Abstract—We present differentiable particle filters (DPFs): a differentiable implementation of the particle filter algorithm with learnable motion and measurement models. Since DPFs are end-to-end differentiable, we can efficiently train their models by optimizing end-to-end state estimation performance, rather than proxy objectives such as model accuracy. DPFs encode the structure of recursive state estimation with prediction and measurement update that operate on a probability distribution over states. This structure represents an algorithmic prior that improves learning performance in state estimation problems while enabling explainability of the learned model. Our experiments on simulated and real data show substantial benefits from end-to-end learning with algorithmic priors, e.g. reducing error rates by ∼80%. Our experiments also show that, unlike long short-term memory networks, DPFs learn localization in a policy-agnostic way and thus greatly improve generalization. Source code is available at https://github.com/tu-rbo/differentiable-particle-filters.

I. INTRODUCTION

End-to-end learning tunes all parts of a learnable system for end-to-end performance—which is what we ultimately care about—instead of optimizing each part individually. End-to-end learning excels when the right objectives for individual parts are not known; it therefore has significant potential in the context of complex robotic systems.

Compared to learning each part of a system individually, end-to-end learning puts fewer constraints on the individual parts, which can improve performance but can also lead to overfitting. We must therefore balance end-to-end learning with regularization by incorporating appropriate priors. Priors can be encoded in the form of differentiable network architectures. By defining the network architecture and its learnable parameters, we restrict the hypothesis space and thus regularize learning. At the same time, the differentiability of the network allows all of its parts to adapt to each other and to optimize their parameters for end-to-end performance.

This approach has been very successful in computer vision. Highly engineered vision pipelines are outperformed by convolutional networks trained end-to-end [8]. But it only works because convolutional networks [15] encode priors in the network architecture that are suitable for computer vision—a hierarchy of local filters shared across the image. Problems in robotics possess additional structure, for example in physical interactions with the environment. Only by exploiting all available structure will we be able to realize the full potential of end-to-end learning in robotics.

But how can we find more architectures like the convolutional network for robotics? Roboticists have captured problem structure in the form of algorithms, often combined with models of the specific task. By making these algorithms differentiable and their models learnable, we can turn robotic algorithms into network architectures. This approach enables end-to-end learning while also encoding prior knowledge from algorithms, which we call algorithmic priors.

Here, we apply end-to-end learning with algorithmic priors to state estimation in robotics. In this problem, a robot needs to integrate uncertain information over time. A single observation can be insufficient to estimate the state, and overestimating uncertainty is beneficial for filtering performance [21, p. 118]. Interestingly, end-to-end learning in DPFs re-discovers what roboticists found out via trial and error: that overestimating uncertainty is beneficial for filtering performance [21] p. 118.

Since DPFs are differentiable, we can learn their models end-to-end to optimize state estimation performance. Our experiments show that end-to-end learning improves performance compared to using models optimized for accuracy. Interestingly, end-to-end learning in DPFs re-discovers what roboticists found out via trial and error: that overestimating uncertainty is beneficial for filtering performance [21] p. 118.

Since DPFs use the Bayes filter algorithm as a prior, they have a number of advantages. First, even with end-to-
end learning, DPFs remain explainable—we can examine the learned models and their interaction. Second, the algorithmic prior regularizes learning, which greatly improves performance in state estimation. Compared to generic long short-term memory networks (LSTMs) [9], DPFs reduce the error rate by \( \sim 80\% \) or require 87% less training data for the same error rate. And finally, the algorithmic prior improves generalization: while LSTMs fail when tested with a different policy than used for training, DPFs are robust to changing the policy.

II. RELATED WORK

There is a surge of recent work that combines algorithmic priors and end-to-end learning for planning and state estimation with histogram-based and Gaussian belief representations.

Planning with known state: Tamar et al. [20] introduced value iteration networks, a differentiable planning algorithm with models that can be optimized for value iteration. Their key insight is that value iteration in a grid-based state space can be represented by convolutional neural networks. Silver et al. [18] proposed the predictron, a differentiable embedding of the TD(\( \lambda \)) algorithm in a learned state space. Okada and Aoshima [16] proposed path integral networks, which encode priors and end-to-end learning for planning and state estimation.

