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De novo creation of fluorescent molecules via adversarial generative modeling

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The development of AI for fluorescent materials design is technologically demanding due to the issue of accurately forecasting fluorescent properties. Besides the huge efforts made in predicting the photoluminescent properties of organic dyes in terms of machine learning techniques, this article aims to introduce an adversarial generation paradigm for the rational design of fluorescent molecules. Molecular SMILES is employed as the input of a GRU based autoencoder, where the encoding and decoding of the string information are processed. A generative adversarial network is applied on the latent space with a generator to generate samples to mimic the latent space, and a discriminator to distinguish samples from the latent space. It is found that the excited state property distributions of generated molecules fully match those of the original samples, with the molecular synthesizability being accessible as well. Further screening of the generated samples delivers a remarkable luminescence efficiency of molecules epitomized by the significant oscillator strength and charge transfer characteristics, demonstrating the great potential of the adversarial model in enriching the fluorescent library.

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Introduction

Organic fluorescent materials, especially small-molecule organic dyes, have been extensively used across several research fields, including in sensors,^{1–4} organic light-emitting diodes (OLED)^{5–8} and bioimaging.^{9–11} It is thus significant to have a rational design framework so that the molecules can possess different functionality to meet various requirements. For instance, a significant luminescence quantum yield with controllable absorption/emission wavelength is generally needed for OLED molecules,¹² while a large Stokes shift and high photo-stability of fluorescent dyes are a prerequisite for reliable bioimaging.¹³ Up to now, most of the fluorescent design work is based on experimental experience in a trial-and-error manner, which is unlikely to give rise to the development of transformative new molecules due to the complexity of chemical space. On the other hand, owing to the complicated transfer mechanism happening in the excited state hypersurface during the photon absorption/emission process, the fluorescent efficiency can be hardly correlated with the molecular structure,¹⁴ posing challenges for the structural design work.

The advent of machine learning brings about a variety of opportunities for the exploration of unknown chemical space and predictions of molecular properties. In particular, numerous efforts have been made to investigate the excited state properties of organic semiconductors,^{15–21} and relevant *de novo* design frameworks^{14,22} have also been proposed to develop new molecules with desired functionalities. For fluorescent materials, machine learning has given successful predictions for the experimental quantum yields and emission energies,^{21,23} by using empirically designed chemical descriptors. The forecasting of oscillator strength and the corresponding fluorescent rate (specifically at the theoretical level) is however less satisfactory,¹⁹ where the machine learning techniques have to be used with caution. It is interesting to see an inverse design case for fluorescent molecules development by employing a reinforcement-like paradigm coupled with ‘*in situ*’ quantum chemical calculations.¹⁴ The generated novel compounds possess the anticipated emission wavelengths and significant Stokes shifts which are subsequently verified experimentally.

Adversarial generation, known as a powerful generative model in machine learning, has attracted great attentions in physical and chemical fields for the innovative design of drug molecules and optoelectronic materials.²⁴ The family of adversarial models basically includes adversarial autoencoder (AAE), generative adversarial network (GAN), reinforcement learning augmented GAN, *etc.*, where the model framework generally consists of a neural network generator and discriminator for the sample generation and discrimination respectively.

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In order to introduce an alternative inverse design paradigm for fluorescent molecules, this article employs an autoencoder based generative adversarial network (AGAN) for the generation of molecular SMILES²⁵ (simplified molecular input line entry system). The architecture is inspired by the LatentGAN,²⁶ in which a heteroencoder is first pre-trained before the training of the GAN. AGAN performs the training in a cohesive manner by optimizing the encoder/decoder and GAN simultaneously. The SMILES strings are fed into the encoder to be transformed to a low-dimensional latent space, which is subsequently decoded back to the character sequences with the reconstruction loss being minimized. A generator is trained to produce a virtual latent space from a Gaussian prior distribution to mimic the real latent samples, while a discriminator is employed to distinguish the real and virtual latent. High throughput quantum chemical calculations are carried out to label the excited state properties of generated molecules. Further fine screening of the generated samples gives rise to molecules with significant luminescence efficiency epitomized by the large oscillator strength and remarkable charge transfer characteristics, indicating the effectiveness of the AGAN model in exploring the fluorescent search space.

The paper is organized as follows. In the Methodology section, the dataset, details of quantum chemical calculations and the model architecture are described. It is then followed by the Results and discussion section which presents the main findings for our fluorescent molecules generation. Finally in section, a Conclusion is drawn.

Methodology

Fluorescent dyes database

The original fluorescent samples are extracted from the database used in Ju *et al.*,²³ where experimental emission wavelength and quantum yield are fitted with machine learning models. 2923 distinct SMILES strings are obtained and the singlet excited state properties are calculated for the labels of molecules. Note that the excited state properties for the 2923 original molecules are acquired in terms of high throughput calculations at the theoretical level, with no experimental data being involved here except for the experimentally synthesized molecular entries (represented as SMILES). All original SMILES are input into the AGAN model for the generative training.

