

NeuFACO: Neural Focused Ant Colony Optimization for Traveling Salesman Problem

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Abstract—This study introduces Neural Focused Ant Colony Optimization (NeuFACO), a non-autoregressive framework designed to address the Traveling Salesman Problem (TSP) by integrating advanced reinforcement learning methods with enhanced Ant Colony Optimization (ACO) techniques. Unlike prior neural-augmented ACO methods that rely on simplistic training schemes, NeuFACO adapts Proximal Policy Optimization (PPO) to train a graph neural network for generating instance-specific heuristic guidance. This learned heuristic is integrated into a highly optimized ACO framework that incorporates several state-of-the-art enhancements, such as candidate lists, focused tour modification, and scalable local search. By jointly exploiting amortized inference and the parallel stochastic exploration characteristic of ACO, NeuFACO delivers efficient and high-quality solutions over a diverse set of TSP instances. Code is available at <https://github.com/shoraaa/NeuFACO>.

Index Terms—Ant Colony Optimization, Traveling Salesman Problem, Proximal Policy Optimization, Graph Neural Networks, Neural Combinatorial Optimization

I. INTRODUCTION

The Traveling Salesman Problem (TSP) is a classical combinatorial optimization problem (COP) that seeks the shortest Hamiltonian tour visiting each city once and returning to the start. Due to its relevance in logistics, production planning, and circuit design, it serves as a benchmark for evaluating optimization algorithms [1].

Exact approaches like integer linear programming guarantee optimality but scale poorly due to exponential complexity [2]. Heuristic and metaheuristic methods offer efficient near-optimal solutions but often lack problem-specific knowledge, rely on handcrafted heuristics, and risk premature convergence to local optima [3], [4].

Deep learning has inspired data-driven approaches to COPs such as TSP [5]. Supervised methods train neural networks to imitate solvers for fast inference but depend on labeled solutions, limiting generalization. Reinforcement learning (RL) avoids this by directly optimizing solution quality through interaction, yet RL-based methods still suffer from weak local refinement, poor scalability, and high sample complexity [6].

To address these issues, hybrid frameworks combine neural models with metaheuristics. Non-autoregressive (NAR) “learn-to-predict” methods use neural networks to generate heuristics

that guide metaheuristics, rather than directly constructing solutions [7], [8]. However, current approaches face high variance and low sample efficiency due to simplistic training, while refinement often relies on standard Ant Colony Optimization (ACO) [9], which exploits TSP structure inefficiently and limits scalability to larger or heterogeneous instances.

A. Paper overview

We introduce Neural Focused Ant Colony Optimization (NeuFACO), a hybrid framework that integrates deep reinforcement learning with a refined variant of ACO. NeuFACO employs Proximal Policy Optimization (PPO) [10] to train a neural policy, offering greater stability and sample efficiency than prior approaches.

For refinement, NeuFACO adapts Focused ACO (FACO) [11], an advanced extension that performs targeted modifications around a reference solution. Unlike conventional ACO, which rebuilds full tours each iteration, FACO selectively relocates a few nodes, preserving strong substructures while improving weaker regions. This focused search narrows exploration to promising areas, enhancing efficiency.

By combining neural priors with restricted refinement, NeuFACO balances global guidance and local exploitation. This reduces disruption to near-optimal tours, accelerates convergence, and scales effectively to large TSP instances.

Experiments show that NeuFACO outperforms neural and classical baselines on random TSPs and TSPLIB benchmarks, solving problems with up to 1,500 nodes. Results also confirm the advantage of PPO in producing high-quality heuristic priors, establishing NeuFACO as a robust and generalizable framework for neural-augmented combinatorial optimization.

