A Dynamic Key Frames Approach to Object Tracking

by

Christopher A. Wilkens

Submitted to the Department of Electrical Engineering and Computer Science in partial fulfillment of the requirements for the degree of Master of Engineering in Computer Science and Engineering

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Abstract

In this thesis, I present a dynamic key frames algorithm for state estimation from observations. The algorithm uses KL-divergence as a metric to identify the frames that contribute the most information to estimation of the system’s current state. The algorithm is first presented in a numerical optimization framework and then developed as an extension to the Condensation algorithm. Finally, I present results from a Matlab simulation of the algorithm.

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Chapter 1

Introduction

Many problems in signal processing and artificial intelligence may be framed as state estimation from past observations. One such task is object tracking. In object tracking, the path of an object is estimated as the object moves through a series of camera frames.

Object tracking is an important task in machine vision. It is useful for a wide variety of tasks from 3D structure-from-motion to video surveillance. In 3D structure-from-motion, we track known points on a 3D object. Then, the 3D structure of those points is inferred from their trajectories. In video surveillance, one could track people of interest and look for suspicious behavior. Similarly, one could track the players in a sporting event. Finally, at a more basic level, object tracking may be used as a means of locating an object in an image during a time when it is occluded, increasing machine intelligence by allowing a computer to understand that an object has not entirely disappeared just because it is not visible.

In this thesis, we will develop a new algorithm for state estimation that provides an efficient way to identify and save relevant historical observations in order to improve tracking robustness. Then we will test our algorithm in simulation.
1.1 Challenges

Many algorithms perform quite well when tracking particular objects under controlled conditions. Unfortunately, object tracking in the general case remains an open problem. Background clutter, partial occlusions, unusual perspectives, lighting, and many other factors make reliable image understanding and tracking extremely difficult.

Some properties that commonly confound state estimation are nonlinear dynamics and complex probability distributions. Unfortunately, both are present in object tracking. A moving object, such as a person, may follow a highly nonlinear dynamic model. Nonlinear dynamics, while not necessarily intractable, make efficient algorithms for state estimation scarce.

The probability distributions present in object tracking are also very complex. For example, in the presence of image clutter, a tracking algorithm may see a plurality of objects that look like the one it was tracking, generating highly multimodal distributions. The problem is magnified when tracking multiple targets, as the image always has a plurality of similar objects.

Occlusions, or frames where the object of interest is hidden behind another object, present another difficult problem. While the temporary loss of data would reflect a failure in many systems, it is a natural occurrence when multiple objects interact in a scene. A successful tracking algorithm must be able to understand the object’s dynamics and predict the location of the object even when it is completely hidden or only partially visible so that it can recover tracking when the object reappears.

Some other difficulties of object tracking are inherent to object detection and image understanding. Varying image conditions may make accurate detection and localization quite difficult and generate a significant amount of noise. The exact location of an object may vary from frame to frame based on subtle changes in its identifying features and even the discretization implicit in the digital image. For example, lighting fluctuations on a shadowed edge of an object may significantly affect the perceived boundary, generating substantial noise in the observed center of the object.
We will demonstrate a general framework that is able to handle these difficulties more effectively than existing algorithms.

1.2 Existing Work

Many solutions to the object tracking problem have been attempted with varying success. In theory, the best approach would use all historical observations in a batch optimization to estimate the object’s current location. Unfortunately, this quickly becomes intractable because the data set grows linearly with time.

To efficiently track objects, many people have developed recursive Bayesian algorithms [10][8][17][6]. These algorithms assume Markovianity of the object’s position process and compute the posterior probability distribution, i.e. the probability distribution over the current location of the object, from the distribution at the last frame and the current image. This gives a recursive form to the solution, rendering the resulting problem tractable.

One of the earliest such algorithms used for object tracking is the Kalman filter [10]. The Kalman filter is an optimal, recursive estimator for linear Gaussian systems. Consequently, if the input observations are best modeled by a linear Gaussian system, this will work very well. Unfortunately, as we have already discussed, dynamics are frequently nonlinear and distributions are often multimodal. The details of the Kalman filter are discussed in section 3.2.1.

The Kalman filter has been extended to handle nonlinear dynamics and subsequently applied to object tracking in more complex situations [9]. The extended Kalman filter (EKF) extends this idea to include nonlinear dynamics by using linear approximations when propagating the covariance. Unfortunately, it is neither an optimal estimator nor proven to converge, so it may be useless in other applications. The Unscented Kalman Filter (UKF) of Julier et al. [9] builds on the EKF by elegantly propagating both the mean and the covariance through the nonlinear system. This addresses some nonlinear dynamics issues, but it still requires Gaussian distributions.

In order to handle image clutter, Shalom et al. developed the probabilistic data
association filter (PDAF) [17][6]. The PDAF is a Bayesian algorithm that can probabilistically handle multiple observations in the vicinity of the object being tracked. The PDAF incorporates the probability that a particular observation was generated by the desired object, giving it more flexibility than all-or-nothing approaches.

A more recent development in recursive Bayesian algorithms adopts the idea of a particle filter [8][15]. Issard and Blake’s Condensation algorithm permits arbitrarily complex posterior distributions and dynamic models at a reasonable cost in accuracy and efficiency [8]. In practice, the Condensation algorithm performs fairly well in many contexts. Its ability to maintain arbitrarily complex distributions allows it to handle the complex posteriors that frequently occur in noisy and occluded object tracking settings. We will extend this algorithm in section 2.2.

However, the Markovian assumption is frequently wrong. To handle these cases, some have attempted to apply semi-Markovian models to other tracking environments [12]. A semi-Markov process is similar to a Markov one, except that the time between state transitions may be distributed according to a more general probabilistic distribution (as opposed to the geometric form inherent in a standard Markov process.) Such models offer significant power but require model-specific recursive algorithms. In contrast, our approach only assumes smoothness.

Some tracking algorithms take a different approach and focus on developing accurate models of the item being tracked [11][1][18]. For example, human motion tracking algorithms will frequently use a model of the human body to improve performance. An extreme example is the SCAPE model which uses a detailed 3D surface model to estimate human pose and track motion accurately [1]. Similarly, the feature-based algorithm of Shi and Tomasi tracks objects by analyzing the motion of model features from frame to frame [18]. These approaches may work well, but like the semi-Markovian models, they are very task- and situation-specific. In particular, they require a good model a priori, and, in many cases, it is difficult or impossible to construct such a model.

Another body of research has tackled the problem of simultaneously tracking multiple targets. One of the biggest challenges with multiple target tracking is the
data association problem: determining the correspondence between observations and targets.

In many ways, the problem is similar to that of tracking objects in clutter. An algorithm must decide whether or not a particular observation corresponds to the current trajectory, with the new constraints that an object may belong to one of many trajectories. Given this similarity, a natural approach to multiple target tracking was to extend the PDAF to handle multiple targets [7]. The resulting joint probabilistic data association filter (JPDAF) has been used and extended in many multi-target tracking applications, including tracking contours with HMMs [3] and mobile robotics [5].

