Learning Trajectory Prediction with Continuous Inverse Optimal Control via Langevin Sampling of Energy-Based Models

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Abstract—Autonomous driving is a challenging multiagent domain which requires optimizing complex, mixed cooperative-competitive interactions. Learning to predict contingent distributions over other vehicles’ trajectories simplifies the problem, allowing approximate solutions by trajectory optimization with dynamic constraints. We take a model-based approach to prediction, in order to make use of structured prior knowledge of vehicle kinematics, and the assumption that other drivers plan trajectories to minimize an unknown cost function. We introduce a novel inverse optimal control (IOC) algorithm to learn other vehicles’ cost functions in an energy-based generative model. Langevin Sampling, a Monte Carlo based sampling algorithm, is used to directly sample the control sequence. Our algorithm provides greater flexibility than standard IOC methods, and can learn higher-level, non-Markovian cost functions defined over entire trajectories. We extend weighted feature-based cost functions with neural networks to obtain NN-augmented cost functions, which combine the advantages of both model-based and model-free learning. Results show that model-based IOC can achieve state-of-the-art vehicle trajectory prediction accuracy, and naturally take scene information into account.

I. INTRODUCTION

Autonomous driving is a challenging multiagent domain which requires optimizing complex, mixed cooperative-competitive interactions. Interacting with other drivers is currently a major impediment to the widespread deployment of driverless autonomous vehicles, and current approaches fall short of a general solution. Rule-based approaches to interaction are too brittle — they require a huge effort to engineer and validate, and they generalize poorly to new scenarios. Learning-based approaches are promising because of the complexity of driving interactions, and the need for generalization. However, learning based systems require a huge amount of data to cover the space of interactive behaviors, and current model-free learning-based methods have not achieved safe, generalizable autonomous interaction thus far.

Learning to predict contingent distributions over other vehicles’ trajectories is a standard approach to simplifying the interaction problem, allowing approximate solutions by trajectory optimization with dynamic constraints. Prediction methods generally fall into two categories: model-based and model-free. Model-based methods use prior knowledge of vehicle kinematics as well as the policies and intentions of other drivers to predict physically realistic maneuvers and trajectories [13]. Learning in these models either learns a space of trajectories or policies directly [2], or does inverse optimal control (IOC), learning a cost function that rationalizes observed behaviors [20], [23], [35]. Because they capture the generative structure of vehicle trajectories, model-based methods can potentially learn more, from less data, than model-free methods. However, good cost functions are challenging to learn, and simple, handcrafted representations may not generalize well across tasks and contexts [16], [28]. In general, model-based methods can be less flexible, and may underperform model-free methods in the limit of infinite data.

Model-free methods [1], [15], [22], [30] take a data-driven approach, aiming to learn predictive distributions over trajectories directly from data. These approaches are more flexible and require less knowledge engineering in terms of the type of vehicles, maneuvers, and scenarios, but the amount of data they require may be prohibitive.

Here we take a model-based approach to prediction, with the goal to achieve the data-efficiency and performance of model-based learning, with the flexibility and generality of model-free learning. Our approach is to combine inverse optimal control (IOC) with model-free deep neural network architectures for cost function learning. Our motivation is that adding structure in the form of kinematic vehicular constraints (which can be modeled very accurately), and the assumption that human drivers optimize their trajectories according to a subjective cost function, provides a powerful inductive bias for learning. Our approach is to focus on learning good cost function representations, and we experiment with various neural network architectures, and model-based and model-free hybrids, to find those which perform the best.

Specifically, we derive a family of algorithms for IOC using energy-based models. We show how training the model is related to a min-max procedure, and how we can use multiple optimal control algorithms for sampling, including Langevin
Sampling, gradient descent and iterative Linear Quadratic Regulation. We design multiple deep neural network architectures for cost function learning, some of which augment a set of hand-crafted features for driving using residual learning [17], and others that use convolutional (CNN) architectures to learn non-Markovian cost functions defined over entire trajectories.

