DNA-Stabilized Silver Nanocluster Design via Regularized Variational Autoencoders

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ABSTRACT

DNA-stabilized silver nanoclusters (Ag$_N$-DNAs) are a class of nanomaterials comprised of 10-30 silver atoms held together by short synthetic DNA template strands. Ag$_N$-DNAs are promising bio-sensors and fluorophores due to their small sizes, natural compatibility with DNA, and bright fluorescence—the property of absorbing light and re-emitting light of a different color. The sequence of the DNA template acts as a "genome" for Ag$_N$-DNAs, tuning the size of the encapsulated silver nanocluster, and thus its fluorescence color. However, current understanding of the Ag$_N$-DNA genome is still limited. Only a minority of DNA sequences produce highly fluorescent Ag$_N$-DNAs, and the bulky DNA strands and complex DNA-silver interactions make it challenging to use first principles chemical calculations to understand and design Ag$_N$-DNAs. Thus, a major challenge for researchers studying these nanomaterials is to develop methods to employ observational data about studied Ag$_N$-DNAs to design new nanoclusters for targeted applications.

In this work, we present an approach to design Ag$_N$-DNAs by employing variational autoencoders (VAEs) as generative models. Specifically, we employ an LSTM-based β-VAE architecture and regularize its latent space to correlate with Ag$_N$-DNA properties such as color and brightness. The regularization is adaptive to skewed sample distributions of available observational data along our design axes of properties. We employ our model for design of Ag$_N$-DNAs in the near-infrared (NIR) band, where relatively few Ag$_N$-DNAs have been observed to date. Wet lab experiments validate that when employed for designing new Ag$_N$-DNAs, our model significantly shifts the distribution of Ag$_N$-DNA colors towards the NIR while simultaneously achieving bright fluorescence. This work shows that VAE-based generative models are well-suited for the design of Ag$_N$-DNAs with multiple targeted properties, with significant potential to advance the promising applications of these nanomaterials for bioimaging, biosensing, and other critical technologies.

CCS CONCEPTS

• Information systems → Data mining. • Computing methodologies → Learning latent representations.

KEYWORDS

nanomaterials design; DNA; variational autoencoders

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1 INTRODUCTION

DNA is a sequence-encoded building block for nanomaterials. By harnessing the well-understood base pairing rules of natural DNA, researchers have developed ways to engineer DNA sequences to fold DNA "origami" [29], build with DNA "bricks" [19], and wire DNA logic circuits [9]. DNA can also imbue sequence-encoded properties to the tiniest of nanoparticles: nanoclusters composed of just a few metal atoms. Of particular interest are DNA-stabilized silver nanoclusters (Ag$_N$-DNAs), which contain 10-30 silver atoms that are stabilized by 1 or 2 short DNA strands [10]. Ag$_N$-DNAs are colloidal nanomaterials that are synthesized in solution by mixing Ag atoms and DNA template strands (Fig. 1, top panel), yielding fluorescent nanoclusters with remarkable sequence-encoded properties. The DNA sequence controls the size and shape of the silver nanocluster, thereby tuning the fluorescence color of Ag$_N$-DNAs from blue wavelengths (~400 nm) to near-infrared (NIR) wavelengths (at least 1,000 nm) [6]. This bright, tunable fluorescence, combined with inherent biological compatibility and sensitivity to the local molecular environment, makes Ag$_N$-DNAs promising for a range of applications, from bioimaging and sensing to nanophotonics.

However, a major challenge faces the development of applications of Ag$_N$-DNAs. Unlike the well-known Watson-Crick base pairing rules of natural DNA, the sequence rules that govern how

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DNA interacts with silver atoms and thereby select for Ag\(_N\)-DNA fluorescence color are not well-understood. Most researchers have used combinatorial screening or intuition to design the DNA template sequences for Ag\(_N\)-DNAs reported in the literature, which is a time-consuming and inefficient process. To enable data-driven approaches to map DNA sequence onto Ag\(_N\)-DNA color, we developed a high-throughput experimental platform for Ag\(_N\)-DNA synthesis and characterization, producing a library linking DNA sequences to the fluorescence colors of Ag\(_N\)-DNAs they stabilize. We previously utilized this library to train classifiers based on subsequence motifs to predict Ag\(_N\)-DNA fluorescence brightness [3] and fluorescence color [5] given an input DNA sequence. (Classification schemes are motivated by the naturally discretized properties of Ag\(_N\)-DNA colors [6].) We then employed the most discriminative subsequence motifs to create new DNA templates. While this approach led to discovery of new Ag\(_N\)-DNAs, it has several limitations: it relies on (i) discriminative as opposed to generative models to sample new DNA templates, (ii) ad hoc feature generation by sub-sequence mining, and (iii) discretization of continuous design properties like brightness and color into balanced classes.

The discovery of Ag\(_N\)-DNAs with NIR fluorescence emission is especially important for bioimaging applications. Biological tissues are much more transparent to NIR light than to visible light, and there is great effort to develop small, nontoxic, and bright fluorescent biolabels in the NIR spectral region. Few NIR Ag\(_N\)-DNAs were reported before 2018, when the discovery of 161 new NIR Ag\(_N\)-DNAs [7, 31, 32] suddenly presented the opportunity to extend machine learning-guided design of Ag\(_N\)-DNAs into the NIR. Because data in the NIR remains scarce, effective approaches to this challenging problem must be sufficiently sensitive to rare data.

