

# SP<sup>2</sup>RINT: Spatially-Decoupled Physics-Inspired ProgRessive INverse Optimization for Scalable, PDE-Constrained Meta-Optical Neural Network Training

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## Abstract

Diffractive Optical Neural Networks (DONNs) harness the physics of light propagation to perform speed-of-light information processing. Training DONN systems to determine the metasurface structures remains a challenging problem. Heuristic methods are fast but oversimplify metasurfaces as element-wise phase masks or convolution units, often resulting in physically unrealizable designs. Accurate simulation-in-the-loop training methods directly optimize metasurfaces using adjoint inverse design methods during end-to-end DONN training, but are inherently computationally prohibitive. To address the DONN training in a physically feasible and scalable manner, we propose a spatially decoupled, progressive training scheme, SP<sup>2</sup>RINT. For the first time, we formulate DONN training as a partial differential equation (PDE)-constrained learning problem, where metasurface responses are relaxed into freely-trainable, banded transfer matrices. We then progressively enforce physical constraints through alternating transfer matrix training and inverse design. It eliminates the need for costly PDE solving per training step while ensuring physical realizability. To further alleviate the runtime bottleneck, we partition the metasurface into independently solvable patches and optimize the transfer matrix of each sub-region in parallel, followed by system calibration to restore global consistency. Across a range of DONN training tasks, SP<sup>2</sup>RINT achieves digital-comparable accuracy while being 1825 $\times$  faster than simulation-in-the-loop approaches. Grounded in a physics-inspired optimization, SP<sup>2</sup>RINT bridges the gap between abstract DONN models and physical metasurface designs and paves the way for scalable training of DONNs with guaranteed physical feasibility and high accuracy.

## 1 Introduction

Diffractive optical neural networks (DONNs) have emerged as a promising technology, leveraging the inherent parallelism for analog

processing at light-speed in diverse tasks such as computer vision, AI inference, scientific computing, sensing, and imaging [3, 5, 12–16, 18, 22, 24–26, 35, 41]. One emerging technology to implement compact, flat optical field manipulation in DONNs is optical metasurfaces. Metasurfaces are engineered planar optical elements, composed of arrays of subwavelength-scale structures known as meta-atoms.

However, training DONNs and designing physically feasible metasurface devices remain notable challenges, rooted in the complexities of optical physics. Traditional approaches to DONN design fall into two categories: *heuristic* approximation methods and *simulation-in-the-loop* training [11]. Heuristic methods simplify the metasurface as a set of element-wise phase masks or convolution kernels, often bypassing rigorous full-wave simulations. These methods translate the desired phase profile into a metasurface layout using a pre-simulated look-up table (LUT) that maps each target phase to a corresponding meta-atom design. Though efficient, this approach often neglects important inter-element interactions and relies heavily on the local periodic approximation (LPA) [35]. The LPA assumes a smooth near-field response, a condition that might hold in some lens designs but is frequently violated in DONNs where strong spatial variations are common. As a result, heuristic methods often produce target phase profiles or transfer matrices that are not physically realizable. When these idealized designs are projected onto feasible metasurface structures, the mismatch leads to significant deviations in optical behavior and degraded DONN performance. Efforts to mitigate this issue, e.g., by applying smoothness regularization [14, 31, 41] to phase profiles to ease hardware design, have generally been inadequate. While smoother profiles are simpler to implement, they significantly limit DONN expressivity and do not ensure true physical feasibility, often leading to impractical designs.

In contrast, simulation-in-the-loop training[11] embeds metasurface optimization directly within the training loop by leveraging adjoint methods. This approach effectively enforces physical feasibility at every training iteration. However, the demanding requirement of solving forward/adjoint Maxwell's equations for each metasurface at each iteration makes it prohibitively expensive and fundamentally unscalable.

To address these bottlenecks in DONN training, we propose SP<sup>2</sup>RINT, a physics-constrained optimization scheme that progressively enforces Maxwell PDE constraints during training. Unlike prior heuristic or simulation-in-the-loop approaches, SP<sup>2</sup>RINT alternates between relaxed DONN training and inverse metasurface projection, ensuring that learned responses remain physically realizable while maintaining efficient design space exploration.

To further reduce the significant computational cost, we develop a patch-wise simulation strategy. This strategy effectively exploits the locality of meta-atom interactions and inherent spatial frequency limits due to diffraction. This enables scalable metasurface inverse design with near-linear complexity, facilitating the design of physically implementable and scalable, large-capacity DONNs.

Our main contributions can be summarized as follows:

- We analyze the core limitations of existing DONN training methods, which suffer from modeling inaccuracies and prohibitive simulation cost. We introduce SP<sup>2</sup>RINT, a scalable PDE-constrained DONN optimization framework that ensures physical feasibility while drastically reducing simulation cost.
- **Spatially-Decoupled Metasurface Simulation:** SP<sup>2</sup>RINT exploits the locality of meta-atom interactions to divide the metasurface into patches, enabling scalable, parallel simulation and reducing complexity from cubic to near-linear.
- **Diffraction-Inspired Transfer Matrix Sketching:** By leveraging the natural smoothness of diffracted fields, we introduce a pulse-based sketching technique that accelerates transfer matrix probing without sacrificing accuracy.
- **Progressive PDE-Constrained Learning:** SP<sup>2</sup>RINT features a novel alternating scheme that interleaves relaxed DONN training with adjoint-based projection onto the Maxwell-constrained subspace, ensuring physical realizability throughout training while maintaining design flexibility and expressivity.
- On multiple DONN benchmarks, SP<sup>2</sup>RINT achieves up to **63.88%** higher accuracy and delivers an **1825×** speed-up compared to state-of-the-art simulation-in-the-loop training methods because of the patched transfer matrices probing and much fewer iterations of inverse design, bridging the gap between analytical DONN training and physically realizable meta-optic hardware.

