\pi_{RL}(\mathbf{a} | \mathbf{s}) = p(\mathbf{a} | f_{RL}(\hat{\mathbf{y}}_{f_0}, \mathbf{e}_{f_0}; \theta_{f_{RL}}))$ . Here the policy network  $f_{RL}$  parameterizes the action distribution p, which in our case is a categorical distribution with probabilities  $\mathbf{s} \in \{s_{f_0}, \ldots, s_{f_m}\}^P : s_{f_i} > 0$ ,  $\sum_{i=0}^m s_{f_i} = 1$ . The entropy  $\mathbf{e}_{f_0}$  is calculated using Monte Carlo dropout (Gal and Ghahramani 2016), but note that other methods for uncertainty quantification can also be supported by PaSeR. Using probabilities s, we sample from a categorical distribution to obtain an action  $\mathbf{a} \in \{0, \ldots, m\}^P$ . For example, if  $\mathbf{a}_k = 2$  for some index k of a, this indicates that  $f_{RL}$  has chosen the  $k^{th}$  patch to be directed to model  $f_2$  for segmentation. Using the sampled action, we pass each patch to its respective model and compute a reward. The reward function is detailed in Eq. 2 and is based on the difference in prediction performance A between the large and small model predictions,  $\hat{\mathbf{y}}_{f_{a_p}}, \hat{\mathbf{y}}_{f_0}$ , as well as a computational cost penalty term C. The action a defines the models run on each patch.

$$R(\mathbf{a} = \{a_1, \dots, a_P\}) = \sum_{p=0}^{P} (1-\lambda) A(\hat{\mathbf{y}}_{f_{a_p}}^{(p)}, \hat{\mathbf{y}}_{f_0}^{(p)}) - \lambda C(f_{a_p})$$
(2)

For our experiments in segmentation we use difference in mean intersection over union the (IoU) as our measure of prediction performance:  $A(\hat{\mathbf{y}}_{f_i}^{(p)}, \hat{\mathbf{y}}_{f_0}^{(p)}) = IoU(\hat{\mathbf{y}}_{f_i}^{(p)}) - IoU(\hat{\mathbf{y}}_{f_0}^{(p)}).$  Note that in Eq. 2, the cost parameter  $\lambda$  parameterizes a convex combination of accuracy and computational cost to provide a simple way to control the influence of each component on the RL policy reward. We design a cost function C in Eq. 3 with range (0,1) as the ratio of the number of learnable parameters in a model to the total number of parameters in all models  $\{f_0, \ldots, f_m\}$ .

$$C(f_i) = \frac{\text{numParams}(f_i)}{\sum_{j=1}^{m} \text{numParams}(f_j)}$$
(3)

Using the reward value R, we compute the policy gradient (Sutton et al. 1999)  $\nabla_{\theta_{f_{RL}}} J = \mathbb{E}[\nabla_{\theta_{f_{RL}}} \log \pi_{RL}(\mathbf{a} \mid \mathbf{s}) * R]$  and update the parameters  $\theta_{RL}$  of the RL policy.

**Fine-Tuning**. The final step of PaSeR is fine-tuning. Here, we jointly update the large models and RL model. The joint training helps the large segmentation models improve their performance on the inputs being directed to them by the RL policy while also further personalizing the RL policy to dis-