In this work, we set out to address the limitations of our earlier Ag\(_N\)-DNA design approaches and employ our new model to enrich the space of known NIR Ag\(_N\)-DNAs. We propose and deploy a regularized variational autoencoder (VAE) model for the design of Ag\(_N\)-DNAs with desired properties summarized in Fig. 1, lower panel. Inputs to our model are sequences for synthesized Ag\(_N\)-DNAs and their measured wavelengths and brightness levels. We train the VAE to encode and decode DNA sequence by employing a bi-directional LSTM architecture. Instead of learning a fully latent space, we regularize a subset of its dimensions to correlate with design properties of interest. Our regularization scheme also accounts for bias in the observations along the design parameters, by compensating for rarer sample Ag\(_N\)-DNAs in the NIR band. We employ the trained model to design Ag\(_N\)-DNA template sequences by truncated sampling from latent space, thus biasing samples towards high wavelength and brightness while obeying the distribution of the remaining latent dimensions. We experimentally test the proposed VAE model on 20 new DNA sequences, finding that all of them produce Ag\(_N\)-DNAs with bright fluorescence and high wavelengths, including a bright NIR Ag\(_N\)-DNA with 845 nm peak fluorescence that has never been observed before.

Our contributions in this work are as follows:

- **Novelty.** We propose, test and deploy the first approach for rational design of Ag\(_N\)-DNAs with multiple continuous properties of interest via a VAE architecture.

- **Generality.** Our framework is general, in that it can extend to more design properties of interest, variable length of DNAs, and for designing other biological sequences with desired properties.

- **Applicability.** We experimentally demonstrate the utility of our approach, employing it to sample and synthesize 20 new Ag\(_N\)-DNAs in the lab, and discover a previously unreported NIR Ag\(_N\)-DNA.

2 RELATED WORK

**Ag\(_N\)-DNA design.** The vast majority of studies on Ag\(_N\)-DNAs employ nanoclusters designed by a combination of combinatorial screening and intuition, which is highly inefficient. To overcome these challenges, we developed high-throughput experimental synthesis and characterization of Ag\(_N\)-DNAs [6], producing a large training dataset that enabled early machine learning approaches based on support vector machine classifiers [4, 5, 7]. These approaches rely on bioinformatics techniques for feature engineering and discretization of a single design property into classes (e.g., high/low fluorescence yield in [4] and color in [5, 7]); as Ag\(_N\)-DNA colors are naturally discretized due to their structural properties, this approach is motivated by physics/chemistry [6]. Perhaps most importantly, these prior approaches rely on discriminative, as opposed to generative, models and ad hoc heuristics to sample from the complex space of all possible DNA sequences. The proposed VAE approach in this work addresses the above limitations: it maps both DNA sequences and multiple design properties into a continuous space from which one can perform truncated sampling to tune properties of interest and decode the samples into DNA sequences.

**Generative models based on VAEs.** Our proposed model is a generative VAE that builds on prior autoencoder (AE) research. AEs have been in use since the mid 1980s, but were initially used for dimensionality reduction and denoising, with little generative ability [11]. In 2014, Kingma and Welling proposed the Variational Autoencoder (VAE) [22], modifying the latent space of the VAE architecture to hold latent distributions, which are then sampled
during the training process. This change allows for VAEs to be used for generative tasks. A drawback of classical VAEs is the inability to control various properties of the features in latent space such as disentanglement, regularization, and monotonicity. Higgins and colleagues proposed the β-VAE framework [16] to control the level of disentanglement in latent space by incorporating a Kullback–Leibler divergence term of the latent distributions from a normal prior. Regularization of VAEs aimed to impose monotonicity of the learned latent space with respect to features of the input was originally introduced in the context of Fader Networks [24] as part of the GLSR-VAE [14] model and later employed in the ARVAE [30] model for image and music datasets. Our proposed model follows a similar property monotonicity approach; however, we apply it to DNA sequences and further consider non-uniform coverage of properties in the training which is inherent to the problem of discovering new Ag₅₋DNA where new samples are both laborious and expensive to obtain.

**Machine learning for biological sequences.** Many sequence embedding approaches build upon word2vec [28], which was designed to represent words as vectors by enforcing low cosine similarity between the vector representations of semantically similar words. FastText [1, 17] is an alternative employing n-grams within words as opposed to whole words. Biological sequence (e.g., RNA, DNA, and proteins) embedding techniques also utilize and extend the above frameworks to obtain representations employed in promoter region [25] and protein [36] classification, taxonomy [34] and neural distance learning [8] and others. Generators for protein or DNA sequences have also been of high interest [35]. Specifically, both Generative Adversarial Network (GAN) [2, 18, 20] and VAE-based [12, 15] generators have been employed for creating nucleotide or amino acid sequences. Gupta and Zou’s FBGAN [13] incorporates an additional feedback component that guides the generator towards desired features, such as peptides with antimicrobial activities. The VAE methods in this group are employed to edit sequences for downstream targets as opposed to direct targeted synthesis [12, 15]. Distinct from our work, the majority of these approaches focus on large biological datasets, both in terms of the available input data as well as the lengths of the encoded biological sequences. Methods tuned for long, information-rich sequences are unlikely to perform as well on shorter strands, like the short 10-base DNA strands that we employ to stabilize Ag₅₋-DNAs. Additionally, the incorporation of additional information is limited to either direct annotation or a semi-supervised editing between runs as in the FBGAN approach.