II. RELATED WORK

A. Traveling Salesman Problem

The Traveling Salesman Problem (TSP) is a classical NP-hard problem in combinatorial optimization [1]. It can be formally defined on a complete graph $G = (V, E)$, where the vertex set $V = \{v_1, v_2, \dots, v_n\}$ corresponds to the n cities and the edge set E denotes all pairwise connections among them. Each edge $(i, j) \in E$ has an associated non-negative cost $d_{i,j}$, often representing distance or travel time. For simplicity, this paper focuses on the two-dimensional Euclidean TSP as

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an illustrative example, where each edge $(i, j) \in E$ has a distance feature $d_{ij} = \|v_i - v_j\|$. The task is to determine a Hamiltonian tour $\pi = (\pi_1, \pi_2, \dots, \pi_n)$ that traverses every city once before returning to the origin, with the goal of minimizing the aggregate travel distance or cost:

$$C(\pi) = \sum_{i=1}^n d_{\pi_i, \pi_{i+1}}, \quad (1)$$

where π is a permutation of V representing the visiting order of the cities and $\pi_{n+1} = \pi_1$.

B. Classical Methods

Classical approaches to the TSP fall into two categories: exact and heuristic-based [3]. Exact methods, such as branch-and-bound [12] and Integer Linear Programming [2], guarantee optimality but suffer factorial time complexity, making them impractical for large instances.

To improve scalability, heuristic and metaheuristic algorithms provide near-optimal solutions in reasonable time [13]. Local search methods like 2-opt, 3-opt, and Lin–Kernighan [14], [15] refine tours by replacing subpaths to reduce cost, while metaheuristics such as Ant Colony Optimization [16] and Genetic Algorithms [17] stochastically explore the solution space by simulating natural processes.

C. Ant Colony Optimization

Ant Colony Optimization is a bio-inspired metaheuristic that emulates the collective foraging behavior of natural ant colonies [16]. In the context of the TSP, artificial ants construct solutions by probabilistically choosing the next city according to two key factors: the pheromone trail τ_{ij} , which encodes the learned desirability of selecting edge (i, j) , and the heuristic desirability $\eta_{ij} = 1/d_{ij}$, which favors shorter edges. The probability p_{ij}^k that ant k moves from city i to city j is typically defined as

$$p_{ij}^k = \begin{cases} \frac{\tau_{ij}^\alpha \eta_{ij}^\beta}{\sum_{l \in \mathcal{N}_i^k} \tau_{il}^\alpha \eta_{il}^\beta} & \text{if } j \in \mathcal{N}_i^k, \\ 0 & \text{otherwise,} \end{cases} \quad (2)$$

where \mathcal{N}_i^k denotes the set of cities not yet visited by ant k , and α and β determine the relative weighting of pheromone intensity and heuristic desirability. Once all ants have completed their tours, pheromone trails are updated both to reinforce high-quality paths and to allow for evaporation, which encourages exploration:

$$\tau_{ij} \leftarrow (1 - \rho)\tau_{ij} + \sum_{k=1}^m \Delta\tau_{ij}^k, \quad (3)$$

where $\rho \in (0, 1]$ is the evaporation rate, and $\Delta\tau_{ij}^k$ represents the pheromone deposited by ant k proportional to the quality of its solution. Over time, ACO converges toward promising regions of the solution space, but it may encounter slow convergence or premature stagnation.

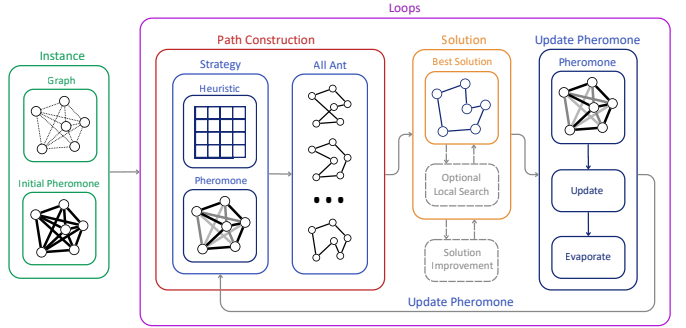


Fig. 1: Original Ant Colony Optimization.

D. Neural Combinatorial Optimization

Neural Combinatorial Optimization (NCO) employs deep learning to derive heuristics for NP-hard problems such as the Traveling Salesman Problem (TSP) [5]. Unlike handcrafted approaches, NCO learns directly from data via supervised imitation of high-quality solutions or reinforcement learning on reward signals such as negative tour length. Architectures explored include pointer networks [18], GNNs [19], and more recently Transformers [20] and diffusion models [21].

