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Machine-learning-based inverse design of wide-band metasurfaces with interdependent unit cells

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E-mail: zernikephaseplate@outlook.com**Keywords:** nanophotonics, metasurface, machine learning, Zernike phase maskSupplementary material for this article is available [online](#)

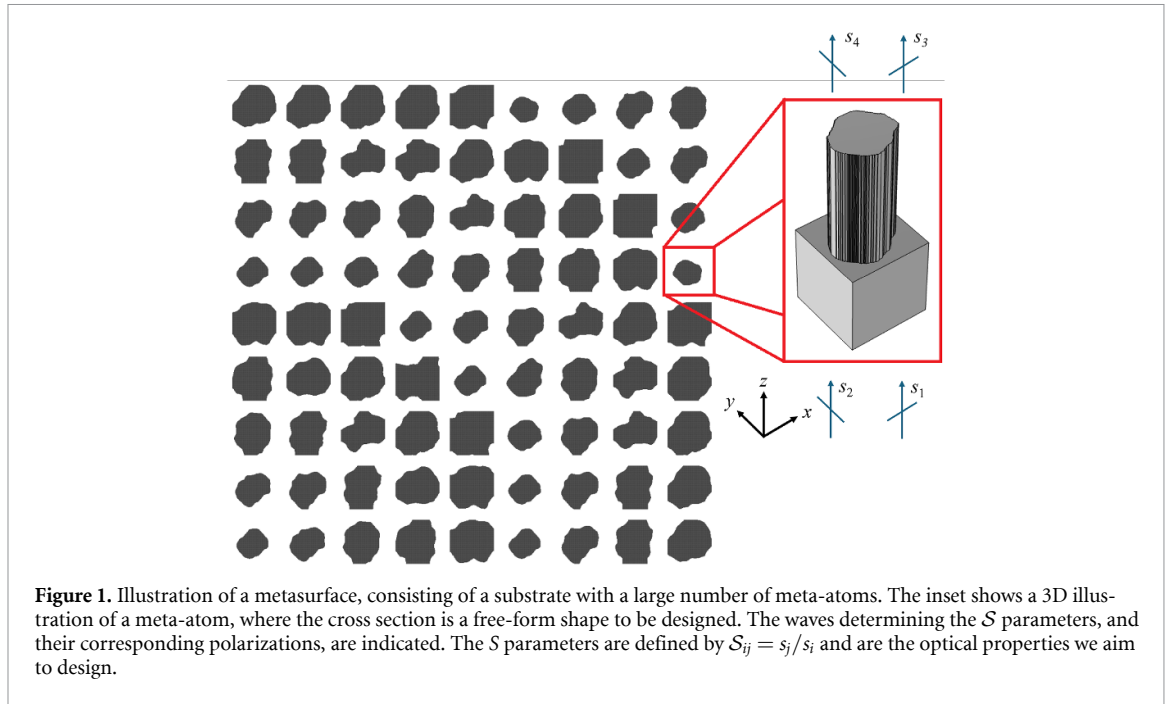
Abstract

Metasurfaces are thin optical materials consisting of tiny meta-atoms that locally change the phase, amplitude, and polarization of light. Such metasurfaces have been shown to exhibit optical responses beyond what is possible with natural materials, e.g. in thin achromatic lenses, holograms, reflectors, and the generation of structured light. A considerable challenge lies in designing the sometimes thousands of meta-atoms that are needed to locally modify the wavefronts of light. Here we use a conditional generative adversarial network to demonstrate for the first time a way to design metasurfaces where the scattering responses of different meta-atoms are interdependent, i.e. some of the individual scattering parameters are unimportant, and it is the relative values between different meta-atoms that matter. We illustrate our approach with a design of a broadband phase mask for a Zernike wavefront sensor, with complex requirements on the phase relationship between the scattering parameters of an inner disk and the surrounding region. Our inverse design method offers increased design freedom, since it does not require specifying absolute scattering parameters, and allows for metasurfaces with large bandwidths.

1. Introduction

Optical metasurfaces are a new class of artificial materials that can manipulate light in unprecedented ways [1–5]. They consist of arrays of nanostructures—also called meta-atoms—that are much smaller than the wavelength of light and can locally modify the phase, amplitude, and polarization of incident light (see figure 1 for an illustration). Optical metasurfaces have many advantages over conventional optical elements, such as ultra-small thickness, high efficiency, and design flexibility. They have enabled the realization of various novel optical devices and functionalities, such as polarization detection [6] and manipulation [7, 8], image processing [9, 10], wavefront control [11, 12], beam steering [13] and focusing [14–16], and the generation and manipulation of beams with optical angular momentum [17]. Optical metasurfaces are expected to have a wide range of applications in optics, photonics, information technology, and biomedicine.

Metasurfaces often act as a phase or amplitude mask, where the phase or amplitude of each meta-atom is related. For example, a quadratic phase profile creates a lens focusing a light beam [18, 19]. The absolute phase of each meta-atom is unimportant; only their relative phase differences matter. In this paper, we will present a way to generate meta-atoms with interdependent properties without fixing the value of phase or amplitude in transmission (or reflection) of any individual meta-atom. This opens up substantial design freedom, which is impossible to achieve with human intuition. However, it is a challenge to determine the shape of the meta-atoms needed for a given desired optical response. Often, thousands of different meta-atoms must be designed and this is especially difficult for multiobjective metasurfaces, as there are no simple forward design techniques available [20–22]. To achieve such inverse design of metasurfaces, we use deep neural networks (DNNs), which are a powerful machine learning