With latent variables: Aoshima [16] proposed path integral networks, which encode information with histogram-based and Gaussian belief representations. We consider the problem of estimating a latent state \( s \) from a history of observations \( o \) and actions \( a \), e.g. a robot’s pose from camera images and odometry. To handle uncertainty, we estimate a probability distribution over the current state \( s_t \) conditioned on the history of observations \( o_{1:t} \) and actions \( a_{1:t} \), which is called belief, \( \text{bel}(s_t) = p(s_t | a_{1:t}, o_{1:t}) \).

A. Bayes Filters

If we assume that our problem factorizes as shown in Fig. 2 the Bayes filter algorithm solves it optimally [21] by making use of the Markov property of the state and the conditional independence of observations and actions. From the Markov property follows that the last belief \( \text{bel}(s_{t-1}) \) summarizes all information contained in the history of observations \( o_{1:t-1} \) and actions \( a_{1:t-1} \) that is relevant for predicting the future. Accordingly, the Bayes filter computes \( \text{bel}(s_t) \) recursively from \( \text{bel}(s_{t-1}) \) by incorporating the new information contained in \( a_t \) and \( o_t \). From assuming conditional independence between actions and observations given the state follows that Bayes filters update the belief in two steps: 1) prediction using action \( a_t \) and 2) measurement update using observation \( o_t \).

1) The prediction step is based on the motion model \( p(s_t | s_{t-1}, a_t) \), which defines how likely the robot enters state \( s_t \) if it performs action \( a_t \) in \( s_{t-1} \). Using the motion model, this step computes the predicted belief \( \bar{\text{bel}}(s_t) \) by summing over all \( s_{t-1} \) from which \( a_t \) could have led to \( s_t \).

\[
\bar{\text{bel}}(s_t) = \int p(s_t | s_{t-1}, a_t) \text{bel}(s_{t-1}) \, ds_{t-1}. \tag{1}
\]

2) The measurement update uses the measurement model \( p(o_t | s_t) \), which defines the likelihood of an observation \( o_t \) given a state \( s_t \). Using this model and observation \( o_t \), this step updates the belief using Bayes’ rule (with normalization \( \eta \)).

\[
\text{bel}(s_t) = \eta \, p(o_t | s_t) \bar{\text{bel}}(s_t). \tag{2}
\]

Any implementation of the Bayes filter algorithm for a continuous state space must represent a continuous belief—and thereby approximate it. Different approximations correspond to different Bayes filter implementations, for example histogram filters, which represent the belief by a histogram, Kalman filters, which represent it by a Gaussian, or particle filters, which represent the belief by a set of particles [21].

B. Particle Filters

Particle filters approximate the belief with particles (or samples) \( S_t = s_t^{[1]}, s_t^{[2]}, \ldots, s_t^{[n]} \) with weights \( w_t^{[1]}, w_t^{[2]}, \ldots, w_t^{[n]} \). The particle filter updates this distribution by moving particles, changing their weights, and resampling them, which duplicates or removes particles proportionally to their weight.
Resampling makes this Bayes filter implementation efficient by focusing the belief approximation on probable states.

The particle filter implements the prediction step (Eq. 1) by moving each particle stochastically, which is achieved by sampling from a generative motion model,

\[ \forall i: s_t^i \sim p(s_t \mid a_t, s_{t-1}^i). \]  

The particle filter implements the measurement update (Eq. 2) by setting the weight of each particle to the observation likelihood—the probability of the current observation conditioned on the state represented by the particle,

\[ \forall i: w_t^i = p(o_t \mid s_t^i). \]  

The particle set is then resampled by randomly drawing particles \( s_t^i \) proportionally to their weight \( w_t^i \) before the filter performs the next iteration of prediction and update.