NCO methods are generally categorized as constructive, incrementally generating solutions, or improvement-based, refining initial ones with techniques like 2-opt [6]. Non-autoregressive (NAR) variants follow a hybrid paradigm: neural encoders extract heuristic signals that metaheuristics exploit for solution construction, contrasting with end-to-end autoregressive models [8].

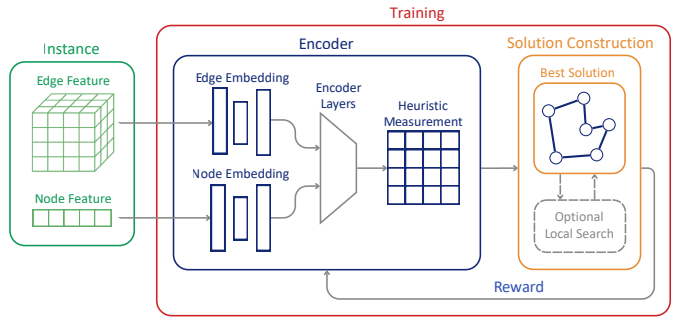


Fig. 2: NAR training procedure.

In Ant Colony Optimization (ACO), classical heuristics are handcrafted and domain-dependent, limiting scalability and adaptability. NAR methods overcome these issues by learning heuristics from data and embedding them into the ACO pipeline. Early implementations relied on REINFORCE [22], but its high variance and slow convergence hindered performance, motivating more stable reinforcement learning techniques.

III. METHODOLOGY

A. Markov Decision Process (MDP) formulation

We formulate the solution construction procedure of TSP as a Markov Decision Process (MDP). At step t , the state

$s_t = (\mathcal{X}, S_t, i_t)$ comprises the instance \mathcal{X} , the set of visited nodes S_t , and the current node i_t . The action is the next city $a_t \in \mathcal{N}_{i_t} \setminus S_t$ sampled from $\pi_\theta(\cdot | i_t)$ under feasibility masking. Transitions deterministically append a_t to the tour and update (S_{t+1}, i_{t+1}) ; the episode terminates after visiting all nodes with a single terminal reward $R = -C(\pi; \mathcal{X})$.

B. Policy Parameterization

A graph neural network f_θ parameterized by θ takes the graph data \mathcal{X} as input and outputs:

- Heuristic matrix $H \in \mathbb{R}^{n \times n}$ representing a learned prior over edge transitions.
- Value prediction $V_\theta(\mathcal{X}) \in \mathbb{R}$ estimating expected return for the given instance.

C. Rollouts and Rewards

For each instance \mathcal{X} , we sample a batch of M tours $\{\pi^{(m)}\}_{m=1}^M$ by chaining the policy over the shrinking feasible set:

$$\pi_{t+1}^{(m)} \sim \pi_\theta(\cdot | \pi_t^{(m)}), \quad t = 1, \dots, n-1.$$

Each tour receives a scalar terminal reward equal to the negative tour length:

$$R^{(m)} = -C(\pi^{(m)}; \mathcal{X}), \quad m = 1, \dots, M. \quad (4)$$

We use a per-instance, per-batch baseline $\bar{R} = \frac{1}{M} \sum_{m=1}^M R^{(m)}$ for variance reduction and to normalize reward and calculated advantage:

$$\tilde{R}^{(m)} = R^{(m)} - \bar{R}, \quad A^{(m)} = \tilde{R}^{(m)} - V_\theta(\mathcal{X}). \quad (5)$$

D. PPO Objective

For each sampled path $\pi^{(m)}$, let $\log p_\theta(\pi^{(m)})$ denote the log probability of sampling under the current policy, and $\log p_{\theta_{\text{old}}}(\pi^{(m)})$ under the previous policy. The log-probability of a sampled tour factorizes as:

$$\log p_\theta(\pi^{(m)}) = \sum_{t=1}^{n-1} \log \pi_\theta(\pi_{t+1}^{(m)} | \pi_t^{(m)}). \quad (6)$$

Then the probability ratio is:

$$r^{(m)}(\theta) = \frac{p_\theta(\pi^{(m)})}{p_{\theta_{\text{old}}}(\pi^{(m)})}. \quad (7)$$