Quantum chemical calculations

High throughput time-dependent density functional theory (TDDFT) calculations are implemented for both the original and generated molecules to obtain the excited state properties. The fluorescent compounds are first processed with the initial geometry optimization using the MMFF94 force field²⁷ in RDKit. The ground state geometries are then optimized using the PBE0 hybrid functional (which is shown to be the most realistic functional in predicting the emissive properties for fluorescent dyes²³) with the def2SVP basis set. TDDFT calculations are then implemented at S_0 geometries at the PBE0/def2SVP level, where the first three singlet transition energies and the corresponding

oscillator strengths are exported. All quantum chemical calculations are performed in gas phase using Gaussian 16,²⁸ with no solvent effect being considered for the sake of simplicity.

During the fine screening stage, the selected fluorescent candidates are optimized at the S_1 state with TDDFT/PBE0/def2TZVP, with the transition energies and oscillator strengths being extracted for the evaluation of emissive properties.

Autoencoder based generative adversarial network

Conventional GAN cannot be used to treat SMILES or other molecular representations due to the nondifferentiability of the input data. Several solutions have been proposed to tackle with this challenge. SeqGAN²⁹ is designed by treating the generator to create actions for sequence generations, while the discriminator is used to give feedbacks. ORGAN³⁰ is built on SeqGAN by incorporating an RNN-like generator and a CNN-like discriminator to generate SMILES strings with a pre-set objective. MolGAN³¹ is developed to directly process the topological graph, with the model architecture consisting of a generator, discriminator and reward network to generate molecular topology with nearly 100% validity. The current sequence-based and graph-based GAN frameworks, however, face the issue of generation quality where the produced compounds have to satisfy the criteria of validity, uniqueness and novelty. It is known that the ORGAN can have a significant non-validity ratio during the generation,³² while MolGAN can suffer from a very low molecular uniqueness. To the best of our knowledge, no model can yet deliver an 'ideal' generation.

In this research, we adopt a SMILES-based GAN architecture for the fluorescent molecules' generation. The SMILES strings are first transformed into a vector-based latent space *via* a GRU encoder (a 1-layer bidirectional Gated Recurrent Unit with 256 hidden dimensions is employed here), which is then transformed back (by using a 3-layer GRU decoder of 512 hidden dimensions) to character sequences for the reconstruction of SMILES. A one-hot embedding layer is applied before the encoder, and the teacher forcing³³ is employed in the decoder to lower the reconstruction loss. As exhibited in Fig. 1, the latent vectors are viewed as the real input for the discriminator. The generator serves to produce virtual latent samples from the Gaussian noise to replicate the real latent, while the discriminator tries to distinguish the two. The training process oscillates between the autoencoder and GAN until the virtual and real latent spaces are indistinguishable. The latent space is of dimension 128. The discriminator is a two-layer neural network with hidden dimensions of 512 and 256, while the generator is a two-layer network as well with hidden dimensions of 512 and 512.

To be in details for the SMILES transformation, the character string is first converted to a one-hot layer with a length set to be the maximal number of characters within the SMILES training data. The 0/1 digital layer is then linearly transformed into an embedded matrix with a dimension of 64, which is subsequently fed into a bidirectional GRU machine. Both forward and backward encodings in GRU are performed to account for the



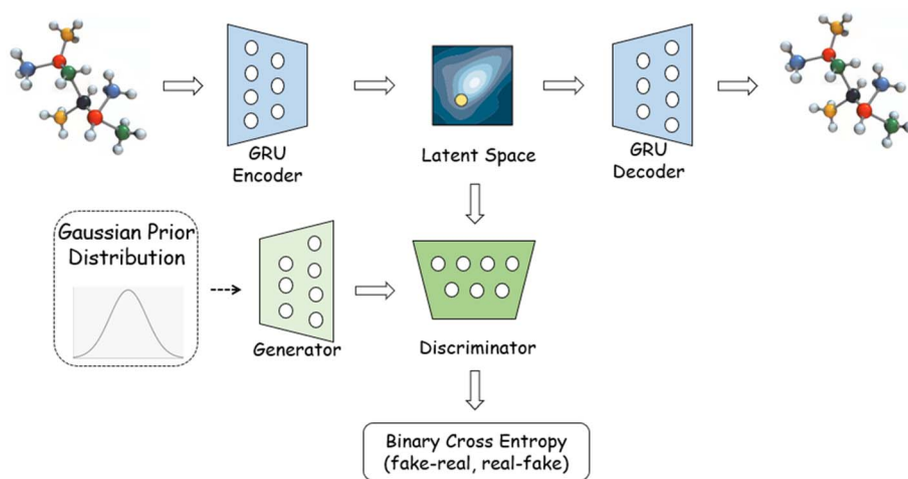


Fig. 1 The architecture of autoencoder based generative adversarial network.

foregoing and following information during the character vectorization. The final hidden state is eventually converted to the latent space *via* a linear transformation. For the decoding process, we perform an almost reverse procedure as conducted in the encoder, to transform the latent vector back into a multiclass probability matrix. The reconstruction is treated as a classification issue with cross entropy being used as the loss function.