IV. DIFFERENTIABLE PARTICLE FILTERS

Differentiable particle filters (DPFs) are a differentiable implementation of the particle filter algorithm with end-to-end learnable models. We can also view DPFs as a new recurrent network architecture that encodes the algorithmic prior from particle filters in the network structure (see Fig. 3a).

With end-to-end learning, we do not mean that every part of a system is learned but that the objective for the learnable parts is end-to-end performance. For efficient end-to-end learning in particle filters, we need learnable models and the ability to backpropagate the gradient through the particle filter algorithm—not to change the algorithm but to compute how to change the models to improve the algorithm’s output.

This section describes our DPF implementation. Our source code based on TensorFlow [1] and Sonnet [4] is available at https://github.com/tu-rbo/differentiable-particle-filters.

A. Belief

DPFs represent the belief at time \( t \) by a set of weighted particles, \( \text{bel}(s_t) = (s_t, w_t) \), where \( S \in \mathbb{R}^{n \times d} \) describes \( n \) particles in \( d \)-dimensional state space with weights \( w \in \mathbb{R}^n \). At every time step, DPFs update the previous belief \( \text{bel}(s_{t-1}) \) with action \( a_t \) and observation \( o_t \) to get \( \text{bel}(s_t) \) (see Fig. 3a).

B. Prediction

The prediction step moves each particle by sampling from a probabilistic motion model (Eq. 3). Motion models often assume deterministic environments; they account for uncertainty by generating noisy versions of the commanded or measured action such that a different version of the action is applied to each particle [21, chap. 5]. We follow the same approach by splitting the motion model into an action sampler \( f \), which creates a noisy action \( \tilde{a}^i \) per particle, and a dynamics model \( g \), which moves each particle \( i \) according to \( \tilde{a}^i \).

\[ \tilde{a}^i = a_t + f_\theta(a_t, e^i) \sim \mathcal{N}, \]  

\[ s_t^i = s_{t-1}^i + g(s_{t-1}^i, a_t^i), \]

where \( f_\theta \) is a feedforward network (see Table 1). \( \theta \) are all parameters of the DPF, and \( e^i \in \mathbb{R}^d \) is a noise vector drawn from a standard normal distribution. Using the noise vector as input for a learnable generative model is known as the reparameterization trick [14]. Here, this trick enables \( f_\theta \) to learn to sample from action-dependent motion noise. The resulting noisy actions are fed into \( g \), which simulates how these actions change the state. Since we often know the underlying dynamics model, we can implement its equations in \( g \). Alternatively, we can replace \( g \) by a feedforward network \( g_\theta \) and learn the dynamics from data (tested in Section V-A3).

C. Measurement Update

The measurement update uses the observation to compute particle weights (Eq. 4). DPFs implement this update and additionally use the observation to propose new particles (see Fig. 3a). The DPF measurement model consists of three components: a shared observation encoder \( h \), which encodes an observation \( o_t \) into a vector \( e_t \), a particle proposer \( k \), which generates new particles, and an observation likelihood estimator \( l_\theta \), which weights each particle based on the observation.
where \( h_\theta, k_\theta, \) and \( l_\theta \) are feedforward networks based on parameters \( \theta \); the input \( \delta_i \) is a dropout vector sampled from a Bernoulli distribution. Here, dropout is not used for regularization but as a source of randomness for sampling different particles from the same encoding \( e_t \) (see Table I).

D. Particle Proposal and Resampling

We do not initialize DPFs by uniformly sampling the state space—this would produce too few initial particles near the true state. Instead, we initialize DPFs by proposing particles from the current observation (as described above) for the first steps. During filtering, DPFs move gradually from particle proposal, which generates hypotheses, to resampling, which tracks and weeds out these hypotheses. The ratio of proposed to resampled particles follows an exponential function \( \gamma^{-1} \), where \( \gamma \) is a hyperparameter set to 0.7 in our experiments. We use 1000 particles for testing and 16 particles for training (to speed up the training process). DPFs implement resampling by stochastic universal sampling [2], which is not differentiable and leads to limitations discussed in Section IV-F.