### 3 PROBLEM FORMULATION

Our goal is to design Ag₅₋-DNAs of specific properties tuned by their stabilizing DNA template sequence. The input to our problem is a training set (S, A) of sequences S and their corresponding properties represented as numeric feature vectors A. Specifically, our training data consist of a set of 10-base DNA sequences annotated by (i) fluorescence emission color quantified as the peak wavelength (WAV) of the emission spectrum of the corresponding Ag₅₋-DNA and (ii) its fluorescence brightness quantified as the local integrated intensity (LII) of a Gaussian fitted to the fluorescence spectral peak. In other words, the input property matrix is 2-dimensional \( A \in \mathbb{R}^{S \times 2} \). Our past work describes the data set acquisition, processing, and curation in detail [5].

Given this input, we aim to learn a generative model for the joint distribution of DNA sequence and properties \( p(S, A) \) based on the training observations (note that we overload the notation and use \( S \) and \( A \) as the corresponding random variables as well). We can then employ \( M \) to sample unobserved sequences \( S' \) with desired properties \( A' \), i.e. \( S' \sim p(S|A = A') \). Specifically, we aim to design DNA templates that stabilize bright Ag₅₋-DNAs with NIR emission, i.e. WAV > 800nm and as high fluorescence yield (LII) as possible. A few such Ag₅₋-DNAs were only recently synthesized for the first time [31, 32]. In this regime (WAV > 800nm) biological tissues become increasingly transparent to light and the Ag₅₋-DNAs can be employed as effective and non-toxic biosensors.

### 4 METHODOLOGY

Estimating the joint distribution of DNA sequences and properties \( p(S, A) \) is challenging with limited training data, since the discrete space of all possible sequences is exponential and testing the properties of all sequences by Ag₅₋-DNA synthesis is impossible. Hence, we seek to learn a joint low-dimensional numeric embedding for sequences and properties that allows for two-way transformation to and from the input space. To this end, we employ the Variational Autoencoders (VAEs) framework [23] which allows for the desired two-way transformation and can flexibly incorporate appropriate encoder/decoder architectures for sequential data such as DNA sequences (Sec. 4.1). To enable sampling from the learned latent space while controlling for WAV and LII of interest, we regularize a subset of the latent dimensions in the VAE to correlate with the observed Ag₅₋-DNA properties from training (Sec. 4.2) and handle imbalanced coverage of property samples (Sec. 4.3). Finally, since we employ a β-VAE architecture that enforces decoupling and normality of the latent space, we can efficiently sample from the conditional latent distribution employing the truncated normal distribution (Sec. 4.4).

#### 4.1 VAE Encoder/Decoder architecture

Our VAE model is composed of two distinct networks, an encoder mapping DNA sequences \( S \) to distributions in latent space \( p(z) \) and a decoder mapping samples from latent space back to sequences (Fig. 2). Observed sequences \( S_i \), \( |S_i| = l \) of length \( l \) are encoded using one-hot encoding into matrices \( X_i \in \mathbb{R}^{l \times 4} \) since our DNA alphabet can take one of 4 possible DNA base values \( \{A,C,T,G\} \).

**Encoder:** The one-hot encoding input matrices \( X_i \) are grouped into training batches of size \( b \) and fed into the first block of the encoder, followed by a many-to-many bi-directional LSTM (Bi-LSTM) with hidden state size \( h \). We select this sequential architecture due to its wide-adoption for sequence learning [27], yet it is among the simplest sequential models with relatively few parameters to tune. The bi-directionalness is essential to capture the context both before and after a given DNA base, which we expect to control the 3D local structure of the DNA strand and its interactions with silver atoms in the Ag₅₋-DNAs. Each Bi-LSTM layer in the block has one Bi-LSTM cell per base position resulting in a total of \( l \) cells. Each cell contains forward and backwards regular LSTM cells. The output of each Bi-LSTM cell is the concatenation of the hidden states of its forward and backward LSTM cell, which is a vector of size \( h \). The
Fig. 2: The architecture of the LSTM encoder and decoder in our VAE model. The Encoder is a sequence of (i) input, (ii) LSTM, (iii) fully connected and (iv) latent mean $\mu_z$ and variance $\sigma_z$ output layers. The Decoder follows a “reversed” architecture with the difference that the LSTM layer is followed by a fully-connected linear layer and a reshape transformation to obtain decoded sequences. Notation: $l$ is the DNA sequence length, $b$ is the training batch size, $w$ is the hidden state size of Bi-LSTM cells, $\nu$ is the linear width of the fully connected layer, and $|z|$ is the dimensionality of the latent space.

LSTM block output is a tensor of shape $R^{bxw}$. We experimented with multiple LSTM layers and by adding dropout layers, but the simplest architecture of one LSTM layer and no dropout resulted in optimal performance for our dataset (details on our hyperparameter search are available in Tbl. 1).