We employ the binary cross entropy (BCE) to classify the real (latent space) and fake (virtual latent space) data in AGAN. Unlike the conventional GAN, we follow a relativistic approach³⁴ to differentiate samples by defining the loss function as,

$$G_loss = -\log[\text{sigmoid}(C(x_r) - C(x_f))] \quad (1)$$

$$D_loss = (\text{discr_real_loss} + \text{discr_fake_loss})/2 \quad (2)$$

$$\text{Discr_real_loss} = -\log[\text{sigmoid}(C(x_r) - C(x_f))] \quad (3)$$

$$\text{Discr_fake_loss} = -\log[1 - \text{sigmoid}(C(x_r) - C(x_f))] \quad (4)$$

where $C(x)$ denotes the discriminator network without the sigmoid layer, x_r and x_f represent the real and fake latent samples respectively. The generator loss (G_loss) can be interpreted as the probability that the given fake data is more realistic than the real data; while the discriminator loss (D_loss) estimates the degree of how realistic the real data is compared to the fake, and how counterfeit the fake data is compared to the real. The model design is claimed³⁴ to be more stable in producing high quality samples than other non-relativistic counterparts, and being computationally cost-effective compared to Wasserstein-GAN (which is frequently used in materials inverse design^{35,36}). It is stated that the relativistic framework is able to taking into account the missing element in the conventional GAN, where the generative network should simultaneously decrease the probability of real data being real when the probability of fake data being real is enhanced. By involving the relativity in the loss function, the model can lead to a better divergence minimization while only requiring

a single discriminator update per generator update (which actually reduces the training time by 400% compared to Wasserstein-GAN), achieving a sufficient computational efficiency along with a high data generation quality.³⁴

The training is conducted by optimizing the reconstruction loss, generator loss and discriminator loss simultaneously after a threshold epoch (set as 20 here), while before that only the autoencoder is trained. 150 training epochs are set, with the learning rate being 0.001 and batch size being 64. For the generation process, the items are randomly sampled from the Gaussian noise and then fed into the generator network, from which the output goes through the GRU decoder to produce the SMILES.

Results and discussion

Generation profiles

Fig. 2 exhibits the loss function convergences of AGAN. It is observed that recon_loss can smoothly converge to 0, demonstrating that an accurate enough mapping between the character string and the latent space is established. The generator and discriminator losses start to oscillate after the pre-set threshold epoch and converge to a level of 5 and 0.2 respectively. The finding is slightly different from the convergence profile in our previous adversarial training²⁴ where the classification loss can approach the theoretical value of BCE for indistinguishable samples. The phenomenon can be attributed to the relativistic approach applied on the discriminator which can possibly lead to a state-of-the-art during the convergence.

The performance of generations is measured with the molecular validity, novelty, uniqueness and overall efficiency, as defined in our previous work.²⁴ To briefly summarize, validity represents the fraction of molecules within the generated samples that can pass the RDKit's sanitization check so that the atomic valency and the consistency of bonds in aromatic rings can be maintained. Novelty is the fraction of the generated molecules that are not present in the training set. Uniqueness gives rise to the percentage of the generated samples that are