E. Supervised Learning

DPF models can be learned from sequences of supervised data \( o_{1:T}, a_{1:T}, s_{1:T}^* \) using maximum likelihood estimation by maximizing the belief at the true state \( s_t^* \). To estimate \( \text{bel}(s_t^*) \) from a set of particles, we treat each particle as a Gaussian in \( E \). Supervised learning and leads to limitations discussed in Section IV-F.

F. Limitations and Future Work

While the LSTM baseline learns localization in a way that stops working when the robot behaves differently (\( \sim 80\% \) error rate), localization with the DPF remains useful with different to supervised learning, where predicting the Markov state at each time step is a useful objective that facilitates future predictions. Differentiable resampling could still improve supervised learning, e.g., by encouraging beliefs to overestimate uncertainty, which reduces performance at the current step but can potentially increase robustness of future state estimates.

Since it is difficult to generate training data that include the true state \( s_t^* \) outside of simulation, we must work towards unsupervised learning, which will require backpropagation through multiple time steps because observations are generally non-Markov. Here are two possible implementations of differentiable resampling that could be the starting point of future work: a) Partial resampling: sample only \( m \) particles in each step; keep \( n-m \) particles from the previous time step; the gradient can flow backwards through those. b) Proxy gradients: define a proxy gradient for the weight of a resampled particle that is tied to the particle it was sampled from; the particle pose is already connected to the pose of the particle it was sampled from; the gradient can flow through these connections.

V. Experiments

We evaluated DPFs in two state estimation problems in robotics: global localization and visual odometry. We tested global localization in simulated 3D mazes based on vision and odometry. We focused on this task because it requires simultaneously considering multiple hypotheses, which is the main advantage of particle filters over Kalman filters. Here, we evaluated: a) the effect of end-to-end learning compared to individual learning and b) the influence of algorithmic priors encoded in DPFs by comparing to generic LSTMs. To show the versatility of DPFs and to compare to published results with backprop Kalman filters (BKFs) [2], we also apply DPFs to the KITTI visual odometry task [5]. The goal is to track the pose of a driving car based on a first-person-view video. In both tasks, DPFs use the known dynamics model \( g \) but do not assume any knowledge about the map of the environment and learn the measurement model entirely from data.

Our global localization results show that 1) algorithmic priors enable explainability, 2) end-to-end learning improves performance but sequencing individual and end-to-end learning is even more powerful, 3) algorithmic priors in DPFs improve performance compared to LSTMs reducing the error by \( \sim 80\% \), and 4) algorithmic priors lead to policy invariance: While the LSTM baseline learns localization in a way that stops working when the robot behaves differently (\( \sim 84\% \) error rate), localization with the DPF remains useful with different

| TABLE I: Feedforward networks for learnable DPF models |
|----------------|--------------------------------------------------|
| \( f_\theta \) | 2 x fc(32, relu), fc(3) + mean centering across particles |
| \( g_\theta \) | 3 x fc(128, relu), fc(3) + scaled by \( E_i [abs(s_t^* - s_{t-1})] \) |
| \( h_\theta \) | conv(3x3, 16, stride 2, relu), conv(3x3, 32, stride 2, relu), conv(3x3, 64, stride 2, relu), dropout(keep 0.3), fc(128, relu) |
| \( l_\theta \) | fc(128, relu), dropout(*keep 0.15), 3 x fc(128, relu), fc(4, tanh) |

fc: fully connected, conv: convolution, *: applied at training and test time
policies (~15% error rate).

In the visual odometry task, DPFs outperform BKFs even though the task exactly fits the capabilities and limitations of Kalman filters—tracking a unimodal belief from a known initial state. This result demonstrates the applicability of DPFs to tasks with different properties: higher frequency, longer sequences, a 5D state instead of a 3D state, and latent actions. The result also shows that DPFs work on real data and are able to learn measurement models that work for visually diverse observations based on less than 40 minutes of video.