We conduct several experiments to evaluate our approach on real-world driving data. We first compare our model with several state-of-the-art prediction techniques on the publicly-available NGSIM dataset [7]. We then evaluate our method on another autonomous highway driving dataset, containing a longer road segment than NGSIM, with more lane curvature as well as multiple highway entrances and exits. We compare our method with the CIOC algorithm [23], and we compare a range of different neural network structures in our framework. Finally, we analyze several synthetic examples to show that our IOC-based method predicts a variety of complex maneuvers, including lane-following, collision-avoidance, and overtaking a slower car ahead.

II. RELATED WORK

Inverse optimal control (IOC) and inverse reinforcement learning (IRL), have seen a variety of methods including maximum margin-based optimization, maximum entropy-based optimization, Bayesian inference, regression, and classification [4]. Most relevant to our work is the maximum causal entropy framework for IRL [15], which assumes an energy-based distribution on trajectories and learns the model parameters by maximizing the entropy over expert data. Wulfmer extended maximum entropy IRL to a deep version [31]. However, these algorithms are only applicable in discrete state spaces (or in continuous state spaces with linear dynamics and quadratic costs [26]), because they use dynamic programming to calculate the precise normalization term which is intractable in the continuous case (detailed in Section 3). For continuous IOC (CIOC), Levine used the Laplace approximation to model the trajectory distribution as a Gaussian [23]. However, the Laplace assumption is not always justifiable, especially when the cost function is complex. Our experiments compare our sampling method with the results of CIOC.

Several model-free methods have been proposed which have a deep relationship to model-based IOC. Finn proposed guided cost learning, a sample-based approach to policy learning which uses energy-based models [11], [12]. Policy search is used to learn a network for trajectories sampling, in order to approximate the normalization term in the energy-based model. Ho [18] and Li [29] synthesized and evaluated full state-control trajectories iteratively using generative adversarial imitation learning. In their setting, the trajectory distribution is captured by the generator in a GAN [14], while the cost function is closely related to the representation learned by the GAN discriminator [11]. Our experiments compare the performance of our model with that of GAIL.

Finally, a recent body of research has applied supervised model-free learning to trajectory prediction. These approaches typically use recurrent encoder-decoder architectures [1], augmented with various techniques such as GANs and aggregation functions such as max-pooling [15], spatio-temporal graphs [30], and attention mechanisms [29], ranking and IOC [22], and convolutional pooling [9]. Our experiments compare our model with the convolutional social pooling approach in [9].

A core concern in energy-based models is how to approximate high-dimensional, non-linear distributions with intractable normalization terms. Xie et al. used Langevin Sampling, an MCMC based sampling method to formulate non-linear energy-based models for a variety of data including image data [32], video data [34] and 3D data [33]. Our algorithmic approach is inspired by this technique.

III. MODEL

A. Task formulation

We first formulate the prediction task as an inverse optimal control problem. Rather than predict the trajectory (x,y position) directly, we use IOC to outputs the control with which we can easily rollout the entire trajectory based on the dynamic model. We define continuous Markov Decision Process (MDP) for a single agent in IOC,

\[
M = < X, U, D, C >,
\]

where \(X \in R^N\) is the state, \(U \in R^n\) is the control, D is the dynamic function \(x_{t+1} = f(x_t, u_t)\), and \(C(x_t, u_t)\) is the unknown cost function. We assume the Dynamic Function \(f\) is known.

A trajectory is a sequence of states and controls with a fixed length \(T\). We then define the cost function of a trajectory as \(C_\theta(\tau)\), linear or non-linear. Normally, \(C_\theta(\tau) = \sum_{i=0}^{T} C_\theta(x_i, u_i)\), but here we explore non-Markov cost functions which formed by a neural network.

B. Energy-based Models

With the goal to infer the distribution of the trajectory, we assume it takes the form of an energy-based model, where the energy term is the cost function.

The probability distribution of the trajectory \(\tau\) is defined as

\[
P(\tau; \theta) = \frac{1}{Z} \exp(-C_\theta(\tau)),
\]

where \(C_\theta(\tau)\) is the cost function and, \(Z = \int \exp(-C_\theta(\tau)) d\tau\) is the normalization term, corresponds \(\int P(\tau) = 1\). The probability of taking a trajectory is small if the corresponding cost is large.