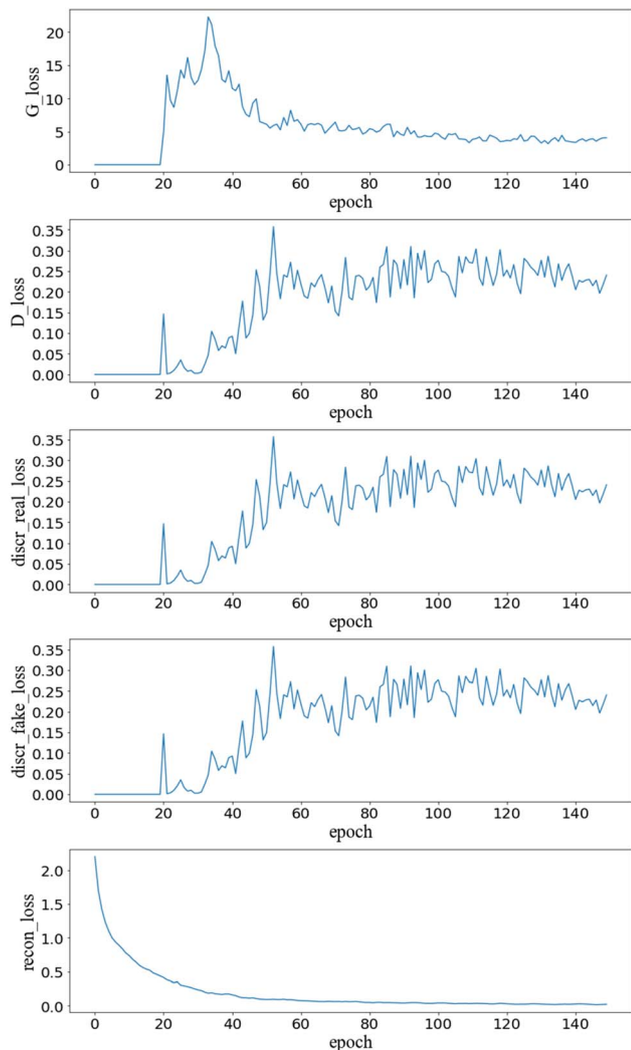


Fig. 2 Convergences of loss functions during the training of AGAN. G_loss , D_loss , $discr_real_loss$, $discr_fake_loss$ are defined in eqn (1)–(4), $recon_loss$ denotes the cross entropy reconstruction loss for the autoencoder.

unique. We compute the overall generation efficiency for molecules that simultaneously fulfill the above three criteria, given different generation sizes.

As shown in Table 1, given the small training set (which is 2923 in this case), generation sizes of 1 K to 20 K are considered. The uniqueness is found to be comparable with the profiles in

Table 1 The generation metrics of AGAN for fluorescent dyes: fraction of valid, novel, unique molecules and the overall generation efficiency fulfilling the above three criteria given different generation sizes

Generation size	Validity	Novelty	Uniqueness	Overall efficiency
1 K	0.2870	0.8600	0.9730	0.1470
5 K	0.2840	0.8632	0.9178	0.1326
10 K	0.2943	0.8618	0.8928	0.1371
20 K	0.2922	0.8677	0.8678	0.1305

Polykovskiy *et al.*³⁷ (approaching 1.0 for LatentGAN at 1 K generation size) and Tan *et al.*²⁴ (0.99 for AAE at 1 K generation size). For the novelty ratio, the current generation result is even better than that in Tan *et al.*²⁴ (where around 0.65 novelty is reported), possibly owing to the addition of the generator network in AGAN which can potentially enhance the probability of creating new molecules not appearing in the original dataset. It is observed that, however, the validity is relatively low. The underlying reason can be tentatively ascribed to the small size of the training data, where there are not enough samples to inform the autoencoder for how to reconstruct a valid SMILES string. The overall generation efficiency is located between 13% and 14%, with 2610 molecules being efficiently produced in case of the generation size of 20 K. We would state that the current generation efficiency is fully acceptable based on the comparability of the efficiently generated size and the input size.

High throughput TDDFT calculations are performed to obtain the singlet transition energies and oscillator strengths for both the generated and original fluorescent dyes, the distributions of which are displayed in Fig. 3(a). It is found that the property distributions of the generation space almost reproduce those of the original space, indicating the effectiveness of the AGAN model. The first excitation energy is in the range of 1.5–4.5 eV (with the generated distribution being slightly broader), covering the full visible spectrum for the light absorption. Both the generated and original oscillator strengths tend to have a distribution near 0, implying the difficulty to find a molecule with large luminescence efficiency.

It is interesting to note that, the fine structures of S_1 distributions differ between the original and generated samples, while the distributions of S_2 and S_3 for the generated dyes seem to have a plausible ‘blue shift’ towards higher energies. The observation is in contrast to our previous work²⁴ where perfect matches of distributions are found when thermally activated delayed fluorescence molecules are generated. Two possible reasons might be announced to address the inconsistencies. One is for the insufficient data within the histogram which can lead to the irregularity of distribution curves. In addition, the diversity in the training set (where samples are known to be experimentally synthesized and collected from different literature), can be another cause for the distribution mismatch, while in Tan *et al.*²⁴ both original and generated sets are of the same molecular type. More details regarding the diversity can be seen in the following analyses.