One fundamental approach is multiple hypothesis tracking (MHT) pioneered by Reid [16]. MHT maintains a set of trajectories based on particular observation-target associations. Unfortunately, the number of possible assignment sets grows exponentially with time. Consequently, much of the ongoing work in MHT revolves around efficient approximations and heuristics. For example, Cox and Hingorani keep only the $k$ best trajectories [4], and Vasquez and Williams exploit correlation effects [20].

A more recent idea applies Markov chain Monte Carlo simulation to the tracking problem [14]. These methods have become possible with the increased power of modern computers and have produced strong results.

While most approaches to multiple target tracking involve data association, the task may be circumvented through alternative statistical techniques. For example, network tomography infers global trajectories from local movements without explicitly identifying objects. Boyd and Meloche use this approach to track pedestrians at a crowded intersection [2].

Finally, another set of tracking algorithms use a key frames approach. A small subset of observations is kept and used to track in a manner similar to batch processing. Vacchetti et al. [19] and Morency et al. [13] use heuristics or hand labeling to identify representative views of an object as their key frames. At any given step, these views are used to estimate the location and perspective of the object being tracked.
When necessary, the key frame set is updated using heuristics.

In contrast, our dynamic key frames algorithm (DKF) keeps the locally optimal frames for estimating both the current location and dynamics of the object. At each step, the algorithm computes and saves the set of key frames most relevant to its understanding of the system’s state. This gives more robustness than either the Bayesian recursive algorithms or the existing heuristic-based key frames approaches.

In the following chapters, we will describe the design and implementation of DKF. Chapter 2 discuss the theory behind the algorithm and formulates it in the particle filter framework of Condensation. Chapter 3 discusses our implementation of both a batch DKF version and the Condensation DKF algorithm. Finally, in chapter 4 we conclude and recommend future work.
Chapter 2

The DKF Algorithm

We propose a general application of the key frames paradigm to estimate state from observations. It will maintain a dynamic active set of key frames chosen to maximize the amount of information known about the current state of the system. At each step, the algorithm will estimate the system’s current state and use Kullback-Leibler divergence (KL-divergence) computations to determine which frames in the active set are most relevant.

Formally, we wish to estimate the trajectory \( X_t = \{x_0, \ldots, x_t\} \) from image observations \( I_0, \ldots, I_t \). A global optimization routine seeks to optimize the full posterior. Namely, it seeks to find the optimal estimate \( \hat{x}_t \) of the current state \( x_t \) given all previous frame data:

\[
\hat{x}_0 \ldots \hat{x}_t = \arg \max_{x_0 \ldots x_t} P(x_0 \ldots x_t | I_0, \ldots, I_t).
\]

In contrast, a recursive Bayesian optimization solution assumes Markovianity, giving the following factorization of the joint posterior:

\[
P(x_0, \ldots, x_t | I_1, \ldots, I_t) = P(x_0) \prod_{i=1}^{k} P(I_i | x_i) \cdot P(x_i | x_{i-1}).
\]

This naturally leads to algorithms of the prediction/update form seen in the Kalman
filter:

\[ P(x_t|I_{t-1}) = \int P(x_t|x_{t-1})P(x_{t-1}|I_{t-1}) \, dx_{t-1} \]

\[ P(x_t|I_t) = \alpha P(I_t|x_t)P(x_t|I_{t-1}) \]

Our approach will take a middle ground, optimizing over a small active set \( AS = \{I^1, I^2, \ldots I^k\} \) of frames, i.e.

\[ \hat{x}_{t_1}, \ldots \hat{x}_{t_k} \hat{x}_t = \arg\max_{x_{t_1}, \ldots x_{t_k}, x_t} P(x_t|AS). \]

Finally, we take the estimate of the current state to be \( \hat{x} \) as computed above. For notational purposes, let \( Z_t \) be the posterior distribution over possible positions \( x_t \) of an object in an image at time \( t \) and \( \hat{x}(Z_t) \) be the best estimate of \( x_t \) from \( Z_t \). Also, \( Z_t(S) \) be the distribution over possible positions \( x_t \) based on dynamic constraints and a set of frames \( S \), and \( \hat{x}(S) = \hat{x}(Z_t(S)) \) as shorthand. In essence,

\[ Z_t(AS) \sim P(x_t|AS). \]

Figure 2-1: At each step, the algorithm keeps the frames most influential to the distribution of \( Z_t \).

In our approach, we maintain an active set of frames \( AS \) and keep only those
frames most relevant to our knowledge object’s current position. At a high level, the algorithm may be summarized as follows:

1. Incorporate the current frame, $f_t$, by adding it to the active set, $AS_t = AS'_{t-1} \cup \{f_t\}$.

2. Estimate $x_t$ from the frames in the active set as $\hat{x}(AS_t)$.

3. Discard frames to keep the active set small, i.e. $AS'_t = SELECT(AS_t)$.

The key to making the algorithm work well is to design $SELECT(AS_t)$ to keep the most useful frames from $AS_t$. Our goal is to demonstrate the superiority of a KL-divergence based $SELECT$. In this thesis, we elect to keep exactly $k$ frames from step to step and compare a few different approaches:

1. Keep the last $k$ frames. This gives

   $$SELECT(AS_t = \{I_{t_1}, \ldots, I_{t_k}, I_{t}\}) = \{I_{t_2}, \ldots, I_{t_k}, I_{t}\}$$

2. Keep the $k$ frames that minimize the Euclidean distance between $\hat{x}(AS_t)$ and $\hat{x}(AS'_t)$, i.e.

   $$SELECT(AS_t = \{I_{t_1}, \ldots, I_{t_k}, I_{t}\}) = AS_t - \arg\min_{I_j \in AS_t} |\hat{x}_t(AS_t) - \hat{x}_t(AS_t - I_j)| .$$

3. Keep the $k$ frames that minimize the KL-divergence between $Z_t(AS_t)$ and $Z_t(AS'_t)$, i.e.

   $$SELECT(AS_t = \{I_{t_1}, \ldots, I_{t_k}, I_{t}\}) = AS_t - \arg\min_{I_j \in AS_t} D_{KL}(Z_t(AS_t - I_j) - Z_t(AS_t))$$

   where $D_{KL}(\cdot)$ is defined as

   $$D_{KL}(Z_t(\tilde{AS}_t)\| Z_t(AS_t)) = \int p_{Z_t(\tilde{AS}_t)}(x) \log \frac{p_{Z_t(\tilde{AS}_t)}(x)}{q_{Z_t(AS_t)}(x)} \, dx$$
or as
\[
D_{\text{KL}}(Z_t(\tilde{A}S_t) \| Z_t(AS_t)) = \sum p_{Z_t(\tilde{A}S_t)}(x) \log \frac{p_{Z_t(\tilde{A}S_t)}(x)}{q_{Z_t(AS_t)}(x)}
\]
for discrete distributions.