The LSTM output is flattened and fed into a fully-connected layer with ReLU activation. The output size of the fully connected layer is $b \times w$, where $w$ is the layer width, i.e., the number of neurons in the layer. The last layer of the encoder includes the latent mean $\mu_z$ and variance $\sigma_z$ dense layers which are traditionally employed in VAEs. Their outputs represent the corresponding distributional parameters of an input’s latent encoding in $z$.

Decoder. The decoder takes as an input a batch of samples from latent space and is trained to reconstruct the DNA sequences in the batch. The first decoder layer is dense and features ReLU activations. Its output size is the same as that of the DNA sequence length. The output from the latter is passed to a bidirectional many-to-many LSTM block with the same architecture as its encoder counterpart. Finally, we transform the LSTM block output into the shape of a one-hot encoded DNA sequence using a dense layer and the output of the latter is transformed in to a batch reconstruction tensor $Y$ of size $b \times 1 \times 4$.

4.2 Property-regularized loss function

The loss function of the basic VAE architecture features a reconstruction $L_{REC}$ and a Kullback-Leibler (KL) divergence $L_{KL}$ term:

$$L_{VAE} = L_{REC}(\phi, \theta) + L_{KL}(\phi, \theta),$$

where $\phi$ and $\theta$ are the parameters of the encoder and decoder respectively. The first term is the reconstruction loss between the input tensor $X$ and the decoded output tensor $Y$, both of shape $b \times 1 \times 4$:

$$L_{REC}(\phi, \theta) = \frac{1}{bl} \sum_{u=0}^{b} \sum_{w=0}^{l} \left[ \frac{4}{j=0} \exp(Y_{uji}) \log \left( \frac{\exp(Y_{uji})}{\sum_{k=0}^{4} \exp(Y_{uik})} \right) \right]$$

VAEs model the latent representation $Z$ as a random variable and hence the end-to-end encoding-decoding process is viewed as a sequence of sampling operations: (i) the input is sampled from the conditional distribution represented by the decoder $X \sim p_{\theta}(X|Z)$ and (ii) $Z$ is sampled from an approximation of the true posterior distribution $Z \sim q_{\phi}(Z|X)$ realized by the encoder and parameterized by the variational parameters $\phi$ [23]. Similar to other variational methods, variational inference in VAEs is performed by maximizing the evidence lower bound (ELBO):

$$\log p(X) \geq \mathbb{E}_{Z \sim q_{\phi}(Z|X)} \left[ \log p_{\theta}(X|Z) \right] - D_{KL}(q_{\phi}(Z|X)||p(Z)),$$

where $p(Z)$ is a prior distribution for the latent representation and $D_{KL}(q_{\phi}(Z|X)||p(Z))$ is the Kullback-Leibler (KL) divergence between the approximation to the posterior distribution and a prior distribution for $Z$. Minimizing the KL divergence improves the tightness of the ELBO bound and how well the approximate posterior aligns with the prior [23] and hence it is the second loss term in Eq. 1, i.e., $L_{KL} = D_{KL}(q_{\phi}(Z|X)||p(Z))$. A typical prior distribution employed for the latent variable $Z$ is multivariate normal with zero co-variances which promotes independence between the dimensions in the latent space. A follow-up model called $\beta$-VAE introduces a weight $\beta$ which multiplies $L_{KL}$ and allows for more control for decoupling of the dimensions of the latent representation [16].

The loss function from Eq. 1 ensures a decoupled latent space and a good reconstruction of the input DNA sequence. Our goal for the latent representation is to also jointly reflect the properties $A_{AGN}$-DNAs (WAV and LII) that are of interest for design. Thus, we extend a property regularization variant of the VAE model [30] that ensures that a subset of the dimensions of $Z$ encode properties, that corresponding latent dimensions monotonically increase in unison with the observed properties $A$. To this end, we add a property regularization term to the basic $\beta$-VAE loss as follows:

$$L_{REC}(\phi, \theta) + \beta L_{KL}(\phi, \theta) + \gamma \sum_{a \in A} L_a,$$

where the last term, which we will also refer to as $L_{REG}$, adds property regularization controlled by a hyper-parameter $\gamma$. The individual summants $L_a$ in $L_{REG}$ enforce alignment between a single property (e.g., WAV, LII) and a corresponding latent dimension:

$$L_a = \text{MAE}(\tanh(5D_a) - \text{sign}(D_a)),$$

where MAE stands for the mean absolute error, $\delta$ is a scaling parameter, $\tanh()$ denotes the hyperbolic tangent function applied element-wise to its argument, $\text{sign()}$ is the sign function also applied element-wise, and $D_a, D_a \in R^{bxb}$ are batch-specific square difference matrices whose elements are defined as follows:

$$D_{r}(i, j) = Z_r(i) - Z_r(j) \text{ and } D_a(i, j) = A_a(i) - A_a(j),$$

where $i$ and $j$ are training instance indices within a given batch, $Z_r(i)$ is the $r$-th dimension of the embedding of instance $i$ that is mapped to attribute index $a$ and $A_a(i)$ is the $a$-th attribute value of instance $i$ provided as input to our model. Intuitively, $L_{REG}$ introduces cost for instance pairs which are ordered differently based on their training attribute with index $a$ and their latent embedding in a corresponding dimension $r$. As a result the VAE will be trained to match the WAV ordering to a WAV proxy latent dimension in $Z$ and similarly the LII to a corresponding LII proxy dimension in latent space. Optimization of the overall objective is performed using standard neural network batch-gradient methods, since all components of the loss function are differentiable with respect to the parameters of the encoder $\phi$ and decoder $\theta$. 
4.3 Handling imbalance in the observations

