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1 Introduction

Photodynamic therapy (PDT) has been extensively investigated for decades as a treatment for tumors due to its minimally invasive nature, ability to preserve normal tissues, and approval for early-stage cancer treatment with relatively low pain.^{1,2} Triplet photosensitizers (PSs) play a key role in determining the efficacy of PDT.^{3,4} However, the traditional trial-and-error efforts to identify successful triplet PSs, which rely on expert knowledge, are time-consuming and intricate processes. This approach involves selecting potential chromophores, introducing reasonable motif derivatization, ensuring molecular synthesizability, achieving long wavelength absorption, and most importantly, securing high intersystem crossing (ISC) efficiency.

Effective generation of heavy-atom-free triplet photosensitizers containing multiple intersystem crossing mechanisms based on deep learning[†]

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Photodynamic therapy (PDT) is a clinically approved therapeutic modality that has demonstrated significant potential for cancer treatment, and triplet photosensitizers (PSs) play a key role in its efficacy. Despite deep learning having emerged as a next-generation tool for material discovery, existing methods mainly target a limited subset of triplet PSs, such as thermally activated delayed fluorescence (TADF) materials, neglecting the critical intersystem crossing (ISC) between the high-lying singlet and triplet states ($\Delta E_{S_{art}}$). To overcome this limitation, we compiled a comprehensive dataset ($\sim 1.90 \times 10^9$) of triplet PSs encompassing various ISC mechanisms. Then, we proposed a novel strategy that incorporates two models: a fragment-based model (Frag-MD) and a character-based model (MD), both integrating a conditional transformer, recurrent neural networks, and reinforcement learning. In silico experiments revealed that the Frag-MD model outperforms the MD model in generating larger conjugated motifs with higher average ring numbers and atom counts; while the MD model generates twice as many unique motifs and excels in novelty and diversity, as evaluated by conditional and MOSES metrics. Therefore, our approach is highly effective for modifying conjugated motifs and designing novel triplet PSs. Notably, the recently reported high-efficiency triplet PSs have been re-identified through ablation experiments using our proposed models, which target $\Delta E_{S_{\alpha}T_{\alpha}}$ and significantly outperform traditional baselines, achieving a prediction accuracy of 73% versus 4%. Our approach holds the potential to establish a new paradigm for discovering novel PSs applicable in PDT.

In recent years, the development of artificial intelligence (AI), especially through machine learning (ML) and deep learning (DL) approaches for molecular design, has significantly transformed the research paradigm in molecular discovery within the medical and materials science communities, sparking considerable interest.⁵⁻⁷ Leveraging advances in computing power and algorithm efficiency, these approaches exhibit an ever-increasing non-linear exploration capability to navigate the vast potential search space and optimize multiple properties simultaneously. They have demonstrated remarkable effectiveness in molecular property prediction, high-throughput virtual screening, and inverse molecular design or molecular generation under given constraints.^{5,8-14} Therefore, these approaches hold great promise for accelerating the discovery of novel triplet PSs.

Given that the ISC process of triplet PSs is a first-order perturbation process, the ISC rate constant can be described by eqn (1), a derivation of equation from the Fermi golden rule, which inevitably depends on quantum chemistry calculations (*e.g.*, semi-empirical methods or density functional theory (DFT) calculations) to obtain the key parameter of the ISC process ($\Delta E_{S,T,r}$, $n \ge 1$, the excited singlet-triplet energy gap).

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$$k_{\rm ISC} \propto \frac{\langle T_n | H_{\rm so} | S_n \rangle^2}{\left(\Delta E_{{\rm S}_n {\rm T}_n}\right)^2} \tag{1}$$

where H_{so} is the Hamiltonian for spin–orbit coupling (SOC) and $\Delta E_{S_nT_n}$ is the energy gap between the excited singlet $(S_n, n \ge 1)$ and triplet $(T_n, n \ge 1)$ states. Thus, it is a common but useful way to design molecules with a smaller energy gap between the S_n and T_n states to facilitate efficient ISC.

It is obvious that the high computational cost of quantum chemistry limits the availability of labeled data for DL-based molecular design approaches. On the other hand, DL-based inverse molecular design, which generates *de novo* molecules with specific and desirable properties rather than relying on high-throughput virtual screening by a molecule prediction model, explores a much broader chemical space. However, it remains a highly challenging task to design successful triplet PSs using DL-based inverse molecular design while simultaneously considering numerous factors such as synthetic accessibility (SA), quantitative estimate of drug-likeness (QED), absorption wavelength, ISC efficiency, and so on.¹⁵ For triplet PSs, heavy-atom-free triplet PSs offer advantages such as low cost, reduced potential toxicity, easier preparation and longer triplet lifetimes compared to triplet PSs that rely on heavy-atom enhanced ISC effects. Recently, heavy-atom-free triplet PSs have gained significance not only in PDT^{3,4} but also in photocatalysis,^{16–20} triplet–triplet-annihilation upconversion (TTA-UC),^{21,22} thermally activated delayed fluorescence (TADF),^{23–25} and other applications. Therefore, our work focuses on designing heavy-atom-free triplet PSs.