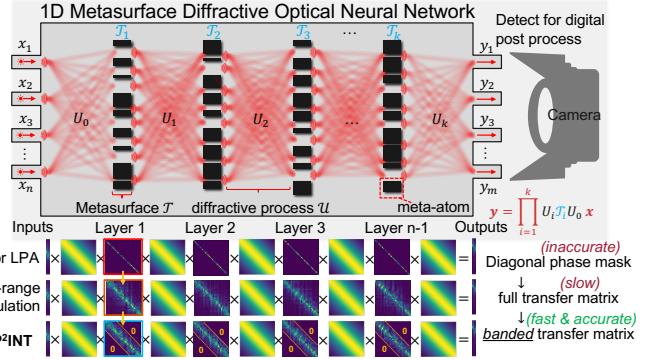
## 2 Background

This section introduces the physical principles of DONNs and existing DONN training methods.

### 2.1 Transfer Matrix Description of DONN Responses

Light propagation in DONNs involves two linear physical processes: free-space diffraction and metasurface-based light modulation, as illustrated in Fig. 1.

**Free-Space Diffraction.** Light diffraction in a homogeneous medium can be precisely modeled using a Green's function formulation, which provides an analytical near-to-far field transformation (N2FF). This representation naturally involves Hankel functions to describe cylindrical wave propagation [1].



**Figure 1: DONN with multi-layer metasurfaces can be modeled as cascaded transformation  $\mathcal{T}$  and diffraction  $U$ . Metasurface transfer matrix modeling comparison among 3 methods. Our SP<sup>2</sup>RINT uses banded transfer matrix probing for fast and accurate modeling.**

**Metasurface-based Modulation.** Metasurfaces modulate the phase and amplitude of incident light through engineered subwavelength structures. The precise response of a metasurface is governed by Maxwell's equations.

In Fig. 1, a  $K$ -layer DONN, comprising alternating layers of modulation and diffraction, can be described by the Transfer Matrix Method (TMM), which naturally enables a layer-wise decomposition of wave propagation,

$$f(\mathbf{x}_{in}; \epsilon) = \left| \left( \prod_{i=1}^K U_i(z) \mathcal{T}_i(\epsilon_i) \right) U_0 \mathbf{x}_{in} \right|^2 = |\mathbf{h}_{out}|^2. \quad (1)$$

where the inputs will be encoded to the intensity or phase of incident light  $\mathbf{x}_{in}$ ,  $U(z)$  is the light diffraction matrix over a distance  $z$ , and  $\mathcal{T}(\epsilon)$  is the metasurface modulation matrix parameterized by its permittivity distribution  $\epsilon$ . In prior work, based on LPA, a metasurface with  $n$  meta-atoms is modeled as an element-wise modulation plate with diagonal response  $\mathcal{T} = \text{diag}([A_1 e^{j\phi_1}, \dots, A_n e^{j\phi_n}])$ . At the end, the light will be converted to electrical signals via photodetector arrays with a square function applied  $|\cdot|^2$ . The metasurface transfer matrix  $\mathcal{T}(\epsilon)$  can be numerically obtained by stimulating the system with orthonormal basis vectors to capture its full impulse response.

### 2.2 Hybrid Diffractive Optical Neural Network

It is a promising trend to hybridize DONN with digital neural networks for adaptive AI inference [19, 34, 38]. Figure 1 shows a hybrid DONN with an optical feature extraction module and a lightweight digital neural head. As the light carrying the input length- $n$  vector propagates through multiple metasurfaces, it forms an intensity distribution on the detector plane. This intensity pattern is coupled out as  $m$ -channel readout signals. This  $m \times n$  linear unit can be used as a convolution with a single input channel, kernel size of  $n$ , and  $m$  output channels. The digital head can be trained for different downstream tasks.

### 2.3 Overview and Analysis of DONN Training Methods

**Heuristic DONN Training Methods.** Heuristic approaches simply the metasurface response, avoiding costly transfer matrix simulations during optimization, and implement the physical design post hoc [3–6, 8, 9, 12–16, 18, 20–22, 24–26, 30, 32, 33, 35, 36, 39, 41]. A widely used method models the metasurface  $\mathcal{T}$  as a diagonal matrix with trainable phase shifts. However, this diagonal assumption neglects

**Table 1: DONN training complexity.**  $E$  denotes the number of training epochs,  $N$  is the size of the training dataset,  $B$  is the inverse design iteration budget for each training epoch ( $B \ll N$ ), and  $n$  represents the spatial dimension of the metasurface.

DONN Training Methods		Algorithmic Complexity
Heuristics-based	LPA[20]	$\mathcal{O}(EN + 1)$
	Convolution[35]	$\mathcal{O}(EN + 1)$
Simulation-based	Simulation-in-the-loop[11]	$\mathcal{O}(ENn^3)$
Simulation-based	SP <sup>2</sup> RINT	$\mathcal{O}(EN + EBn)$

inter-element optical coupling, making the resulting  $\mathcal{T}$  physically unrealizable.