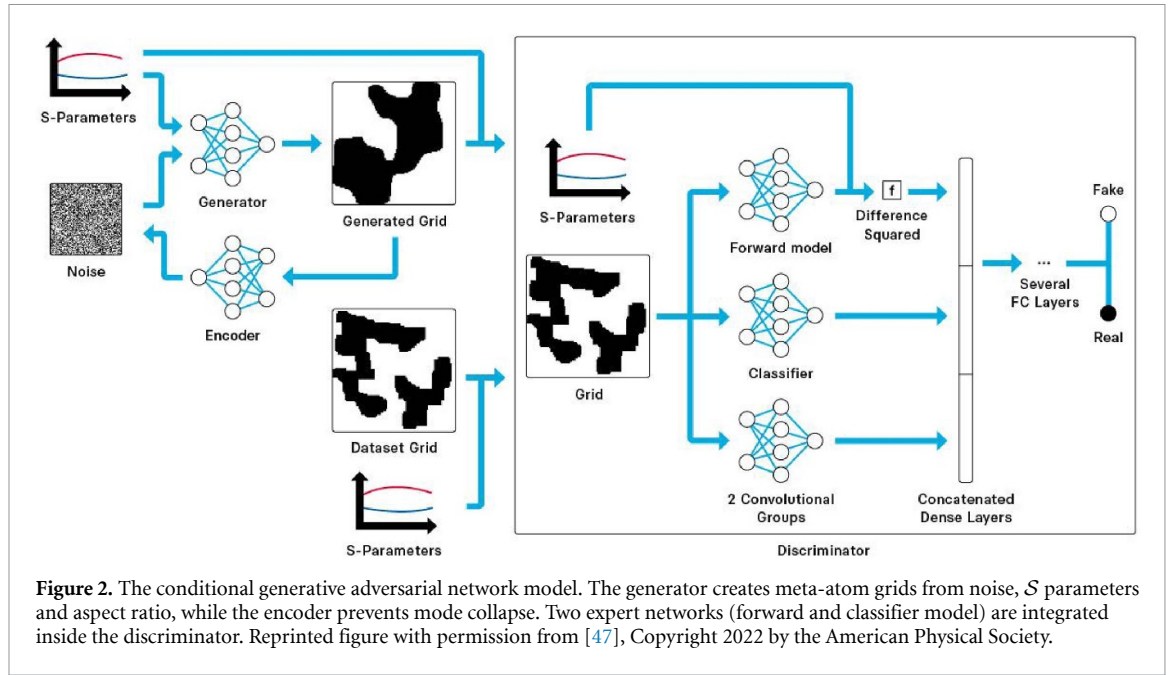
architecture [23–25] that can be used to learn complex patterns and functions from data, including to learn laws of physics [26, 27]. They have been widely used for various tasks such as image recognition [28, 29], natural language processing [30], and recommendation systems [31]. More recently, neural-network models have been used to achieve inverse design. This is often implemented by constructing a surrogate model that predicts physical properties for a device with arbitrary geometry [32–34], potentially aided by dimensionality reduction [35], manifold learning [36], or physics-informed networks [37–39]. Such models can then be inverted using a tandem network [40–42], an autoencoder [43, 44], or a conditional generative adversarial network (CGAN) [45–49]. A key advantage of DNNs in the design of physical devices is their ability to quickly generate designs for devices with any desired physical response. For example, in an optical metasurface we can design meta-atoms with any desired transmission amplitude and phase in the subset of physically realizable scattering parameters once trained, eliminating the need to run an optimization for each meta-atom. The training of neural networks requires a large number of simulations of the forward problem, but these simulations are independent, meaning they can be run simultaneously on high-performance computer clusters.

In this paper, we study a novel conditional generative adversarial network (CGAN) graph configuration for the inverse design of metasurface unit cells with interdependent properties, i.e. with design requirements including relative differences between scattering parameters of different meta-atoms. This implies that we must co-design all the meta-atoms, often thousands of meta-atoms, simultaneously. This provides several advantages over other inverse design techniques: (1) We get more design freedom, since we do not need to arbitrarily fix an absolute value of the \mathcal{S} parameters at a specific location, which is important, e.g. for lenses where only relative phase differences are really required. (2) It allows us to design metasurfaces with large bandwidths, which we will demonstrate for a Zernike phase mask. (3) We can find designs that were untractable until now.

We will first present a graph for a CGAN scheme that can co-design all meta-atoms. Subsequently, we will discuss the training of the CGAN model. After we have presented our CGAN model, we will illustrate our CGAN on a problem that was untractable until now: a Zernike phase mask. This problem requires relative phase differences between meta-atoms in two different regions and also has a complex set of design requirements that cannot be solved by a simple parameter map or by human intuition.

2. Results

Our graph is based on a CGAN for the inverse design of individual meta-atoms [47], shown in figure 2, which is then adapted to enable the codesign of many meta-atoms. The machine learning method works as follows. Initially, a forward model \mathcal{F} is established to approximate the optical properties of meta-atoms as a function of the grid representing the lithographic mask. This forward model aids in accelerating the assessment of optical properties compared to direct full-wave solutions. A binary grid is utilized



to represent the lithographic mask U of a meta-atom, indicating where the top layer is etched away and where not. Due to similarities with pixelated images, a DenseNet [50] is used to model the scattering parameters. This model is trained with about 80 000 semi-random grids created and loaded into a finite-element simulator to obtain the scattering parameters. We also use a classifier network \mathcal{C} to estimate the fabrication feasibility as described in reference [47]. The two expert models (\mathcal{F} and \mathcal{C}) are embedded into the discriminator of a CGAN by feeding the unit cell U to the two networks and their output to a hidden layer of the discriminator after computing the metrics of the model, such as evaluating the difference and difference squared of the desired/ground truth S parameters and the by the forward model approximated S parameters $\mathcal{F}(U)$.