The goal of IOC is to find a distribution that best fits the expert control. In other word, we maximize the log-likelihood on expert trajectories \((\tau_i \in Traj_{obs})\).

\[
l(\theta) = \frac{1}{n} \sum_{i=1}^{n} \log P(\tau_i; \theta) = \frac{1}{n} \sum_{i=1}^{n} (-C_\theta(\tau_i) - \log(Z)).
\]
C. Learning Algorithm

In the energy-based model, the normalization term $Z$ is intractable. In CIOC, it uses Laplace approximation that models the trajectory distribution as a Gaussian distribution. Hence, the likelihood can be approximated by the Taylor expansion on cost function around control $u$,

$$l(\theta) = \frac{1}{2} C_U C_U^T - C_U + \frac{1}{2} \log | - C_U | - \frac{d_u}{2 \log (2 \pi)},$$

where $C_U = \frac{\partial C}{\partial \theta}$, $C_U = \frac{\partial^2 C}{\partial \theta^2}$.

However, the assumption is not always justifiable. As an alternative, we use sample-based approach to calculate the normalization term, and the gradient is,

$$\frac{\partial}{\partial \theta} l(\theta) = \frac{1}{n} \sum \frac{\partial}{\partial \theta} C_\theta(\tau_t) = E_{P(\tau; \theta)}[\frac{\partial}{\partial \theta} - C_\theta(\tau_t)],$$

because

$$\frac{\partial}{\partial \theta} \log(Z) = \frac{1}{Z} \frac{\partial}{\partial \theta} \int \exp(-C_\theta(\tau))d\tau$$

$$= \int \frac{1}{Z} \frac{\partial}{\partial \theta} \exp(-C_\theta(\tau)) \frac{\partial}{\partial \theta} - C_\theta(\tau)d\tau$$

$$= \int \frac{\partial}{\partial \theta} C_\theta(\tau) P(\tau; \theta)d\tau$$

$$= E_{P(\tau; \theta)}[\frac{\partial}{\partial \theta} - C_\theta(\tau_t)].$$

The expectation term can be approximated by sampling,

$$\frac{\partial}{\partial \theta} l(\theta) = \frac{1}{n} \sum \frac{\partial}{\partial \theta} C_\theta(\tilde{\tau}_t) - \frac{1}{n} \sum \frac{\partial}{\partial \theta} C_\theta(\tilde{\tau}_t),$$

where $\tilde{\tau}$ is the sampled trajectories and $n$ is the number of samples. Different sampling approaches are discussed in Section 4.2.

D. Sampling Method

In order to approximate the normalize term $Z$ in EBM, we introduce two sampling methods.

The first one is Langevin Sampling, a widely used method in Markov chain Monte Carlo sampling.

The other one is iterative linear quadratic regulation (iLQR) which is common in optimal control field. iLQR can compute the control sequence with the lowest cost

1) Langevin Sampling: The state in our model can be divided into two parts, vehicle status $x_v$ and environment $x_e$. Modifying control affects the status but not the environment.

We sample the trajectories based on the input environment information. For each expert trajectory, we synthesize one trajectory based on the associated environment. In other words, we sample from the conditional distribution with fixed environment and maximize the conditional likelihood,

$$l(\theta) = \frac{1}{n} \sum_{i=1}^N \log P(\tau_i | X e_i = x_e; \theta).$$

Our sampling algorithm only updates the control, which leads to a change in status $x_v$.

The iterative process for Langevin Sampling is

$$U_{i+1} = U_i - \frac{\delta^2}{2} \frac{\partial}{\partial U} C_\theta(U_i) + \delta \text{Noise}_i.$$

The Langevin dynamics consists of a deterministic part, which is the gradient descent for control $u$, and a stochastic part, which is a Brownian motion that helps the chain to escape spurious local minima.

For each sampling sequence, we sample exactly one trajectory.