Another heuristic method accounts for meta-atom interactions by modeling the metasurface as a learned convolution kernel [35], still relying on LPA. The reliance on LPA still limits accuracy, especially in regimes where strong inter-element interference or non-periodic layouts cause the approximation to break down.

**Simulation-based Training Methods.** To ensure physical fidelity, simulation-in-the-loop methods embed full-wave simulations directly into the DONN training process [11]. At each iteration, gradients are computed via the adjoint method, which requires both forward and adjoint simulations[10]. While this approach provides accurate modeling and guarantees physical feasibility throughout training, its computational cost scales poorly with system size.

Table 1 compares the computational complexity of different DONN training paradigms, decomposed into two components: training cost (related to  $E, N$ ) and design cost (related to  $B, n$ ). SP<sup>2</sup>RINT offers a favorable trade-off by decoupling training from simulation. Its design cost scales linearly with metasurface size, thanks to the use of localized, patched transfer matrix probing. As a result, SP<sup>2</sup>RINT retains physical fidelity while remaining computationally tractable.

## 2.4 Metasurface Design Methods

Various methods have been proposed for metasurface design. A widely adopted approach is the LUT-based method [2, 7], where phase elements are independently selected from a pre-simulated library/LUT and simply ensembled to approximate a target phase profile. Adjoint method-based inverse design [23, 28, 29, 40] directly optimizes the design variables in high-dimensional space to maximize a certain objective function using adjoint gradients. However, adjoint-based methods rely on multiple computationally intensive full-wave optical simulations, which are prohibitively time-consuming. To clarify, DONN training is fundamentally more challenging than metasurface inverse design tasks. Inverse design solves a deterministic optimization problem for a specific figure-of-merit. In contrast, DONNs must learn metasurface designs on thousands of input-label pairs on the training dataset and generalize to the test set, making it a PDE-constrained learning task.

## 3 Proposed DONN Training Scheme SP<sup>2</sup>RINT

We aim to train DONNs under physically realistic conditions that obey Maxwell's equations. We formulate DONN training as a PDE-constrained subspace optimization problem. To address both physical realizability and scalability challenges, we introduce SP<sup>2</sup>RINT, a progressive training framework that alternates between relaxed DONN learning and adjoint-based inverse design. SP<sup>2</sup>RINT integrates two key innovations: **progressive soft projection**, which gradually enforces physical constraints while enabling efficient design space exploration,

and **patched transfer matrix probing**, which enables scalable modeling of large metasurfaces without resorting to full-system simulations.

## 3.1 Problem Formulation

Instead of being formulated as a deterministic optimization problem as in conventional inverse design, DONN training is formulated as a PDE-constrained learning problem:

$$\begin{aligned} \boldsymbol{\epsilon}^*, \mathbf{w}^* = \operatorname{argmin}_{\boldsymbol{\epsilon}, \mathbf{w}} \mathbb{E}_{(\mathbf{x}, \mathbf{y}) \sim \mathcal{D}} [\mathcal{L}(g_w \circ f(\mathbf{x}; \boldsymbol{\epsilon}), \mathbf{y})], \quad f(\mathbf{x}; \boldsymbol{\epsilon}) = |\mathbf{h}|^2, \\ \text{s.t. } (\nabla \times (\boldsymbol{\epsilon}_{\text{tot}}^{-1} \nabla \times) - \omega^2 \mu_0 \epsilon_0) \mathbf{h} = \mathbf{b}(\mathbf{x}) \rightarrow A(\boldsymbol{\epsilon}) \mathbf{h} = \mathbf{b}(\mathbf{x}). \end{aligned} \quad (2)$$

Here,  $f(\mathbf{x}; \boldsymbol{\epsilon})$  models the diffractive optical system, its output being the light intensity detected on photodetector arrays (for TM polarized light, this intensity is proportional to  $|\mathbf{h}|^2$ , where  $\mathbf{h}$  is the magnetic field). The function  $g_w(\cdot)$  represents the subsequent digital processing head, encompassing operations like normalization, biasing, and the final digital classification layers, with  $w$  as its learnable variables. The primary trainable parameters for the optical system are the  $k$  metasurface structures, characterized by their permittivity distributions  $\boldsymbol{\epsilon} = (\epsilon_1, \dots, \epsilon_k)$ . Input data and label pairs  $(\mathbf{x}, \mathbf{y})$  are drawn from the training dataset  $\mathcal{D}$ , and  $\mathcal{L}$  is the chosen loss function. The magnetic field  $\mathbf{h}$  follows Maxwell's equation, which sets a challenging Maxwell PDE constraint to the optimization problem.  $\boldsymbol{\epsilon}_{\text{tot}}$  describes the entire optical system, including multiple cascaded metasurfaces and their spacing cladding. For simplicity, we use a linear equation for the PDE constraint  $A(\boldsymbol{\epsilon}) \mathbf{h} = \mathbf{b}(\mathbf{x})$ , where  $\mathbf{h}$  is the vectorized magnetic field and  $\mathbf{b}(\mathbf{x})$  is the vectorized input source related to  $\mathbf{x}$ .