The synthetic accessibility (SA)³⁸ is also examined which is measured in terms of the fragment contributions and the molecular complexity. The SA score is simply estimated as the difference between the fragment score and the complexity penalty, where the fragment score can be understood as the frequency of the current molecular fragments appearing in the empirical chemical fragment database, and the complexity penalty characterizes the presence of complex structural features that may lead to overcounts of molecular substructures. Note that the SA score spans from 1 to 10 with higher values denoting enhanced difficulties to synthesize a molecule. A major peak of SA around 2–3 is found, indicating that these



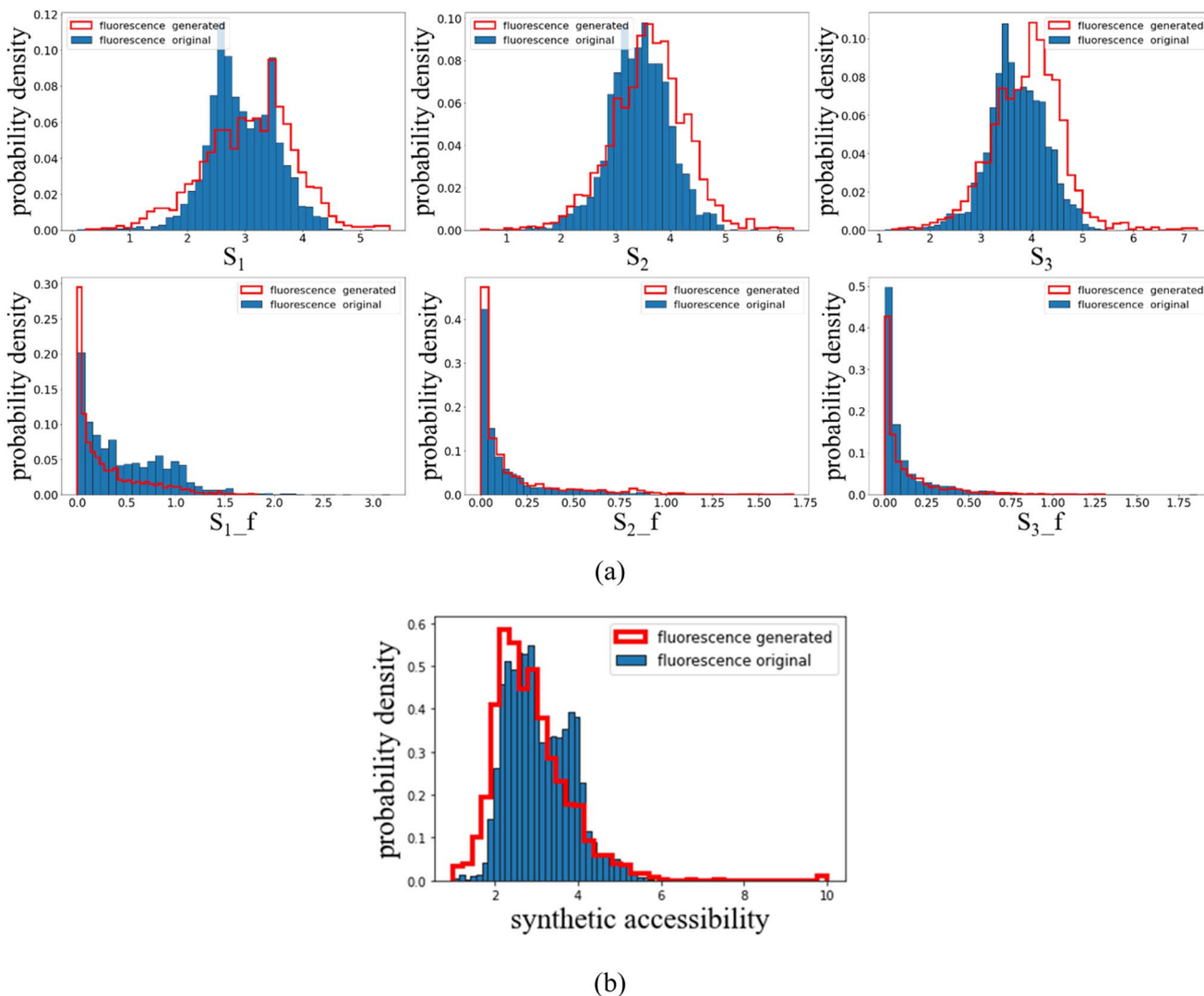


Fig. 3 (a) Distributions of singlet excited state energies and oscillator strengths for the generated and original fluorescent dyes. Note that energies are in eV and oscillator strengths are dimensionless. (b) Distributions of synthetic accessibility for the generated and original fluorescent molecules.

fluorescent dyes are easily synthesizable. Since the original molecules are recorded to be synthesized experimentally,²³ the generated samples are supposed to have a wide experimental accessibility.

Regarding the diversity of the original and generation sets, internal diversities (IntDiv₁ and IntDiv₂, as defined to be one minus the average Tanimoto similarity within a molecular set,²⁴ with higher values denoting higher diversity) are computed. IntDiv₁ and IntDiv₂ (the indices represent the order of the diversity) for the original samples are 0.8845 and 0.8618, while those for the generated samples are 0.8854 and 0.8696. The analysis points towards an enhanced diversity compared to the values in Tan *et al.*,²⁴ where a level around 0.82 is reported. The discrepancy makes sense as the molecular entries in the current training set are collected from different experimental literature, while only a focused library of donor–acceptor molecules is used in our previous work.