Notice that the state changes as the control is changed; at the same time, the change of control in the previous frame affects each cost later. Thus the derivative is calculated by chain rule,

$$\frac{\partial}{\partial u_i} C_\theta(U_T) = \sum_{i=t}^T \frac{\partial}{\partial u_i} C_\delta(U_T) = \sum_{i=t}^T \frac{\partial}{\partial x_i} \frac{\partial}{\partial u_i} \prod_{j=t}^{i-1} \frac{\partial}{\partial x_j} + \frac{\partial}{\partial u_i} C_\delta(U_T).$$

We can also use gradient descent to find the local mode of the energy function by eliminating the Brownian motion term in Langevin Sampling.

$$U_{T+1} = U_T - \frac{\delta^2}{2} \frac{\partial}{\partial U} C_\theta(U_T).$$

2) Iterative Linear Quadratic Regulation: Iterative Linear Quadratic Regulation (iLQR) is a variant of Differential dynamic programming (DDP) [24] [4]. Given an initial trajectory, it updates the trajectory by repeatedly solving for the optimal policy under linear quadratic assumptions.

Let $(x^i_t, u^i_t)$ be the i-th iteration trajectory. The dynamic is known, $x_{t+1} = f(x^i_t, u^i_t)$. Define $\Delta x_t = x_{t+1} - x_t, \Delta u_t = u_{t+1} - u_t$, then,

$$\Delta x_{t+1} \approx f(x^i_t, \Delta x_t + f(u^i_t) \Delta u_t)$$

$$C_\theta(x^i_t, u^i_t) \approx \Delta x_t c_{x_t} + \Delta u_t c_{u_t} + \frac{1}{2} \Delta x_t c_{x_x} \Delta x_t$$

$$+ \frac{1}{2} \Delta u_t c_{u} \Delta u_t + \Delta u_t c_{u_x} \Delta x_t + C_\theta(\Delta x_{t-1} - u_{t-1}).$$

where the subscripts denote the Jacobians and Hessians of the dynamic $f$ and cost function $C$.

E. Min-max Interpretation

We now show that the training procedure can be approximately interpreted as a min-max game. [33] Rewrite the deviation into

$$\frac{\partial}{\partial \theta} l(\theta) = \frac{\partial}{\partial \theta} \left[ \frac{1}{n} \sum C_\theta(\tilde{\tau}_t) - \frac{1}{n} \sum C_\theta(\tilde{\tau}_t) \right].$$

Let,

$$V_\theta(\tilde{\tau}) = \frac{1}{n} \sum C_\theta(\tilde{\tau}_t) - \frac{1}{n} \sum C_\theta(\tilde{\tau}_t)$$

Thus, minimum $V$ is equivalent to maximum the likelihood function.

Mode Shifting The learning step attempts to increase the value $V_\theta(\tilde{\tau})$ by updating $\theta$. It shifts the low energy mode from the current trajectories $\{\tilde{\tau}_i\}$ towards the expert trajectories $\{\tau_i\}$. In other word, with fixed $\tilde{\tau}_t$, and $\tau_t$, we want $\sum C_\theta(\tilde{\tau}_t)$
to be as small as possible comparing to \( \sum C_\theta(\tilde{\tau}_i) \) by changing \( \theta \).

**Mode Seeking** Then we decrease the value \( V_\theta(\tilde{\tau}) \) by selecting new \( \tilde{\tau} \). By sampling we can generate trajectories \( \{\tilde{\tau}_i\} \) which are optimal or near-optimal (close to the modes of the cost function).

As a result, our training process is
\[
\theta = \arg\min_\theta \max_{\tilde{\tau}} V_\theta(\tilde{\tau}).
\]

**F. Algorithm Flow**

The training algorithm of energy-based model learning is presented as follows,

**Algorithm 1** Inverse Optimal Control by EBM.

1: **input** expert trajectories \( \tau_i \).
2: **output** the cost function parameter \( \theta \), the synthesized trajectories.
3: Let \( t \leftarrow 0 \), initialize \( \theta \).
4: **repeat**
5:   **Mode Seeking:** Use current cost function to synthesis trajectories by iLQR or Langevin Sampling.
6:   **Mode shifting:** Use synthesized trajectories to update \( \theta by \theta(t+1) = \theta(t) + \gamma L'(\theta(t)) \).
7:   \( t \leftarrow t + 1 \).
8: **until** \( t = T \).

