LEARNING ENERGY DECOMPOSITIONS FOR PARTIAL INFERENCE OF GFLOWNETS

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Abstract

This paper studies generative flow networks (GFlowNets) to sample objects from the Boltzmann energy distribution via a sequence of actions. In particular, we focus on improving GFlowNet with partial inference: training flow functions with the evaluation of the intermediate states or transitions. To this end, the recently developed forward-looking GFlowNet reparameterizes the flow functions based on evaluating the energy of intermediate states. However, such an evaluation of intermediate energies may (i) be too expensive or impossible to evaluate and (ii) even provide misleading training signals under large energy fluctuations along the sequence of actions. To resolve this issue, we propose learning energy decompositions for GFlowNets (LED-GFN). Our main idea is to (i) decompose the energy of an object into learnable potential functions defined on state transitions and (ii) reparameterize the flow functions using the potential functions. In particular, to produce informative local credits, we propose to regularize the potential to change smoothly over the sequence of actions. It is also noteworthy that training GFlowNet with our learned potential can preserve the optimal policy. We empirically verify the superiority of LED-GFN in five problems including the generation of unstructured and maximum independent sets, molecular graphs, and RNA sequences.

1 INTRODUCTION

Generative Flow Networks (Bengio et al., 2021a, GFlowNets or GFNs) are frameworks to sample objects through a sequence of actions, e.g., iteratively adding nodes to a graph. Their key concept is training a policy that sequentially selects the actions to sample the object from the Boltzmann distribution (Boltzmann, 1868). Such concepts enable discovering diverse samples with low energies, i.e., high scores, as an alternative to reinforcement learning (RL)-based methods which tend to maximize the return of the sampled object (Silver et al., 2016; Sutton & Barto, 2018).

To sample from the Boltzmann distribution, GFlowNet trains the policy to assign action selection probability based on energy of terminal state (Bengio et al., 2021a;b; Malkin et al., 2022a), e.g., a high probability to the action responsible for the low terminal energy. However, such training has fundamental limitations in credit assignment, as it is hard to identify the action responsible for terminal energy (Pan et al., 2023a). This limitation stems from solely relying on the terminal energy associated with multiple actions, lacking the information to identify the contribution of individual actions, akin to challenges in RL with sparse reward (Arjona-Medina et al., 2019; Ren et al., 2022).

An attractive paradigm to tackle this issue is *partial inference* (Pan et al., 2023a) that trains flow functions with local credits, e.g., evaluation of the intermediate states or transitions. Such local credit identifies individual action contributions to the terminal energy before reaching the terminal state. To this end, Pan et al. (2023a) proposed a forward-looking GFlowNet (FL-GFN), which assigns the local credit based on the energy of incomplete objects associated with intermediate states.

However, FL-GFN crucially relies on two assumptions that may not hold in practice. First, FL-GFN requires evaluating the energy of the intermediate state in the trajectories. However, the energy function can be expensive or even impossible to evaluate. Next, FL-GFN assumes the energy of intermediate states to provide useful hints for the terminal energy. However, this may not be true when the intermediate energy largely fluctuates along the sequence of states, e.g., low intermediate energies may lead to a terminal state with high energy. We illustrate such a pitfall in Figure 1.



Figure 1: **The local credit evaluation in bag generation (Example 1)**. (**first row**) The task is to generate a bag of entities, where the seven same entities yields a high score. (**second row**) Left-to-right indicates the state transitions over a given trajectory. (**third row**) The energy-based evaluation fails to produce informative local credits since every intermediate state has zero energy, whereas our potential function produces informative credits by enforcing the potentials to be uniformly distributed.

Contribution. We propose learning energy decomposition for GFlowNet, coined LED-GFN. Our key idea is to perform partial inference by decomposing terminal state energy into a sum of learnable potentials associated with state transitions and use them as local credits. In particular, we show how to regularize the potential function to preserve the ground-truth terminal energy and minimize variance over the trajectory to yield informative potentials. Figure 1 highlights how LED-GFN provides informative local credits compared to the existing approach.

To be specific, our energy decomposition framework resembles the least square-based return decomposition for episodic reinforcement learning (Efroni et al., 2021; Ren et al., 2022). We parameterize potential function with a regression model that is constrained to be equal to the terminal energy when aggregated over the entire action sequence. We also regularize the potential function to minimize the variance along the trajectory, so that the potential function provides dense local credits in training GFlowNet. Such potentials associated with intermediate transitions provide informative signals, as each of them is enforced to be correlated with the terminal energies. The training of potential function is online, which uses samples collected during GFlowNet training.