To address the challenges of insufficient labeled data in high-throughput virtual screening²⁶⁻²⁸ and inverse molecular design,^{5,8-10,14,29-33} various strategies have been developed,

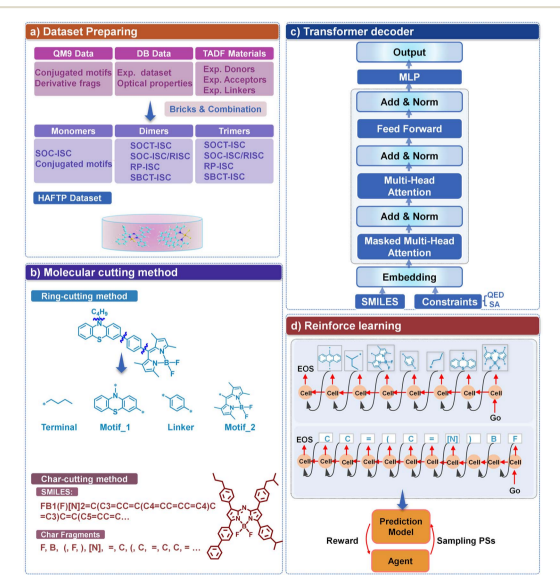


Fig. 1 Workflow of Frag-MD with a fragment-based method and MD with a character-based method. (a) Dataset preparation details, (b) ringcutting method and char-cutting method, (c) transformer decoder, and (d) fragment-based RNN or character-based RNN for reinforcement learning.

including constructing chemically reasonable and constrained molecules, selectively labeling a subset of molecules, and employing DFT functionals with lower computational cost. For example, Gómez-Bombarelli et al.5 built a space of 1.6 million molecules by combining 110 donor, 105 acceptor and 7 bridging moieties. More specifically, Buglak et al.30 focused on 70 heavyatom-free boron-dipyrromethene (BODIPY) dyes in three different solvents to predict singlet oxygen generation using a multiple linear regression model within a DL framework. To improve the model's prediction accuracy of ΔE_{S1T1} and reduce its dependence on labeled data, Wang et al. developed a smart strategy that combines a heuristic algorithm (active learning) with a deep neural network.34 This resulted in an efficient and useful self-improving discovery system from a space of more than 7 million molecules. Similarly, Nigam et al. developed a DL workflow to find novel organic emitters with desired ΔE_{S1T1} and appreciable fluorescence rates,13 using a genetic algorithm as the heuristic algorithm. More recently, Blaschke et al.35 constructed a framework that connects the REINVENT reinforcement learning framework to discover acceptors in TTA-UC.14 However, a key point that has been overlooked is the ISC process arising from higher excited singlet and triplet states $(\Delta E_{S_nT_n}, n > 1)$, leading to limited studies on the ΔE_{S1T1} value³⁶ and TADF molecules ($\Delta E_{S1T1} \leq 0.2 \text{ eV}$).³⁷ While the ISC process of TADF molecules includes the SOC-ISC and the revised ISC (RISC) process, which extend the triplet state lifetime and enhance their applicability in PDT, TADF molecules represent only a small subset of ideal triplet PSs. Notably, some monomers achieve high ISC efficiency purely through the SOC-ISC mechanism, without requiring RISC. For example, porphyrin derivatives have been used as commercial reagents in PDT.38,39 Similarly, other donor-acceptor systems can be effective candidates through a spin orbital charge transfer ISC (SOCT-ISC)40 or radical pair ISC (RP-ISC) mechanism,41 which are modulated by solvent polarity, electronic coupling, and other related parameters.42

In this study, we first developed a comprehensive dataset $(\sim 1.90 \times 10^9)$ of triplet PSs with multiple ISC mechanisms by combining linkers, donors and acceptors. The ISC efficiency metric was set within a $\Delta E_{S_nT_n}$ range of 0–0.3 eV. Beyond traditional molecular-level metrics, we adopted conjugatedmotif-level metrics as a more reasonable evaluation approach. We then proposed a novel strategy for designing heavy-atomfree triplet PSs, which mimics the design process of a chemistry expert and can be divided into two main tasks. Task 1 is the de novo design aimed at discovering new conjugated motifs, and Task 2 employs a conjugated-motif-based method focused on conjugated motif derivation. To accomplish these two tasks, we utilized a character-based molecular generation architecture (MD/GD)43 for Task 1 and a fragment-based molecular generation architecture (Frag-MD/Frag-GD)⁴⁴ for Task 2 (Fig. 1). The general reinforcement learning (RL) workflow for target-directed molecular generation consists of a recurrent neural network (RNN) and scoring function models.35,45 Frag-GD and GD only consist of general modules, while Frag-MD and MD consist of mixture modules which introduce a conditional transformer for molecule dataset