4. Keep all the frames.

We expect the KL-divergence to have the best performance of the tractable SELECT implementations. The KL-divergence measures the distance between the \(Z_t\) distributions, thereby approximating the information from all the frames in \(AS_t\) as well as possible.

In contrast, keeping the last \(k\) frames fails to consider the relevance of the observations. For example, occlusions frequently happen for significant, contiguous blocks of time. If an object is occluded for more than \(k\) frames, keeping the last \(k\) frames is of little added benefit. One might employ a heuristic to determine that the current frames were empty, but the other approaches solve the problem more generally.

Minimizing the Euclidean distance provides some measure of relevance when selecting the active set. Its weakness is that it only propagates the mode of \(Z_t\). At each step, the algorithm is forced to commit to a particular trajectory assignment and picks frames that will, in the future, bias it toward that choice. Thus, while it may handle occlusions well, it will likely fail when it is important to maintain the multimodality of \(Z_t\).

Finally, as previously noted, keeping the entire frame history in the active set is computationally intractable. However, it is a useful performance benchmark that we will use later.

### 2.1 Multiple Target Tracking

One of our goals in this work is to demonstrate the performance of DKF when simultaneously tracking multiple targets. To simultaneously track, we augment \(X_t\) (and
therefore \( x_t \) to include the position of all objects being tracked, e.g.

\[
x_t = \begin{bmatrix}
x_1^t \\
x_2^t \\
\vdots
\end{bmatrix}
\]

where \( x_j^t \) is the location of object \( j \) at time \( t \). This sacrifices efficiency for accuracy, as it significantly increases the dimensionality of the tracking space. In the future, when we refer to \( X, x_t, \) and \( \hat{x} \), we are referring to the concatenation of all the locations and estimates for all objects being tracked.

This complicates our computation of \( \hat{x}_t \). In general, using the instantaneous mode of a multimodal distribution is potentially unstable because the estimate may jump between modes. In the case of multiple targets, this manifests itself as \( \hat{x}_t \) jumping between trajectories.

### 2.2 Condensation-DKF

In order to practically implement dynamic key frames, we need to give efficient ways to compute \( \hat{x} \) and \( \text{SELECT}(AS_t) \). Instead of seeking an analytic solution, we will seek a numerical approximation using a particle filter. Specifically, we will extend the Condensation algorithm. This will give us arbitrary (theoretical) precision in a simple implementation.

Issard and Blake’s Condensation algorithm [8] is a standard, weighted particle filter resampled at each timestep. At time \( t \), we start with the weighted samples from the previous step, \( \{ s_{t-1}^{(n)}, \pi_{t-1}^{(n)} \} \) where \( s_{t-1}^{(n)} \) corresponds to sample \( n \) and \( \pi_{t-1}^{(n)} \) is the weight for sample \( n \). We sample from this distribution and do a randomized prediction to get the next set of samples, \( \{ s_t^{(n)} \} \). Finally, we compute new weights based on the current image.

The algorithm can be summarized as follows:

1. **Sample.** Given \( \{ s_{t-1}^{(n)}, \pi_{t-1}^{(n)} \} \), sample from the weighted particle distribution to
get a set of particles \( \{ s^{(n)}_{t-1} \} \).

2. **Drift and diffuse.** Deterministically predict the next location for each particle and add noise to each sample according to the dynamic model. This gives \( \{ s^{(n)}_t \} \).

3. **Weight.** Compute new weights \( \{ \pi^{(n)}_{t-1} \} = \{ Pr(s^{(n)}_t | I_t) \} \), i.e. the conditional probability of the sample given the current frame.

At the sampling stage, the particles will be distributed according to \( Pr(s^{(n)}_t | Z_{t-1}) \).

In the drift and diffusion stage, the probability of getting a particular sample \( s^{(n)}_t \) can be written as \( Pr(s^{(n)}_t | s^{(n)}_{t-1}) \). The algorithm is implemented such that this conditional probability matches the system dynamics. This gives samples \( \{ s^{(n)}_t \} \) distributed according to

\[
Pr(s^{(n)}_t | s^{(n)}_{t-1}) \cdot Pr(s^{(n)}_{t-1} | Z_{t-1}) = Pr(s^{(n)}_t | Z_{t-1}).
\]

Finally, the weights are then chosen to be \( \{ \pi^{(n)}_{t-1} \} = \{ Pr(s^{(n)}_t | f_t) \} \). This implies that the final weighted particle distribution is a representation of

\[
Pr(s^{(n)}_t | Z_{t-1} \cap f_t).
\]

In the limit as the number of particles grows, the collection will approach the true distribution, thus making the Condensation algorithm a correct recursive estimator.

The only remaining task is to estimate \( \hat{x}_t \) from \( Z_t \). For this thesis, we compute \( \hat{x}_t \) as the mode of \( Z_t \). As noted earlier, this is slightly problematic.

Now, we are ready to define Condensation-DKF. We extend our samples to include a trajectory over the frames in the active set. Previously, condensation samples contained only position and velocity information for the object at time \( t \). Now, samples will also include position information for each of the frames contained in the active set.

Conveniently, the computation of \( \hat{x} \) does not change from a traditional condensation implementation. It is computed as the instantaneous mode of \( Z_t \) by ignoring the historical trajectory information to get the marginal distribution over \( x_t \).
Now, we need to show how to implement DKF. First, in order to compare two potential active sets, we need a way to represent $Z_t(\tilde{AS}_t)$ where $\tilde{AS}_t \subset AS_t$. For this, we keep the same particles, discard information for frames not in $\tilde{AS}_t$, and compute new weights $\pi^n_{\tilde{AS}}$. This gives us an approximation of $Z_t(\tilde{AS}_t)$.

When using the Euclidean distance metric, we can compute $\hat{x}(\tilde{AS}_t)$ and pick the best subset $\tilde{AS}_t$.