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Algorithm 1: PaSeR Algorithm
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**Data:**  $\mathcal{D}_{PT}, \mathcal{D}_{RL}, \mathcal{D}_{FT}$  **Parameters:**  $\theta_{f_{RL}}, \theta_{f_0}, \dots, \theta_{f_m}$  **Hyp:**  $\lambda, \eta, \beta$ **Models:** RL policy  $f_{RL}$ , small/efficient model  $f_0$  and m large task models  $\{f_1, \dots, f_m\}$ 1 for  $f_i \in \{f_1, \ldots, f_m\}$ # Pretrain each large task model 2 do for  $\mathbf{x}^{(p)}, \mathbf{y}^{(p)} \in \mathscr{P}(\mathcal{D}_{PT})$ 3 # For each data point in pre-training dataset do 4  $\begin{vmatrix} \hat{\mathbf{y}}_{f_i}^{(p)}, \hat{\mathbf{z}}_{f_i}^{(p)} \leftarrow f_i(\mathbf{x}^{(p)}) \\ l \leftarrow \mathcal{L}(\hat{\mathbf{z}}_{f_i}^{(p)}, \mathbf{y}^{(p)}) \end{vmatrix}$ # Get task predictions and logits from model  $f_i$ 5 # Compute loss (cross entropy) 6  $\theta_{f_i} \leftarrow \theta_{f_i}^{j_i} - \eta \nabla_{\theta_{f_i}} l$ # Update model parameters 7 8 end 9 end # Pretrain small/efficient model with KD loss 10 for  $\mathbf{x}, \mathbf{y} \in \mathcal{D}_{PT}$ 11 do  $\hat{\mathbf{y}}_{f_0}, \hat{\mathbf{z}}_{f_0} \leftarrow f_0(\mathbf{x})$ # Get *small* model  $(f_0)$  prediction 12  $\hat{\mathbf{y}}_{f_m}, \hat{\mathbf{z}}_{f_m} \leftarrow f_m(\mathbf{x})$ # Get largest model prediction 13  $l \leftarrow \mathcal{L}(\hat{\mathbf{z}}_{f_0}, \mathbf{y}) + \beta \mathcal{L}_{KD}(\hat{\mathbf{z}}_{f_0}, \hat{\mathbf{z}}_{f_m})$  # Compute loss with KD 14  $\theta_{f_0} \leftarrow \theta_{f_0} - \eta \nabla_{\theta_{f_0}} l$ # Update model parameters 15 16 end 17 for  $\mathbf{x}, \mathbf{y} \in \mathcal{D}_{RL}$ # Train RL policy model 18 do # Get small model prediction and entropy 19  $\hat{\mathbf{y}}_{f_0}, \mathbf{e}_{f_0} \leftarrow f_0(\mathbf{x})$  $\begin{aligned} \mathbf{s} &\leftarrow f_{RL}(\hat{\mathbf{y}}_{f_0}, \mathbf{e}_{f_0}) \\ \mathbf{a} &\sim \pi_{RL}(\mathcal{A} \mid \mathbf{s}) \end{aligned}$ 20 # Get probabilities of actions from RL model # Sample action from RL policy distribution 21 for  $a_p \in \mathbf{a}$ # For each model and patch in action 22 do 23  $\begin{aligned} \hat{\mathbf{y}}_{a_p}^{(p)} \leftarrow f_{a_p}(\mathbf{x}^{(p)}) & \text{# Get model prediction} \\ R += (1 - \lambda) A(\hat{\mathbf{y}}_{f_{a_p}}^{(p)}, \hat{\mathbf{y}}_{f_0}^{(p)}) - \lambda C(f_{a_p}) & \text{# Compute accuracy+cost-based reward} \end{aligned}$ 24 25 26 end  $\begin{aligned} \nabla_{\theta_{RL}} J &= \mathbb{E}[\nabla_{\theta_{RL}} \log \pi_{RL}(\mathcal{A}|s) * R] \\ \theta_{f_{RL}} &\leftarrow \theta_{f_{RL}} - \eta \nabla_{\theta_{f_{RL}}} J(\pi_{RL}) \end{aligned}$ # Compute policy gradient 27 # Update RL model 28 29 end 30 for  $\mathbf{x}, \mathbf{y} \in \mathcal{D}_{FT}$ # Finetune models 31 do Repeat Lines: 5-7 for each large model Repeat Lines: 19-28 for RL model 32 end

cern the strengths and weaknesses of each constituent segmentation model for each input patch.

#### **Experimental Setup**

We train three UNet segmentation models  $f_0, f_1, f_2$  with 16571, 1080595, and 17275459 parameters respectively on  $\mathcal{D}_{PT}$  for 200 epochs, followed by training our RL model  $f_{RL}$  with 14736 parameters on  $\mathcal{D}_{RL}$  for 200 epochs. Finally, we fine-tune all models on  $\mathcal{D}_{FT}$  for 200 epochs. PaSeR trains with a batch size of 32 using the Adam optimizer (Kingma and Ba 2014) with  $\eta = 0.0001$ . In our battery segmentation experiment we set  $\beta = 0.01$  using grid search and  $\lambda = 0.5$  which corresponds to an even balance between performance and cost. For the noisy MNIST dataset we set  $\lambda = 0$ , see section R4. Adaptability to Complementary Models for more details.