This PDE-constrained formulation dictates that each iteration requires solving the forward Maxwell equation to obtain  $\mathbf{h}$ , followed by solving an adjoint Maxwell equation to compute the gradients  $\frac{d\mathcal{L}}{d\boldsymbol{\epsilon}}$  with respect to the metasurface parameters:

$$A(\boldsymbol{\epsilon})^\top \mathbf{h}_{\text{adj}} = -\frac{d\mathcal{L}}{d\mathbf{h}}, \quad \frac{d\mathcal{L}}{d\boldsymbol{\epsilon}} = -\mathfrak{R}(\mathbf{h}_{\text{adj}}^\top \mathbf{h}). \quad (3)$$

However, rigorous full-wave simulation of the entire optical system is prohibitively time-consuming, making it impractical to embed this simulation within the outer training loop.

## 3.2 Understanding Difficulty in Designing Implementable DONNs

**3.2.1 The Essence of the PDE Constraint.** As shown in Fig. 1 and Eq. (1), each metasurface with permittivity  $\epsilon_i$  has a transfer matrix  $\mathcal{T}_i(\epsilon_i)$  that maps the input  $\mathbf{x}_i$  to the output field near the metasurface. However, not every arbitrary transfer matrix  $\mathcal{T}$  can find its physical metasurface implementation. **Therefore, training a DONN that adheres to Maxwell's equations inherently becomes a constrained optimization problem, where the learned transfer matrices  $\widehat{\mathcal{T}}_i(\epsilon_i)$  must reside within the subspace of physically implementable transformations**, which we refer to as the *implementable subspace*.

**3.2.2 Training Difficulties.** We identify **three key difficulties** that hinder the design of physically implementable and high-performance DONNs: **1 Characterizing the Valid Subspace:** The intricate optical physics make a precise analytical characterization of the implementable subspace unavailable, hence it is difficult to guide the optimization effectively. **2 Prohibitive Simulation Cost:** Directly enforcing physical realizability by repeatedly solving PDEs for each metasurface design during training is computationally formidable. **3 Non-Convex Optimization Landscape:** The subspace of physically

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**Algorithm 1:** Progressive Projected Training for DONN

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**Input:** DONN model  $\mathcal{M}_{\widehat{\mathcal{T}}, w}$ , use  $\mathcal{M}$  for short; Training set  $\mathcal{D}(x, y)$ ; Uniform metasurface init  $\epsilon^0$ ; Initial and final binarization sharpness:  $s_0, s_T$ ; Inverse design iteration budget per epoch  $B$ , number of inverse design iteration per projection  $I$ ; Number of metasurface layers  $K$ ; patch size  $P$