Computing the KL-divergence is slightly more difficult. The distribution $Z_t$ is ideally continuous. This gives us the following form for the KL-divergence:

$$D_{KL}(Z_t(\tilde{AS}_t) \| Z_t(AS_t)) = \int p_{Z_t(\tilde{AS}_t)}(x) \log \frac{p_{Z_t(\tilde{AS}_t)}(x)}{p_{Z_t(AS_t)}(x)} \, dx.$$  

However, we cannot easily compute this function in continuous space, so we would like to use the discrete version as an approximation. The easy approach is to perform the sum such that each sample trajectory $s^n_t$ corresponds to a point in the sum, i.e.

$$D_{KL}(Z_t(\tilde{AS}_t) \| Z_t(AS_t)) = \sum_n \alpha^n_{\tilde{AS}} \log \frac{\beta^n_{\tilde{AS}}}{\gamma^n_{AS}}$$

for some appropriate $\alpha$, $\beta$, and $\gamma$. We observe that the total probability of a trajectory selected this way is $\delta^n \cdot \pi^n_S$, where $S$ is the active set being used and $\delta$ is the probability implicitly encoded in the particle distribution. This suggests that we take

$$\alpha = \beta = \delta^n \cdot \pi^n_{\tilde{AS}_t}.$$  

However, if we add over all samples, a multiplicative factor of $\delta^n$ is implicitly present. Thus, we actually want

$$\alpha = \pi^n_{\tilde{AS}_t},$$

$$\beta = \delta^n_{t-1} \cdot \pi^n_{\tilde{AS}_t},$$

and

$$\gamma = \delta^n_{t-1} \cdot \pi^n_{AS_t}.$$
This gives the final form

$$D_{KL}(Z_t(\tilde{A}S_t)\|Z_t(AS_t)) = \sum_n \pi_{\tilde{A}S_t}^n \log \frac{\delta_n^t \cdot \pi_{\tilde{A}S_t}^n}{\delta_{t-1}^n \cdot \pi_{AS_t}^n}.$$ 

which conveniently simplifies to

$$D_{KL}(Z_t(\tilde{A}S_t)\|Z_t(AS_t)) = \sum_n \pi_{\tilde{A}S_t}^n \log \frac{\pi_{\tilde{A}S_t}^n}{\pi_{AS}^n}.$$

We will use this form to compute the Condensation-DKF.
Chapter 3

Implementation and Results

To test the DKF approach, we implemented it in Matlab and tested it against simulated data sets. The simulation trajectories were designed to have the following difficult properties related to object tracking:

1. **Occlusion.** In each trial, one trajectory has a significant (≈ 10 frame) occluded region.

2. **Clutter.** In cluttered environments, there may be many candidate observations near a given trajectory. In our tests, one trajectory is cluttered, containing multiple observations near the object.

3. **Multiple targets.** Each trial contains two separate intersecting trajectories. The trajectories are intended to represent similar objects, so the recognition stage cannot differentiate between them.

4. **Noise.** We add Gaussian noise to simulate standard measurement noise.

For the test trajectories, an “image” is simply a collection of locations \((x_i, y_i)\) corresponding the locations of an object. Such data could presumably be obtained by running an object detection scheme first.
3.1 Data Sets

Tests were primarily conducted on an *occluded intersection* data set of 100 trials. In this data set, the first trajectory is occluded as it passes by the second trajectory (see figure 3-1.) The second trajectory is “cluttered” at each timestep with an average of one extra observation near the trajectory. Finally, Gaussian noise is added to each trajectory.

Figure 3-1: In this run, two objects move from left to right through an image. The ground truth positions are denoted with an ‘x’ while the observations are denoted with a ‘.’. The red object is occluded for a long period of time and reappears just on the other side of the blue object, making successful tracking difficult.
3.2 Benchmarks

As a benchmark, we implemented the standard Kalman filter and the Condensation algorithm.

3.2.1 Kalman Filter

As mentioned earlier, the Kalman filter is an optimal recursive estimator for linear Gaussian systems. It may be derived fairly easily, but we will just summarize the results. First, one must model their system using linear state update equations as

\[ x_t = A \cdot x_{t-1} + w_{k-1} \]

\[ y_t = C \cdot x_t + v_t \]

where \( x_t \) is the internal state at time \( t \), \( y_t \) is observed, and \( w_t \sim N(0, Q) \) and \( v_t \sim N(0, R) \) are Gaussian noise variables. The resulting Kalman filter consists of prediction and update equations for both the mean, \( \hat{x} \), and the covariance, \( P \).

**Prediction:**

\[ \hat{x}_{t|t-1} = A_k \hat{x}_{t-1|t-1} \]

\[ P_{t|t-1} = A P_{t-1|t-1} A^T + Q. \]

**Update:**

\[ K_t = P_{t|t-1} C^T (C P_{t|t-1} C^T + R)^{-1} \]

\[ \hat{x}_{t|t} = \hat{x}_{t|t-1} + K_t (y_t - C \hat{x}_{t|t-1}) \]

\[ P_{t|t} = (I - K_t C) P_{t|t-1} \]

The prediction equations predict the mean \( \hat{x}_{t|t-1} \) and covariance \( P_{t|t-1} \) of the distribution of \( x_t \) given the distribution at time \( t - 1 \). Then, the update equations adjust this prediction based on the actual observations at time \( t \).

The only special feature of our Kalman filter implementation is the extension of
the state space to include the locations of two objects and to identify observations with a particular trajectory. To enable multiple target tracking, observations were assigned to trajectories before updating $\hat{x}$ based on the current frame. To do this, we used a nearest-neighbors approach and assigned observations to the closest trajectory. A sample trial using the Kalman filter is in figure 3-2.

![Figure 3-2: Tracking with a Kalman filter.](image)

Figure 3-2: Tracking with a Kalman filter. The solid red and blue lines represent the estimated trajectories computed by the Kalman filter as the objects move down and to the right. The data association used for the Kalman filter forced it to recover the red trajectory; however, it switched the two objects.

A key weakness of the Kalman filter in these tests was its inability to represent multimodal distributions. At the intersection, the Kalman filter could not represent the multimodality that appears in $Z_t$. (I.e. did the paths cross or did the objects turn away from each other at the last minute?) In combination with the early assignment of observations, this causes the Kalman filter to follow the cluttered trajectory instead of waiting for the other trajectory to stop being occluded. The success rate for the
Kalman filter was a mere 24% (see table 3.1.)

### 3.2.2 Condensation

Similar to the Kalman filter, Condensation was implemented using an extended state space to store the position of both objects. In our implementation, each sample also stored the velocity of the object for prediction purposes. To compute $\hat{x}$ from the samples, we used an implementation of the mean-shift algorithm to find the mode of $Z_t$.

As noted earlier, the mode of $Z_t$ is not necessarily stable in a particle filter. This can be clearly seen in the results as the hypothesis hops between trajectories.

![Figure 3-3: Tracking with Condensation. The two solid lines represent a failed run of the condensation filter. The occlusion is too long and the red trajectory cannot recover, while the blue hypothesis goes halfway between the red and blue objects.](image)
3.3 Numerical Optimization

Our first implementation of DKF used Matlab’s numerical optimization methods. Specifically, we wrote a function

$$\alpha(x_{t_1}, \ldots, x_{t_k}, x_t) = Pr(x_{t_1}, \ldots, x_{t_k}, x_t|AS_t)$$

where

$$AS_t = \{I_{t_1}, \ldots, I_{t_k}, I_t\}.$$ 

We use Matlab’s built-in Levenberg-Marquardt optimizer to find

$$\tilde{X}_t = \arg \max_{x_{t_1}, \ldots, x_{t_k}, x_t} \alpha(x_{t_1}, \ldots, x_{t_k}, x_t).$$

Unfortunately, $\alpha$ is symmetric: as defined, trajectories may be arbitrarily interchanged while maintaining the same $\alpha$ value. Consequently, after finding $\tilde{X}_t$, we must match the trajectories. For this, we take the Euclidean distance between the overlapping points in $X_{t-1}$ and $\tilde{X}_t$ and compute the minimal matching. Finally, we append the final value of $\tilde{X}_t$ to $X_{t-1}$ to get $X_t$.