**IV. Experiments**

We quantitatively and qualitatively evaluate the trajectory predictions of our inverse optimal control framework on two driving datasets and several synthetic examples.

In autonomous driving, the state \( x \) consists of vehicle status (position, steering, speed) and environment information (lane, signs and other vehicle states). In our datasets the environment information is provided as the position of lane, the speed limit, the road boundary and other vehicle states which is estimated through cameras and other sensors. Control \( u \) has two dimensions: steering and acceleration. A vehicle trajectory is a sequence of states and controls with a constant interval between frames, typically 0.1s.

For our cost function learning procedure, our input is a large amount of observed trajectories. The full trajectories of the ego vehicle and other vehicles are provided, as is the lane which the vehicle sees at the first time step (represented as a cubic polynomial).

We assume only the starting point of a trajectory is known for each vehicle. Thus, we need to predict the future trajectories of other vehicles before predicting that of a given vehicle. For simplicity, these other-vehicle trajectory predictions use a constant velocity model. For evaluation, we compare the predicted trajectories with the ground truth in the testing set.

**A. NGSIM Dataset**

We first experiment on the publicly available NGSIM US-101 dataset \([7]\). This dataset consists of real highway traffic captured at 10Hz over a time span of 45 minutes. We preprocess it into a 5 second / 50 frame blocks. There are 831 total scenes with 96,000 5-second vehicle trajectories. There are no control values provided. Thus we need to infer the control of each vehicle given the vehicle state, which includes the position, speed and heading angle. Assuming bicycle model \([27]\) dynamics, we perform an inverse-dynamics optimization using gradient descent to minimize the distance between the trajectory reconstructed with the inferred controls, and the real one.

1) **Evaluation Metrics:** We use Root-Mean-Square-Error (RMSE) between the predicted trajectory and ground truth to evaluate the performance of our model.

RMSE for \( i \)-th position is defined as:
\[
RMSE = \sqrt{\frac{1}{N} \sum_k E_{\text{err}}(\tau_{\text{pre}}^k, \tau_{\text{obs}}^k, i)}
\]

A small RMSE is desired.

2) **Model Comparison:** We compare the following models:
   - Constant Velocity: simplest baseline, generates trajectories with constant velocity and zero steering.
   - C-VGMM + VIM: Uses maneuver-based variational Gaussian mixture models with a Markov random field based vehicle interaction module described in \([8]\).
   - GAIL-GRU: Considers a GRU model based on generative adversarial imitation learning described in \([21]\).
   - M-LSTM: Uses the maneuver-based LSTM approach described in \([10]\).
   - CS-LSTM: Updates V-LSTM by introducing convolutional Social Pooling, as described in \([9]\).

| Method                  | 1s  | 2s  | 3s  | 4s  |
|-------------------------|-----|-----|-----|-----|
| MEIOC via Langevin     | 0.318 | 0.644 | 1.149 | 2.138 |
| MEIOC via iLQR         | 0.351 | 0.603 | 0.969 | 1.874 |
| Constant Velocity      | 0.484 | 1.500 | 2.911 | 4.718 |
| C-VGMM + VIM \([8]\)   | 0.66 | 1.56 | 2.75 | 4.24 |
| GAIL-GRU \([21]\)      | 0.69 | 1.51 | 2.55 | 3.65 |
| M-LSTM \([10]\)        | 0.58 | 1.26 | 2.12 | 3.24 |
| CS-LSTM \([9]\)        | 0.61 | 1.27 | 2.09 | 3.10 |

**TABLE I**

NGSIM DATASET RESULTS.