The graph setup of this CGAN model is given by

$$\begin{aligned} \mathcal{G}[\mathcal{N}, \mathcal{S}, \mathcal{A}] &= U_G, \\ \mathcal{D}[U, \mathcal{S}, \mathcal{A}] &= \mathcal{D}[U, f(\mathcal{F}(U), \mathcal{S}), t(\mathcal{C}(U), \mathcal{A})] = O, \end{aligned} \quad (1)$$

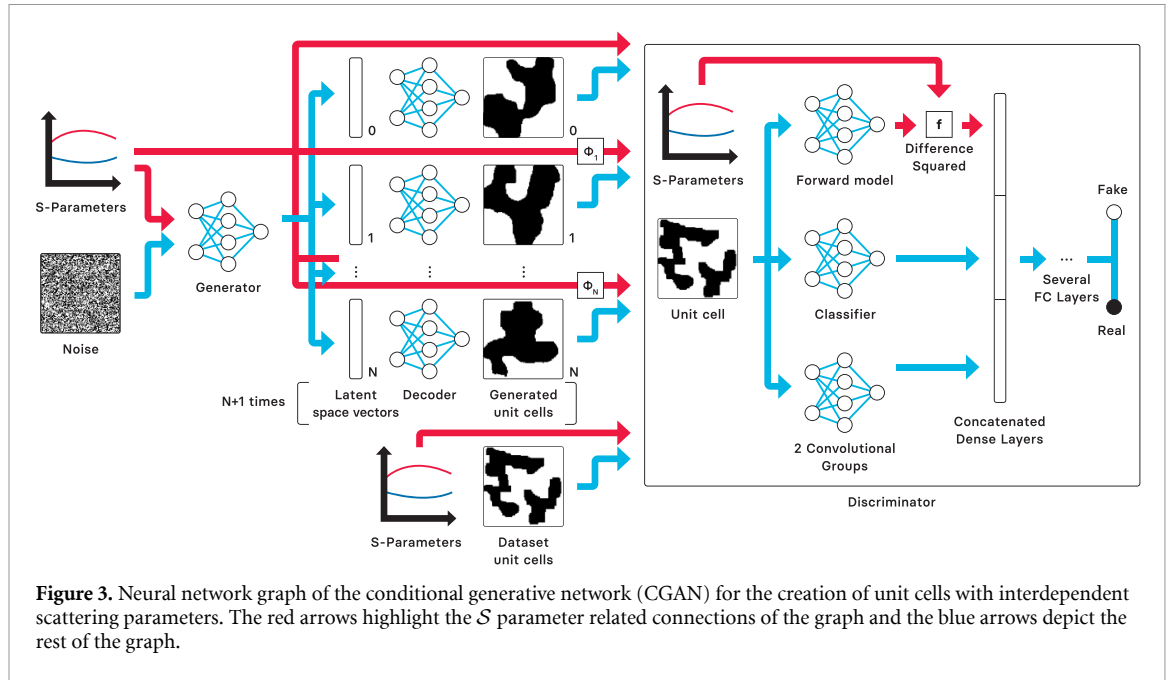
where \mathcal{G} is the generator, \mathcal{D} is the discriminator, \mathcal{S} are the scattering parameters (optical response), \mathcal{A} is the aspect ratio of the meta-atom, \mathcal{N} is noise, U is the lithographic mask of the meta-atom, and O is the discriminator output. The loss functions of the discriminator and generator are given by the usual cross-entropy loss:

$$\begin{aligned} \mathcal{E}_D(U_D, \mathcal{S}, \mathcal{A}, \mathcal{N}) &= -\ln(\mathcal{D}(U_D, \mathcal{S}, \mathcal{A})) - \ln(1 - \mathcal{D}(\mathcal{G}(\mathcal{N}, \mathcal{S}, \mathcal{A}), \mathcal{S}, \mathcal{A})), \\ \mathcal{E}_G(\mathcal{S}, \mathcal{A}, \mathcal{N}) &= \ln(1 - \mathcal{D}(\mathcal{G}(\mathcal{N}, \mathcal{S}, \mathcal{A}), \mathcal{S}, \mathcal{A})), \end{aligned} \quad (2)$$

where U_D is a unit cell from the data set.

2.1. Interdependent \mathcal{S} parameters

We now discuss how the above neural-network model for inverse design can be adapted to not only create one meta-atom U_0 with desired scattering parameters \mathcal{S}_0 , but also $N \in \mathbb{N}$ additional meta-atoms $U_{1..N}$ with interdependent \mathcal{S} parameters $\mathcal{S}_{1..N}$, where $\mathcal{S}_n = \phi_n(\mathcal{S}_0)$ with $n \in [1, N]$ (e.g. $\phi_n(\mathcal{S}_0)$ could be $\phi_n^1(\mathcal{S}_0)$, a phase rotation with constant amplitude, for beam steering). Generally, \mathcal{S}_0 has to be in the manifold \mathbb{S} of \mathcal{S} parameters spanned by the training data (used to train the discriminator and the forward model) to facilitate that the output of the generator is not a hallucination and the generated unit cell U_0 actually approximates the desired \mathcal{S} parameters (\mathcal{S}_0) sufficiently well. For the same reason, the function ϕ_n must be a transformation that projects the set of desired \mathcal{S} parameters \mathcal{S}_0 back onto \mathbb{S} ($\phi_n(\mathcal{S}_0) \in \mathbb{S}$). This is not a strict condition in the sense that the generator will still provide output if $\phi_n(\mathcal{S}_0) \notin \mathbb{S}$, but, in the same way that the output U_0 is most likely a hallucination when $\mathcal{S}_0 \notin \mathbb{S}$, the generated unit cells U_n will not approximate the projected \mathcal{S} parameters \mathcal{S}_n if ϕ_n is not an endomorphism on \mathbb{S} . Nevertheless, this can be guaranteed by constructing a suitable training data set.