**Output:** Trained model with implementable metasurfaces

```

1  $\epsilon^0 \leftarrow [\epsilon^0] \times K; n_p \leftarrow B/I; S \leftarrow \text{Sched}(s_0, s_T, B)$ 
2 for epoch  $t \leftarrow 1$  to  $n_{\text{epochs}}$  do // Sec. 3.3
3    $\epsilon^t \leftarrow \epsilon^{t-1};$ 
4    $\widehat{\mathcal{T}}^t \leftarrow \text{ProbeTM}(\epsilon^t, s_T, P);$  // Sec. 3.4
5    $\mathcal{M}.\text{SetTM}(\widehat{\mathcal{T}}^t, k), \forall k \in [K];$  // Load  $\mathcal{T}$  for  $k^{\text{th}}$  layer
6   foreach batch index  $j, (x, y)$  in  $\mathcal{D}$  do
7     TrainStep( $\mathcal{M}, x, y$ ); // Train  $\mathcal{M}$ 
8     if  $j \bmod b_p = 0$  and  $j \neq 0$  then
9       |  $\epsilon_i^t \leftarrow \text{Proj}(\widehat{\mathcal{T}}_i^t), \forall i \in [K];$  // Solve Eq.(6)
10       $\epsilon^t \leftarrow \text{Proj}(\widehat{\mathcal{T}}_{\text{tot}} \text{ of } \mathcal{M});$  // Solve Eq. (6) and (7)
11       $S.\text{reset}();$  // Sec. 3.5

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implementable transfer matrices  $\mathcal{T}_i$  is highly non-convex, which significantly increases the risk of the training process prematurely converging to bad local optima. This, in turn, curtails effective exploration of the vast design space and impedes the discovery of high-expressivity metasurface designs.

To effectively address these challenges, several fundamental **questions** must first be answered:

**Q1. How to effectively restrict transfer matrices within the implementable subspace?** While *penalty methods* require unavailable analytical subspace characterization and *reparametrization methods* require unaffordable PDE solving per iteration, we choose to use **progressive projection** (i.e., **periodically perform inverse design**) to balance efficiency and PDE constraint compliance.

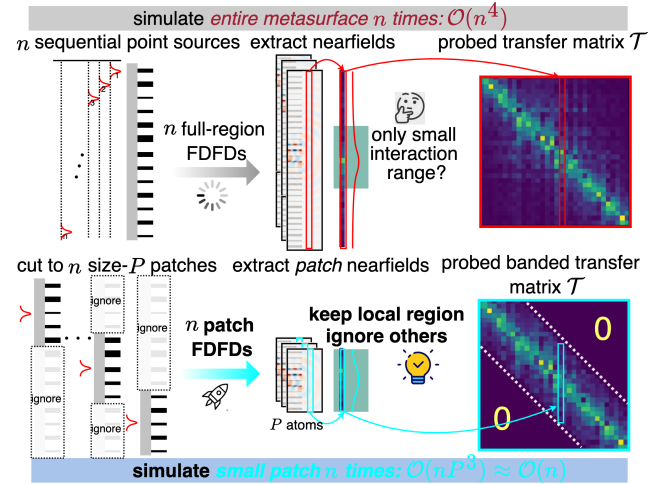
**Q2. How can we avoid per-iteration, unscaleable PDE solving?** While TMM allows us to decompose full-system simulation into layer-wise subproblems, each layer still requires expensive forward/adjoint simulations on the whole metasurface. To make training scalable, we must **decouple field evaluation  $h_i$  from design updates on  $\epsilon_i$  and solve PDEs only when necessary**.

**Q3. How to perform projection to balance exploration and efficiency?** Since metasurface inverse design is inherently a slow and difficult binary optimization problem, effective training requires a carefully scheduled projection strategy. *Hard projection* strictly enforces feasibility but limits exploration and increases simulation cost, while *soft projection* improves efficiency and optimization flexibility but risks constraint violation. **To ensure fast convergence and maintain physical realizability, we need to balance these trade-offs through progressive projection that gradually tightens constraints during training.**

### 3.3 Overview of Proposed DONN Training Flow

To efficiently solve Eq. (2), we propose a *variable separation* method to decouple the transfer matrix  $\widehat{\mathcal{T}}$  used in DONN training from the metasurface response  $\mathcal{T}$  that requires  $n$  full-metasurface simulations to extract. The formulation is rewritten as

$$\begin{aligned} \widehat{\mathcal{T}}^*, w^* = & \underset{\widehat{\mathcal{T}}, w}{\text{argmin}} \mathbb{E}_{(x, y) \sim \mathcal{D}} [\mathcal{L}(g_w \circ f(x; \widehat{\mathcal{T}}), y)], \\ \text{s.t. } A(\epsilon_i) \widehat{\mathcal{T}}_i &= B, \forall i \in [K]; \widehat{\mathcal{T}}_i = \mathcal{T}_i, \forall i \in [K]. \end{aligned} \quad (4)$$



**Figure 2: Proposed spatially-decoupled transfer matrix probing method cuts the metasurface into small patches for patch simulation that reduces complexity from cubic to linear.**

To solve this constrained optimization problem, the projected gradient descent can be adopted, as shown in Alg. 1, where we alternatively solve two subproblems. The first subproblem is **1 Relaxed DONN Training**:

$$\widehat{\mathcal{T}}^{t+1}, w^{t+1} = \underset{\widehat{\mathcal{T}}, w}{\text{argmin}} \mathbb{E}_{(x, y) \sim \mathcal{D}} [\mathcal{L}(g_w \circ f(x; \widehat{\mathcal{T}}), y)]. \quad (5)$$

The second subproblem is **2 Metasurface Inverse Design**: optimizing the metasurface permittivity  $\epsilon$  to match the target matrix  $\mathcal{T}$ :

$$\begin{aligned} \epsilon^{t+1} = & \underset{\epsilon}{\text{argmin}} \sum_{i=1}^K \|\mathcal{T}_i - \widehat{\mathcal{T}}_i^{t+1}\|_F^2, \\ \text{s.t. } A(\epsilon_i) \mathcal{T}_i &= B, \forall i \in [K]. \end{aligned} \quad (6)$$

This subproblem is very time-consuming. The following section discusses how we efficiently solve **2**.