#### Baselines

We compare PaSeR to six baselines with complementary strengths to illustrate how we improve upon each of these baselines in either IoU performance and/or IoU per GigaFlop efficiency. (1) IDK Cascade (Wang et al. 2017): We implement the IDK-Cascade model with a cost aware cascade using the same segmentation models in PaSeR. For the IDK loss and cost function we use cross entropy loss and our previously defined cost function (Eq. 3), while tuning this baseline with an exhaustive grid search. (2) PaSeR-**RandPol.**: We setup PaSeR with a random policy for actions drawn uniformly from a categorical distribution. We call this method PaSeR-RandPol. (3) MatPhase (Tabassum et al. 2022): We also compare PaSeR to a state of the art (SOTA) model specialized for the task of battery material phase segmentation. The MatPhase model is an ensemble method which combines UNet segmentation models with pixel level IDK classification and a convolutional neural network. (4) DeepLabV3+ (Chen et al. 2018b): To put PaSeR in context with modern DL models, we compare it to DeepLabV3+, a SOTA segmentation model which uses atrous convolutions alongside an encoder-decoder. (5) Seg-Former (Xie et al. 2021): We also compare our method to SegFormer, a recent SOTA segmentation model which combines transformers with small multi-layer perceptron decoders. (6) EfficientViT (Cai et al. 2022): We also compare to the lightweight EfficientViT, which uses linear attention.

#### **Evaluation Metrics**

(1) Intersection-Over-Union (IoU): We employ IoU (aka. Jaccard index), a popular and effective metric used to evaluate performance on image segmentation tasks. (2) Flops (F): We profile the number of floating point operations per instance for PaSeR and baselines in inference mode when run on the full test set. This gives us the raw computational cost of each model. (3) IoU Per GigaFlop  $\left(\frac{IoU}{GigaFlop}\right)$ : While Flops measures compute required per model, we introduce a new metric called IoU per GigaFlop which is defined by the ratio  $\frac{IoU}{GigaFlop}$ . This metric enables a unified understanding of performance effectiveness and computational cost.

#### **Dataset Description**

**Battery Material Phase Segmentation**. Our battery material phase segmentation dataset consists of 1,330 images (1270 training images, 20 validation, and 40 test images) obtained from low-res microtomography (inputs), each of size (224,256) along with pixel level labels of 3 classes (obtained from high-res computational tomography): pore, carbon, and nickel. We split these images into 16 equal size patches of size (56, 64) each.

**Noisy MNIST**. The standard MNIST dataset (Deng 2012) consists of 70,000 grayscale images (50,000 training, 10,000 validation and 10,000 test images). We create three different versions of this dataset for foreground/background segmentation with three noise types respectively: Gaussian blur with radius 1, Gaussian blur with radius 2 and a box blur with a fixed convolutional filter. See Fig. 3 for examples of each noise type.

#### **Results & Discussion**

In line with our goal of designing a computationally parsimonious framework, we investigate PaSeR performance in the context of the following research questions.

**R1.** How does the task performance and computational efficiency of PaSeR compare with the IDK-Cascade paradigm? **R2.** How well does PaSeR balance IoU and efficiency relative to SOTA segmentation models?

**R3.** How adaptable and robust is the PaSeR decision policy to noisy data?

**R4.** How adaptable and robust is the PaSeR decision policy to task models with complementary strengths?

**R5.** What are the effects of the various components of PaSeR, ( $\lambda$ , MC-Sampling) on achieving an effective balance between computational cost and task performance?

## **R1:** Task Performance and Computational Efficiency vs. IDK-Cascade

To evaluate model task performance, we compare our PaSeR model to the cost-aware IDK cascading decision baseline, and a variant of PaSeR (i.e., PaSeR-RandPol.) with

the same segmentation models as PaSeR except with a random policy instead of a learned RL policy. The performance results are depicted in Table 1. Looking at the battery dataset, we see that PaSeR outperforms the IDK Cascade model by 6.28% in terms of the IoU metric. PaSeR also achieves the highest IoU/GigaFlop, outperforming IDK-Cascade by 196%.

Note that the IDK-Cascade model currently underperforms PaSeR on the Battery dataset. Hence, for a fair comparison with our method, we tune the IDK Cascade model to match the IoU performance of PaSeR and denote this model as IDK-Cascade (IoU Match). We achieve this by adjusting the entropy thresholds used in each stage of the cascade until we obtain a least-upper-bound performance (i.e., within a tolerance of  $10^{-3}$  of IoU) compared to PaSeR on the same test set. In Table 2, comparing the flops of both models (for the same IoU performance), we see that the PaSeR model requires 90% fewer flops compared to IDK-Cascade (IoU Match) to achieve similar performance. This is further corroborated by the IoU/GigaFlop metric in Table 2 wherein we see that PaSeR achieves a 923% improvement on this metric thereby indicating that PaSeR is able to yield good performance at much lower computational cost compared to the IDK cascading modeling paradigm.