We extensively validate LED-GFN on various tasks: set generation (Pan et al., 2023a), bag generation (Shen et al., 2023), molecular discovery (Bengio et al., 2021b), RNA sequence generation (Jain et al., 2022), and the maximum independent set problem (Zhang et al., 2023). We observe that LED-GFN (1) outperforms FL-GFN when the assumption of intermediate energy does not hold, (2) excels in practical domains compared to GFlowNets and RL-based baselines, and (3) achieves similar performance to FL-GFN even when intermediate energy provides the "ideal" local credit.

2 PRELIMINARIES

In this section, we describe generative flow networks (Bengio et al., 2021a, GFlowNets or GFNs) and their partial inference algorithm. We describe additional related works in Appendix A.

2.1 GFLOWNETS

GFlowNets sample from discrete space \mathcal{X} through a sequence of actions from the action space \mathcal{A} that make transitions in the state space \mathcal{S} . For each complete trajectory $\tau = (s_0, s_1, \ldots, s_T)$, the terminal state is the object $x = s_T \in \mathcal{X}$ to be generated. The state transitions are determined by the action sequence (a_1, \ldots, a_{T-1}) , e.g., a_t determines $s_t \to s_{t+1}$. The policy $P_F(s'|s)$ selects the action a to transition from the current state s to the next state s' and induces a distribution over the object x.

The main objective of GFlowNet is to train the policy $P_F(\cdot|\cdot)$ that samples objects from the Boltzmann distribution with respect to a given energy function $\mathcal{E} : \mathcal{X} \to \mathbb{R}$ as follows:

$$P_F^{\top}(x) \propto \exp(-\mathcal{E}(x)),\tag{1}$$

where $P_F^{\top}(x)$ is the distribution of sampling an object x induced from marginalizing over the trajectories conditioned on $x = s_T$. We omit the temperature for simplicity. To this end, GFlowNet trains with auxiliary objectives based on state transition, trajectory, or sub-trajectory information.

Detailed balance (Bengio et al., 2021b, DB). The DB utilizes the experience of state transitions to train GFlowNet. It trains the GFlowNet with a forward policy model $P_F(s'|s)$, a backward policy $P_B(s|s')$, and a state flow estimator $F(\cdot) : S \to \mathbb{R}^+$ by minimizing the following loss function:

$$\mathcal{L}_{\text{DB}}(s, s') = (\log F(s) + \log P_F(s'|s) - \log F(s') - P_B(s|s'))^2$$

where the flow F(s) of the terminal state $s_T = x$ is defined to be identical to the energy $\exp(-\mathcal{E}(x))$.

Trajectory balance (Malkin et al., 2022a, TB). The TB aims to learn the policy faster by training on full trajectories. To this end, TB requires a forward policy model $P_F(s'|s)$, a backward policy $P_B(s|s')$, and a learnable scalar Z to minimize the following loss function:

$$\mathcal{L}_{\text{TB}} = \left(\log Z + \sum_{t=0}^{T-1} \log P_F(s_{t+1}|s_t) - \mathcal{E}(x) - \sum_{t=0}^{T-1} \log P_B(s_t|s_{t+1}) \right)^2.$$

This objective is resilient to the bias from inaccurate flow estimator $F(\cdot)$ used in DB, since it directly propagates the terminal energy to train on intermediate states. However, the TB suffers from the high variance of the objective over the collected trajectories (Malkin et al., 2022a).

Sub-trajectory balance (Madan et al., 2023, subTB). The subTB trains forward and backward policies $P_F(s'|s)$, $P_B(s'|s)$ and a learnable scalar Z similar to the TB. However, it trains on flexible length of sub-trajectory $s_U \rightarrow s_{U+1} \cdots \rightarrow s_{U+L}$ to minimize the following loss function:

$$\mathcal{L}_{\text{subTB}} = \left(\log F(s_U) + \sum_{t=U}^{U+L-1} \log P_F(s_{t+1}|s_t) - \log F(s_{U+L}) - \sum_{t=U}^{U+L-1} \log P_B(s_t|s_{t+1})\right)^2.$$

This objective enables controlling the bias-variance trade-off by interpolating between the DB that trains on a single state transition and the TB that trains on a complete trajectory.

2.2 PARTIAL INFERENCE FOR GFLOWNETS

The GFlowNet training is often challenged by limitations in credit assignment, i.e., identification and promotion of the action responsible for the observed low energy. This limitation stems from relying solely on the terminal state energy as the training signal. The terminal energy lacks information to identify the contribution of individual action, akin to how reinforcement learning with sparse reward suffers from credit assignment (Arjona-Medina et al., 2019; Ren et al., 2022).