generation, followed by general modules,⁴⁴ which bias the molecular space towards the desired region. The final label 'D' means the scoring function model of the studied models trained from diverse molecule types and multiple ISC mechanisms. In contrast, Frag-MB/Frag-GB and MB/GB are regarded as 4 baseline models, where the scoring function model is trained on data labeled with ΔE_{S1T1} . In silico experiments demonstrate the practical viability of our method in accomplishing both tasks. Notably, recently reported high-efficiency triplet PSs have been re-identified through ablation experiments, further demonstrating the molecular design ability of our proposed models. Thus, Frag-MD/MD has the potential to discover novel PSs applicable in PDT.

2 Methods

The workflow presented here is developed based on a policy gradient model with a modified version of REINVENT³⁵ by introducing a conditional transformer. For de novo molecular design in Task 1, we implemented a conventional char-cutting method,⁴⁶ referred to as the character-based method, to facilitate free molecular derivation without the constraint of preserving conjugated motifs. For Task 2, which focuses on designing conjugated-motif-based molecules, we split molecules into fragments using a ring-cutting method to retain conjugated motifs (referred to as the fragment-based method).47,48 As shown in Fig. 1, the char-cutting method splits molecules into atoms or functional groups, which serve as tokens for a molecule generation model (e.g., an RNN model). In contrast, the ring-cutting method splits molecules into fragments based on single bonds outside the ring. These fragments include terminal groups (R-groups), conjugated motifs (scaffolds), and linkers. Specifically, the wavy lines represent the cleavage sites, and the star symbols represent the combination sites.

2.1 Dataset

To design triplet PSs with multiple ISC mechanisms, a wellcurated dataset is crucial. Hence, we constructed our dataset by combining fragments from conventionally studied chromophores,⁴⁹ custom donors or acceptors in TADF materials,³⁶ and molecules from the QM9 dataset.⁵⁰ It has the following advantages: (1) conventionally studied chromophores, which are typically large conjugated molecules, exhibit long-wavelength absorption and practical derivatization, aligning well with the desired molecular space; (2) molecular scale and molecular diversity can be further expanded by introducing A–D–A, A–A, and D–D systems to the conventional D–A and D–A–D types (TADF molecules); (3) the incorporation of novel fragments from the QM9 dataset (124 K), which is widely used in quantum chemistry models, significantly enhances molecular diversity while maintaining computational feasibility.

Detailed information on dataset construction is provided in ESI Part 1,† and its basic characteristics are summarized in Table 1. The final dataset contains $\sim 1.90 \times 10^9$ molecules, with a subset of 3.5×10^6 molecules (500 K molecules for each

Molecule types	Molecule numbers	Dataset components	Labeled molecules	Descriptions of combination
Monomers	${\sim}1.90 imes10^{9}(1898972708)$	50 0000	${\sim}1.40 imes10^4~(14005)$	Con* + termi-frag
Dimers1 (directly connected)		50 0000		Con* + Con*
Dimers2 (alkyl linkers)		50 0000		2 Con* + alkyl**
Dimers3 (aryl linkers)		50 0000		$2 \operatorname{Con}^* + \operatorname{aryl}^{**}$
Trimers1 (directly connected)		50 0000		2 Con* + Con**
Trimers2 (alkyl linkers)		50 0000		3 Con* + alkyl***
Trimers3 (aryl linkers)		50 0000		3 Con* + aryl***

^a Con, star, alkyl and aryl stand for conjugated motifs, reaction site, alkyl linker and aryl linker, respectively. The Con recovery rate of dataset is 1.0.

molecule type) selected in our study to balance computational costs. To the best of our knowledge, this dataset is substantially larger and contains a broader range of molecular types compared to previously reported datasets $(10^3-10^5, \text{ specifically TADF})$ molecules in the initially studied dataset).^{1,29,34}

2.2 Agent in reinforcement learning

In RL, the agent for all models adopted an RNN architecture to generate molecules tailored to desired properties through score functions. To constrain the RNN-generated molecules to a reasonable range of SA and QED properties while enabling subsequent target optimization of ΔE_{ST} and E_{abs} and improving the fine-tuning efficiency, a lightweight RNN architecture (MD/ Frag-MD) was derived from a heavier transformer decoder model through knowledge distillation.⁵¹

The conditional transformer decoder was trained on the selected dataset (3.5×10^6) . Its architecture is illustrated in Fig. 1, where the next token in a Simplified Molecular Input Line Entry System (SMILES) sequence is generated based on the preceding token. To mitigate the risk of information leakage from tokens within the decoder portion of the sequence, a masked multi-head self-attention layer was implemented in this model. As the core component of the transformer decoder, this layer integrates multiple scaled dot-product attention mechanisms, enhancing the model's ability to effectively extract and prioritize critical information from sequential data. The attention mechanism is concisely represented by eqn (2):

Attention
$$(Q, K, V) = \operatorname{softmax}\left(\frac{QK^T}{\sqrt{d_k}}\right)V$$
 (2)

where *Q*, *K* and *V* are the query, key, and value vectors, respectively, and d_k is the dimension of *Q* and *K*.