Fig. 4 gives rise to a two-dimensional principal components analysis (PCA) for the latent space, to further investigate the diversity of the original and generation sets. The original and generated SMILES are separately fed into the autoencoder where the models are trained with the reconstruction loss being minimized respectively. The converged autoencoder (after 150 epochs) can therefore deliver the latent space as robust numerical representations for character strings. A dimension reduction *via* PCA is then performed to extract the main features of the chemical space. As shown in Fig. 4, the reduced original space exhibits a certain degree of aggregation which may correspond to the similar topologies in the experimental synthesis. A broader coverage of the reduced generation space is found compared to the original one, implying a possible higher diversity of the generated samples. It is interesting that the generated space can fully capture the major characteristics of the original space including the structural aggregation,



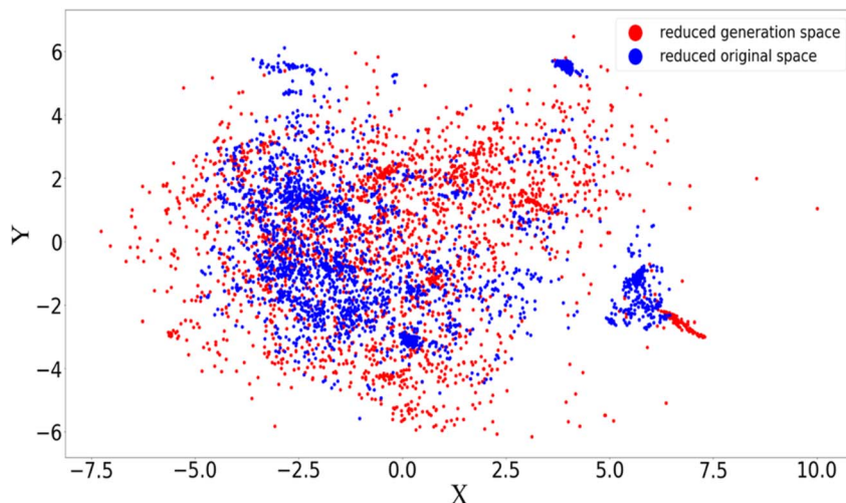


Fig. 4 Two-dimensional PCA analysis of latent space for the autoencoder used in AGAN. The X and Y axes are the principal components selected from the PCA analysis.

demonstrating the reliability of the adversarial algorithm in producing mirrored spaces. The detected augmentation of diversity in the generation set might partially explain the energy distribution mismatch issue mentioned above.

Fine screening

In order to explore the emissive properties of the generated molecules, we select 10 dyes with the largest absorptive oscillator strengths from the generation space to perform the excited state optimization. The S_1 geometry optimization is conducted at the TDDFT/PBE0/def2TZVP level (where the PBE0 is justified to be the most accurate functional in describing the fluorescent properties according to Ju *et al.*²³), and the emissive energies and oscillator strengths are displayed in Table 2. Significant emissive oscillator strengths are observed which is comparable

with the absorptive ones, implying a remarkable fluorescent rate for these new molecules. The emission energies lie in the visible region and being slightly smaller than the absorption energies, exhibiting moderate Stokes shifts as calculated from 5 nm to 33 nm. The molecular structures of the fine screened dyes are shown in Fig. 5, where π -conjugated, polycyclic, benzonitrile-like systems are observed, as usually present in the fluorescent compounds.³⁹

The optimized S_0 and S_1 geometries for the two representative molecules from Table 2 are exhibited in Fig. 6. No substantial structural displacement is found between the ground state and the excited state, which is consistent with the small Stokes shift where relaxations on the hypersurface can readily happen. The highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) are both highly delocalized with a notable charge transfer

Table 2 The emissive and absorptive S_1 energies and corresponding oscillator strengths for the finally screened fluorescent candidates