Partial inference is a promising paradigm to resolve this issue by incorporating local credits (Pan et al., 2023a). Specifically, the partial inference aims to evaluate individual transitions or sub-trajectories, i.e., local credits, and provide informative training signals for identifying the specific contributions of actions. To this end, Pan et al. (2023a) proposed Forward-Looking GFlowNet (FL-GFN), which evaluates intermediate state energy as a local credit signal for partial inference.

Forward-Looking GFlowNet (Pan et al., 2023a, FL-GFN). To enable partial inference, the FL-GFN defines a new training objective that incorporates an energy function $\mathcal{E} : \mathcal{X} \to \mathbb{R}$ for intermediate states. To this specific, FL-GFN modifies the DB as follows:

$$\mathcal{L}_{\rm FL}(s,s') = (\log \tilde{F}(s) + \log P_F(s'|s) - \mathcal{E}(s) + \mathcal{E}(s') - \log \tilde{F}(s') - \log P_B(s|s'))^2, \quad (2)$$

where $\tilde{F}(s) = F(s) \exp(\mathcal{E}(s))$ is the re-parameterized flow function and $\mathcal{E}(s') - \mathcal{E}(s)$ is the energy gain associated with the transition from s to s'. Note that the energy function is defined only on the discrete space \mathcal{X} , hence FL-GFN evaluates an intermediate energy $\mathcal{E}(s)$ as the energy of an incomplete object associated with the intermediate state s. Pan et al. (2023a) shows that optimum of Equation (2) induces a policy $P_F(\cdot|\cdot)$ that samples from Boltzmann distribution. While Equation (2) is associated with DB, FL-GFN is also applicable to subTB with energy gains in the level of sub-trajectory.

3 LEARNING ENERGY DECOMPOSITION FOR GFLOWNET

While the FL-GFN is equipped with partial inference capabilities, it relies on the energy function to assign local credits, which can be expensive to evaluate, or lead to sub-optimal training signals (details are described in Section 3.1). In this paper, we propose learning energy decomposition for GFlowNets (LED-GFN) to achieve better partial inference. In what follows, we describe our motivation for better partial inference (Section 3.1) and the newly proposed LED-GFN (Section 3.2).



Figure 2: **Illustration of energy decomposition for partial inference of GFlowNet.** LED-GFN enables partial inference with potentials which (a) approximate the object energy via summation, and (b) minimize variance along the action sequence.

3.1 MOTIVATION FOR BETTER PARTIAL INFERENCE

Our motivation stems from the limitations of FL-GFN, which performs partial inference based on evaluating the energies of intermediate states with respect to a single transition (for DB) or sub-trajectory (for subTB). In particular, we are inspired by how the energy gain may yield a sub-optimal local credit signal due to the following pitfalls (see Figures 1 and 9(b)).

- **A.** The energy evaluation for an incomplete object, i.e., intermediate state, can be non-trivial. In addition, the cost of energy evaluation can be expensive, which can bottleneck efficient training when called for all visited states.
- **B.** The energy can exhibit sparse or high variance on intermediate states within a trajectory, even returning zero for most states, which is non-informative for partial inference.



Figure 3: Negative relative mean error (\downarrow) for estimating the true Boltzmann distribution on Example 1-type task (Bag).

We further provide a concrete example for the pitfall **B**. of FL-GFN, which is illustrated in Figure 1 and Figure 3^1 for both conceptual and empirical purposes, respectively:

Example 1. Consider adding objects from $\{A, B, C, D, E\}$ to a bag with a maximum capacity of nine. Define the energy as -1 when the bag contains seven identical objects and 0 otherwise.

For Example 1, the intermediate energy (which is always 0) does not provide information for the terminal energy. However, the number of the most frequent elements in the bag is informative even at intermediate states since greedily increasing the number leads to the best terminal state.

Our observation in Example 1 hints at the existence of a partial inference algorithm to provide better local credit signals. We aim to pursue this direction with a learning-based approach. That is, we parameterize the class of potential functions that decompose the terminal energy to provide local credit signals for GFlowNet training. Our key research direction is to understand about what conditions of the potential functions are informative for partial inference.

3.2 Algorithm description

In this section, we describe our framework, coined learning energy decomposition for GFlowNet (LED-GFN), which facilitates partial inference using learned local credit. To this end, we propose to decompose the terminal energy into learnable potentials defined on state transitions. Similar to FL-GFN, we reparameterize the flow model with local credits, i.e., potentials. In contrast to FL-GFN, we optimize the local credits to enhance partial inference by minimizing variance of potentials along the action sequence. See Figure 2 for an illustration of LED-GFN.