In order to output multiple unit cells U_N with interdependent properties, the original generator architecture is split into two networks

$$\begin{aligned} \mathcal{G}_l[\mathcal{S}_0, \mathcal{A}, \mathcal{N}] &= [l_0, \dots, l_N], \\ \mathcal{L}[l_n] &= U_n, \\ \mathcal{L}[\mathcal{G}_l[\mathcal{S}_0, \mathcal{A}, \mathcal{N}]] &= [U_0, \dots, U_N], \end{aligned} \quad (3)$$

where \mathcal{G}_l is a multiple latent vector generator, which outputs $N + 1$ latent vectors and \mathcal{L} is a latent vector decoder used to decode each latent vector l_n into a unit cell U_n . All these unit cells get individually fed to the discriminator with ϕ_n -transformed \mathcal{S} parameters (ϕ_0 is the identity operation):

$$\mathcal{D}[U_n, \phi_n(\mathcal{S}_0), \mathcal{A}] = O_n. \quad (4)$$

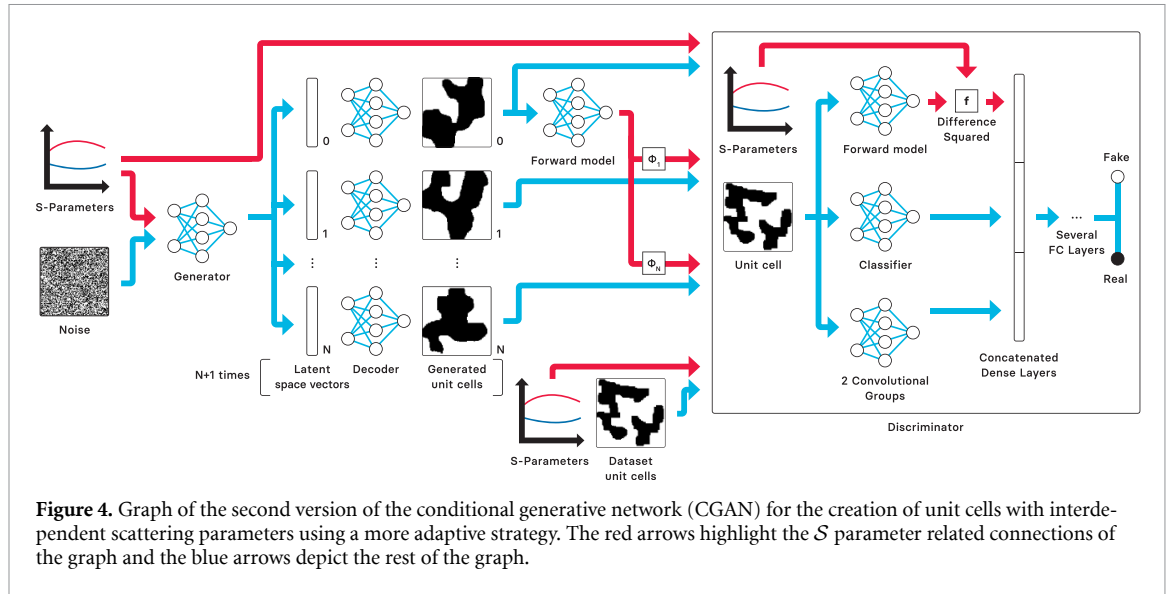
The neural network graph of this adapted CGAN set up is visualized in figure 3. Subsequently, a weighted sum of the cross entropy losses of the O_n outputs is used in the loss function:

$$\begin{aligned} \mathcal{E}_{\mathcal{D}}(U_D, \mathcal{S}_0, \mathcal{A}, \mathcal{N}) &= -\ln(\mathcal{D}(U_D, \mathcal{S}_0, \mathcal{A})) - \frac{\sum_{n=0}^N a_n \ln(1 - O_n)}{\sum_{n=0}^N a_n}, \\ \mathcal{E}_{\mathcal{G}}(\mathcal{S}_0, \mathcal{A}, \mathcal{N}) &= \frac{\sum_{n=0}^N a_n \ln(1 - O_n)}{\sum_{n=0}^N a_n}, \end{aligned} \quad (5)$$

where the factors a_n are the weights of the generator output U_n (and the respective discriminator output O_n) in the loss function.

2.2. Model training

While $a_0 = 1$ for all epochs, a_1 to a_N start out at 0.001 at epoch 0. Thereby, the generator outputs U_1 to U_N play almost no role in the early stages of training, making this effectively a standard inverse design CGAN setup (comparable to the graph in equation (2)). This improves the generator output U_0 and, in the process, minimizes the \mathcal{S} parameter error $|\mathcal{S}_0 - \mathcal{F}[U_0]|$. Once the \mathcal{S} parameter error sinks under a certain threshold, a_1 to a_N increase as the \mathcal{S} parameter error of U_0 decreases (and approach $0.5/N$ in the limit of the error going to 0). In this way, the generator first learns to output l_0 . Additionally, the latent decoder is trained to output U_0 from l_0 , while the discriminator learns the wide range of characteristics that have to be fulfilled for a meta-atom mask to be a suitable design. Once this works so well that most of a set of training instances would converge to a useful generator, the weight factors a_1 to a_N are increased in order to gradually shift the focus and learning of the generator to output l_1 to l_N (note that the same latent decoder network \mathcal{L} is used to produce U_0 to U_N from l_0 to l_N). More details about the model training is provided in the Methods section.