The regularized VAE model imposes a penalty in $L_{REG}$ for pairs of instances whose embeddings in regularized dimensions are ordered differently than the reference values of their properties. Specifically, the regularization loss for a given attribute $a$ within a batch of size $b$ instances is computed as an average of all instance pairs:

$$L_a = \frac{2}{b(b+1)} \sum_{i=1}^{b-1} \sum_{j=i+1}^{b} \left| \tanh(\delta D_a(i,j)) - \sign(D_a(i,j)) \right|.$$  

The above definition assumes that attribute values in batches and in the overall dataset are uniformly randomly distributed. In settings where the training dataset contains non-uniformly distributed attribute values, the regularization, and consequently the trained VAE model, will over-represent intervals of attribute values with high support and neglect rare values. Note that this imbalance is especially relevant to employing VAEs for Ag$_N$-DNA design. In particular, we have many fewer NIR Ag$_N$-DNAs in the training data, while at the same time our goal is to design DNAs for NIR Ag$_N$-DNAs. Given the relative scarcity of training instances for desired attribute values, how can we "focus" the training on representing that attribute value region well?

To this end, we propose a weighted MAE alternative in which differences for rare pairs induce higher penalties. Specifically, if $v(i,j)$ is an instance pair score function, we define a weighted MAE attribute loss as follows:

$$L_{\text{reg}} = \text{WMAE}(\text{tanh}(\delta D_a(Z^a)) - \sign(D_a^p(Z^a))))$$

$$= \frac{1}{v} \sum_{i=1}^{b-1} \sum_{j=i+1}^{b} v(i,j) \left| \tanh(\delta D_a(i,j)) - \sign(D_a(i,j)) \right|,$$  

where $v = \sum_{i=1}^{b-1} \sum_{j=i+1}^{b} v(i,j)$. To over-represent rare pairs we consider scores that are inversely proportional to the probability of such pairs. In particular we employ the exponential function:

$$v(i,j) = e^{-\alpha p^a_i p^a_j},$$

where $p^a_i$ is the probability of observing the attribute value of the $i$-th sample and $\alpha \geq 0$ is a parameter controlling the rate of score decrease with increasing pair probability. Note that as $\alpha \to 0$ the score of all pairs approaches 1, regardless of their probability and the weighted MAE loss reduces to the original unweighted version. In our experiments, small values of $\alpha = 0.01$ improve the representation of rare attribute instances without significant impact on the overall reconstruction accuracy.

To estimate the probability of attribute values empirically, we compute a fixed-bin-width frequency histogram from the attributes of training samples and normalize each bin by the total number of training instances. The probability of a specific instance $p^a_i$ is the probability of the bin corresponding to its attribute value.

4.4 Truncated VAE sampling for DNA design

Recall that we proposed the regularized VAE as an approach to represent the joint distribution $p(S,A)$ of DNA sequences and the corresponding Ag$_N$-DNA properties and our goal is to design DNAs with specific properties, i.e., sample $S' \sim p(S|A \in [A_{ib, A_{ub}])$, where $[A_{ib}, A_{ub}]$ specifies some property ranges of interest for design. In our specific case, we would like to design bright NIR Ag$_N$-DNAs, so the range of interest is high WAV and high LII.

The process of sampling from our VAE is demonstrated in Fig. 3. Since we cannot control directly WAV and LII, we sample $Z'$ from the latent space conditional on the property-regularized proxy dimensions being in specified bands. Since the latent distribution is regularized (via the KL divergence loss term $L_{KL}$) to approximate a normal prior distribution $p(Z)$ and retain samples that fall in the bands of interest, we do not need rejection sampling. Due to the Gaussian assumption for the latent embeddings, we can employ truncated normal sampling—an efficient sampling approach that does not require rejection [26].

Given a sample in latent space $Z'$ we employ the trained decoder to obtain an output approximation for a one-hot encoding $Y' \in \mathbb{R}^{1 \times 1}$ (Fig. 3).

5 EXPERIMENTAL EVALUATION

This section reports on new Ag$_N$-DNAs we experimentally synthesized based on sampled sequences from a trained VAE model and discusses implications of tuning, training and deploying the model, including effect of hyper-parameters and lessons learned from this first deployment for design of new Ag$_N$-DNAs.

5.1 Experimental setup

Data. Our training dataset consists of $|S| = 2661$ DNA sequences of length $l = 10$ together with the properties WAV and LII of their corresponding stabilized Ag$_N$-DNAs. The distribution of property values in the training set can be seen in Fig. 4 (grey bars). To visualize results and to select truncation points for sampling and synthesis of new Ag$_N$-DNAs from higher wavelengths, we adopt the same color class definitions from past work, motivated by the
physical properties of Ag$_N$-DNAs [6]. We define DNA sequences with WAV < 580 nm to be Green, 600 nm < WAV < 660 nm to be Red, and 660 nm < WAV < 800 nm to be Very Red (details in [5]). We also introduce a new NIR class with WAV > 800 nm, which is the particularly rare class (Fig. 4a) that we aim to target with sequence generation. These color definitions play no role in VAE training but are useful for comparing to past work on Ag$_N$-DNAs.