To generate molecules with desired properties, a conditional character P is incorporated as an input during the molecule generation process. This conditional character P is a Boolean string of length 2, where each bit (1 or 0) indicates whether the molecule meets the QED and SA conditions. Once embedded, the conditional character and the molecular SMILES strings are concatenated, enabling the model to learn from both sources concurrently during training. Specifically, during the training of the conditional transformer decoder model, the following loss function is minimized given a conditional character P, as shown in eqn (3).

$$L = -\sum_{i=1}^{n} \log p(S_i | S_{
(3)$$

To generate SMILES under a conditional character *P*, the model learns the conditional distribution p(S|P), as shown in eqn (4).

$$p(S|P) = \prod_{i=1}^{n} p(S_i|S_{
(4)$$

Using the trained conditional transformer decoder, constrained desired molecules were generated based on a conditional character *P* (QED and SA), resulting in a final new dataset of molecules (3.5×10^6) . During the training stage of the RNN with this dataset, the loss function aims to minimize the negative log-likelihood or cross-entropy function, as shown in eqn (5).

$$L = -\sum_{i=1}^{n} \log p(x_i | x_{< i})$$
(5)

In the generation stage of the trained RNN, the autoregressive method was used to generate SMILES strings in units of molecular fragments or characters as shown in eqn (6).

$$p(x) = \prod_{i=1}^{n} p(S_i | S_{
(6)$$

2.3 ΔE_{ST} and E_{abs} prediction models as scoring functions in reinforcement learning

The labeled dataset for the prediction model contains ~14 K molecules for multiple molecule types that favor multiple ISC mechanisms (Table 1). To determine the energy gap between high excited singlet and triplet states ($\Delta E_{S_nT_n}$, $1 \le n \le 6$) and the energy level of the absorption wavelength (E_{abs}), ten excited singlets and ten exited triplets were calculated based on the ground state geometries optimized using DFT//B3LYP/6-31G. Including more excited states enhances the accuracy of the first six excited energy levels. These calculations were performed using the Gaussian 16 program package.⁵² The prediction model employed is a typical molecular graph convolutional neural (GCN) network.⁵³ The molecular graph is an undirected graph G = (V, E), where nodes V and edges E represent atoms

and chemical connection bonds between atoms, respectively. The ConvMolFeaturizer function was used to encode the molecular fingerprint into a two-dimensional representation, which facilitates the prediction of quantum properties.54 Hyperparameters such as the number of graph convolutional layers, dense layers, dropout, and learning rate were optimized using a grid search method, which systematically evaluates all possible combinations of specified hyperparameters via crossvalidation. An early stopping strategy was implemented during the training stage. The relationship between training epochs and the mean absolute error (MAE) values of ΔE_{ST} and E_{abs} is presented in ESI Fig. S1,[†] and the MAE of ΔE_{ST} in the subset of the dataset was less than 0.3 eV, indicating that our model is accurate enough for RL applications in triplet PS design. Further details about the prediction model are provided in ESI Part 2.†

2.4 Loss function in reinforcement learning

In this study, we employed the RL strategy implemented in REINVENT to refine our RNN model (agent). The loss function was optimized using the policy gradient descent optimizer method to fine-tune the agent model. The RL fine-tuning procedure is outlined as follows: (1) copy the trained fragment/ character-based RNN model as the prior model, keeping all parameters fixed during fine-tuning, and use the original fragment-based or character-based RNN model as the agent model; (2) sample molecules with the agent model and save the likelihood of each sample molecule as log $p(S)_{Agent}$, and use the prior model to estimate log $p(S)_{Prior}$ for the generated SMILES strings; (3) calculate the reward score from the prediction models, multiply it by a coefficient σ , and define the loss function as follows:

$$Loss = [\log p(S)_{prior} + \sigma Score(S) - \log p(S)_{agent}]^2$$
(7)