Training with potential functions. To be specific, we decompose the energy function \mathcal{E} associated with the terminal state into learnable potential functions ϕ_{θ} as follows:

$$\mathcal{E}(x) \approx \Phi_{\theta}(\tau) = \sum_{t=0}^{T-1} \phi_{\theta}(s_t \to s_{t+1}), \tag{3}$$

where $\tau = (s_0, s_1, \dots, s_T)$, $x = s_T$, and the potential functions are defined on state transition $s_t \rightarrow s_{t+1}$. Similar to FL-GFN, we use the potential function to train the forward and backward

¹The detailed experiment settings are described in Section 4.1.

Algorithm 1 Learning energy decomposition for GFlowNet

- 1: Initialize the buffer \mathcal{B} , forward and backward policy P_F, P_B , state flow F, and model ϕ_{θ} .
- 2: Update the model ϕ_{θ} to minimize ℓ_{ED} if the generation trajectories are given in advance.
- 3: repeat
- 4: Sample a batch of trajectories $\{\tau_b\}_{b=1}^{B_1}$ from forward policy P_F .
- 5: Update buffer $\mathcal{B} \leftarrow \mathcal{B} \cup \{\tau_b\}_{b=1}^{B_1}$.
- 6: **for** n = 1, ..., N **do**
- 7: Sample a batch of trajectories $\{\tau_b\}_{b=1}^{B_2}$ from the buffer \mathcal{B} .
 - Update the model ϕ_{θ} to minimize ℓ_{LS} with $\{\tau_b\}_{b=1}^{B_2}$.
- 9: end for

8:

10: Compute intermediate energy gains $\phi_{\theta}(s_i, a_i)$ for all $(s_i, a_i) \in \tau$.

11: Update the GFlowNet P_F, P_B, \tilde{F} to minimize \mathcal{L}_{LED} with τ and $\phi_{\theta}(s_i, a_i)$ for all $(s_i, a_i) \in \tau$. 12: **until** converged

▷ Energy decomposition learning

GFlowNet policies P_F , P_B and flow model F to minimize the following loss:

$$\mathcal{L}_{\text{LED}}(s,s') = (\log F(s) + \log P_F(s'|s) + \phi_{\theta}(s \to s') - \log F(s') - \log P_B(s|s'))^2.$$
(4)

Given a sub-trajectory $(s_0, \ldots, s_u = s, s_{u+1} = s')$, one can derive this objective from Equation (2) by replacing $\mathcal{E}(s), \mathcal{E}(s')$ with $\sum_{t=0}^{u} \phi(s_t)$ and $\sum_{t=0}^{u+1} \phi(s_t)$, respectively. That is, it is evident that Equation (4) preserves the optimal policy of GFlowNet when $\mathcal{E}(x) = \Phi_{\theta}(\tau)$ is satisfied for all trajectories τ terminating with x.

Our objective becomes equivalent to that of FL-GFN when $\phi_{\theta}(s \rightarrow s') = \mathcal{E}(s') - \mathcal{E}(s)$, but our key idea is to *learn* the potential function ϕ_{θ} instead of the energy gain which can be expensive and may exhibit sparsity or high variance, as pointed out in Section 3.1. Note that one can also introduce an approximation error $\mathcal{E}(x) - \Phi_{\theta}(\tau)$ as an additional correction term to preserve the optimal policy of GFlowNet even when the potential function ϕ_{θ} is inaccurate. In Appendix B.1, we describe how LED-GFN consistently induces the optimal policy that samples from the Boltzmann distribution.

Training potentials with squared loss. In training the potential function, our key motivation is twofold: (a) accurately estimating the true energy through summation and (b) providing dense and informative training signals by minimizing variance along the action sequence.

To this end, given a trajectory $\tau = (s_0, \dots, s_T = x)$, we train the potential functions ϕ_{θ} to minimize the loss function for (a) achieving $\mathcal{E}(x) \approx \Phi_{\theta}(\tau)$ with (b) dropout-based regularization:

$$\ell_{\rm LS}(\tau) = \mathbb{E}_{\boldsymbol{z} \sim {\rm Bern}(\lambda)} \left[\left(\frac{1}{T} \mathcal{E}(s_T) - \frac{1}{C} \sum_{t=0}^{T-1} z_t \phi_{\theta}(s_t \to s_{t+1}) \right)^2 \right].$$
(5)