**Metrics.** Intuitively, a well-trained VAE for targeted Ag$_N$-DNA design should (i) reconstruct DNA sequences well and (ii) impose ordering in regularized (proxy) latent dimensions similar to that of their corresponding property observations from training. To quantify sequence reconstruction *Accuracy*, we measure the fraction of correctly recovered DNA bases after decoding, namely:

$$\text{Accuracy} = 1 - d_H(S, S')/l,$$

where $S$ is the input DNA sequence, $S'$ is its reconstructed DNA sequence after taking the maximum loadings from the VAE output encoding $Y'$ (See Fig. 3), $l$ is the length of sequences and $d_H(\cdot, \cdot)$ is the Hamming distance between the two argument sequences. To quantify the alignment of proxy dimensions with their observed attribute values, we compute the *Correlation* between the regularized latent dimension embedding of training/validation instances and their corresponding properties. Note that the above measures have counterparts in the loss function, but we use these more interpretable measures to select hyper-parameter configurations for synthesis. In tuning the model, we also reserve a random subset of instances for validation to gauge test experimentally following our sampling approach (Sec. 4.4). We employ a VAE trained on all instances (no validation set) and with hyper-parameters $a = 0.01, \beta = 0.007, y = 1, \delta = 1, |z| = 19, h = 13, w = 16$, single LSTM layer, and no dropout. We tune hyper-parameters by performing a grid search and select the model with both high Accuracy and Correlation for properties (details in the Supplement). We generate 1000 samples of DNA templates and rank them by their re-encoded WAV proxy $Z''_{WAV}$. Specifically, each sample $Z'$ is first decoded and translated to a DNA sequence $S'$, which is then re-encoded by the encoder to obtain the re-encoded WAV proxy. The top-20 sequences of highest $Z''_{WAV}$ are selected for synthesis.

We experimentally synthesized Ag$_N$-DNAs using the selected 20 strands and measured their fluorescence properties, finding that all 20 sequences yield brightly fluorescent Ag$_N$-DNAs with WAV between 695nm and 845nm. One Ag$_N$-DNA falls into our targeted region of WAV > 800nm, a 240% increase in NIR frequency compared to the training data. Notably, the other generated sequences form fluorescent Ag$_N$-DNAs very close to the NIR WAV threshold, without a single nonfluorescent sample or Ag$_N$-DNA at Green or Red WAV values (Fig. 4a). Furthermore, the distribution of fluorescence brightness values (LII) also increases substantially compared to training data (Fig. 4b).

### 5.3 Training, latent space and sampling

We next provide more insight into the best model employed for synthesis (Sec. 5.2). Fig. 5 summarizes various metrics of the model during training over $e = 2000$ epochs, using 85% of the data for training to also allow characterization of validation statistics. Fig. 5(a) shows the break-down of the loss components. The reconstruction $L_{REC}$ and regularization $L_{REG}$ loss components monotonically decrease as the VAE is learning to both encode-decode sequences and also training properties in their corresponding proxy dimensions in $Z$. This effect is also evident from Accuracy (Fig. 5(b)) and Correlation (Fig. 5(c),(5(d)) profiles. It is important to note that both proxy dimensions tend to retain significant correlation with both WAV and LII. This is because the training WAV and LII are inherently correlated and so are their proxies, regardless of the KLD loss that “works” to de-correlate the latent space.

Fig. 6 presents a visualization of the learned latent representations in terms of the Ag$_N$-DNA color classes defined in Sec. 5.1. Note that our VAE models the properties (WAV and LII) in continuous space, and we introduce this natural (and physically-motivated) binning into classes only to aid the visualization. The latent embeddings of training samples (both centroids and individual samples) follow the natural WAV order of classes (Fig. 6(a)). Note that the

**Figure 4:** Wet lab synthesis results: Probability density distributions of (a) WAV (units of wavelength in nanometers, representing Ag$_N$-DNA color) and (b) LII (brightness) for training (grey) and newly synthesized Ag$_N$-DNAs.

**Figure 5:** Reconstruction (a) and regularization (b) loss components during training over 2000 epochs (grey) and newly synthesized Ag$_N$-DNAs.

**Figure 6:** Visualization of the learned latent representations in terms of the Ag$_N$-DNA color classes defined in Sec. 5.1.
Figure 5: Training profiles of the VAE model we employ for de novo AgN₄-DNA synthesis over ε = 2000 epochs. (a): Break-down of the loss components for training (LREC, LLD, LREG) and the overall training LTRA and validation LVAL loss; (b): Training and validation accuracy; (c): WAV 5(d): LII correlation. The grey dashed curves (other) in the last two figures show the correlation of the remaining latent dimensions with the training properties.