2.5 Experiment setting

The objective of our model is to design heavy-atom-free triplet PSs containing multiple ISC mechanisms. A successful triplet PS needs to satisfy the following four constraints: $\Delta E_{ST} \leq 0.30$, $E_{\rm abs} \leq 2.48$, QED ≥ 0.38 , and SA ≤ 4.0 . The threshold of $\Delta E_{\rm ST} \leq$ 0.30 captures not only TADF molecules with the SOC-ISC/RISC mechanism but also more general triplet PSs encompassing multiple ISC mechanisms, such as SOCT-ISC, RP-ISC and SOC. The E_{abs} value above 0.38 eV signifies an absorption wavelength greater than 500 nm, guiding the model toward molecules with extended conjugated systems for enhanced applicability in PDT. $QED \ge 0.38$ was chosen based on the average QED value of the reported triplet PS distribution, and this value of PDT drugs significantly differs from the typical QED of 0.6 for conventional drugs, highlighting the distinct chemical space of triplet PSs. The SA value below 4.0 generally indicates that a molecule is synthetically accessible. Molecules that satisfy these constraints are defined as "desired molecules", while "successful molecules" are further distinguished by their uniqueness. During the RL process, the batch size for sampling was set to 64. These

molecules were labeled with their four scores as the rewards, and the desired molecules were saved during sampling.

2.6 Baselines and training setting

Frag-MB/Frag-GB and MB/GB are regarded as four baselines for comparison with the studied generative models. All workflows and datasets remain consistent across models, except that the prediction models in the baseline models were trained on data labeled with ΔE_{S1T1} ³⁴ which favors the generation of TADF molecules. For each model, the fine-tuning process of RL was conducted for 5000 iterations. To mitigate experimental variability, each RL fine-tuning procedure was repeated three times. This study employed two distinct sets of evaluation metrics as benchmarks to rigorously assess the performance of the generative models. The first standard metrics were outlined by Jin et al.,55 while the second metrics were adapted from the MOSES benchmarking platform,⁵⁶ a widely recognized standard in *de novo* molecular generation. Besides, to evaluate the fragments of generated molecules more effectively, we used conjugate motif diversity, which reflects the hierarchical organization of molecular motifs and better guides models toward generating triplet PSs.

3 Results and discussion

3.1 Generation efficiency

To assess the generation efficiency of each model, we analyzed the relationship between the number of steps and the cumulative number of unique desired molecules (referred to as successful molecules for simplicity). As shown in Tasks 1 and 2 of Fig. 2, the number of successful molecules increased with the generation steps, indicating that the model had effectively learned the association between conditional inputs and generated molecules. However, the generation efficiency differed across models. In Task 1 (left panel of Fig. 2), MD/MB exhibited a faster generation rate of successful molecules compared to GD/GB. Similarly, in Task 2 (right panel of Fig. 2), Frag-MD/ Frag-MB demonstrated a faster generation rate than Frag-GD/ Frag-GB. This suggests that, the RNN, derived from knowledge distillation of the conditional transformer decoder, enhanced the generation efficiency of successful molecules. More specifically, in Task 1 (left panel of Fig. 2), MD significantly outperformed MB, whereas GD showed a marginal improvement over GB. This finding demonstrated that our scoring function, as a more accurate prediction tool, also contributed to enhancement of generation efficiency in the knowledge distillation-based RNN model. A similar result was observed in Task 2, as shown in the right panel of Fig. 2. When comparing the number of successful molecules generated by MD (the top model in Task 1) with those generated by Frag-MD (the top model in Task 2), we found that MD produced substantially more successful molecules. Additionally, the average number of tokens for MD (88.9) was significantly higher than that for Frag-MD (7.6), indicating a more diverse range of modification for MD. In conclusion, the efficient generation and diverse modification for the character-based model significantly facilitate the exploration of novel molecules.

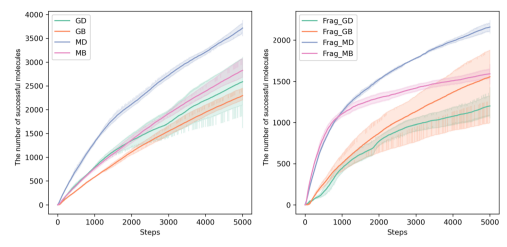


Fig. 2 Relationship between the number of successfully generated molecules and the steps in RL of Task 1 (left) and Task 2 (right).

3.2 Diversity and size of conjugated motifs

To further assess the diversity of conjugated motif level (rather than molecular level) generation by the MD model in Task 1, we collected the conjugated motifs in the generated molecules. This involved splitting the generated motifs, selecting unique instances, and removing duplicates. Encouragingly, the MD model exhibited 18 836 unique conjugated motifs, nearly twice the number (9754) observed in Frag-MD, further indicating that MD is well-suited for *de novo* triplet PS design in Task 1 by exploring novel conjugated motifs rather than relying solely on molecule modification or derivation. To examine the size of conjugated motifs, the distributions of ring numbers and atoms numbers of conjugated motifs for the successfully generated molecules were analyzed (Fig. 3). Frag-MD exhibited slightly higher average ring numbers (1.53 *vs.* 1.28) and atom counts (13.42 *vs.* 11.16), suggesting that the fragment-based generation method is more advantageous for creating larger motifs, which may exhibit longer wavelength absorption and demonstrate better applicability in PDT. The results for other models are provided in ESI Table S2.[†]

3.3 Benchmark performance

As part of this analysis, we collected the total number of molecules generated in each cycle to evaluate the performance of all models against the benchmarks. The average values are summarized in ESI Table S2.† Regarding conditional metrics, the MD model performed the best in terms of Novelty, whereas the Frag-MD model ranked the last. Additionally, all character-based methods exhibited superior performance compared to their fragment-based counterparts across the Novelty and Div metrics.