Id	Smiles	Emission		Absorption	
		E_S1 (eV)	E_S1_f	A_S1 (eV)	A_S1_f
mol_1	<chem>CCN1/C(C2=CC=C(C3=CC=C(N(C4=CC=CC=C4)C5=CC=CC=C5)C=C3)S2)=NG6=C1C7=C(C8=C6C=CC=N8)N=CC=C7</chem>	2.969	1.4451	3.0344	1.4718
mol_2	<chem>FC(C=CC=C1)=C1C=CC2=CC=C(NC(C3=CC=CC=C3C)=O)C=C2</chem>	3.5919	1.4546	3.6645	1.4568
mol_3	<chem>F[B-]1(F)[N+]=2C(=C#Cn1c(cc3)\C=C\c1oc(cc1)-c1ccc([N+](=O)[O-])cc1)C=CC=2</chem>	2.3355	1.6269	2.3926	1.612
mol_4	<chem>OC1=CC(O2)=C(C=C1)C=C2/C=C/C3=CC=C(N4CCCC4)C=C3</chem>	3.2705	1.6294	3.3375	1.6566
mol_5	<chem>OC1=CC=C(C#CC2=CC(C#N)=C(C#CC3=CC=CC(C#N)=C3)C=C2)C=C1</chem>	3.2075	1.5565	3.2493	1.5583
mol_6	<chem>F[B-]1(F)[N+]=2C(=C#Cn1c(cc3C)\C=C\C1c1oc(cc1)-c1oc(cc1)-c1oc(cc1)-c1ccc([N+](=O)[O-])cc1)C=CC=2</chem>	1.9596	1.5718	2.0132	1.5316
mol_7	<chem>N#CC1=CC=C(C#CC2=CC=C(C#CC3=CC=CC=C3C#N)C=C2C#N)C=C1</chem>	3.2363	1.8274	3.2864	1.8329
mol_8	<chem>F[B-]([N+])1=C2C=C(C3=CC=CC=C3)OC2=CC1=C4(F)N5C4=CC(O6)=C5C=C6C7=CC=CC=C7C(C)C</chem>	2.2242	1.8164	2.364	1.7839
mol_9	<chem>FC1=CC=C(C2=CC=C(C#CC3=CC=CC=C3)C=C2)C=C1</chem>	3.8249	1.4566	3.8856	1.4618
mol_10	<chem>C1(N(C2=CC=CC=C2)C3=CC=CC=C3)=CC=C/C=C/C4=CC=C(N(C5=CC=CC=C5)C6=CC=CC=C6)C=C4)C=C1</chem>	2.945	1.5698	3.0107	1.6149



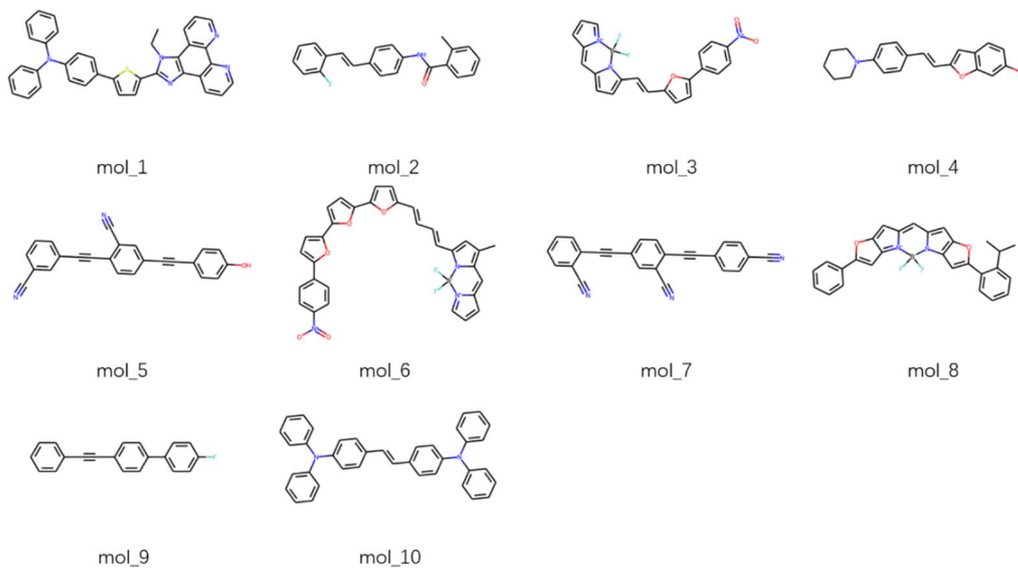


Fig. 5 Topological structures of fluorescent molecules from the generation space with the largest 10 absorptive oscillator strengths.