Finally, on the MNIST dataset PaSeR outperforms IDK-Cascade (IoU Match) by 6.1% and 88.4%% on IoU and IoU/GigaFlop metrics respectively. Here IDK-Cascade (IoU Match) underperforms on the IoU metric vs PaSeR because the entropy based threshold of IDK-Cascade (IoU Match) is not nuanced enough to determine the correct model assignment for a given input. In fact, the accuracy of model assignment by the IDK-Cascade (IoU Match) is only 80% while PaSeR has a model assignment accuracy of 92.7%.

# **R2:** Performance Comparison with SOTA Segmentation Models

The problem of battery material phase segmentation has been investigated by a few previous efforts (see Sec. ). The most recent and best model of this group of efforts is MatPhase. We characterize the performance of PaSeR with respect to this SOTA battery material phase segmentation model as well as the recent monolithic SOTA segmentation models DeepLabV3+, SegFormer and EfficientViT. The distributed nature of PaSeR vs monolithic architectures such as SegFormer allows PaSeR to be deployed in an EFC system where monolithic SOTA models would not satisfy computational edge constraints.

In Table 1 we see that although MatPhase (Tabassum et al. 2022) outperforms PaSeR in terms of segmentation performance, it does so employing significantly more computation. Specifically, MatPhase employs **1297**% more computation than PaSeR to obtain a **9.7**% performance improvement. Further, we notice that PaSeR achieves a minimum improvement of **174**% over all baselines on the IoU/GigaFlop metric. This is a significant result showing the usefulness of PaSeR relative to SOTA models like MatPhase in computationally constrained environments.

When comparing to DeepLabV3+, SegFormer and EfficientViT on the Battery dataset, we see that PaSeR is within

Model	Battery			Noisy MNIST		
Model	IoU	Flops	IoU/GigaFlop	IoU	Flops	IoU/GigaFlop
Matphase (Tabassum et al. 2022)	0.8144	$2.11\times10^{12}$	$0.39 \times 10^{-3}$		_	
DeepLabV3+ (Chen et al. 2018b)	0.7817	$1.55 \times 10^{12}$	$0.51 \times 10^{-3}$	0.8459	$2.07 \times 10^{13}$	$4.08 \times 10^{-5}$
SegFormer (Xie et al. 2021)	0.7692	$5.84 \times 10^{11}$	$1.32 \times 10^{-3}$	0.8448	$7.56 \times 10^{12}$	$1.12 \times 10^{-4}$
EfficientViT (Cai et al. 2022)	0.7765	$4.34 \times 10^{11}$	$1.79 \times 10^{-3}$	0.8344	$3.72 \times 10^{14}$	$2.24 \times 10^{-6}$
IDK-Cascade (Wang et al. 2017)	0.6987	$4.20 \times 10^{11}$	$1.66 \times 10^{-3}$	0.7750	$1.15 \times 10^{13}$	$6.73 \times 10^{-5}$
PaSeR-RandPol.	0.7234	$5.33 \times 10^{11}$	$1.36 \times 10^{-3}$	0.6376	$7.05 \times 10^{12}$	$9.05 \times 10^{-5}$
PaSeR (ours)	0.7426	$1.51 \times 10^{11}$	$4.91 imes10^{-3}$	0.8231	$6.51\times10^{12}$	$1.27 imes10^{-4}$

Table 1: Battery material phase segmentation and Noisy MNIST results comparison between PaSeR and SOTA models.

Model		Battery		Noisy MNIST		
	IoU	Flops	IoU/GigaFlop   IoU	Flops	IoU/GigaFlop	
IDK-Cascade (IoU Match)	0.7444	$1.54 \times 10^{12}$	$0.48 \times 10^{-3} \mid 0.7755$		$6.74 \times 10^{-5}$	
PaSeR (ours)	0.7426	$1.51 \times 10^{11}$	$4.91 \times 10^{-3}$ 0.8231	$6.51\times10^{12}$	$1.27 imes10^{-4}$	

Table 2: Battery material phase segmentation and Noisy MNIST IoU Match Results for PaSeR and IDK-Cascade .