This observation further demonstrated that character-based molecule generation models are more likely to yield diverse

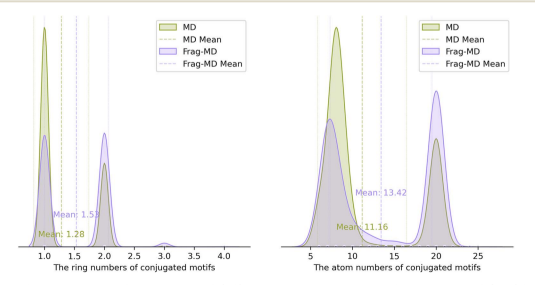


Fig. 3 Distribution of the ring numbers of conjugated motifs (left) as well as the atom numbers of conjugated motifs (right) for the successful molecules.

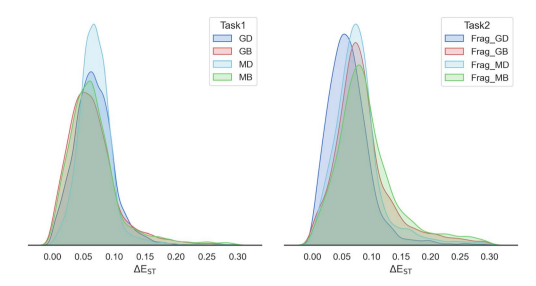


Fig. 4 Distribution of ΔE_{ST} for the successful molecules generated by all models for Task 1 based on the character-based method (left) and Task 2 based on the fragment-based model (right).

derivatives, consistent with the findings presented in Fig. 2. However, the MG model exhibited the highest diversity, slightly surpassing MD. This result can be attributed to the constraints imposed by the chemical space limitations of the transformer decoder.⁴⁴ With respect to the MOSES metrics, Frag-MD performed the best in terms of Unique and Frag, despite the overall poor performance of all models. Hence, Frag-MD is proficient in fragment combination and fragment derivation for identifying desired triplet PSs. The MOSES metrics, including SNN and IntDiv, showed similar trends to Novelty and Div observed in the conditional metrics. Although the metrics for the four studied models were compared with their baseline counterparts in both Task 1 and Task2, not all models showed better performance. For example, Frag-MD performed notably worse than its baseline, Frag-MB, in Novelty metrics, and MD exhibited substantially lower diversity compared to its baseline, MB. All studied models consistently showed improved diversity and larger conjugated motifs, as detailed in ESI Table S2.[†]

We visualized the property distributions of the total successfully generated molecules across all models in both Task 1 and Task 2. As depicted in Fig. 4 and 5 for Task 1 and Task 2, the distributions of $\Delta E_{\rm ST}$ and $E_{\rm abs}$ varied among four models, and a uniform distribution was observed across all models. Notably, the left-skewed distribution curves of $E_{\rm abs}$, generated by the MD/GD and Frag-MD/Frag-GD models, when compared to their respective baselines, exhibited their capability to generate larger conjugated molecules (Fig. 5). The underlying logic is that a lower $E_{\rm abs}$ energy level corresponds to longer absorption wavelengths, making these molecules better candidates for triplet PSs. However, the desired left-skewed

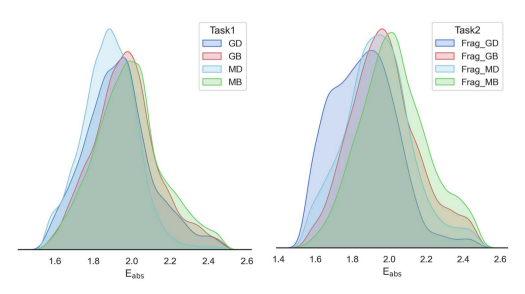


Fig. 5 Distribution of E_{abs} for the successful molecules generated by all models for Task 1 (left) and Task 2 (right).

distribution curves of $\Delta E_{\rm ST}$ are not observed in Fig. 4, indicating a limitation in the models' ability to accurately reproduce the $\Delta E_{\rm ST}$ property. To explore potential reasons behind this discrepancy, we performed DFT calculations to obtain groundtruth data.