The *T*-length random vectors z promotes the dropout, where $z_t = 0$ sampled from the Bernoulli distribution with probability $1 - \lambda$. Dividing by *T* and $C = \sum_{t=0}^{T-1} z_t$ aligns the scales to compensate for the scale reduction induced by dropout. When $\lambda = 1$ and the loss function is minimized, i.e., $\ell_{LS}(\tau) = 0$ for all τ , the potential function decomposes the energy function without error. When $\lambda < 1$, dropout prevents heavy reliance on specific potentials to satisfy Equation (3), thereby reducing the variance and sparsity of the potentials along the action sequence.² Note that our intuition is similar to recent works on learning return decomposition to alleviate sparse reward problems in reinforcement learning (Arjona-Medina et al., 2019; Gangwani et al., 2020; Ren et al., 2022).

To train the potential function, we define its training as online learning within GFlowNet training, i.e., learning from trajectories obtained during GFlowNet training. We describe the overall algorithm in Algorithm 1. Such an alternative training of the potential function and the policy is similar to model-based reinforcement learning algorithms (Luo et al., 2018; Sun et al., 2018; Janner et al., 2019) for monotonic improvement of policies. Note that LED-GFN can also be implemented with subTB, where the details are described in Appendix B.2.

²Ren et al. (2022) provide a formal proof that Equation (5) serves as a surrogate objective to satisfy Equation (3) while reducing the variance and sparsity of the potentials along the action sequence.



Figure 4: **The performance on bag generation.** The solid line and shaded region represent the mean and standard deviation, respectively. The LED-GFN shows superiority compared to the considered baselines on both DB-based and subTB-based implementations.



Figure 5: **The performance on molecule generation.** The solid line and shaded region represent the mean and standard deviation, respectively. The LED-GFN shows superiority compared to the considered baselines in generating diverse high reward molecules.

4 EXPERIMENT

We extensively evaluate LED-GFN on various domains, including bag generation (Shen et al., 2023), molecule generation (Bengio et al., 2021a), RNA sequence generation (Jain et al., 2022), set generation (Pan et al., 2023a), and the maximum independent set problem (Zhang et al., 2023). Following prior studies, we consider the number of modes, i.e., samples with energy lower than a specific threshold, and the average top-100 score as the base metrics, which are measured via samples collected during training. We report all the performance using three different random seeds.

4.1 BAG GENERATION

First, we consider a bag generation task (Shen et al., 2023). The action is adding an object from seven distinct entities to a bag with a maximum capacity of 15. The bag exhibits low energy when including seven or more repeated entities of the same type. In this task, we compare our method with GFN and FL-GFN. We consider both DB-based (Bengio et al., 2021b) and subTB-based (Malkin et al., 2022a) implementations. The detailed experimental settings are described in Appendix C.

Results. We present the results in Figure 4. Here, one can observe that our method excels in bag generation compared to GFN and FL-GFN on both DB and subTB. In particular, FL-GFN fails to make improvements on the subTB-based implementation, since most states do not provide informative signals for partial inference (as illustrated in Figure 1). In contrast, LED-GFN consistently improves performance by producing informative potentials to enhance partial inference.

4.2 MOLECULE GENERATION

Next, we validate LED-GFN in a more practical domain: the molecule generation task (Bengio et al., 2021a). This task aims to find molecules with low binding energy to the soluble epoxide hydrolase protein. In this task, a molecule is generated by constructing junction trees (Jin et al., 2018), with the



Figure 6: **The performance on RNA sequence generation.** The solid line and shaded region represent the mean and standard deviation, respectively. The LED-GFN shows superiority compared to the considered baselines in generating diverse high reward RNA sequences.

actions of adding molecular building blocks. The binding energy between the molecule and the target protein is computed using a pre-trained oracle (Bengio et al., 2021a).

In this experiment, we consider PPO (Schulman et al., 2017), and three GFN models: DB, TB, and subTB (Madan et al., 2023) as the baselines. Additionally, we compare our approach with GAFN (Pan et al., 2023b) and FL-GFN. For FL-GFN and LED-GFN, we consider a subTB-based implementation. The overall implementations and experimental settings follow prior studies (Bengio et al., 2021a; Pan et al., 2023a), which are described in Appendix C.

Results. The results are presented in Figure 5. One can see that LED-GFN outperforms the considered baselines in enhancing the average score of unique top-100 molecules and the number of modes found during training. These results highlight that LED-GFN is also beneficial for real-world generation problems with a large state space.