3) **Results:** Table 1 and Figure 1 show the RMSE results for different methods. Figure 5 shows randomly selected trajectory predictions. Notice that the results of C-VGMM + VIM, GAIL-GRU, GAIL-GRU and CS-LSTM are retrieved from the original paper, yet we calculate constant velocity to make sure it is a fair comparison. The obtained constant velocity is 4.718, which is close to that originally reported at around 4.78.

For both optimal control methods, we set the parameter to make sure their speed is no less than 0.1s/sample, which is the lowest requirement for prediction.

Though the testing data typically should not include the scene in training data, such division is hard for the NGSIM dataset, because it includes the same scene through different times. We tried random splits, but there is no major difference.
between the training and testing data. As a result, we do use NGSIM data as a whole to train and test, the same as most baseline methods used for comparison.

4) Discussion: Our method achieved a substantial improvement compared with a state-of-art model-free prediction method. This suggests that representing vehicle kinematics and controls is fundamentally advantageous for prediction.

Comparing MEIOC via iLQR and langevin, iLQR result is better. We found it is probably due to the speed varies a lot in NGSIM dataset. Both method predict good on steering while following the lane. Comparing to steering, acceleration is always changed more rapidly (i.e. It is rare to have sudden turn in highway while a sudden stop is more usual) Langevin sampling tend to produce a smooth control making it hard to predict sudden change for acceleration.

It is true that there are some differences between the two methods. Our method relies on an accurate measure of speed and heading angle, so that the initial acceleration and steering can be predicted and used in other-vehicle prediction and ego-rollout. The result is very sensitive to initialization, while the NGSIM dataset is collected by a fixed camera so the relative position and speed calculation are quite accurate.

B. Autonomous Driving Dataset

1) Dataset: In order to evaluate our model in a more diverse range of scenario, we use a dataset collected from an autonomous car during repeated drives on a several-mile stretch of highway, which includes both car states and environment information. In this dataset, we compared our IOC algorithm with CIOC [23], the current state-of-art continuous IOC algorithm discussed in section 4. Compared to NGSIM dataset, this dataset is more challenging for two reasons:

- The lane curvature of this dataset is 3x larger than NGSIM.
- The data is noisy – especially for other vehicles. For NGSIM, the camera is set up in a fixed position while the autonomous driving dataset uses car-mounted cameras and LIDAR to estimate the states of other vehicles.

The advantage of this dataset is that it contains a more realistic driving scenario, as the control of the autonomous vehicle is collected by hardware on the car, which is more accurate than inferred controls.

The car state consists of four parts: x, y position, vehicle orientation and velocity. For the autonomous vehicle, the control is provided by two scalars: the steering and acceleration. Environment information consists of all lanes, represented as cubic polynomials. To solve the problem of noisy GPS signal, Kalman filtering is used to denoise the data. We use rollout data for both training and testing. The number of expert trajectories is 44,000, and each has a length of $T = 30$ frames with 0.1s intervals for a total of 4 seconds.

2) Evaluation Metrics: Other than RMSE, we introduce likelihood between predicted trajectory and ground truth. The reason is that, RMSE gives the same penalty for both x axis and y axis, while we try to give the y-axis (latitude) a bigger penalty, because even a small lateral shift may trigger a collision with an adjacent vehicle.

Assume,

$$\tau_1 = [(x_{11}, y_{11}), \ldots, (x_{1n}, y_{1n})],$$

$$\tau_2 = [(x_{21}, y_{21}), \ldots, (x_{2n}, y_{2n})].$$

The likelihood is defined as:

$$L(\tau_1, \tau_2) = \prod_{i=1}^{T} \frac{\phi(x_{1i}; x_{2i}, 1)\phi(y_{1i}; y_{2i}, 1)}{\phi(x_{1i}; x_{2i}, 1)\phi(y_{1i}; y_{2i}, 1)},$$

where $\phi(a; \mu, \sigma)$ is the probability under a Gaussian distribution with mean $\mu$ and standard deviation $\sigma$.