3.4 Theoretical validation by DFT calculation

We first collected two datasets of the successful molecules generated by all studied models (dataset 1) and their baseline models (dataset 2), respectively. One key difference between these datasets lies in the ground truth used in the prediction model: $\Delta E_{S_{s,T_s}}$ for the studied models and $\Delta E_{S_{s,T_s}}$ for the baseline models. Initially, each generated dataset was labeled exclusively with its own prediction function, implying that each molecule just has one prediction value. Subsequently, we incorporated the counterpart's prediction model to add a secondary prediction value. Ultimately, two sub-datasets (sub-dataset 1 and subdataset 2) were formed by selecting molecules with prediction values for both $\Delta E_{S_{a}T_{a}}$ and ΔE_{S1T1} below 0.3 eV, thereby classifying these molecules as successful molecules in both the baseline and studied models. To further validate these molecules, the ground truth of the two sub-datasets was obtained by DFT calculation. Encouragingly, the ground truth distribution of $\Delta E_{S_nT_n}$ predominantly fell below 0.3 eV, whereas the ground truth distribution of ΔE_{S1T1} largely exceeded 0.3 eV, with some values even surpassing 1.0 eV (Fig. 6). This observation highlights the superior prediction accuracy of the investigated models compared to the baselines. According to the ground truth distribution of ΔE_{ST} in sub-dataset 1 (left panel in Fig. 6), we observed that the baseline models (ΔE_{S1T1} as ground truth) exhibited a low accuracy rate of 4%, implying a high error rate of 96% in generating successful molecules. Conversely, the studied models ($\Delta E_{S_nT_n}$ as groundtruth) exhibited a high accuracy of 73%, resulting in a low error rate of 27% in generating successful molecules. More interestingly, the ground truth distribution of ΔE_{ST} below 0.3 eV exhibited a Gaussian-like

pattern rather than being concentrated around zero, which is conducive to multiple ISC mechanisms. Similar results were observed in the distribution of ΔE_{ST} in sub-dataset 2 (right panel in Fig. 6).

3.5 Ablation studies

Boron dipyrromethene (BODIPY) compounds have been recognized as one of the most promising types of PSs for PDT, according to their excellent photophysical properties, including high extinction coefficients of visible light, superior photostability, and robust chemical stability, as well as practical and efficient derivatization capabilities. Novel BODIPY derivatives functioning as triplet PSs are continuously discovered based on expert knowledge.⁵⁷ To demonstrate the effectiveness of our DLbased methods to discover heavy-atom-free triplet PSs, we conducted ablation studies by systematically excluding the SMILES representations of BODIPY derivatives from the training and validation datasets, evaluated by Tanimoto similarity (Tanimoto coefficient > 0.4).⁵⁸

To generate BODIPY-constrained molecules, we employed a scaffold-constrained method⁵⁹ for both the *de novo* design scenario and the fragment-based molecular generation scenario. In the de novo design scenario, Fig. 7a demonstrates that our character-based method is capable of not only generating structure-similar published compounds but also discovering novel, unpublished compounds. Specifically, M2, a structure-similar published compound, is a fluorescein probe,⁶⁰ while M1, a derivative molecule of M2, probably exhibits strong absorption in the visible region and shows different triplet properties, which indicates M1 is a potential triplet PS. On the other hand, M3 and M4 can represent de novo compounds, with the chrysene motif in M3 and the phenanthrene motif in M4 that never exist in the training and validation datasets, which are different from the following unpublished molecules by the fragment-based method. Thus, the character-based method demonstrates robust capability to

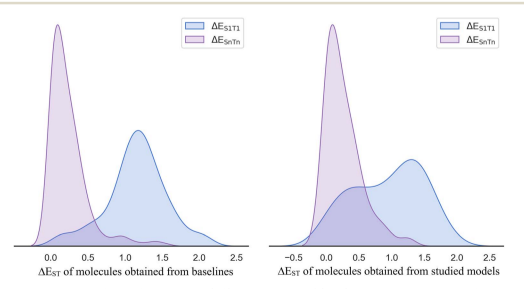


Fig. 6 The ground truth distributions of ΔE_{ST} of sub-dataset 1 (left) and sub-dataset 2 (right); $\Delta E_{S_nT_n}$ is the ground truth for the studied models and ΔE_{S1T1} is the ground truth for the baseline models.