4.3 RNA SEQUENCE GENERATION

We also consider a RNA sequence generation task for discovering diverse and promising RNA sequences that bind to human transcription factors (Barrera et al., 2016; Trabucco et al., 2022; Jain et al., 2022). The action is appending or prepending an amino acid to the current sequence. The energy is pre-computed based on wet-lab measured DNA binding activity to Sine Oculis Homeobox Homolog 6 (Barrera et al., 2016). We consider the same baselines as in the molecule generation task.

Results. The results are presented in Figure 6. One can observe that LED-GFN outperforms the considered baselines. Furthermore, FL-GFN only makes minor differences compared to GFN, while LED-GFN makes noticeable improvements. These results highlight that energy-based partial inference can fail to improve performance in practical domains, while the potential learning-based approach consistently leads to improvements.

4.4 COMPARISON WITH IDEAL LOCAL CREDITS

In these experiments, we demonstrate that LED-GFN can achieve similar performance compared to FL-GFN, even when the intermediate state energy is sufficient to identify the contribution of the action, i.e., ideal local credit (Zhang et al., 2023). Note that this tasks are idealized, since designing such an energy function requires a complete understanding of the domain. Especially, we focus on two tasks: set generation (Pan et al., 2023a) and the maximum independent set problem (Zhang et al., 2023). For these tasks, we compare our method with GFN and FL-GFN.

Set generation. The set generation task is similar to the bag generation. The actions are adding an objects from 30 distinct objects to a set with a maximum capacity of 20. The energy is evaluated by accumulating the individual energy of each entity, so the intermediate energy gain has complete information to identify the contribution of each action (Pan et al., 2023a). We describe the detailed experiment settings in Appendix B.



Figure 7: **The performance comparison with ideal local credits.** The solid line and shaded region represent the mean and standard deviation, respectively. The LED-GFN shows similar performance to the FL-GFN, even when the intermediate energy is sufficient to identify the action contribution.



Figure 8: (a-b) The negative relative mean error comparison (lower is better). Our LED-GFN better approximates the target distributions. (c) The Tanimoto similarity with respect to the average scores (upper-right is better). Our LED-GFN produces more diverse and promising molecules.

Maximum independent set problem. This task aims to find the maximum independent set by selecting nodes, and the energy is evaluated based on the current size of the independent set (Zhang et al., 2023). Here, we compare the performance on validation graphs following Zhang et al. (2023). At each step, we sample 50 solutions for each validation graph to measure the average scores and the number of mode founds (greater than 18.5). The overall implementations and hyper-parameters follow prior studies (Zhang et al., 2023).

Results. As illustrated in Figure 7, one can see that our approach achieves similar performance to FL-GFN, even though the intermediate state energy provides ideal local credit for partial inference. These results highlight that the potentials of LED-GFN can be as informative as ideal local credit, which provides complete identification of the action contributions.

4.5 Ablation studies

Goodness-of-fit to the true Boltzmann distribution. We show that how well our algorithm make a good sampling distribution to sample from the target distribution, i.e., Boltzmann distribution. Following Shen et al. (2023), we measure the relative mean error between the empirical generative distribution and target distribution following. In Figures 8(a) and 8(b), one can observe that LED-GFN achieves a better approximation to target distribution compared to considered baselines.

Diversity vs. high score. Next, we further verify that our algorithm not only generates high-scoring samples but also diverse molecules. Specifically, we analyze the trade-off between the average score of the top-100 samples and the diversity these samples. To measure diversity, we compute the average pairwise Tanimoto similarity (Bajusz et al., 2015). In Figure 8(c), we illustrate the Tanimoto similarities with respect to the average of top-100 scores. Here, one can observe that LED-GFN achieves better diversity with respect to the average scores.



Figure 9: (a) The benefits of regularizing variance of potentials. The regularization improves the performance. (b) The performance over the number of energy evaluation. The FL-GFN can not make improvement with respect to the number of energy evaluation.



Figure 10: **Comparison between potential learning methods for partial inference.** The least-square based energy decomposition shows most promising results for partial inference.

Regularized vs. non-regularized potentials. We also analyze how reducing the variance of potentials benefits the improvement in performance. To this end, we compare energy decomposition learning with variance regularization and its counterpart without regularization, i.e., set $\lambda = 1$ in Equation (5). In Figure 9(a), one can see that applying regularization yields more promising performance. These results highlight that inducing dense potentials plays a significant role in improving performance.

Number of energy evaluation vs. performance We analyze the performance with respect to the number of energy evaluation, which can be expensive. In Figure 9(b), one can see that FL-GFN can not improve performance since it requires evaluating the energy for every visited states. In contrast, LED-GFN uses a potential function without energy evaluation for intermediate states.

Energy decomposition learning vs. alternative potentials learning. We further investigate the following alternative potential learning schemes.