In this case, computing the posterior probability of a path includes associating observations with trajectories. This gives it a significant advantage, allowing it do isolate the two trajectories with near perfection when they were both visible.

Unfortunately, the myriad local minima of the optimization space made this approach somewhat difficult. Matlab was highly effective at optimizing paths once the observations were associated with the correct trajectory; however, all of Matlab’s built-in optimization algorithms rarely changed more than a couple observation associations from their values in the initialization vector. As a result, we needed to carefully select the initialization vector.

We finally used dynamic programming to construct the initialization trajectories. We used a modified construction of the Viterbi algorithm that evaluated the total posterior (instead of multiplying the conditional by the previous probability) to con-
struct two trajectories entirely from observation points and interpolated occlusions. (Note that these trajectories are not optimal in any overall sense - they are merely a good starting point.)

Fortunately, due to the method by which trajectories were matched to the past, this implementation did not suffer from the instability seen in the Condensation algorithm. The only exception is the second frame, where the optimizer almost always switched the two trajectories briefly.

Ultimately, when the optimization trackers erred, they swapped trajectories at the intersection. For this problem, the optimizer was largely at the mercy of the initialization vector.

Figure 3-4: DKF keeping all frames. The solid red and blue lines represent the trajectories estimated by the algorithm and the rainbow colored lines represent the optimal partial trajectories found at each timestep. The temporary crossover at the beginning occurs in step 2 because estimating two trajectories from four points is ambiguous.
As one would expect, the approach that kept the entire history in the active set performed the best, successfully tracking both objects 68% of the time (see figure 3-4 and table 3.1.) On the other hand, the routine that kept only the last four frames (see figure 3-5) performed significantly worse, successfully tracking only 55% of the frames. It is worth noting that the implementation of the batch processor was extremely robust to the replicated trajectory. It was rarely fooled more than temporarily by the replicated trajectory in the same way as the Kalman filter. Consequently, the difference in the DKF implementations lies in their ability to track objects through the occluded intersection and recover the correct path in the end. Thus, the baseline for an algorithm that randomly guessed is 50% and, therefore, the difference between 55% and 68% is much larger than it might seem at first glance.

The two intelligent DKF schemes demonstrated a significant improvement. Selecting frames based on the Euclidean distance metric kept frames that gave more information about the occluded intersection point (see figure 3-6.) On the other hand, it did not always keep the most logical frames. For example, in the blue example trajectory, the first frame is kept for a long time. (This is evidenced by the rainbow colored partial paths that can be seen reaching from the first observation past the intersection point.) The KL-divergence metric made more reasonable decisions - the partial paths show that it generally kept more recent observations, as one would expect. The success rates corroborate the intuitive notions, with a 60% success rate for the Euclidean metric and a 64% success rate for the KL-divergence metric.

Overall, the results from numerical optimization were quite promising. On this type of data set, at least, DKF with the KL-divergence metric stands to provide a significant enhancement over existing algorithms.

### 3.4 Condensation DKF

For our final tests, we implemented the Condensation variant of DKF as described in chapter 2. In the Condensation filter, the performance difference was less obvious. The raw success rate was about the same, though the DKF variant generally
produced smoother, more accurate trajectories (the results in table 3.1 do not reflect this because they use a boolean quantity for success.)

The discrepancy is likely due to the incomparability of particle counts and the method of measuring success. The added dimensionality in the DKF variant requires more particles to accurately handle the distribution; however, it is not clear that this makes a fair comparison. For the purposes of this thesis, we used the same number of particles (5000) for both the standard and DKF versions of Condensation.
Figure 3-6: DKF using four frames and the Euclidean distance metric. The dashed red and blue lines represent the trajectories estimated by the algorithm and the rainbow colored lines represent the optimal partial trajectories at each timestep. With this metric, the algorithm may elect to keep older observations, giving rise to longer partial trajectories than when only the last four frames were kept (see figure 3-5.) The Euclidean distance metric likes to keep the first observation around for a long time, which generates the long rays seen around the intersection.

3.5 Results

The results (see table 3.1) were generally as anticipated. The simple benchmark approaches could not reliably handle the occlusions and replications. These were handled much more effectively by batch DKF implementations. Success across the different DKF implementations varied as expected, and the KL-divergence metric yielded performance amazingly close to the version that kept the entire history.

Unfortunately, the Condensation DKF version did not perform as well as expected, likely because of incomparable particle counts. In general, the DKF version produced
Figure 3-7: DKF using four frames and the KL-divergence metric. The solid red and blue lines represent the trajectories estimated by the algorithm and the rainbow colored lines represent the optimal partial trajectories found at each timestep. Like the Euclidean distance metric, the KL-divergence metric elects to keep observations before the occlusion; however, it is more intelligent and does not simply keep the first observation.

smoother, more accurate trajectories but was still susceptible to getting lost when run with comparatively low particle counts.

We defined a successful run as one in which the data association was correct for 80% of the run. When the batch methods erred, they handled the occlusion but switched trajectories at the intersection. Consequently, they were well above or below the 80% cutoff.

In contrast, the Kalman filter and condensation algorithms were closer to the cutoff. The trajectories would frequently merge around the occluded intersection and split again a while later.
Figure 3-8: Condensation DKF using four frames and the KL divergence metric. The solid red and blue lines represent the trajectories estimated by the algorithm. Compared to the standard Condensation filter on the same data in figure 3-3, the DKF implementation has less of a tendency to get confused as it approaches the blue trajectory and is able to pick up the red object when it reappears.

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Kalman Filter</td>
<td>24%</td>
</tr>
<tr>
<td>Condensation</td>
<td>50%</td>
</tr>
<tr>
<td>All-frames</td>
<td>68%</td>
</tr>
<tr>
<td>Most recent</td>
<td>55%</td>
</tr>
<tr>
<td>Euclidean</td>
<td>60%</td>
</tr>
<tr>
<td>KL-Divergence</td>
<td>64%</td>
</tr>
<tr>
<td>DKF (</td>
<td>AS</td>
</tr>
<tr>
<td>Condensation-DKF (</td>
<td>AS</td>
</tr>
</tbody>
</table>

Table 3.1: The DKF approaches performed well, with performance among the DKF variants commensurate with expectations.
Chapter 4

Conclusion and Further Work

The simulation results demonstrated that batch DKF using the KL-divergence metric in the simulation environment is a good approximation of full batch processing, even with an active set of only four items. It was able to successfully track objects through occlusions, intersections, and duplications that more frequently fooled other algorithms.