A. Global Localization Task

The global localization task is about estimating the pose of a robot based on visual and odometry input. All experiments are performed in modified versions of the navigation environments from DeepMind Lab [3], where all objects and unique wall textures were removed to ensure partial observability. Data was collected by letting the simulated robot wander through the mazes (see Fig. 4). The robot followed a hand-coded policy that moves in directions with high depth values from RGB-D input and performs 10% random actions. For each maze, we collected 1000 trajectories of 100 steps with one step per second for training and testing. As input for localization, we only used RGB images and odometry, both with random disturbances to make the task more realistic. For the observations, we randomly cropped the rendered 32 × 32 RGB images to 24 × 24 and added Gaussian noise (σ = 20, see Fig. 4d-f).

As actions, we used odometry information that corresponds to the change in position and orientation from the previous time step in the robot’s local frame, corrupted with multiplicative Gaussian noise (σ = 0.1). All methods were optimized on short trajectories of length 20 with Adam [13] and regularized using dropout [19] and early stopping. We will now look at the results for this task.

1) Algorithmic priors enable explainability: Due to the algorithmic priors in DPFs, the models remain explainable even after end-to-end learning. We can therefore examine a) the motion model, b) the measurement model, and c) their interplay during filtering. Unless indicated otherwise, all models were first learned individually and then end-to-end.

\begin{align*}
\text{Motion Model:} & \quad \text{Obs. likelihood estimator} \\
\text{Measurement Model:} & \quad \text{Obs. likelihood estimator}
\end{align*}

Fig. 5: Learned motion model. (a) shows predictions (cyan) of the state (red) from the previous state (black). (b) compares prediction uncertainty in \( x \) to true odometry noise (dotted line).

\begin{align*}
\text{Obs. likelihood estimator} & \quad \text{Obs. likelihood estimator} \\
\text{Particle proposer} & \quad \text{Particle proposer}
\end{align*}

Fig. 6: Learned measurement model. Observations, corresponding model output, and true state (red). To remove clutter, the observation likelihood only shows above average states.

a) Motion Model: Fig. 5a shows subsequent robot poses together with predictions from the motion model. These examples show that the model has learned to spread the particles proportionally to the amount of movement, assigning higher uncertainty to larger steps. But how does this behavior depend on whether the model was learned individually or end-to-end?

Fig. 5b compares the average prediction uncertainty using models from different learning schemes. The results show that individual learning produces an accurate model of the odometry noise (compare red and the dotted black lines). End-to-end learning generates models that overestimate the noise (green and orange lines), which matches insights of experts in state estimation who report that “many of the models that have proven most successful in practical applications vastly overestimate the amount of uncertainty” [21, p. 118].

b) Measurement Model: Fig. 6 shows three example observations and the corresponding outputs of the measurement model: proposed particles and weights depending on particle position. Note how the model predicts particles and estimates high weights at the true state and other states in locally symmetric parts of the maze. We can also see that the data
distribution shapes the learned models, e.g. by focusing on dead ends for the second observation, which is where the robot following the hand-coded policy will look straight at a wall before turning around. Similar to motion models, end-to-end learned measurement models are not accurate but effective for end-to-end state estimation, as we will see next.

c) Filtering: Figure 7 shows filtering with learned models. The DPF starts by generating many hypotheses (top row). Then, hypotheses form clusters and incorrect clusters vanish when they are inconsistent with observations (second row). Finally, the remaining cluster tracks the true state.

2) End-to-end learning improves performance: To quantify the effect of end-to-end learning on state estimation performance, we compared three different learning schemes for DPFs: individual learning of each model (ind), end-to-end learning (e2e), and both in sequence (ind+e2e). We evaluated performance in all three mazes and varied the amount of training trajectories along a logarithmic scale from 32 to 1000. We measured localization performance by error rate, where we consider a prediction erroneous if the distance to the true state, divided by $E_t[\text{abs}(s_t - s_{t-1})]$, is greater than 1.