- One may train a proxy model φ_θ : X → ℝ to predict the terminal energy, and utilize it to compute potentials φ_θ(s → s') = φ_θ(s') φ_θ(s). This approach can be interpreted as extension of model-based GFlowNet (Jain et al., 2022) for partial inference.
- Based on the LSTM-based decomposition method (Arjona-Medina et al., 2019), one can design the potential $\phi_{\theta}(s_t \rightarrow s_{t+1})$ as the difference between two subsequent predictions for $(s_0, a_0, \ldots, s_t, a_t)$ and $(s_0, a_0, \ldots, s_{t+1}, a_{t+1})$ using an LSTM.

In Figure 10, we compare each method in molecule and set generation tasks with a DB-based implementation. Here, one can see that the least square-based approach shows the most competitive performance due to its capabilities in producing dense and informative potentials.

5 CONCLUSION

In this paper, we propose learning energy decomposition for GFlowNets (LED-GFN). Experiments on various domains show that LED-GFN is promising compared to existing partial inference methods for GFlowNet. An interesting avenue for future work is developing new partial inference techniques using learnable local credit, other than the flow reparameterization considered in our work.

Reproducibility. We describe experimental details in Appendix C, which provides the base implementation references, environments, and detailed hyper-parameters. In the supplementary materials, we also include the codes for molecule generation tasks based on the official implementation codes of the prior study (Pan et al., 2023a).

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A RELATED WORKS

Generative augmented flow network (Pan et al., 2023b, GAFN). The GAFN is a learning framework that incorporates intermediate rewards for exploration purposes. Specifically, GAFN measures the novelty score of a given state, which provides an intrinsic signal to facilitate better exploration towards unvisited states. To compute the novelty score, this method also incorporates online training of random network distillation (Burda et al., 2019), which assigns lower scores to unseen states compared to more frequently observed states.

Model-based GFlowNet. Jain et al. (2022) propose model-based training of GFlowNet for discovering diverse and promising biological sequences. They train a proxy model of the energy function to mitigate the expensive cost of evaluating biological sequences, such as wet-lab evaluation. Additionally, they introduce an active learning algorithm for the model-based GFlowNet, leveraging the epistemic uncertainty estimation of the model to improve exploration.

Return decomposition learning. Our LED-GFN approach is inspired by return decomposition learning of reinforcement learning in sparse reward settings. Their goal is to decompose the return into step-wise dense reward signals (Arjona-Medina et al., 2019; Gangwani et al., 2020; Ren et al., 2022). They have studied various return decomposition methods. First, Arjona-Medina et al. (2019) utilize an LSTM-based model to produce step-wise proxy rewards. Next, Gangwani et al. (2020) propose a simple approach that uniformly redistributes the terminal reward over the trajectory. Ren et al. (2022) train a proxy reward function using randomized return decomposition learning which is contrained to produces dense and informative proxy rewards.

B DETAILS OF LED-GFN

B.1 PRESERVING OPTIMAL POLICY OF GFLOWNET

Although the potential function is inaccurate, we show that optimum of \mathcal{L}_{LED} can induce an optimal policy that samples from Boltzmann distribution. We give a simple proof by reduction to the optimum of DB objective (Bengio et al., 2021b). Suppose the parameterization $\phi_{\theta}(s \to s') = \phi_{\theta}(s') - \phi_{\theta}(s)$, and $\log \hat{F}(s) = -\phi_{\theta}(s) + \log \tilde{F}(s)$. Then, we can reformulate \mathcal{L}_{LED} as follows:

$$\mathcal{L}_{\text{LED}}(s, s') = \left(\log \hat{F}(s) + \log P_F(s'|s) - \log \hat{F}(s') - P_B(s|s')\right)^2.$$

If we add an additional correction term $-\mathcal{E}(x) + \Phi_{\theta}(\tau) = -\mathcal{E}(x) + \phi_{\theta}(x)$ to the terminal flow such that $\log \hat{F}(x) = -\mathcal{E}(x)$, this objective becomes to be equivalent to the reparameterization of DB, where the optimum induces a policy sampling from a Boltzmann distribution (Bengio et al., 2021b). Therefore, the optimum of LED-GFN can still induce the policy that samples from the Boltzmann distribution. We refer to this correction-based approach as LED-GFN*.

However, our implementation follows a prior study in return decomposition learning (Ren et al., 2022), which uniformly redistributes the decomposition error over the transitions within the given trajectory (we denote this approach as LED-GFN in experiments). In Figure 11, we empirically observe that this approach further improves the training of GFlowNets. We assume that uniformly redistributed decomposition error partially provides more dense and informative local credit signals correlated with future energy.