And the clipped PPO surrogate objective is:

$$\begin{aligned} \mathcal{L}_{\text{policy}}(\theta) = -\frac{1}{M} \sum_{m=1}^M & \min \left(r^{(m)} A^{(m)}, \right. \\ & \left. \text{clip}(r^{(m)}, 1 - \varepsilon, 1 + \varepsilon) A^{(m)} \right). \end{aligned} \quad (8)$$

The value losses calculated using mean squared error (MSE):

$$\mathcal{L}_{\text{value}}(\theta) = \frac{1}{M} \sum_{m=1}^M \left(V_\theta(\mathcal{X}) - \tilde{R}^{(m)} \right)^2 \quad (9)$$

To promote exploration, entropy is computed on the normalized heuristic distribution: $\tilde{H}_{ij} = H_{ij}/(\sum_k H_{ik})$ and

$$\mathcal{H}(\tilde{H}) = -\frac{1}{n} \sum_{i=1}^n \sum_{j=1}^n \tilde{H}_{ij} \log(\tilde{H}_{ij} + \varepsilon). \quad (10)$$

Then we define the entropy loss as:

$$\mathcal{L}_{\text{entropy}} = -\beta \mathcal{H}(\tilde{H}), \quad (11)$$

and we minimize $\mathcal{L} = \mathcal{L}_{\text{policy}} + \mathcal{L}_{\text{value}} + \mathcal{L}_{\text{entropy}}$.

IV. SOLUTION SAMPLING

The ACO constructs a probability distribution over transitions:

$$p_{ij} \propto \tau_{ij}^\alpha \cdot H_{ij}, \quad (12)$$

where τ_{ij} is the pheromone and H_{ij} is the learned heuristic, and α is a weighting parameter.

Our method incorporates multiple state-of-the-art enhancements aimed at improving both the computational efficiency and solution quality of ACO algorithms. These improvements address three key aspects of solution construction: efficient node selection, focused modifications to high-quality tours, and scalable local search. Figure 3 demonstrates the final algorithm for solution sampling.

A. Incorporating the Min-Max Ant System

To enhance solution quality and stabilize the search process, NeuFACO adopts the Min-Max Ant System (MMAS) [23] for pheromone updates. MMAS modifies the classical Ant System in two key aspects: (1) pheromone trails are restricted to the interval $[\tau_{\min}, \tau_{\max}]$ to prevent premature convergence, and (2) updates rely solely on the best-so-far or iteration-best solution, rather than aggregating contributions from all ants.

In our implementation, pheromone updates follow the standard MMAS rule:

$$\tau_{ij} \leftarrow \begin{cases} (1 - \rho)\tau_{ij} + \Delta\tau_{ij}^{\text{best}} & \text{if edge } (i, j) \text{ chosen,} \\ (1 - \rho)\tau_{ij} & \text{otherwise,} \end{cases} \quad (13)$$

$$\tau_{ij} \leftarrow \text{clamp}(\tau_{ij}, \tau_{\min}, \tau_{\max}). \quad (14)$$

This mechanism is particularly critical when combined with neural-guided heuristics, as it prevents the learned model from prematurely dominating the search dynamics.

B. Node Selection via Candidate and Backup Lists

Candidate lists are widely used in ACO to restrict node selection to the k -nearest neighbors, thereby reducing complexity. Formally, for current node i , the candidate set is

$$\mathcal{C}_i = \{u_{i,j} \mid j \leq k, \text{visited}(u_{i,j}) = 0\}. \quad (15)$$