### 3.4 Spatially-Decoupled Metasurface Simulation for Scalable Transfer Matrix Probing

In subproblem **2**, we need to extract  $\mathcal{T}_i$  for all  $k$  metasurfaces by solving  $nk$  full-metasurface Maxwell equations in total. Based on a strong physics prior that meta-atom responses often have restricted interaction distance, we can observe a clear **banded diagonal** structure on the transfer matrix  $\mathcal{T}$ . In other words, the light source shone on one meta-atom will not scatter to far-away locations in the near field, such that we only need to simulate a small region and ignore all light transmission in the exterior region up to a certain acceptable approximation error, as shown in Fig. 2. Hence, we propose a spatially-decoupled metasurface simulation approach by cutting a large metasurface into overlapping patches of  $P$  meta-atoms and a stride of 1 meta-atom. For a metasurface with  $n$  meta-atoms, we only need to simulate  $n$  small size- $P$  patches, which significantly reduces the cost when extracting the transfer matrix. More importantly, the simulation cost now **scales linearly with the size of the metasurface** if patches are simulated sequentially, or even remains constant runtime if patches are simulated in parallel as they are fully decoupled, transforming an otherwise cubic complexity into a much more scalable approach.

### 3.5 Progressive PDE-Constrained Training

It is critical to carefully schedule the alternating frequency between **1** and **2** as well as the binarization sharpness in **2**. As shown in

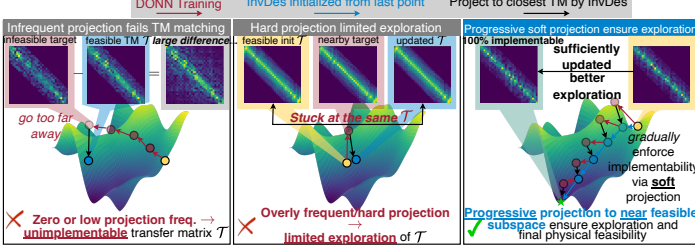


Figure 3: Our proposed SP<sup>2</sup>RINT framework enables both exploitation and physical feasibility.

Fig. 3, a full epoch of unconstrained training often causes  $\widehat{\mathcal{T}}_i$  to drift too far from the subspace, such that the subsequent projection step can no longer find a physically realizable metasurface that accurately recovers the target response. This results in a severe performance drop and even divergence. Increasing the projection frequency helps prevent  $\widehat{\mathcal{T}}_i$  from drifting too far from the subspace. However, due to the highly non-convex nature of the subspace, as discussed in Sec. 3.2.2, overly frequent hard projection often causes the optimization to get stuck near the initial point without meaningful progress, shown in Fig. 3. To resolve this, we introduce a *progressive PDE-constrained training* strategy. Rather than enforcing hard binarization from the start, we allow the inverse design module to initially project to relaxed, continuous-valued patterns that lie near the subspace. As training proceeds, we gradually tighten the binary constraint, ensuring the final metasurfaces are both expressive and physically realizable.

### 3.6 Extra System-Level Fine-Tuning for Enhanced Optimality

As shown in Alg. 1 Line 10, besides matching  $\mathcal{T}_i$  of each individual metasurface (Eq.(6)), we introduce an extra fine-tuning stage to further calibrate the response of the entire optical system. This is achieved by optimizing the metasurface  $\epsilon$  to match the end-to-end system-level transfer matrix,

$$\epsilon^{t+1} = \arg \min_{\epsilon} \left\| \mathcal{T}_{\text{tot}} S - \widehat{\mathcal{T}}_{\text{tot}}^{t+1} S \right\|_F^2, \quad \mathcal{T}_{\text{tot}} = \prod_{i=1}^K U_i \mathcal{T}_i(\epsilon_i), \quad (7)$$

$$\text{s.t. } A(\epsilon_i) \mathcal{T}_i = B, \quad \forall i \in [K].$$

$\mathcal{T}_{\text{tot}}$  represents the total transfer matrix of the multi-layer DONN system. The target total transfer matrix,  $\widehat{\mathcal{T}}_{\text{tot}}^{t+1}$ , is constructed from the layer-wise target modulation matrices  $\widehat{\mathcal{T}}_i^{t+1}$  obtained from the DONN training stage shown in Eq. (5), typically as  $\widehat{\mathcal{T}}_{\text{tot}}^{t+1} = \prod_{i=1}^K U_i \widehat{\mathcal{T}}_i^{t+1}$ . The matrix  $S$  consists of a set of one-hot probe stimuli. This system-level fine-tuning enables improved alignment with the desired global optical response and ultimately enhances end-to-end DONN performance.

## 4 Evaluation

### 4.1 Evaluation Settings

The hybrid DONN for classification tasks comprises an optical feature extractor and a lightweight digital head. The optical feature extractor has 2 cascaded metasurfaces and operates as a  $3 \times 3$  convolution (9 in-ports) with square nonlinearity, a single input channel, and 4 output channels (4 out-ports). For RGB images, we convert them to grayscale. Input pixels within the convolution window are phase-encoded ( $[0-1]$  maps to  $[0, \pi]$ ). For simplicity, we adopt a one-dimensional (1D) metasurface configuration, where each metasurface consists of 32 Si meta-atoms with air cladding, a period of 300 nm and a pillar height of 750 nm. The pillar widths of the meta-atoms are trainable parameters. The diffraction distances  $z$  between layers are all set to 4  $\mu\text{m}$ . The

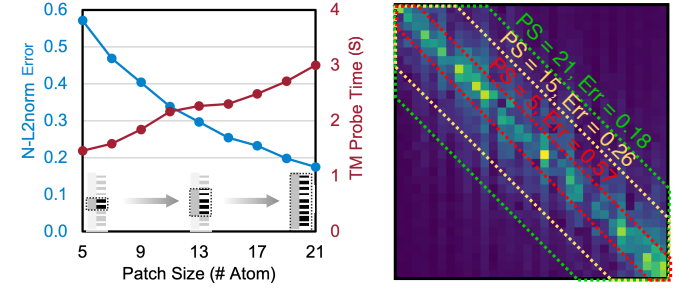