Figure 11: The performance on bag generation.

B.2 TRAINING ON SUBTB

The LED-GFN can also be implemented on subTB by incorporating the potentials within subtrajectories. To this specific, one can modify subTB as follows:

$$\mathcal{L}_{\text{LED-subTB}} =$$

$$\left(\log \tilde{F}(s_U) + \sum_{t=U}^{V-1} \log P_F(s_{t+1}|s_t) + \sum_{t=U}^{V-1} \phi_\theta(s_t \to s_{t+1}) - \log \tilde{F}(s_V) - \sum_{t=U}^{V-1} \log P_B(s_t|s_{t+1})\right)^2$$

which is based on a sub-trajectory $s_U \to s_{U+1} \cdots \to s_V$. This objective is equivalent to FL-GFN on the subTB when $\phi_{\theta}(s \to s') = \mathcal{E}(s') - \mathcal{E}(s)$ (Pan et al., 2023a), but we replace it with the potentials for better credit assignment.

C EXPERIMENTAL DETAILS

For all experiments, the neural network architecture of potential function is identical to that of GFlowNet policy. We set the learning rate for potential functions to 0.001. The dropout is applied to potentials in least square-based energy decomposition learning Equation (5). We set the dropout probability as 10% for tasks with a trajectory length less than 10 and 20% for others.

Bag generation (Shen et al., 2023). The experiment settings, implementations, and hyper-parameters are based on prior studies (Shen et al., 2023). The bag generation task is to generate a bag with a maximum capacity of 15. There are seven types of entities, where each action includes one of them in the current bag. If it contains seven or more repeats of any items, it has reward 10 with 75% chance, and 30 otherwise. The threshold for determining the mode is 30.

In each round, we generate $B_1 = 32$ bags from the policy. The GFlowNet model consists of two hidden layers with 16 hidden dimensions, which is trained with a learning rate of 1e-4. We use an exploration epsilon of 0.01. In addition, their implementation involves a buffer for enabling backward policy-based learning (Shen et al., 2023; Malkin et al., 2022b). We utilize this buffer for energy decomposition learning. The mini-batch size B_2 is same as B_1 . We set the number of iterations in energy decomposition learning N = 8 for each round. Note that reducing N still leads to promising results compared to baseline.

Molecule generation (Bengio et al., 2021a). The experiment settings, implementations, and hyperparameters are based on prior studies (Bengio et al., 2021a; Pan et al., 2023a). The maximum trajectory length is 8, with the number of actions varying between around 100 and 2000 which is depending on the state. The threshold for determining the mode is 7.5.

In each round, we generate four molecules, i.e., $B_1 = 4$. The model consists of Message Passing Neural Networks (Gilmer et al., 2017) with ten convolution steps and 256 hidden dimensions, which is trained with a learning rate of 5e-4. We rescale the reward so that the maximum reward is close to one, and exponent of it is set to 8.0. We use an exploration epsilon of 0.05. In energy decomposition learning, we do not use a buffer and immediately use the molecules that are sampled in each round. For PPO, we set the entropy coefficient to 1e-4 and do not apply the reward exponent because it causes a gradient exploding.

RNA sequence generation (Shen et al., 2023). The experiment settings, implementations, and hyper-parameters are based on prior studies (Shen et al., 2023). The action is defined as prepending or appending an amino acid to the current sequence. The maximum length is 8 and the number of actions is 4. The mode is determined based on whether it is included in a predefined set of promising RNA sequences (Shen et al., 2023).

In each round, we generate $B_1 = 16$ sequences. The GFlowNet model consists of two hidden layers with 128 hidden dimensions, which is trained with a learning rate of 1e-4. The reward exponent is set to 3.0. We use an exploration epsilon of 0.01. In addition, their implementation involves a buffer for enabling backward policy-based learning iteration (Shen et al., 2023). The hyper-parameters for energy decomposition learning is the same as that of bag generation. For PPO, we set the entropy coefficient to 1e-2.

Set generation (Pan et al., 2023a). The experiment settings, implementations, and hyper-parameters are based on prior studies (Pan et al., 2023a). The set generation task is to generate a set which involves 20 entities. There are 30 types of entities, where each action includes one of them in the current set. We set the threshold for determining the mode as 0.25.

In each round, we generate $B_1 = 16$ sets. The GFlowNet model consists of two hidden layers with 256 hidden dimensions, which is trained with a learning rate of 0.001. The hyper-parameters for energy decomposition learning is the same as that of molecule generation.