When $\mathcal{C}_i = \emptyset$, classical ACO scans all unvisited nodes, an $O(n)$ operation. To avoid this, we adopt the backup strategy of [24]: each node maintains a precomputed backup list BKP_i , from which the first unvisited neighbor is selected. Only if both \mathcal{C}_i and BKP_i are empty do we fall back to

$$J = \arg \min_{u: \text{visited}(u)=0} d(i, u). \quad (16)$$

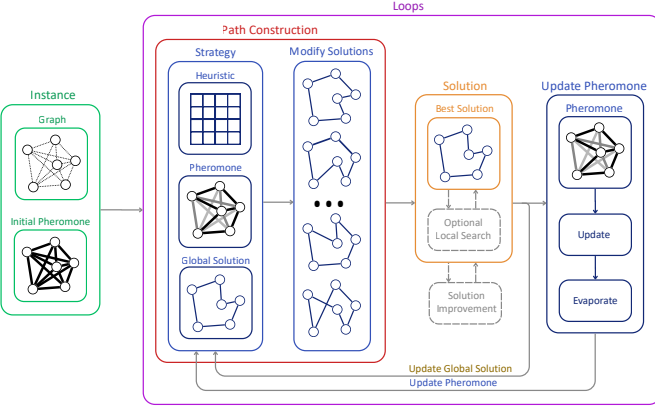


Fig. 3: State-of-the-art Ant Colony Optimization.

This hierarchical rule preserves stochastic exploration early on while substantially reducing overhead in the later stages of tour construction.

C. Node Relocation to maintain path structure

We adopt the focused modification strategy introduced in [11], which aims to explore the solution space by making minimal impactful changes to existing high-quality tours. At the start of construction, each ant stochastically copies a complete tour from either the iteration-best or the global-best candidate and subsequently modifies it by introducing new nodes based on the standard transition rule.

From the selected edge (u, v) and the original tour π , a relocate move produces a new tour π' by removing v from its current position and reinserting it immediately after u . Let $p = \text{pred}(v)$ be the predecessor of v in π , $s = \text{succ}(v)$ be the successor of v , and s_u be the successor of u . Then the cost change from performing such a move is

$$\Delta C = -d_{p,v} - d_{v,s} - d_{u,s_u} + d_{p,u} + d_{u,v} + d_{v,s_u}. \quad (17)$$

To maintain structural similarity to the reference tour, modification is limited: once a predefined minimum number of new edges (MNE) is introduced (we set MNE=8), the ant stops altering the route and continues following the remaining portion of the original tour. This strategy restricts the number of relocated nodes, thereby concentrating the search effort on promising subregions of the solution space while avoiding the computational burden of constructing tours entirely from scratch. Figure 4 demonstrates an example of a node relocation operation.

D. Scalable Local Search

Local search, particularly the 2-opt heuristic, is highly effective for improving ACO-generated solutions but is often applied sparingly due to its computational cost.

To enable frequent refinement, our algorithm incorporates two optimizations from [11]. First, it tracks modified edges in each constructed tour and applies 2-opt only to those differing from the previously optimized tour. Second, it restricts 2-opt to candidate edges, thereby reducing the search space.

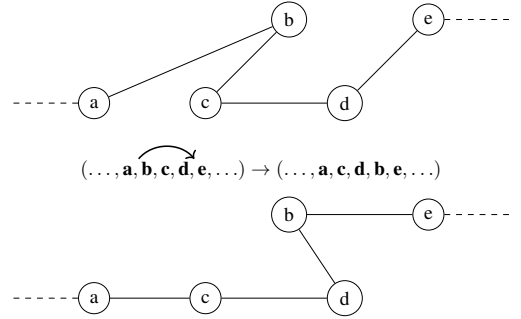


Fig. 4: Illustration of node relocation.

With a constant-size candidate list and a limited set of modified edges, this optimized 2-opt can be applied to every solution efficiently, fully exploiting high-quality tours without prohibitive overhead.

V. EXPERIMENTS

This section presents experimental results to validate the effectiveness of our algorithm. We compare it against other ACO-based methods as well as state-of-the-art neural approaches for solving the TSP. Our source code is built upon codebases of previous ACO-based methods: DeepACO (NeurIPS 2023) [7] and GFACS (AISTATS 2025) [25].

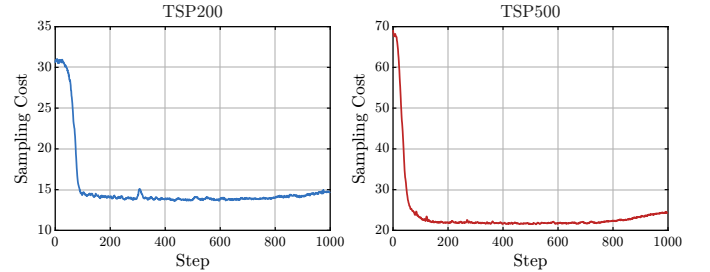


Fig. 6: Test objective cost over training step.