The prefect prediction has likelihood of 1. A bigger likelihood is desired.

| Method         | CIOC | iLQR | Langevin |
|----------------|------|------|----------|
| Training Likelihood | 0.795 | 0.745 | 0.799    |
| Testing Likelihood   | 0.653 | 0.745 | 0.810    |
| Training RMSE        | 0.789 | 0.871 | 0.767    |
| Testing RMSE         | 0.987 | 0.786 | 0.660    |

TABLE II

Comparison result with hand-crafted cost function.

3) Model Comparison: On this dataset, we focus on comparison between different IOC based prediction methods. The Continuous Inverse Optimal Control (CIOC) is used as a baseline. Table 2 shows our sample-based method is better and more stable than COIC. The training likelihood curve in Figure 7 also suggests a lower stability for CIOC. Langevin Sampling and gradient descent sampling yield similar results.

Due to the limited amount of data, the number of corner cases is relatively small. As a result, the testing result can be better than training. However, we could still observe that higher likelihood leads to better prediction in corner cases based on analysis of the predicted trajectories.

4) Cost Function Comparison: Langevin sampling method is capable to handle non-linear cost function. We evaluate different NN-augmented cost functions in Table 3. As for other neural network structures, a marginal improvement is
Table III

| Method          | Human-defined | NN as transformer | NN as residual | NN as residual to each | CNN    |
|-----------------|---------------|-------------------|---------------|------------------------|--------|
| Training Likelihood | 0.800         | 0.799             | 0.80          | 0.773                  | 0.814  |
| Testing Likelihood  | 0.810         | 0.794             | 0.804         | 0.772                  | 0.733  |
| Training RMSE     | 0.767         | 0.668             | 0.823         | 0.802                  | 0.626  |
| Testing RMSE      | 0.660         | 0.576             | 0.700         | 0.713                  | 0.778  |

TABLE III

Comparison result using Langevin Sampling with different cost function settings.

Fig. 2. Comparing different methods (Left: Baseline CIOC, Middle: iLQR, Right: Langevin Sampling).

Fig. 3. Predicted Trajectories from Autonomous Driving dataset.

achieved by ‘NN as transformer’. This structure provide non linear connection layer as a transformer of the original input sub-cost. This imply that there are some internal connection between each sub-cost. e.g. if there is a car in front, the collusion cost will increase. At this time, the lane keeping cost will be neglected or even negative impact.

‘NN as residual’ decreases the accuracy. It seems that the raw input will interfere the cost function, even argumented non-linear neural network.

CNN is the only method which have overfitting problem, with small training error but large testing error. Although we have a large training set, there are too much similar trajectories (i.e. following lane) making them not enough to fit a good parameter. More diversity of dataset, more corner cases are needed.

Detailed NN structure are provided in appendix. In this experiment we showed that the EBM model is able to handle the neural network design. Current designs are small nets and not well turned. We believe neural network structs will achieve better results.

C. Synthetic Examples

Corner cases are important for model evaluation. Therefore, we construct 4 typical corner cases to test our model. Figure 9 shows the predicted trajectory for several synthetic examples.

Fig 4(a)(b) are sudden-brake, where the orange vehicle in front of the ego car is braking. There is no car alongside in (a), so the predicted control is to change the lane. There are cars alongside in (b), which is why the ego car trigger braking.

(c) is another car trying to cut-in to the current lane. Our model plans a trajectory to avoid the collision, and even performs an overtake maneuver.

(d) shows a large lane curvature, yet our model still performs good lane following.

For all graphs, green points represent the predicted trajectory of the Langevin Sampling method, orange points show other vehicles and grey lines show the lane.

In short, our IOC algorithm via Langevin Sampling is capable of learning a reasonable cost function to avoid collision and handle cut-in situations.
V. CONCLUSION AND FUTURE WORK

In this paper, we implement continuous IOC with energy-based models in order to learn trajectory prediction. In comparison to other prediction methods, IOC-based methods show a large improvement by taking vehicle kinematics into account. In our experiments with an Autonomous Driving dataset, our method with Langevin Sampling is shown to generate better predictions with higher stability.

We design multiple neural network structures to augment a hand-designed cost function. Our experiments show energy-based model along with Langevin Sampling is capable to handle complex cost function. However, current results do not show great improvement thus introduce instability.