Figure 3: Examples of types of noise added to MNIST data.

4% of the IoU that those models achieve. Despite their slightly better performance on IoU, PaSeR is much more efficient on the IoU/GigaFlop metric by 863%, 272% and 174% for DeepLabV3+, SegFormer and EfficientViT respectively. On the Noisy MNIST dataset, we see the same pattern again. For the DeepLabV3+ model, PaSeR has an 211% higher IoU/GigaFlop while also outperforming the SegFormer model by 13.4% on IoU/GigaFlop. The EfficientViT model performs poorly on this dataset because it is designed for high resolution images and downscales the image by a factor of 8 when outputting segmentation maps. To compensate for this downscaling, we upscale our 32x32 MNIST images to 256x256 for this model.

**Cityscapes.** To demonstrate PaSeR on a modern segmentation task while also integrating pretrained models, we train PaSeR on the Cityscapes dataset (Cordts et al. 2016) using three task models: our small UNet, SegFormer-B0, and SegFormer-B5 with  $\lambda = 0.10$  achieving a test set IoU of 0.8163 which is comparable with SOTA model performance.

### **R3:** Adaptability to Unseen Contexts (Battery Data)

Data and products in real-world (IoT-based) manufacturing pipelines are often plagued by process noise leading to instances from unseen input data distributions. It is in such contexts that the true effectiveness of pipelines such as PaSeR come to the fore in terms of being able to adapt in unseen data contexts.

To investigate the adaptability of our RL policy based

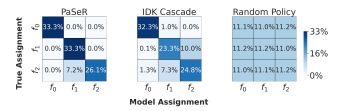


Figure 4: Model assignment confusion matrices for PaSeR, IDK-Cascade and PaSeR-RandPol.

Model	IoU (Noisy)	Degradation
PaSeR-RandPol.	0.5864	-18.94%
PaSeR	0.7322	-1.4%

Table 3: PaSeR vs PaSeR-RandPol. on noisy datasets. Note that PaSeR-RandPol. fails to adapt in the case of noisy data.

PaSeR and demonstrate its effectiveness relative to the random policy in PaSeR-RandPol., we create a variant of our battery segmentation dataset injected with salt and pepper noise. This is done to simulate data quality degradation of the input to the segmentation pipeline, due to equipment / process noise. Further, we create pre-trained variants of all segmentation models  $\{f_1, \ldots, f_m\}$  (except  $f_0$  i.e., the small U-Net) on a combination of clean and noisy data. Finally, we just replace (without fine-tuning  $f_0$ ,  $f_{\rm RL}$ ) the models  $\{f_1, \ldots, f_m\}$  in the fully-trained PaSeR model, with variants trained on noisy as well as clean data.

We then investigate performance of PaSeR and PaSeR-RandPol. (both augmented with same set of segmentation models) on a noisy held-out set of data. Note that by leaving  $f_0$  and RL policy  $f_{\rm RL}$  unaware of the noisy data, we have created a scenario which is unseen w.r.t the RL policy (and model  $f_0$  on whose predictions and entropy the RL policy decisions are conditioned). Table 1 showcases IoU segmentation results (on the battery dataset) of PaSeR and PaSeR-RandPol. in the clean data context while Table 3 showcases corresponding IoU results in a noisy context. From these results, we notice that both models experience degradation under the unseen noisy context. However, the degradation in IoU performance experienced by PaSeR is minimal (1.4%), owing to the RL policy being able to adapt, unlike in PaSeR-RandPol. which shows significant performance degradation (18.94%). We find that PaSeR sends 5.7% more patches to the larger models (that have been exposed to the noisy data) than in the clean data case, thereby showcasing strong evidence of adaptability in unseen contexts. This **advantage of adaptability in noisy**, **unseen scenarios with minimal degradation** is also a significant advantage of PaSeR and its cost-aware RL model.

### **R4.** Adaptability to Complementary Models (Noisy MNIST)

We demonstrate robustness of PaSeR to utilize models with complementary strengths, on the Noisy MNIST dataset. We train each segmentation model  $(f_0, f_1, f_2)$  on the task of foreground/background segmentation on each noisy dataset respectively, training  $f_0$  on the Gaussian blur with radius 1,  $f_1$  on Gaussian blur radius 2 and  $f_3$  on box blur data. Examples of the three noise types are shown in Fig. 3. Each segmentation model learns how to denoise its own noise type and thereby has a unique strength relative to other models.