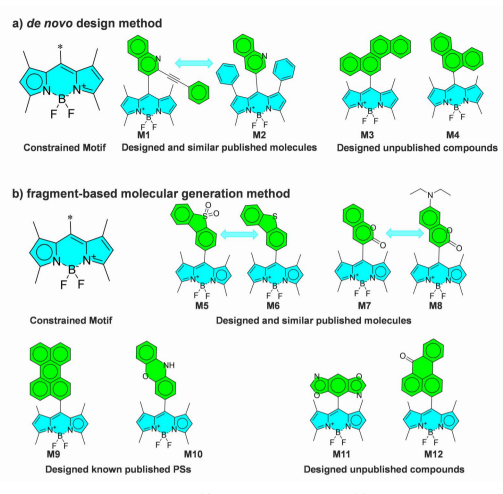


Fig. 7 Sampled generated molecules by different methods. (a) *De novo* design method and (b) fragment-based molecule generation method. Note: M1, M5, and M7 represent designed compounds, while M2, M6, and M8 correspond to structure-similar published compounds.

design novel triplet PSs, showing its potential for advancing PDT. In the fragment-based molecular generation scenario, Fig. 7b illustrates that, in addition to structure-similar compounds (M5 as a derivatization of M6 (ref. 61) and M7 as a de-derivatization of M8,62 represented as potential triplet PSs) and unpublished compounds (M11 and M12, represented as a novel fragment combination), known published molecules (M9 (ref. 63) and M10 (ref. 64)) were successfully designed. The ISC process of M10 has been meticulously investigated, and its underlying mechanism is elucidated as the SOCT-ISC mechanism. Similarly, the high triplet quantum yield ($\Phi_{\rm T} = 0.60$) of M9 also comes from the SOCT-ISC mechanism. Furthermore, M9 shows red-shifted spin-allowed charge transfer absorption. These two published molecules highlight the ability of our fragment-based method in fragment combination and fragment derivation in searching for desired triplet PSs. Similarly, other case studies about naphthalimide (NI) derivatives and perylenemonoimide (PMI) derivatives are presented in ESI Fig. S8 and S9.† Our fragment-based method successfully reidentified four well-known published molecules, including two TADF and two non-TADF molecules for NI derivatives. Moreover, published perylene (Pery)-phenothiazine (PTZ) triplet

photosensitizers (PSs) were obtained through both Pery-constrained and PTZ-constrained case studies, as detailed in ESI Fig. S10 and S11.[†]

4 Conclusion

In this study, we have successfully developed DL-based methods for de novo design and motif optimization of heavy-atom-free triplet photosensitizers (PSs), which incorporate multiple intersystem crossing (ISC) mechanisms. In particular, we developed a comprehensive dataset ($\sim 1.90 \times 10^9$) of triplet PSs for the first time, which included a variety of molecular types, favoring multiple ISC mechanisms. Subsequently, we trained a prediction model with the high excited singlet-triplet gap $(\Delta E_{S_nT_n} \leq 0.3 \text{ eV} \text{ as the ground truth, } n \geq 1)$ for the scoring function. To search de novo conjugate motifs (Task 1) as well as perform a conjugated-motif-based modification (Task 2) for desired triplet PSs, a reinforcement learning (RL) network was employed. The silicon experimental results revealed that the Frag-MD model outperformed the MD model in generating larger conjugated motifs, with higher average ring numbers (1.53 vs. 1.28) and atom counts (13.42 vs. 11.16), which illustrated the advantages of PS modification in Task 2 by creating larger motifs with longer wavelength absorption. In contrast, the MD model generated twice as many unique conjugated motifs and performed better in terms of the conditional and MOSES metrics for novelty and diversity, which showed the *de novo* design ability of the character-based method in Task 1. Impressively, theoretical calculations indicated that our proposed models (Frag-MD/MD and Frag-GD/GD) outperformed their baselines (Frag-MB/MB and Frag-GB/GB) in which the label data of prediction model based on ΔE_{S1T1} has a significantly higher prediction accuracy (73% vs. 4%). Highefficiency triplet PSs reported in recent studies were re-identified through ablation experiments. Thus, our method proves to be highly practical and effective, holding the promise of establishing a new paradigm for the discovery of novel PSs applicable in PDT.

Data availability

The code and datasets used in the study are publicly available from the GitHub repository: https://github.com/Kepeng2019/ Fragment_and_char_based_PSs_generation. The detailed procedures for constructing the dataset of photosensitizers, prediction models for score function in reinforcement learning, conjugated motif diversity for all models, distribution of QED and SA properties for the unique desired molecules generated by all models for Task 1 and Task 2, as well as DFT/TDDFT calculation results of selected compounds.

Author contributions

Yu Kang and Tingjun Hou organized and supervised the research and guided the manuscript writing and experimental design. Kepeng Chen and Jike Wang conceived the overall framework of the character-based and fragment-based models for triplet photosensitizer design. Kepeng Chen constructed the DFT-labeled dataset and performed model validation. Xiaoting Zhang and Wenbo Yang evaluated the generated molecules and proposed practical optimization strategies. Kepeng Chen drafted the manuscript, with critical revisions by Dan Li, Tingjun Hou, Wenbo Yang, and Yu Kang. Yu Kang and Tingjun Hou contributed their expertise in computational chemistry to improve the paper in terms of triplet photosensitizer design. Jike Wang provided specialized guidance on neural network adaptation for this task.

Conflicts of interest

There are no conflicts to declare.

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