Figure 6: Visualization of the learned latent representation Z for training data instances partitioned into WAV classes. (a) shows the means μ, of latent representation of samples in 3D space, where the horizontal axes are the first two principal components of the 19-dimensional embeddings and the vertical axis is the WAV proxy dimension. Individual samples are depicted as class-colored circles, while class-based centroids are shown as diamonds. (b): Color class-based distributions of training samples in latent space along the WAV proxy dimension.

Figure 7: WAV (a) and LII (b) proxy distributions of 1000 latent samples and their corresponding distributions after re-encoding the sequences (c),(d). Samples and re-encoded samples are superimposed over the training distributions depicted as wider black-bar histograms.

5.4 Ablation analysis
In this section, we remove essential components of the VAE model to test their impact on model quality (Fig. 8). Particularly, we are interested in answering the following questions: Is property regularization necessary for the VAE architecture to "isolate" the LII and WAV proxies into interpretable latent dimensions? Does the weighting of rare property samples improve the representation of corresponding samples in latent space?

We first characterize the latent space learned by an Unregularized β-VAE, i.e., γ = 0 and all other hyper-parameters set in the same way as in Sec. 5.2,5.3. Fig. 8(a) shows the correlation of all its latent dimensions with the WAV property. Unlike its regularized counterpart (training profile in Fig. 5(c)), none of this model’s latent dimensions correlate as strongly with WAV. A side-by-side comparison of the color class distributions of training samples of our Regularized model and its Unregularized counterpart is in Fig. 8(b) (first latent dimensions to Unregularized). We observe a similar behavior for the other regularized property, LII (figures not included). Without regularization, the β-VAE architecture cannot learn to isolate the AgN₄-DNA properties in its latent representation.

We also investigate the effect of weighted regularization (Sec. 4.3) on the representation of rare property values. Fig. 8(c) shows centroids of latent samples in the WAV proxy dimension for the four WAV classes as a function of α—the hyper-parameter controlling importance of rare pairs of properties in the regularization. We aim to "well-separate" rare property values. As α (and hence relative
5.5 Varying sequence length

Ideally, our model should be generalizable to other sequence lengths. Thus, we ask: Can the model trained for sequences of length \( l = 10 \) be employed for other values of \( l \)? To study this, we employ a small sample of \( \text{Ag}_N \)-DNAs stabilized by sequences of lengths \( l = 8, 12, 16 \) with measured WAV and LII from our recent work \[7\]. We investigate the quality of embedding for different length sequences in our model trained for \( l = 10 \). For this, we must first choose how to represent variable length sequences within a \( l = 10 \) VAE model.

For \( l = 8 \), we pad the sequence to increase length to \( l = 10 \). The padding characters can be placed on either side of the sequence (8-FB), the front only (8-F) and in the back only (8-B). Padding positions feature uniform distributions in the one-hot encoding (i.e. all four positions have a value of 0.25). For sequences of lengths longer than \( l = 10 \), we apply a sliding window approach and represent a single long sequence as a set of its sliding size-10 windows, all sharing the same WAV and LII properties.

We encode the sequences with our trained \( l = 10 \) model to characterize how the model embeds them. Fig. 9 presents the accuracy and correlation values for all \( l \) and padding options. While accuracies are comparable to validation results for sequence \( l = 10 \) (note that the expected Accuracy of \( l = 8 \) sequences is 0.8 as successful matching of padding characters is random), the WAV and LII correlations, however, are significantly lower than validation results for \( l = 10 \). This outcome suggests that simple padding and sliding window approaches are insufficient to generalize to varying \( l \). It may be necessary to consider alternatives in which \( l \) and “do-not-matter” positions are explicitly modeled.

6 DISCUSSION

Compared to past machine learning models for \( \text{Ag}_N \)-DNA design \[4, 5, 7\], the VAE generative model presented here has several new advantages. First, past models learned only a single \( \text{Ag}_N \)-DNA property (WAV or LII), while the generative model here can distinctly target multiple \( \text{Ag}_N \)-DNA properties, namely both WAV and LII. It is advantageous to design DNA sequences correlated with multiple \( \text{Ag}_N \)-DNA properties (e.g. fluorescence color, brightness, chemical stability, and sensitivity to an analyte of interest). Thus, multi-objective design methods like the models introduced here are critically needed to advance \( \text{Ag}_N \)-DNA applications. Second, this method does not require strictly defined \( \text{Ag}_N \)-DNA “color classes” to learn \( \text{Ag}_N \)-DNA color; this is particularly ideal for rare \( \text{Ag}_N \)-DNAs in the newly explored NIR spectral range, where little chemical information exists to motivate learning \( \text{Ag}_N \)-DNA color as a classification problem. Also of note is that our VAE model is more successful in targeting the high WAV space for generated sequences (despite no explicit “class” targeting), and in particular the high LII space. Finally, future examination of the latent space may provide new insights into how DNA sequence selects for \( \text{Ag}_N \)-DNA properties, advancing fundamental science of these nanomaterials.

While the current implementation of our model yields strong experimental performance, expansions for future work can lead to further improvements. Among these are expanding the attributes used to yield further classifying information, and more aggressive truncation of the sampling distribution to more strongly target the higher end of the WAV scale.