The unit cells U_1 to U_n will generally approximate the \mathcal{S} parameters $\phi_1(\mathcal{S}_0)$ to $\phi_N(\mathcal{S}_0)$, because these are fed to the discriminator, which is punishing deviations. Thereby, the following sum is minimized

$$|\mathcal{S}_0 - \mathcal{F}[U_0]| + \sum_{n=1}^N |\phi_n(\mathcal{S}_0) - \mathcal{F}[U_n]|. \quad (6)$$

2.3. Model without reference phase/amplitude

In the neural network graph above, we still fix the scattering parameters \mathcal{S}_0 of one of the meta-atoms. This is often undesirable as it unnecessarily restricts the design freedom, e.g. for a lens, the transmission phase of the center meta-atom can be arbitrary as long as the relative phase compared to the center meta-atom has a quadratic phase as a function of the distance to the center meta-atom. If the forward model \mathcal{F} has the necessary accuracy, then we can modify equation (4) by replacing $\phi_n(\mathcal{S})$ with $\phi_n(\mathcal{F}[U_0])$ for $n > 0$:

$$\begin{aligned} \mathcal{D}[U_0, \mathcal{S}_0, \mathcal{A}] &= O_0 \\ \mathcal{D}[U_n, \phi_n(\mathcal{F}[U_0]), \mathcal{A}] &= O_n \quad \forall n > 0. \end{aligned} \quad (7)$$

The network graph of this setup is shown in figure 4. In this way, it is possible to give more design freedom to the generator to find a set of unit cells where

$$|\mathcal{S}_0 - \mathcal{F}[U_0]| + \sum_{n=1}^N |\phi_n(\mathcal{F}[U_0]) - \mathcal{F}[U_n]| \quad (8)$$

is minimized. In the training process, we will generally feed \mathcal{S}_0 parameters to the generator and discriminator, although, by utilizing an input mask (\mathcal{S}_m) for the \mathcal{S}_0 parameters, it is possible to feed only a subset of all parameters to the network or none at all. Feeding a wide variety of $\mathcal{S}_0 (\in \mathbb{S})$ parameters to the networks helps generalize their capabilities, and feeding none ($\mathcal{S}_m = 0 \rightarrow |\mathcal{S}_m(\mathcal{S}_0 - \mathcal{F}[g_0])| = 0$) enables the maximal design freedom of the generator to minimize

$$\sum_{n=1}^N |\phi_n(\mathcal{F}[U_0]) - \mathcal{F}[U_n]|. \quad (9)$$

2.4. Zernike phase masks

We subsequently use our trained CGAN for the design of a broadband phase mask for a Zernike wavefront sensor (ZWFS) [51–54]. A ZWFS can be used to detect deviations of a detected wavefront from a plane wavefront, which can then be used to correct the incoming light signal. Therefore, it is a very powerful technique for ground-based telescopes, where the incoming light is distorted by the Earth's atmosphere, and eliminating this effect can significantly improve a telescope's ability to view distant

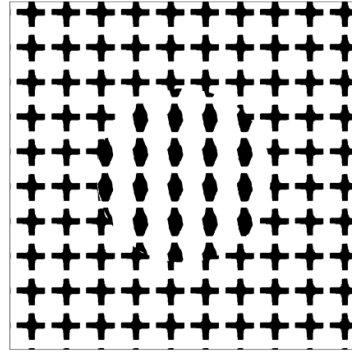


Figure 5. A sketch of the 2D cross section of the ZWFS metasurface (meta-atoms not to scale), unit cells from figure 7(a). A Zernike phase mask has a central circular region and a surrounding region; there are phase relationships between the S parameters of both regions.

objects. In addition, segmented aperture telescopes, e.g. at the Keck observatory or the recently launched James Webb Space telescope, require accurate phasing between the different segments in order to form the precise images required. Recent work at the Keck II telescope demonstrated the utility of metasurface masks by improving the observed Strehl ratios of imaged distant stars [55].

A ZWFS implements the phase contrast technique, where light is passed through a focal plane mask consisting of two regions, imparting different phases on impinging light. The light then interferes at an exit pupil to create intensity variations that can be readily imaged with a camera. A phase mask can, for instance, be created by etching a shallow dimple into a piece of glass to create a difference in optical path length between the etched and non-etched regions. However, such an implementation leads to a limited dynamic range in terms of measurable phase errors. The dynamic range can be improved by using polarization as a degree of freedom and simultaneously impart two different phases on two orthogonal polarization states, which can later be separated to create two images containing different phase information. Here, we utilize our CGAN to design metasurfaces that not only impart different phases on two polarization states, but also do so in a highly achromatic manner. This achromatic behavior is important for broadband functionality of metasurface devices in general, and in the ZWFS context this enables the use of more of the incident light signal as compared to a highly chromatic device.

We design a Zernike phase mask that consists of two regions, as illustrated in figure 5, with the following interdependent S parameters over the entire frequency band of interest:

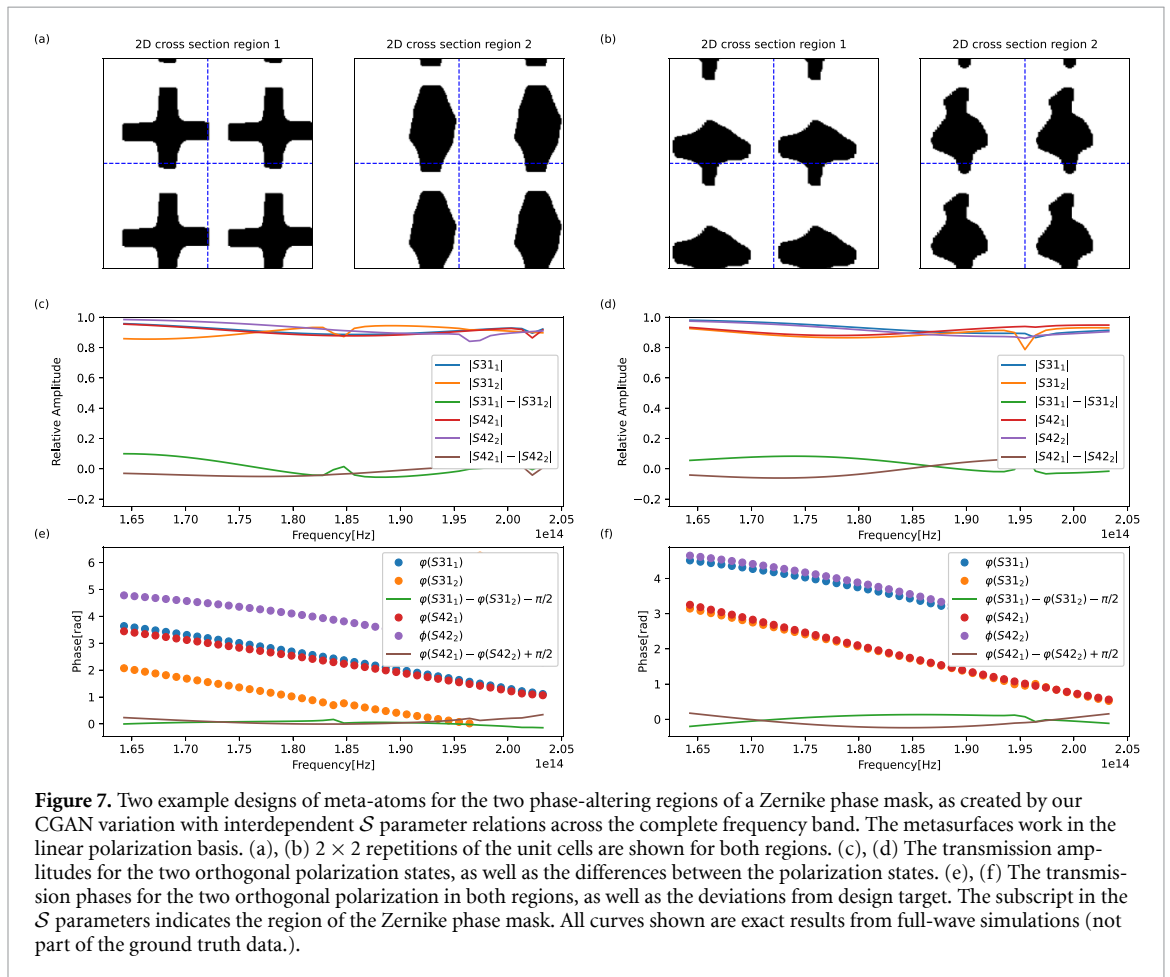
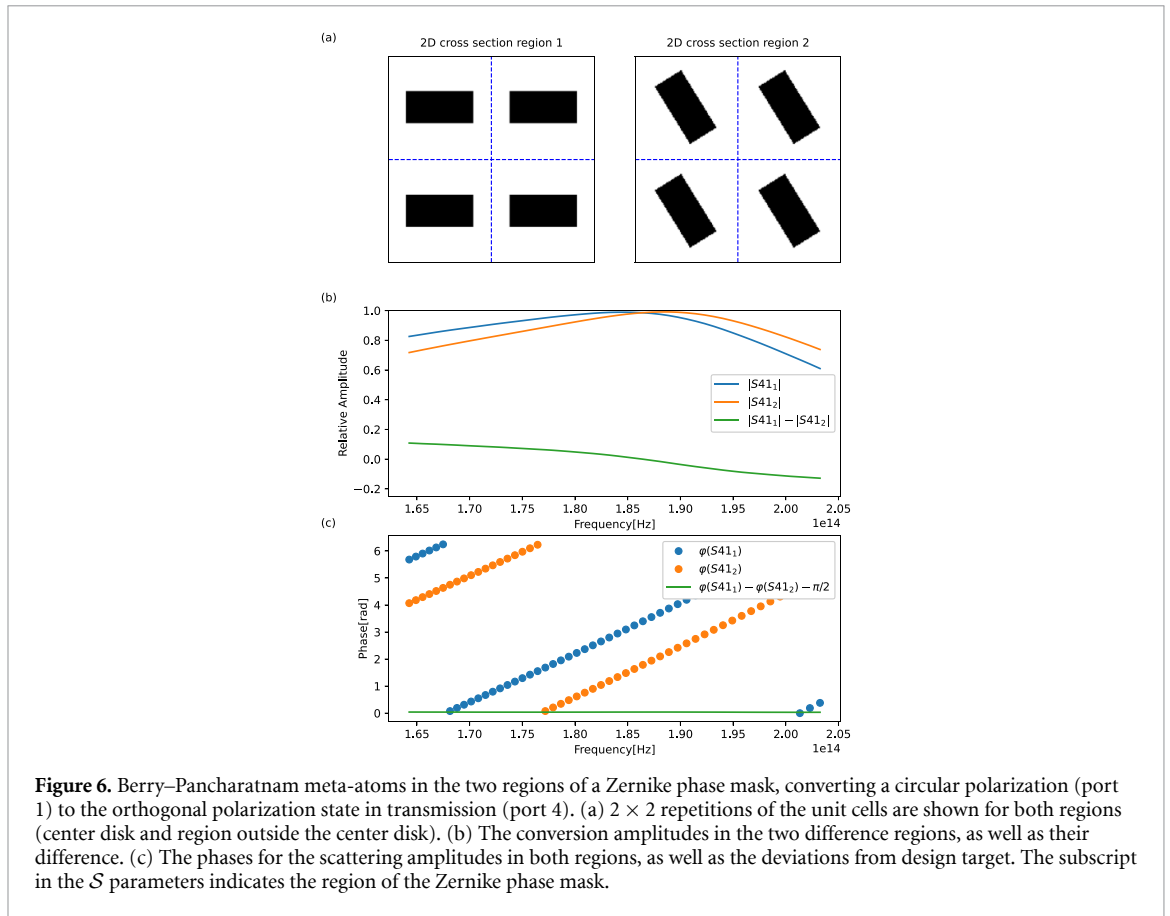
- a relative phase difference of $\pi/2$ between the S_{31} parameters in both regions;
- a relative phase difference of $-\pi/2$ between the S_{42} parameters in both regions;
- high and approximately constant amplitudes of S_{31} and S_{42} in both regions;

where ports 1 and 2 are two orthogonal polarization states at the input plane and ports 3 and 4 are two orthogonal modes at the output plane. Therefore, the transformation ϕ , mapping S parameters from the central region to the surrounding region, is given by

$$\phi_1(S_{31}) = S_{31} e^{\frac{\pi}{2}}, \phi_1(S_{42}) = S_{42} e^{-\frac{\pi}{2}} \quad (10)$$

in our example of a Zernike phase mask.