Figure 4: Different patch size trades off transfer matrix probing error and runtime.

Table 2: Comparison across different DONN training methods on different benchmarks, our proposed SP<sup>2</sup>RINT consistently achieves best performance. *Sim-in-the-loop* method has low accuracy after epoch 1 and takes 2 years to finish 100 epochs.

Benchmark	Baselines	Train CE	Train Acc	Test CE	Test Acc
Fashion-MNIST[37]	SP <sup>2</sup> RINT	9.00E-02	96.85%	0.45	88.44%
	LPA[20]	1.52E-01	94.77%	2.25	58.79%
	conv LPA [35]	1.69E-01	94.11%	6.40	13.45%
	smoothed metasurface [14]	2.39E-02	99.45%	22.61	16.21%
	sim-in-the-loop [11]	Time out	Time out	Time out	Time out
avg. improv.		+66.67%			
SVHN[27]	SP <sup>2</sup> RINT	3.16E-01	91.03%	0.76	81.61%
	LPA[20]	2.99E-01	91.70%	6.71	23.92%
	conv LPA [35]	2.21E-01	93.67%	2.51	11.51%
	smoothed metasurface [14]	5.98E-02	98.55%	5.04	7.23%
	sim-in-the-loop [11]	Time out	Time out	Time out	Time out
avg. improv.		+82.58%			
Darcy Flow[17]		Train N-L2norm		Test N-L2norm	
	SP <sup>2</sup> RINT	0.37		0.35	
	LPA[20]	0.32		0.60	
	conv LPA [35]	0.33		0.87	
	smoothed metasurface [14]	0.33		0.48	
avg. improv.		+42.40%			
total avg. improv.		+63.88%			

wavelength is fixed at 850 nm. The digital head is simply a 4-channel 1x1 convolution followed by a pooling layer and 2 linear layers.

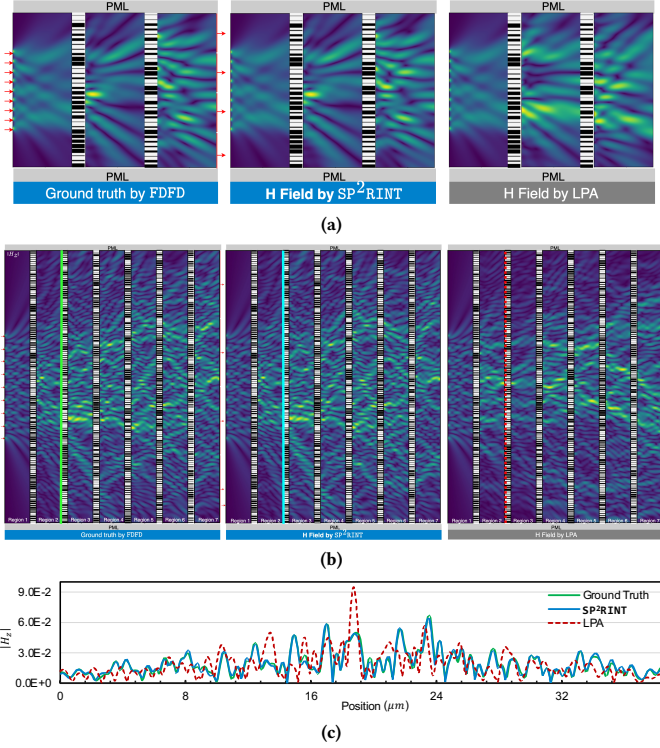
### 4.2 Patch Size Selection: Efficiency vs. Accuracy Trade-off

We first determine a critical hyperparameter: the patch size in patched TM probing. In Fig. 4, larger patch sizes improve the fidelity of transfer matrix probing by capturing more near-field interactions, but this comes at the cost of increased simulation time. To balance probing accuracy and efficiency, we set the patch size to 17 meta-atoms in all experiments. It is also important to note that the optimal patch size depends on factors such as the wavelength, the meta-atom material, and the diffraction distance between the metasurface and the observation plane.

### 4.3 Main Result

Table 2 compares different DONN training methods. Fashion-MNIST[37] and SVHN[27] represent image recognition (metric is *accuracy* and *cross-entropy (CE) loss*), while Darcy Flow[17] shows scientific machine learning for solving PDEs (metric is *normalized L2-Norm*). For each method, we evaluate the test set performance using the final trained DONNs with real simulated responses of implemented metasurfaces.

We compare SP<sup>2</sup>RINT with three heuristic methods (LPA, conv LPA, and smoothed) and simulation-in-the-loop training. The *Smoothed Metasurface* method uses a regularization term to enforce similar meta-atom sizes across neighbors [41].



**Figure 5:  $|H_z|$  fields comparison on the 9-in-4-out metasurface system. (a) 2-layer 32-metaatom, (b) 6-layer 128-metaatom, (c) Magnetic field  $H_z$  slice comparison. Our  $SP^2RINT$  can accurately capture the metasurface transfer matrix, much more accurate than simple phase mask modeling using LPA, yielding almost the same  $H$  field amplitude calculated by FDFD.**

To clarify, train-test performance gap is mainly due to *metasurface modeling error* and *NN generalization error*. Since all heuristic methods oversimplify the complex metasurface responses, the optimized transfer matrices are *not physically realizable*, hence they exhibit significant performance degradation when mapped to real metasurfaces during inference. The huge drop is mostly due to inaccurate metasurface modeling. In contrast, the optimized transfer matrices produced by  $SP^2RINT$  are guaranteed to lie within the implementable subspace. Hence, the training accuracy evaluated using  $\hat{\mathcal{T}}$  remains almost the same when mapped to real metasurface responses  $\mathcal{T}(\epsilon)$ . Our method eliminates the metasurface modeling error (our main goal) and achieves the best test accuracy up to the inevitable train-test generalization gap (not the issue we aim to solve in this work). Figure 5 visually compares the estimated and simulated optical fields within the diffractive feature extractor during inference.  $SP^2RINT$  accurately reproduces the field calculated by FDFD simulation, whereas LPA shows a significant field approximation error.

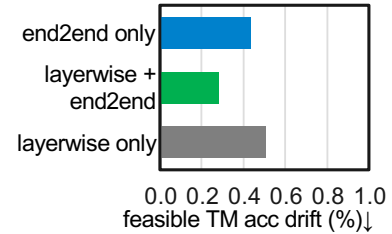
For the simulation-in-the-loop method[11], we estimate the runtime for a single epoch to be  $50k/32 \times (2 \text{ metasurfaces}) \times (32 \text{ sims/TMProb}) \times 6.4s/\text{sim} = \sim 178$  hours for Fashion-MNIST with batch size 32, due to the large number of full-wave simulations involved, making it prohibitively time-consuming. It needs 2 years to finish 100 epochs. As a result, we consider this intractable and do not report results for this baseline.  $SP^2RINT$  only takes 5.85 min per epoch, which directly shows 1825 $\times$  speedup.