After training the segmentation models, we train PaSeR 's RL policy with  $\lambda = 0$  such that it learns the optimal policy without regard for computational cost. We have the dataset containing equal proportions of each noise type, so the optimal policy will send one-third of the images to each segmentation model. Then we fine-tune the pre-trained RL model assuming it has learned an optimal policy. We do this by linearly increasing  $\lambda$  while measuring the total variation distance (TVD) from the optimal policy which was previously learned. Once this TVD hits a pre-specified threshold, we stop fine-tuning.

To understand the robustness of the PaSeR RL policy, we examine the model assignment confusion matrices in Fig. 4. Here, PaSeR (with a TVD threshold of 10%) has nearly perfect assignment of images to the  $f_0$  and  $f_1$  task models, while only sending 7.2% of images which should have gone to the  $f_2$  model to the  $f_1$  model. This occurs because of the 10% TVD threshold, which gives PaSeR the flexibility to send a small percentage of images to the  $f_1$  model instead of  $f_2$ . Comparing this to the model assignment of IDK-Cascade, we see that it sends 10% of  $f_1$  model images to  $f_2$ , while also incorrectly sending 7.2% of  $f_2$  model images to  $f_1$ . This is why IDK-Cascade cannot match the performance of PaSeR. The IDK-Cascade with entropy as the gating mechanism is not adaptable enough to accurately assign images to the best model. Finally, note that the PaSeR-RandPol. assigns images at random to each task model and thereby has the poorest performance across all metrics.

#### **R5:** Sensitivity to Hyperparameters

We now investigate how  $\lambda$  (cost parameter) and entropy map estimation affect PaSeR performance.

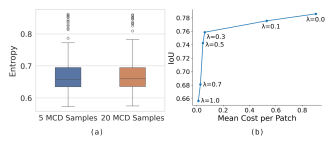


Figure 5: (a) Distribution of entropy estimates with 5 and 20 Monte Carlo Dropout (MCD) samples. (b) PaSeR IoU vs Mean Cost as  $\lambda$  changes on battery material phase segmentation dataset.

**Performance vs Cost Trade-off.** In Fig. 5(b), we show PaSeR's performance/cost trade-off curve as  $\lambda$  decreases for the battery segmentation task. The mean cost is calculated using Eq. 3. This cost function is based on the number of parameters in each task model with  $f_2$  having a significantly higher cost than  $f_1$ . As expected, as  $\lambda$  increases, mean cost falls and performance decreases. The sharp drop in cost between  $\lambda = 0.0$ . and  $\lambda = 0.3$  occurs because of the high difference in the cost of using the large task model  $f_2$  vs using the smaller models. As  $\lambda$  increases in this range, PaSeR uses  $f_2$  less, leading to a quick drop in mean cost.

Number Effect of of **MCDropout** Samples. PaSeR computes entropy maps using Monte Carlo (MC) dropout sampling which requires taking multiple samples of each prediction. To test the sensitivity of estimation of entropy to the number of MC samples taken, we show a box plot of the entropy distributions in Fig. 5(a). Comparing 5 MC dropout samples to 20 MC dropout samples shows no significant difference between the distributions of entropies. A t-test between these distributions gives a p-value of 0.6986, allowing us to safely assume these distributions are the same and use 5 MC samples in PaSeR for entropy estimation. We account for these 5 MCD samples in all our previous flops calculations.

#### Conclusion

In this work, we have developed a computationally parsimonious and more effective alternative to the IDK cascading decision pipeline and demonstrated that our proposed model PaSeR outperforms SOTA models on the task of battery material phase segmentation. We also propose a new metric IoU per GigaFlop which is useful for characterizing effectiveness of models to yield good predictions at low computational cost. Through various qualitative and quantitative results, we demonstrate that PaSeR yields a minimum performance improvement of 174% on the IoU/GigaFlop metric with respect to compared baselines. We also demonstrate PaSeR's adaptability to complementary models trained on the noisy MNIST dataset, where it outperforms all baselines on IoU/GigaFlop by a miniumum 13.4%. In the future, we shall extend PaSeR to incorporate other sophisticated cost metrics and test it in the context of multi-model pipelines comprised of data-driven and scientific simulation models.

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