7 CONCLUSIONS

In this paper we proposed, evaluated and deployed a \( \beta \)-VAE generative model for the design of \( \text{Ag}_N \)-DNA nanomaterials. Our model was able to learn a joint representation for stabilizing DNA templates and \( \text{Ag}_N \)-DNA properties, including fluorescence color and fluorescence brightness, from a highly imbalanced training data set by regularizing the latent space to correlate with \( \text{Ag}_N \)-DNA properties. To counteract imbalanced training samples, our model
employed weighting scheme to over-represent such instances. To test the model’s efficacy, we targeted the design of DNA template sequences for especially rare NIR-fluorescent Ag\textsubscript{N}\textsubscript{N}-DNAs, which represent only 2% of training instances. Our experiments showed that out of 20 DNA template sequences generated by the VAE-based model, all succeeded in producing Ag\textsubscript{N}\textsubscript{N}-DNAs with both bright and high wavelength fluorescence, including a new NIR-emissive Ag\textsubscript{N}\textsubscript{N}-DNA with 840 nm peak fluorescence. The successful selection of a NIR Ag\textsubscript{N}\textsubscript{N}-DNA in this test set represents a 240% increase in the target Ag\textsubscript{N}\textsubscript{N}-DNA color class, an improvement upon past machine learning models for Ag\textsubscript{N}\textsubscript{N}-DNA design despite a significantly imbalanced training data. In addition to enhanced predictive power, our model is the first to learn multiple Ag\textsubscript{N}\textsubscript{N}-DNA properties, with significant implications for the advancement of Ag\textsubscript{N}\textsubscript{N}-DNA applications in bioimaging and biosensing. Our results show that VAE-based generative models are highly promising for the design of nanomaterials whose properties are encoded by biomolecular sequence and for which only sparse experimental observations may be available. As the fields of DNA and protein nanotechnology [29, 33] continue to expand, such computational models may be crucial in the advancement of biomolecule-based nanotechnologies.

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REFERENCES

SUPPLEMENTAL MATERIAL

Our model includes multiple hyperparameters (listed in Table 1) that can be grouped into two categories:

1. Architectural hyperparameters: \([|z|, L_w, L_d, h/2, w]\) controlling the shape and function of layers in the architecture; and

2. Loss hyperparameters: \((\alpha, \beta, \gamma, \delta)\) controlling the behavior of the loss function.

Hyperparameters have varying impacts on different metrics. We perform a grid search across all our hyperparameters to optimize our model for both accuracy and latent space correlation. Tested value of each parameter are listed in Table 1 and optimal parameters denote with bold font.

<table>
<thead>
<tr>
<th>Hyper-parameter</th>
<th>Values for grid search</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\alpha)</td>
<td>0.005, 0.006, 0.007, 0.008, 0.009, 0.01, 0.02</td>
</tr>
<tr>
<td>(\beta)</td>
<td>1.0, 3.0, 5.0, 10.0</td>
</tr>
<tr>
<td>(\gamma)</td>
<td>0.0, 0.3, 0.5</td>
</tr>
<tr>
<td>(\delta)</td>
<td>1.0, 5.0, 10.0</td>
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<tr>
<td>Latent Dimensions (</td>
<td>(z</td>
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<tr>
<td>LSTM Layers ((L_w))</td>
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<tr>
<td>LSTM Dropout ((L_d))</td>
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</tr>
<tr>
<td>LSTM Info (h/2)</td>
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</tr>
<tr>
<td>Encoder Width ((w))</td>
<td>12.0, 16.0, 20.0</td>
</tr>
</tbody>
</table>

Table 1: Table of hyperparameters that were used during the model testing phase, highlighted values represent the chosen hyperparameters. Note: \(h\) is the size of the concatenated output of both the forward and backward LSTM cells, hence the size of each LSTM cell hidden state (LSTM info) is \(h/2\).

To illustrate the effect of individual hyperparameters on the model’s reconstruction accuracy and correlations, we plot changes of these metrics in an interval around the optimal hyperparameters value while keeping the rest of the parameters set to their optimal values (bold in Tbl. 1).

We present results from this experiment in Figure 10, and indicate the optimal hyperparameter values by red squares. Consider first the figures on reconstruction accuracy as a function of hyperparameters. For \(\alpha\), \(\gamma\) and \(\delta\) it is evident that, as these hyperparameters increase in value, both training and validation accuracy decrease (\(w\) has a similar trend, as the change from 12 to 16 starkly increases training and validation accuracy, though stays stagnant upon further increase). \(|z|\) has the opposite effect: as we increase the values of this hyperparameter, training and validation accuracy both increase. Finally, our chosen \(\beta\) results in marginally smaller training and validation accuracy when compared to the other values of \(\beta\), and our chosen value for LSTM Info (h/2) results in larger training and validation accuracy than its counterpart values. The corresponding correlation figures, however, (WAV and LII Proxies represented by solid and dashed lines here, respectively) demonstrate that increases in reconstruction accuracy often results in deteriorated correlation. To achieve a good balance between reconstruction accuracy and latent correlation, we therefore choose our optimal hyperparameters indicated in Table 1.

Figure 10: Effect of hyper parameters on reconstruction accuracy and correlations. All parameters, apart from the one varied in each figure, are set to the optimal regimes denoted by bold values in Tbl. 1.