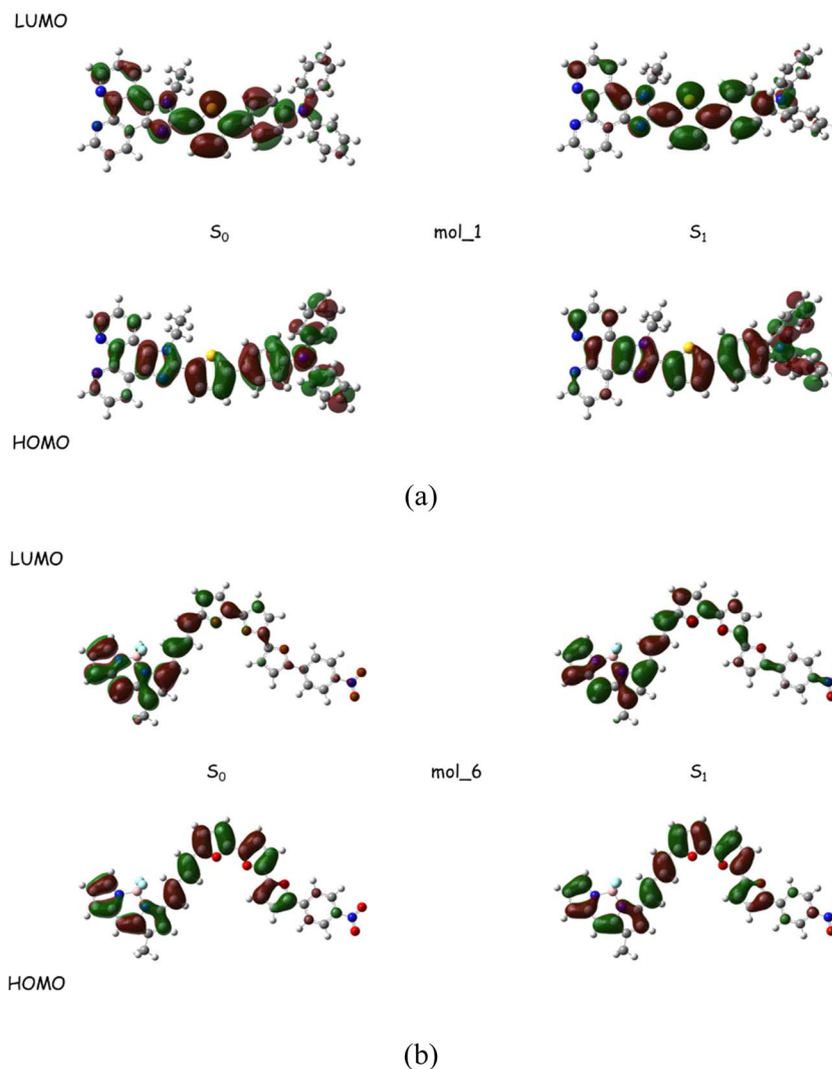


Fig. 6 Frontier orbitals for two of the fine screened fluorescent dyes at S_0 and S_1 geometries.



characteristics being detected. The electronic structure of the generated samples can imitate most of the known fluorescent dyes,^{40,41} further validating the effectiveness of the adversarial model in creating molecules with reliable photophysical properties.

Conclusions

For a short conclusion, the article has proposed an efficient framework by employing the autoencoder based generative adversarial network to generate novel fluorescent compounds. The GRU encoder and decoder are utilized to map the string-based data onto the low-dimensional latent space. A GAN architecture is then applied to create a virtual latent space from the generator so that it can fully mimic the real latent based on a relativistic loss mechanism. The new samples are produced from the Gaussian noise and eventually decoded to molecular SMILES with the validity, novelty, uniqueness being examined. Given the current generation efficiency (around 13%), the model can deliver more than 2000 new molecules with satisfactory luminescence properties. Considering the small training input size (2923 fluorescent molecules that are experimentally synthesized), it is equivalent to doubling the fluorescent library in a very low computational cost (where the model cost-effectiveness is stated in contrast to the empirical trial and error method used in conventional molecular design which may take decades to develop thousands of fluorophores). High throughput TDDFT calculations are performed for nearly 5500 samples to obtain the excited state properties. It is found that the distributions of singlet transition energies and oscillator strengths for the generated and original molecules are highly comparable. The SA of the generation space is also similar with the original one, further validating the experimental accessibility of the novel compounds. By implementing the excited state optimization for the molecules with the largest absorptive strengths, the generated dyes are proved to possess a remarkable fluorescent rate along with a moderate Stokes shift. The charge transfer characteristics are detected in the frontier orbitals analysis, demonstrating the efficiency of AGAN in producing anticipated electronic structures.

The current adversarial methodology has displayed a great potential in designing molecules with the desired fluorescent properties. It can be easily extended to the design of a large family of luminescent materials with different functionalities, with one of the most attractive design cases possibly being to develop molecules with large Stokes shifts particularly for bioscience applications. Additional improvements of the model are expected including the enhancement of the generation validity given the small input size, or by involving a predictor-augmented reinforcement algorithm⁴² to further elevate the generation efficiency, and the works are subject to the future research.

Data availability

All of the codes associated with the research work are made available at [https://github.com/Xiyuan-Quantitative-](https://github.com/Xiyuan-Quantitative-Technology/relativistic-GAN)

Technology/relativistic-GAN. The data that support the findings of this study are obtainable from the corresponding author upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

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