**Table 3: Compare different projection sharpness schedules.** Progressively tightening the projection from a soft to a hard-binariization device ensures both exploration and physical feasibility. **CE** is Cross-Entropy.

Sharpness Schedule	Per Projection	Per Epoch	Per Train
CE Loss $\downarrow$	0.579	<b>0.428</b>	0.954
Accuracy $\uparrow$	86.19%	<b>88.32%</b>	80.82%
Transfer Matrix			

**Table 4: Comparison between the projection schedule in one training epoch.** By default, we define 1 unit budget as 20 iterations. Each iteration includes 2 times of forward transfer matrices probing and corresponding adjoint simulation.

Cost	Proj. frequency	Proj. iters	CE $\downarrow$	Accuracy $\uparrow$
budget $\times 1$	1 / epoch	20	1.190	56.04%
	2 / epoch	10	0.939	71.68%
	4 / epoch	5	0.360	87.59%
	5 / epoch	4	0.444	84.99%
	<b>10 / epoch</b>	<b>2</b>	0.352	89.54%
	20 / epoch	1	0.354	88.64%
budget $\times 1/2$	5 / epoch	2	0.514	87.64%
budget $\times 1/2$	10 / epoch	1	0.406	88.76%
budget $\times 2$	10 / epoch	4	0.452	83.91%

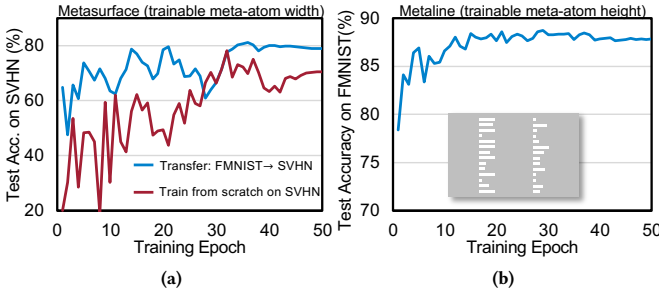


**Figure 6: Layerwise+end-to-end inverse design gives the best projection accuracy.**

#### 4.4 Discussion on Projection Configurations

**4.4.1 Progressive Soft Projection Schedule.** As shown in Table 3, when strict binarization is enforced at every projection step, the transfer matrix of metasurfaces stagnates with minimal updates compared to initialization. If the binarization is only enforced at the end of the training process, it over-relaxes the inverse design process (i.e., projection) and fails to adjust model weights to tolerate the permittivity binarization effects, thus leading to low accuracy after convergence. Our proposed progressive soft projection gradually increases the binarization sharpness over time, allowing the metasurface to evolve meaningfully and contribute to higher classification accuracy, while still ensuring implementability through the final binarization.

**4.4.2 Projection Frequency.** Table 4 compares different projection frequencies under different optimization runtime budgets. Increasing the projection frequency while proportionally reducing the number of iterations per projection maintains the overall runtime but significantly improves test accuracy. In our setting, we use 20 inverse design iterations per epoch with 10 projections per epoch, i.e., 2 adjoint gradient updates per projection.



**Figure 7: (a) Transfer learning of DONNs from Fashion-MNIST to SVHN shows faster convergence and higher performance than training on SVHN from scratch. (b) SP<sup>2</sup>RINT can be generally applied to other devices, such as metalines with trainable meta-atom height.**

**4.4.3 Inverse Design Projection Objective.** Figure 6 shows how different projection objectives impact the performance drift from unconstrained target performance to the feasible performance. Layer-wise projection with our extra system-level fine-tuning effectively mitigates the performance drift caused by the inverse design imperfection.

**4.4.4 Transfer Learning from Fashion-MNIST to SVHN.** To demonstrate the effectiveness of the learned optical feature extractor, we performed transfer learning from Fashion-MNIST to SVHN. Despite the visual differences between the datasets, one consisting of grayscale images of clothing and the other of colorful house numbers, the transferred model shows faster convergence and improved performance compared to training from scratch, as shown in Fig. 7(a).

**4.4.5 Generalization to Other Types of Meta-Optic Structures.** Beyond metasurfaces, where light modulation is realized by varying the width of meta-atoms with fixed height, alternative approaches exist to control light behavior. For example, metalines achieve light modulation by fixing the width of meta-atoms and varying their height. Our proposed SP<sup>2</sup>RINT framework is compatible with such architectures and can be used to train DONNs implemented with metalines. This is demonstrated in Fig. 7(b), which shows the training dynamics of a metiline-based DONN.

## 5 Conclusion

We propose SP<sup>2</sup>RINT, a scalable and physically grounded framework for training DONN. SP<sup>2</sup>RINT combines PDE-constrained optimization with spatially decoupled transfer matrix probing and progressive projection to ensure physical realizability without requiring per-iteration full-wave simulations. Evaluation shows that SP<sup>2</sup>RINT achieves average test accuracy improvements of 63.88% over widely used heuristic training methods and delivers 1825 $\times$  speed-up compared to the simulation-in-the-loop training. These results highlight SP<sup>2</sup>RINT’s potential to bridge the gap between high-performance deep learning and physically implementable nanophotonic hardware, enabling scalable, generalizable, and deployable optical neural systems.

## 6 Acknowledgment

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