The resulting learning curves in Fig. 8a-c show that end-to-end learned DPFs (orange line) consistently outperform individually trained DPFs (red line) across all mazes. Individual training is worst with few training trajectories (less than 64) but also plateaus with more data (more than 125 trajectories). In both cases, the problem is that the models are not optimized for state estimation performance. With few data, training does not take into account how unavoidable model errors affect filtering performance. With lots of data, the models might be individually accurate but suboptimal for end-to-end filtering performance. End-to-end learning consistently leads to improved performance for the same reasons.

Performance improves even more when we sequence individual and end-to-end learning (green line in Fig. 8a-c). Individual pretraining helps because it incorporates additional information about the function of each model into the learning process, while end-to-end learning incorporates information about how these models affect end-to-end performance. Naturally, it is beneficial to combine both sources of information.

3) Algorithmic priors improve performance: To measure the effect of the algorithmic priors encoded in DPFs, we compare them with a generic neural network baseline that replaces the filtering loop with a two-layer long-short-term memory network (LSTM) [9]. The baseline architecture uses the same convolutional network architecture as the DPF—it embeds images using a convolutional network $h_{\theta}$, concatenates the embedding with the action vector and feeds the result into $2\times\text{lstm}(512), 2\times\text{fc}(256, \text{relu}), \text{and fc}(3)$—and is trained end-to-end to minimize mean squared error.

The comparison between DPF (ind+e2e) and the LSTM baseline (blue) in Fig. 8a-c shows that the error rate of DPF (ind+e2e) is lower than for LSTM for all mazes and all amounts of training data. Also in all mazes, DPF (ind+e2e) achieve the final performance of LSTM already with 125 trajectories, $\frac{1}{8}$ of the full training set.

We performed a small ablation study in maze 2 to quantify the effect the known dynamics model on this performance. When the dynamics model is learned, the final error rate for DPFs increases from 1.6% to 2.7% compared to 6.0% error rate for LSTMs. This shows that knowing the dynamics model is helpful but not essential for DPF’s performance.
Algorithmic priors lead to policy invariance: To be useful for different tasks, localization must be policy-invariant. At the same time, the robot must follow some policy to gather training data, which will inevitably affect the data distribution, add unwanted correlations between states and actions, etc.

We investigated how much the different methods overfit to these correlations by changing the policy between training and test, using two policies A and B. Policy A refers to the heuristic exploration policy that we used for all experiments above (see Sec. V-A). Policy B uses the true pose of the robot, randomly generates a goal cell in the maze, computes the shortest path to the goal, and follows this path from cell to cell using a simple controller mixed with 10% random actions.

The results in Fig. 9 show that all methods have low error rates when tested on their training policy (although DPFs improve over LSTMs even more on policy B). But when we use different policies for training and test, LSTM’s error rate jumps to over 80%, while DPF (ind+e2e) still works in most cases (5% and 26% error rate).

The LSTM baseline is not able to generalize to new policies because it does not discriminate between actions and observations and fits to any information that improves state estimation. If the training data includes correlations between states and actions (e.g. because the robot moves faster in a long hallway than in a small room), then the LSTM learns this correlation. Put differently, the LSTM learns to infer the state from the action chosen by the policy. The problem is that this inference fails if the policy changes. The algorithmic priors in DPFs prevent them from overfitting to such correlations from the Bayes filter algorithm reduce variance without adding bias—that these algorithmic priors capture some true structure about the problem, which data does not help to improve upon.

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The results in Fig. 9 show that all methods have low error rates when tested on their training policy (although DPFs improve over LSTMs even more on policy B). But when we use different policies for training and test, LSTM’s error rate jumps to over 80%, while DPF (ind+e2e) still works in most cases (5% and 26% error rate).