The key to the success of DKF is its selection of the most relevant frames. By using the KL-divergence to compare posterior distributions, it saves the locally optimal set of key frames from step to step and uses those frames to find the optimal estimate.

Unfortunately, the Condensation DKF implementation demonstrated weaker results than the batch implementation. It generally produced smoother estimates that were closer to the ground truth than the traditional Condensation algorithm, but it did not demonstrate the extra resilience to occlusions and intersections that the batch version did.

One of the key benefits to our approach is its simple generality. The algorithm requires minimal assumptions about the structure of the input. One need only specify the posterior function \( \alpha \) to track in entirely different contexts. In a more general way, this methodology can be easily applied to any problem involving state estimation from observations with minimal effort.

The next stage of this work would be to test DKF in a real system and to try a faster implementation. A good source of test data would be a soccer match or traffic
scene, where many identical targets move around the field with fairly complicated dynamics.

One problem with the current implementation is its speed. The Condensation DKF implementation requires about 20 seconds to process each frame on an Intel 2.6GHz Core2 processor when run with 5000 particles. In large part, this is because it was implemented inefficiently in Matlab. As a result, the runtime is impractical for any real-time application. To legitimately evaluate Condensation-DKF, it should be implemented in a manner that ensures reasonable performance.

Finally, on a theoretical level, it would be interesting to see if any process structure will give a convenient, closed-form solution for optimization of \( \hat{x}_t \) and \( AS'_t \).

Overall, while our approach has not yet reached practicality, it is a useful framework that may be easily and powerfully applied to many different situations to improve tracking performance.
Appendix A

Matlab Code

This appendix contains the Matlab code used for the simulation. The code used to generate the datasets follows immediately in section A.1. The code for the testing framework is in section A.2. After that, section A.3 contains code for computing the posterior probabilities from observations. Sections A.4, A.5, and A.6 contain the implementations of the batch, Condensation, and Kalman algorithms respectively, and, finally, section A.7 contains for various utility functions.

A.1 Generating Data

genData_3D.m

% Generates ground truth data with an occluded intersection:
% - two, intersecting trajectories
% - one trajectory is occluded around the intersection
% - the other trajectory exhibits multiple data points for a given
% truth point.
%
%
% gt - the ground truth data
% obs - the "observations" of the ground truth
% tstamps - the time stamps for the observations
% init0 - the starting locations of the trajectories
function trajectory = genData_3D(L, dyn_mode)


tstamps = generateTimestamps(L);

x10 = [0 5 2]'; % the location of intersection for the two trajectories
param1 = [0.2 0 0]'; % the velocity of the first trajectory at the intersection
param2 = [0.1 -0.1 .1]'; % the velocity of the second trajectory at the intersection

gt1 = simulateTrajectory(dyn_mode, L, x10, param1); % ground truth
gt2 = simulateTrajectory(dyn_mode, L, x10, param2); % ground truth

gt = [gt1 gt2];

[dim one] = size(x10);
num_targ = 2;

obs1 = addNoise(gt1, 0.1); % observations
obs2 = addNoise(gt2, 0.1); % observations

init0 = [obs1{1};obs2{1}];

o1split = 4;%ceil(3*rand());
o1splitlvl = .15*rand();
o2occs = ceil(L/5)+ceil(L/3*rand());
o2occf = ceil(L/3*rand())+o2occs;

data_stat = [o1split o1splitlvl o2occs o2occf];

obs1f = split(obs1,L,o1splitlvl,o1split);%occlude(obs1, L, 10, 15);
ob2f = occlude(obs1, L, o2occs, o2occf);%obs2;
obs = mergeObservations(obs1, obs2);

% obs = addBackgroundNoise(obs, .25, -5, 5, 0, 10);

trajectory = struct('L', L, 'dim', dim, 'num_targ', num_targ,
    'dyn_mode', dyn_mode, 'gt', {gt}, 'obs', {obs}, 'tstamps', {tstamps},
    'init0', {init0}, 'data_stat', {data_stat});
end

genData_NM.m

function trajectory = genData_NM(L, dyn_mode)

tstamps = generateTimestamps(L);

x10 = [0 5]'; % the location of the midpoint of the first trajectory
x20 = [0 6.25]'; % the location of the midpoint of the second trajectory
param1 = [0.2 0]'; % the velocity of the first trajectory at the intersection

[dim one] = size(x10);
num_targ = 2;

gt1 = simulateTrajectory(dyn_mode, L, x10, param1); % ground truth
gt2 = simulateTrajectory(dyn_mode, L, x20, param1); % ground truth
gt = [gt1 gt2];

obs1 = addNoise(gt1, 0.1); % observations
obs2 = addNoise(gt2, 0.1); % observations
init0 = [obs1{1}; obs2{1}];
% o1split = 4;%ceil(3*rand());
% o1splitlvl = .15*rand();
o2occs = ceil(L/5)+ceil(L/3*rand());
o2occf = ceil(L/3*rand())+o2occs;
%
% data_stat = [o1split o1splitlvl o2occs o2occf];

obs1f = obs1;%split(obs1,L,o1splitlvl,o1split);%occlude(obs1, L, 10, 15);
obs2f = occlude(obs2, L, o2occs, o2occf);%obs2;

obs = mergeObservations(obs1f,obs2f);

%obs = addBackgroundNoise(obs,.25,-5,5, 0, 10);

data_stat = [];

trajectory = struct('L', L, 'dim', dim, 'num_targ', num_targ,...
    'dyn_mode', dyn_mode, 'gt', {gt}, 'obs', {obs}, 'tstamps', {tstamps},...
    'init0', {init0}, 'data_stat', {data_stat});

end

genData_OI.m

% Generates ground truth data with an occluded intersection:
% - two, intersecting trajectories
% - one trajectory is occluded around the intersection
% - the other trajectory exhibits multiple data points for a given
% truth point.
%
function trajectory = genData_OI(L, dyn_mode)

% gt - the ground truth data
% obs - the "observations" of the ground truth
% tstamps - the time stamps for the observations
% init0 - the starting locations of the trajectories

tstamps = generateTimestamps(L);

x10 = [0 5]'; % location of intersection for the two trajectories
param1 = [0.2 0]'; % velocity of the first trajectory at intersection
param2 = [0.1 -0.1]'; % velocity of the second trajectory at intersection

[dim one] = size(x10);
num_targ = 2;

gt1 = simulateTrajectory(dyn_mode, L, x10+[0 0]', param1); % ground truth
gt2 = simulateTrajectory(dyn_mode, L, x10, param2); % ground truth
gt = [gt1 gt2];

obs1 = addNoise(gt1, 0);%0.1); % observations
obs2 = addNoise(gt2, 0);%0.1); % observations

init0 = [obs1{1};obs2{1}];

o1split = 4;%ceil(3*rand());
o1splitlvl = .15*rand();
o2occs = ceil(L/6)+ceil(L/4*rand());
o2occf = ceil(L/3*rand())+o2occs;

data_stat = [o1split o1splitlvl o2occs o2occf];
obs1f = obs1;%split(obs1,L,o1splitlv1,o1split);%occlude(obs1, L, 10, 15);
obs2f = occlude(obs2, L, o2occs, o2occf);%obs2;

obs = mergeObservations(obs1f,obs2f);