A. Setup

1) *Training*: We train and evaluate NeuFACO against two baseline models, DeepACO and GFACS. NeuFACO is trained with 20 PPO steps per epoch, a batch size of 20, a learning rate of 0.001, a clip ratio of 0.2, and an entropy coefficient of 0.01. All experiments are conducted on a machine with an AMD RX6800 GPU and an Intel i5-13400 CPU at 2.50 GHz. For fair comparison, DeepACO and GFACS are re-trained using their originally reported parameters, with the sole adjustment being the number of epochs, aligned to ours. Figure 8 presents the model sampling cost over training steps on the validation and test datasets, respectively.

2) *Benchmark*: We evaluate NeuFACO on two benchmark datasets: randomized TSP instances with size 200, 500 and 1000, each with 128 instances [7], and TSPLib [26] with up to 1500 nodes. Each experiment is repeated $R = 10$ times, and results are reported as averages. For other baseline methods,

Method	TSPLib200		TSPLib500		TSPLib1000	
	Gap(%)	Time	Gap(%)	Time	Gap(%)	Time
DeepACO	1.50	13.70s	3.43	143.92s	4.86	1047.59s
GFACS	1.44	13.75s	2.46	150.49s	3.99	1141.24s
NeuFACO	1.39	0.22s	2.06	0.48s	2.98	1.03s

Fig. 5: Different prior comparison.

results are taken directly from their respective papers. Errors are computed as $E = (\text{cost} - \text{optimal})/\text{optimal} \times 100\%$. The optimal values for gap calculation are obtained using an exact solver.

B. Comparison with other NAR solvers

We compare NeuFACO with previously proposed non-autoregressive methods, DeepACO and GFACS, both of which outperform the classical ACO without neural guidance [7], [25]. For all compared NAR models, the number of ants is fixed at $M = 100$, and each run consists of $I = 100$ ACO iterations. Unlike these methods, which employ neural-guided perturbation, NeuFACO does not rely on this technique; instead, it applies scalable local search at every iteration, providing effective refinement without additional perturbation mechanisms.

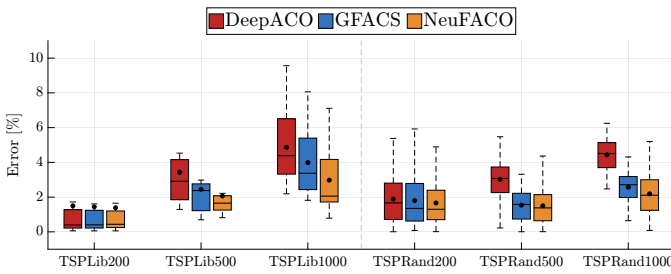


Fig. 7: Minimum, maximum and average errors comparison.

Figure 5 shows that among DeepACO, GFACS, and NeuFACO, our model achieves the best solution quality across nearly all datasets. NeuFACO delivers these improvements with a $5\text{--}100\times$ runtime reduction relative to prior methods, primarily due to the refined sampler that leverages neural heuristics while incurring minimal quality loss. Moreover, the use of PPO with entropy regularization alleviates policy degradation, ensuring solution quality on par with or superior to existing models. The gains are particularly pronounced on large-scale instances, such as TSPLib graphs with up to 1500 nodes, demonstrating NeuFACO’s suitability for efficiently solving large TSP instances without sacrificing accuracy.

Figure 8 further illustrates the evolution of average objective costs over iterations, confirming NeuFACO’s rapid convergence compared to other baselines.

C. Comparison with other RL solvers

In this section, we compare NeuFACO with problem-specific RL approaches for the TSP, including established baselines (AM [20], POMO [27], Pointerformer [28]), heatmap-based

Method	TSP200		TSP500		TSP1000	
	Gap(%)	Time	Gap(%)	Time	Gap(%)	Time
DeepACO	1.89	16.86s	3.03	119.58s	4.32	659.43s
GFACS	1.80	17.36s	1.54	136.14s	2.48	730.56s
NeuFACO	1.66	0.28s	1.50	0.74s	2.00	1.98s

Fig. 5: Different prior comparison.