The LSTM baseline is not able to generalize to new policies because it does not discriminate between actions and observations and fits to any information that improves state estimation. If the training data includes correlations between states and actions (e.g. because the robot moves faster in a long hallway than in a small room), then the LSTM learns this correlation. Put differently, the LSTM learns to infer the state from the action chosen by the policy. The problem is that this inference fails if the policy changes. The algorithmic priors in DPFs prevent them from overfitting to such correlations from the Bayes filter algorithm reduce variance without adding bias—that these algorithmic priors capture some true structure about the problem, which data does not help to improve upon.

4) Algorithmic priors lead to policy invariance: To be useful for different tasks, localization must be policy-invariant. At the same time, the robot must follow some policy to gather training data, which will inevitably affect the data distribution, add unwanted correlations between states and actions, etc.

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because DPFs cannot directly infer states from actions.

DPFs generalize better from A to B than from B to A. Since generalization from B to A is equally difficult for DPFs with individually learned models, the error increase cannot come from overfitting to correlations in the data through end-to-end learning but is most likely because the states visited by policy A cover those visited by policy B but not vice versa.

The alternative approach to encoding policy invariance as a prior is to learn it by adding this variance to the data. Our results show that if we train on combined training data from both policies (A+B), all methods perform well in tests with either policy. This approach in the spirit of domain randomization and data augmentation helps DPFs because it covers the union of the visited states and (additionally) helps LSTM by including state-action correlations from both policies. But to make the LSTM localization truly policy invariant such that it would work with any new policy C, the training data has to cover the space of all policies in an unbiased way, which is difficult for any interesting problem.

### B. Visual Odometry Task

To validate our simulation results on real data, we applied DPFs on the KITTI visual odometry data set, which consists of data from eleven trajectories of a real car driving in an urban area for a total of 40 minutes. The data set includes RGB stereo camera images as well as the ground truth position and orientation of the car in an interval of $\sim$0.1 seconds. The challenge of this task is to generalize in a way that works across highly diverse observations because the method is tested on roads that are never seen during training. Since the roads are different in each trajectory, it is not possible to extract global information about the car’s position from the images. Instead, we need to estimate the car’s translational and angular velocity from the stream of images and integrate this information over time to track the car’s position and orientation.

We tackle this problem with a DPF in a five dimensional state space, which consists of the position, orientation, forward velocity and angular velocity. DPFs learn to perform visual odometry from a known initial state using a simple first-order dynamics model $g$ and a learnable action sampler $f_a$. Since there is no information about the action of the driver, the action sampler produces zero mean motion noise on the velocity dimensions, which is then evaluated with the measurement model. For a fair comparison, we used the same network architecture for the observation encoder $h_\theta$ as in the backprop Kalman filter paper [7], which takes as input the current image and the difference image to the last frame (see Fig. 10). Our observation likelihood estimator $l_\theta$ weights particles based on their velocity dimensions and the encoding $h_\theta(o_t)$. Since, the initial state is known, we do not use a particle proposer. We train the DPF individually and end-to-end, using only the velocity dimensions for maximum likelihood estimation.

We evaluated the performance following the same procedure as in the BKF paper. We used eleven-fold cross validation where we picked one trajectory for testing and used all others for training with subsequences of length 50. We evaluated the trained model on the test trajectory by computing the average error over all subsequences of 100 time steps and all subsequences of 100, 200, 400, and 800 time steps.

Table II compares our results to those published for BKFs [7]. DPFs outperform BKFs, in particular for short sequences where they reduce the error by $\sim$30%. Any improvement over BKFs in this task is surprising because Gaussian beliefs seem sufficient to capture uncertainty in this task. The improvement could come from the ability of particles to represent long tailed probability distributions. These results demonstrate that DPFs generalize to different tasks and can be successfully applied to real data.

### VI. Conclusion

We introduced differentiable particle filters to demonstrate the advantages of combining end-to-end learning with algorithmic priors. End-to-end learning optimizes models for performance while algorithmic priors enable explainability and regularize learning, which improves data-efficiency and generalization. The use of algorithms as algorithmic priors will help to realize the potential of deep learning in robotics. The components of the DPF implementation, such as sample generation and density estimation, will be useful for producing differentiable versions of other sampling-based algorithms.

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