Utilizing two polarization states with opposite phase difference between the regions, it is possible to improve the dynamic range of Zernike wavefront sensing [56]. Designing broadband meta-atoms for such a Zernike phase mask is challenging, because there are requirements on phase and amplitude simultaneously, for two different polarization states. It can be achieved in the circular polarization basis with Berry–Pancharatnam meta-atoms [57–59], but such an implementation suffers from leakage which ultimately affects the Zernike sensing. Furthermore, subsequent polarization splitting in a circular basis requires additional optical components, which complicates the system. As a reference, we plot in figure 6 the efficiency of a Zernike phase plate with Berry–Pancharatnam meta-atoms; we see that the efficiency is high at the center frequency, but then gradually drops towards the edges of the operational bandwidth, which is a fundamental limitation of this type of metasurfaces [60]. We now ask our CGAN for interdependent meta-atoms to design the metasurface for a Zernike phase plate and, in figures 7(a) and (b), we plot the meta-atom pairs for two different designs returned by the CGAN. The resulting etching masks, shown in figures 7(a) and (b), show that our CGAN can find designs in a much larger design



space, with designs that are difficult to find with traditional approaches. In figures 7(c) and (d), we plot the transmission efficiency of the meta-atoms in both regions. The CGAN-derived designs have efficiencies that are constant over a very large bandwidth; they are slightly fluctuating, but do not systematically drop off towards the edges of the operational bandwidth. In figures 7(e) and (f), we observe that the transmission phases satisfy the requirement to a very high degree, also constant over the very large bandwidth. We point out that the desired phase is here imprinted for both polarization states in linear polarization basis (\mathcal{S}_{31} and \mathcal{S}_{42}), eliminating the need for additional optical components needed for the classical designs using Berry–Pancharatnam metasurfaces. We plot also the other \mathcal{S} parameters in the supplementary material, showing amongst others that the cross polarization leakage is negligible. This confirms that CGAN-based inverse design can find designs beyond the current state of the art for meta-atoms with interdependent optical properties.

3. Discussion

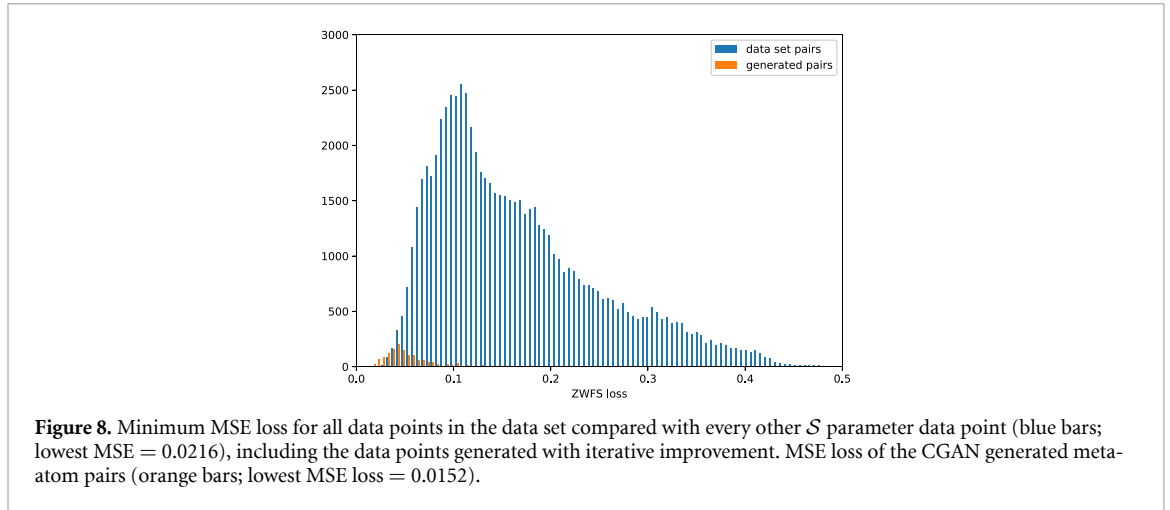
In this article, we have demonstrated a CGAN-based inverse design method for metasurfaces consisting of meta-atoms with interdependent optical properties. This is a very common situation in optics, since an overall phase shift of the wavefronts is often unimportant, but leads to intractable designs without the use of machine learning. We demonstrated our CGAN with a metasurface for the phase mask of a ZWFS, which contains two regions of meta-atoms with multiple, relative requirements (phase and amplitude) on multiple \mathcal{S} parameters of the meta-atom pairs. The CGAN is able to design etching masks for such phase plates with much lower leakage than existing solutions. In addition, our CGAN has built-in functionality to enforce that its resultant designs are fabricationally feasible. The training time of our CGAN is about 12 to 56 h on 2 GPUs. The inference time is a few milliseconds. In figure 8, we estimate the learning ability by plotting the mean square error (MSE) of every pair of meta-atoms from our training set (blue bars), which is a measure of how a Zernike phase plate with random meta-atoms chosen from the training data set perform, and we compare it with the MSE for designs created with different generators (orange bars). In this plot, the training data set pairs already include the samples that were generated by iterative improvement by using the CGAN to find training samples in sparsely populated regions of the design space. We see that the CGAN is able to find designs for meta-atom pairs with a significantly lower MSE than present in the training data set. Our Zernike phase masks demonstrate that machine learning can help with optical design of devices that cannot be designed with conventional methods.