EBM is very promising for inverse optimal control, because as a descriptive model it is capable of representing the whole trajectory distribution in theory. We believe it can also be extended to multi-agent situations.

In autonomous driving, there are different moving objects including other vehicles, pedestrians, etc. Currently, we only predict the individual trajectories, while we assume other vehicles are driving by constant velocity. However, there is a major problem that vehicles are actually interacting to each other. For example, a vehicle behind may try to overtake if the front one trigger the break. In this case, the ego car should not change the lane. Therefore, multi-agent planning is a direction to explore. One possible implementation is to calculate the joint trajectory distribution for all moving agents, and sample multiple trajectories at the same time. Assume we have K agents and each of them has a trajectory $\tau_i$, then,

$$P(\tau_1, ..., \tau_K|\theta) = \frac{1}{Z}e^{\sum_{i=1}^{K} C_\theta(\tau_i)}.$$  

The cost function of each agent shares the same parameters. Notice that the cost function for one vehicle is dependent on the information on the others. We plan to focus on various generalizations of energy-based models in future work, with the belief that it has great potential for application to trajectory prediction and optimal control.

APPENDIX

Detailed definition of hand-crafted cost function

1) Feature-based Cost Function: Typically, the final cost function for a trajectory is defined as the sum of the cost for each frame with a state-control pair,

$$Cost_\theta(\tau) = \sum_{(x,u) \in \tau} Cost_\theta(x,u).$$

For feature-based cost function, we design multiple features and combine them linearly to obtain the final cost function. IOC can be used to learn the linear weight for each feature cost,

$$Cost_\theta(x,u) = \sum_{k=1}^{K} \theta_k f_k(x,u).$$

where $f_k(x,u)$ is hand crafted based on human expertise. See appendix for detailed settings.

The human-crafted cost function are defined as 10 components,

- The distance to the goal (x and y).
- The distance to the center of the lane.
- The penalty to collision to other vehicle. It is inversely proportional to distance to other vehicle.
- The L2-norm of acceleration and steering.
- The L2-norm for difference of acceleration and steering between two frames.
- The heading angle to lane.
- The difference to speed limit.

The goal is set as keeping the lane by the speed of speed limit. For initialization, we calculate all these components in training sample and normalized them in to same scale.

2) Neural Network Designs: Neural network is a powerful function approximator. It is capable of approximating arbitrarily complex nonlinear function given sufficient training data, and is flexible in incorporating prior information, which in our case is the manually designed features. We design three different network structures as an add-on to the feature-based function in the experiments.

Figure 1 shows the network structure. The red block stands for the human defined feature and the blue block the neural network layer, which uses fully connected layers. (a) stand the linear function.

(b) is 'NN as transformer'. Instead of linearly combining $J$ functional costs, we input the feature costs into a 2 layer fully-connected neural network and use the output as the final cost. Basically, we introduce nonlinearity to the human defined features.

(c) is 'NN as residual'. We feed the raw data into a two layer fully-connected neural network and output a scalar. The final cost is then the sum of the scalar output and the original CIOC cost. In this design, neural network serves as a residual to correct the cost, so it does not affect the result by much.

(d) is 'NN as residual to each'. The idea is similar to 'NN as residual', yet we output the $J$ scalars corresponding to $J$ features rather than one single scalar. We add each of the $J$ scalars to each feature as residual and output the linear combination. The experiments indicate that this method may distract the human defined features and decrease the accuracy.

Fig. 5. (a) hand-crafted Cost (b) nn as transformer (c) NN as residual (d) NN as residual to each.

A. Convolutional Neural Network Setting

To use some DDP (differential dynamic programming) methods including iLQR, we need to specify the cost function for each state. However, Langevin Sampling does not have
this constraint. We use a convolutional structure to connect the temporal information between states. That is, \( \text{Cost}_t = F(X, U(0)) \), where \( F \) is a fully-convolution neural network. The input to \( F \) is the single frame cost. Figure 2 displays the structure of the CNN.

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