% obs = addBackgroundNoise(obs,.25,-5,5, 0, 10);

trajectory = struct('L', L, 'dim', dim, 'num_targ', num_targ,...
    'dyn_mode', dyn_mode, 'gt', {gt}, 'obs', {obs}, 'tstamps', {tstamps},...
    'init0', {init0}, 'data_stat', {data_stat});

end

generateTimestamps.m

function T = generateTimestamps(N)
for i=1:N,
    T(i) = i;
end

addBackgroundNoise.m

function obs_out = addBackgroundNoise(obs, level, xmin,xmax,ymin,ymax)
[one L] = size(obs);
obs_out = obs;

xdelta = xmax-xmin;
ydelta = ymax-ymin;

for i=3:L
    cur_obs = obs{i};
num_pts = floor(abs(2*level*randn()));
pts = ones(num_pts,1)*[xmin ymin] + ...
    rand(num_pts,2).*(ones(num_pts,1)*[xdelta ydelta]);
cur_obs = [cur_obs;pts];
obs_out{i} = cur_obs;
end
end

addNoise.m

% add noise to data
function Y = addNoise(X, noise_magn)
[N, dimX] = size(X);
Xnoisy = X + noise_magn*randn(size(X));
for i=1:N,
    Y{i} = Xnoisy(i,:);
end

loadDatasetFile.m

function dataset = loadDatasetFile(name)
load([name '.mat']);
end

makeDatasetFile.m

function dataset = makeDatasetFile(name)
N = 50;
L = 45;
dim = 2;
num_targ = 2;
dyn_mode = 2;
dataset = struct('L', cell(1,N), 'dim', cell(1,N), 'num_targ', cell(1,N),
    'dyn_mode', cell(1,N), 'gt', cell(1,N), 'obs', cell(1,N),
    'tstamps', cell(1,N), 'init0', cell(1,N), 'data_stat', cell(1,N));

for i=1:N
    trajectory = genData_OI(L, dyn_mode);
    dataset(i) = trajectory;
end

save(['name '.mat'],'dataset');

end

occlude.m

function obs_out = occlude(obs_in, L, t_start, t_end)
obs_out = obs_in;
[a b] = size(obs_in{t_start});
for i=t_start:t_end
    obs_out{i}=ones(0,b);
end
end

simulateTrajectory.m

% returns the entire trajectory of a particle in 2D
% mode == 0 -> linear..param = velocity
% mode == 1 -> continuous velocity..param = initial velocity
% mode == 2 -> continuous velocity, through x0 at time L/2..param =
% velocity at step L/2
function X = simulateTrajectory(mode, L, x0, param)
switch mode
case 0
    curr_x = x0;
    v = param;
    for i=1:L,
        X(i,:) = curr_x;
        curr_x = curr_x + v;
    end

case 1
    curr_x = x0;
    v = param;
    max_a = 0.02;% *ones(size(v));
    for i=1:L,
        X(i,:) = curr_x;
        a = max_a*randn(size(v));
        v = v + a;
        curr_x = curr_x + v;
    end

case 2
    l = ceil(L/2);
    before = simulateTrajectory(1, l, x0, -param);
    after = simulateTrajectory(1, L-l+1, x0, param);
    before = before(end:-1:2,:);
    X = [before; after];
end
split.m

function obs_out = split(obs_in, L, noise_magn, max_replicas)
obs_out = {};
for i=1:L
    cur_obs = obs_in{i};
    num_reps = ceil(max_replicas * rand());
    cur_obs = repmat(cur_obs, num_reps, 1);
    cur_obs = cur_obs + noise_magn*randn(size(cur_obs));
    obs_out = [obs_out, cur_obs];
end
end

A.2 Testing Framework

batch_sim.m

function [error bad_trial results] = batch_sim(dataset_file,output_tag,indices)

dataset = loadDatasetFile(dataset_file);
[one N] = size(dataset);

if isempty(indices)
    indices = 1:N;
end
[one num_trials] = size(indices);

error = [];
bad_trial = [];
%% This allows me to comment out a particular test without generating
%% errors.
Yb = []; Yib = []; Yk = []; Yc = [];

Eb = []; Eib = []; Ek = []; Ec = [];

results = struct('dataset',cell(1,num_trials),
                 'index',cell(1,num_trials),
                 'data',cell(1,num_trials),
                 'Yb',cell(1,num_trials),
                 'Yib',cell(1,num_trials),
                 'Yk',cell(1,num_trials),
                 'Yc',cell(1,num_trials),
                 'error',cell(1,num_trials));

trial_num = 1;
for i=indices
    tic;
    disp(i);
    fig = figure(i); clf;

    trajectory = dataset(i);
    drawData(trajectory);

    gt = trajectory.gt;
    obs = trajectory.obs;
    tstamps = trajectory.tstamps;
init0 = trajectory.init0;
num_targ = trajectory.num_targ;

disp('Batch');
Yb = runIntelBatch(obs, tstamps, init0, 0);
Eb = calc_error(gt,Yb);

display('Intelligent Batch');
Yib = runIntelBatch(obs, tstamps, init0, 3);
Eib = calc_error(gt,Yib);

disp('Kalman');
Yk = runKalman(obs, tstamps, init0);
Ek = calc_error(gt,Yk);

disp('Condensation');
Yc = runCondensation(obs, tstamps, init0);
Ec = calc_error(gt, Yc);

E = [Eb Eib Ek Ec];
bad = E>.55;
bad = add_cols(bad,num_targ);
bad = bad>0;
disp(bad);
error = [error;E];
bad_trial = [bad_trial;bad];

filename = sprintf([dataset_file '\%i_' output_tag],i);
saveas(fig,filename,'jpg');
close;
result = struct('dataset',{dataset_file},'index',{i},... 'data',{trajectory},'Yb',{Yb},'Yib',{Yib},'Yk',{Yk},... 'Yc',{Yc},'error',{error});
results(trial_num) = result;
trial_num = trial_num+1;
toc;
end