Machine-learning-based designs always come at the cost of a large number of training data samples that need to be simulated, but this cost can be acceptable if no other design method is known. The cost is also reduced if the CGAN can be reused for multiple designs, e.g. in gradient-phase metasurfaces where thousands of different meta-atoms are needed. For example, another possible application of our CGAN for metasurfaces with interdependent unit cells could be a metasurface lens without chromatic aberrations. Since the phase profile of a lens is wavelength-dependent, the individual unit cells need to have interdependent \mathcal{S} parameters—the required phase profile would be given by $\varphi(r, \lambda) = -\frac{2\pi}{\lambda}(\sqrt{r^2 + f^2} - f)$, where f is the focal point distance of the lens and r is the distance of a unit cell from the center of the lens. Consequently, the shift in phase from unit cell to unit cell also depends on the wavelength. Hence, the \mathcal{S} parameters (at each and every frequency) of the individual unit cells in a metasurface lens without chromatic aberration must obey interdependent relations. If the network is trained with a sufficiently diverse training set, it can be reused for other applications using the same materials platform; otherwise the networks will need to be retrained. Another possibility is to reuse the networks in a transfer learning setting, where the trained networks are reused with additional layers at the beginning and end of the neural-network structure. We therefore anticipate that our CGAN-based inverse design method for meta-atoms with interdependent optical properties will lead to new designs for a multitude of optical devices. While it is a black box model that allows to find solutions beyond human intuition on physical insight, it can also be combined using recent techniques to incorporate the physics in the surrogate/forward model [61]. Furthermore, since our method is transparent to the physics (the physics is contained in the training data), our method can be straightforwardly transferred to systems in other paradigms of physics, such as acoustics, mechanics, and fluid dynamics.

4. Methods

4.1. Training of neural networks

The models were trained on NVIDIA Tesla V100 SXM2 GPUs with 32 GB RAM with the following optimizers and training durations:



- forward network
 - optimizer: Adamax(learning rate = 0.001, $\beta_1 = 0.9$, $\beta_2 = 0.999$, $\epsilon = 10^{-7}$)
 - typical training time: 200 epochs in 4 h
- classifier
 - optimizer: Adamax(learning rate = 0.001, $\beta_1 = 0.9$, $\beta_2 = 0.999$, $\epsilon = 10^{-7}$)
 - typical training time: 150 epochs in 5 h
- generator optimizer: Adamax(learning rate = 0.001, $\beta_1 = 0.9$, $\beta_2 = 0.999$, $\epsilon = 10^{-7}$)
- discriminator optimizer: Adam(learning rate = 0.000 002, $\beta_1 = 0.5$, $\beta_2 = 0.999$, $\epsilon = 10^{-7}$)
- encoder optimizer: Adamax(learning rate = 0.001, $\beta_1 = 0.9$, $\beta_2 = 0.999$, $\epsilon = 10^{-7}$)
- typical CGAN training time: 80 to 400 epochs in 12 to 56 h (2 GPUs).

The following python packages were used:

- tensorflow
- numpy
- pandas
- multiprocessing.

Validation loss curves for the forward network, the classifier network, the generator, and the discriminator can be found in the supplementary material. The generator and the discriminator are competing in a zero-sum game and, therefore, their validation loss curves have information on the stability of the training process rather than the quality of the outputs.

4.2. Initial training data generation

Our training data set consists of a set of 100×100 binary grids, where 0s mark the areas in which the material is etched away and 1s mark the areas where the lithographic mask prohibits the etching process and consequently the material stays intact. In order to create a data set of semi-random pixelated images representing the lithographic masks of experimentally feasible structures, the binary grids were grown from random noise by filling too small holes or erasing too small islands of increasing sizes, since too small holes and islands cannot be realized by current lithographic techniques. In this way, random noise was transformed successively from iteration to iteration into structures with courser features until the fabrication constraints were met. As a result of this process, the allowed geometries of our meta-atoms are only limited by the resolution of the grid (in practice our grid pixels are so much smaller than the wavelength of light and slightly smaller than the estimated fabrication error that further reducing the pixel size is pointless) and by fabrication feasibility—and not by any other constraints arising from the computational method.

4.3. Training strategy and iterative improvement of training data

To improve the quality of the training data set, we have iteratively improved the training data using the following algorithm:

1. Expert models training: We train the classifier and forward networks on the current set of grids, labels, and \mathcal{S} parameters.
2. CGAN training: We train several CGAN generators with the classifier and forward models trained in the previous step.
3. Classifier data enhancement: From all generators of this iteration, we produce grids for various different kinds of target \mathcal{S} parameters and noise, round them to values of 0 and 1, save and label them for the classifier, and then add the data to the classifier training data.
4. Forward model data enhancement: The quality and accuracy of grids generated by the generator is directly dependent on the accuracy of the forward model in the regime of interest. Since the number of randomly generated grid pairs that satisfy the design criteria for a Zernike phase mask is rather low, we increase the size of the training set iteratively. To this end, a diverse selection of grids in the regime of interest is created by the generators of this iteration, corrected for fabricationability if necessary, simulated by COMSOL, and then added to the training data of the forward model. In this way, we gradually create more and more training data pairs that are relevant for the design of a Zernike phase mask.

By applying this iterative process, we are able to increase the accuracy of all models, reduce the number of corrections to the grids generated by the generator necessary to satisfy feasibility constraints, and obtain \mathcal{S} parameters of the generated grids that better approximate the target parameters.

4.4. Electromagnetic simulations

The electromagnetic simulations of the meta-atoms (training sample set) are performed using the finite-element method with the package COMSOL. Scattering parameters are calculated using absorbing boundary conditions (ports) for a plane wave incident from the top of the metasurface. The errors on the \mathcal{S} parameters in the finite-element simulations are approximately 1%.

Data availability statement

All data that support the findings of this study are included within the article (and any supplementary files).

Supplementary material available at <https://doi.org/10.1088/2632-2153/ae365e/data1>.

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