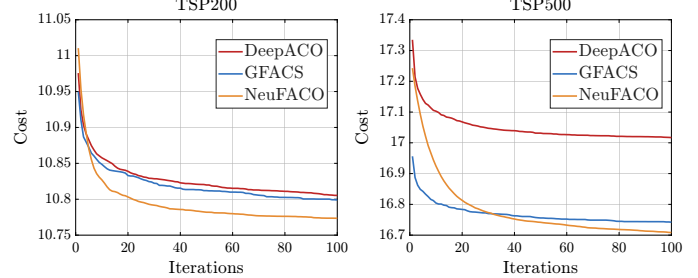


Fig. 8: Objective cost over iterations between different priors.

Results for other methods are taken from [25], [31], [32]. We reported the averaged objective value over 128 instances of TSP, originally from [38].

Method	TSP500		TSP1000			
	Obj.	Gap(%)	Time	Obj.	Gap(%)	Time
Concorde	16.55	–	10.7s	23.12	–	108s
AM	21.46	29.07	0.6s	33.55	45.10	1.5s
POMO	20.57	24.40	0.6s	32.90	42.30	4.1s
DIMES	17.01	2.78	11s	24.45	5.75	32s
SO	16.94	2.40	15s	23.77	2.80	26s
Pointerformer	17.14	3.56	14s	24.80	7.90	40s
H-TSP (AAAI 23)	17.55	6.22	0.18s	24.65	6.62	0.33s
GLOP (AAAI 24)	16.91	1.99	1.5m	23.84	3.11	3.0m
INViT-3V+Greedy (ICML 24)	16.78	1.56	5.48s	24.66	6.66	9.0s
LEHD+Greedy (NeurIPS 23)	16.78	1.56	0.13s	23.84	3.11	0.8s
BQ+Greedy (NeurIPS 23)	16.72	1.18	0.36s	23.65	2.30	0.9s
SIGD+Greedy (NeurIPS 23)	16.71	1.17	0.23s	23.57	1.96	1.2s
SIT+Greedy (NeurIPS 25)	16.7	1.08	14.95s	23.57	1.95	0.2s
DeepACO	16.84	1.77	15s	23.78	2.87	1.1m
GFACS	16.80	1.56	15s	23.72	2.63	1.1m
NeuFACO	16.43	1.33	0.91s	23.34	1.16	0.98s
			+14.95s			+40.03s

Fig. 9: Performance on TSP500 and TSP1000 with $M = 256$ ants and $I = 1000$ iterations in NeuFACO. Reported times include heuristic inference and solution sampling.

DIMES [29], improvement-based SO [30], and more recent methods such as H-TSP [31], GLOP [32], INViT [33], LEHD [34], BQ [35], SIGD [36], and SIT [37].

Results demonstrate that NeuFACO consistently achieves superior or highly competitive performance across all baselines. It surpasses every established methods and remains competitive with recently proposed approaches, often delivering better solution quality at the cost of slightly longer runtimes. The experiments also highlight a key bottleneck of NAR methods, including ours: solution sampling remains CPU-bound, leading

to higher runtime despite comparable amortized inference. Nevertheless, NeuFACO marks a significant advance for both NAR and RL-based approaches to the TSP.

VI. CONCLUSION

In conclusion, NeuFACO integrates deep reinforcement learning with a refined ACO framework to address limitations of non-autoregressive models for the TSP. By combining PPO-based policy learning with targeted refinement around high-quality solutions, it achieves a strong balance between global guidance and local exploitation. This synergy preserves solution structure, accelerates convergence, and scales effectively to large instances. Experiments show that NeuFACO consistently achieves superior or highly competitive performance compared to state-of-the-art neural baselines on both randomized and benchmark datasets. While runtimes are longer due to CPU-bound sampling, the method delivers higher solution quality, underscoring the effectiveness of PPO-guided priors in enhancing ACO and establishing NeuFACO as a robust and generalizable framework for neural-augmented combinatorial optimization.

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