% Compute some statistics
bad_count = sum(bad_trial);
bad_count

save([dataset_file '\" output_tag '.mat'], 'results', 'bad_count');
end
calc_error.m

function error = calc_error(gt, Y)
delta = abs(Y-gt);

delta = delta.*delta;

dist = add_cols(delta,2); %REVISE FOR GENERIC
dist = sqrt(dist);
error = mean(dist);
drawData.m

function drawData(trajectory)
style = ['b' 'r' 'g'];
dim = trajectory.dim;
for i=1:trajectory.num_targ
    drawTrajectory(trajectory.gt(:,((i-1)*dim+1:i*dim)), ...
        ['x' style(i)]); hold on;
end

drawObservation(trajectory.obs, '.k'); hold on;
end

drawObservation.m

% draw complete trajectory
function drawObservation(obs, sym)
    [n,m] = size(obs);
    [junk dim] = size(obs{1});
    x = [];
    for i=1:m,
        x = [x; obs{i}];%obs{i}(1) obs{i}(2];
    end
    if dim == 2
        plot(x(:,1), x(:,2), sym);
    else
        plot3(x(:,1), x(:,2), x(:,3), sym);
    end

drawTrajectory.m

% draw complete trajectory
function drawTrajectory(X, sym)
    [N dim] = size(X);
    if dim == 2
        plot(X(:,1),X(:,2), sym);
else
    plot3(X(:,1),X(:,2),X(:,3), sym);
end

mean_error.m

function e = mean_error(gt, est)
e = abs(gt-est);
e = e.*e;
e = e*[1;1];
e = sqrt(e);
e = mean(e);

plotRun.m

function plotRun(results, runs, algorithms)

for run=runs
    figure(run);
    result = results(run);
    data = result.data;
    gt = data.gt;
    obs = data.obs;
    Y = {result.Yk result.Yc result.Yb result.Yib};
    for i=1:length(obs)
        plot(obs{i}(:,1),obs{i}(:,2),'k.'); hold on;
    end
    plot(gt(:,1),gt(:,2),'bx'); hold on;
    plot(gt(:,3),gt(:,4),'rx'); hold on;
    for i=1:4
        if algorithms(i)
            plot(Y{i}(:,1),Y{i}(:,2),'b-'); hold on;
        end
    end
end
plot(Y{i}(:,3),Y{i}(:,4),'r-'); hold on;
end
end
end

plot_frame.m

function plot_frame(gt, obs, Yb, Yk, Yc, frame)

if (frame>1)
    drawTrajectory(gt(1:frame-1,1:2), 'xk'); hold on;
    drawTrajectory(gt(1:frame-1,3:4), 'xk'); hold on;
    drawObservation(obs(1:frame-1), '.k'); hold on;

    styles = [ 'b-';'r-';'g-'];
    for i=0:(2-1)
        drawTrajectory(Yb(1:frame-1,(2*i+1):(2*(i+1))), 'k-');
        drawTrajectory(Yb(frame-1:frame,(2*i+1):(2*(i+1))), styles(i+1,:));
    end
end

drawTrajectory(gt(frame,1:2), 'xb'); hold on;
drawTrajectory(gt(frame,3:4), 'xr'); hold on;
drawObservation(obs(frame), '.g'); hold on;
end

A.3 Posterior Probabilities

compute_dStates.m
targetLogLikelihood.m

function [v, logll] = targetLogLikelihood(state, obs)

num_obs, dim] = size(obs);
[junk, N] = size(state);
num_targ = N/dim;

%obs_rep = repmat(obs,1,num_targ);
obs_rep = zeros(num_obs,N);
for i=0:(num_targ-1)
    obs_rep(:,(dim*i+1):(dim*(i+1)))=obs;
end

%state_rep = repmat(state,num_obs,1);
state_rep = state(ones(1, num_obs),:);
err = obs_rep - state_rep;

dist = err.*err;
dist = add_cols(dist,dim);
[min_err idx_vec] = min(dist,[],2);
easy_state = reshape(state,dim,num_targ)';
err = abs(obs - easy_state(idx_vec,:));
v = reshape(err,1,num_obs*dim);

logll = sum(v.*v);%logll = dot(v,v);%Substitution made for speed

totalPosterior.m

% defines the total posterior on all the data
% - allStates is a [Nxdim] matrix
% - allObs is an array of size N
% - timestamps: list of times associated with the observations
function post = totalPosterior(allStates, allObs, timestamps, mode_dynmodel)
    [N, dimStates] = size(allStates);
    % compute likelihood for current sta/obs
    log_llk = 0;
    post = [];
    for idx=1:N,
        sta = allStates(idx,:);
        obs = allObs{idx}; % set current observation
        [v, log_llk(idx)] = targetLogLikelihood(sta, obs);
        post = [post v];
    end

    % - compute time derivates of states
    if (mode_dynmodel==0)
        avg_v_rep = repmat(avg_v,N-1,1);
        err = dStates-avg_v_rep;
    elseif (mode_dynmodel==1)
        err = dStates(1:N-2,:)-dStates(2:N-1,:);
    end
    dyn = reshape(err,1,numel(err));
    post = [post 3*dyn];

%above constant was 2 for best fit to curve, 20 provided best tracking
% when square roots were not taken on likelihood, ~60 when taking sqrt
### A.4 Batch DKF

batchPredict.m

% Selects an initialization point for the maximization.
% y is the optimum found at the last step
function init = batchPredict(num_targ, allObs, tstamps, mode_dynmodel, init_obs)
  % [a b] = size(y);
  % if a>1
  %   prediction = 2*y(a,:)-y(a-1,:);
  % else
  %   prediction = y(a,:);
  % end
  % init = [y;prediction];

  % We use the viterbi algorithm...ish...

  init_obs = allObs{1};

  [junk dim] = size(init_obs);
  total_dim = num_targ*dim;
  [one N] = size(allObs);
  [num_end_obs dim] = size(allObs{end});

  init_obs = [init_obs; NaN*ones(1,dim)];
  [num_init_obs dim] = size(init_obs);

  init_perms = nchoosek(1:num_init_obs,num_targ);
  [num_init_perms dim] = size(init_perms);
  opt = cell(1,num_init_perms);
  logic_opt = cell(1,num_init_perms);
for i=1:num_init_perms
    cur_start = init_obs(init_perms(i,:),:);
    opt{i} = reshape(cur_start',1,total_dim);
    logic_opt{i} = init_perms(i,:);
end

for i=2:N
    cur_obs = [allObs{i};NaN*ones(1,dim)];

    [num_obs dim] = size(cur_obs); % the dim assignment is redundant
    [one num_old_options] = size(opt);
    num_new_options = npermk(num_obs,num_targ);
    options = vpermk(1:num_obs,num_targ);
    new_opt = cell(1,num_new_options);
    new_logic_opt = cell(1,num_new_options);

    err = zeros(1,num_old_options);
    for new_choice=1:num_new_options
        new_obs_idx = options(new_choice,:);
        cur_state = cur_obs(new_obs_idx,:);
        cur_state = reshape(cur_state',1,total_dim);
        new_state = cell(1,num_old_options);
        for old_choice=1:num_old_options
            old_state = opt{old_choice};
            save_state = [old_state;cur_state];
            for j=1:num_targ
                targ_idx = (dim*(j-1)+1):(dim*j);
                if any(isnan(old_state(end,targ_idx)))
                    if any(isnan(save_state(targ_idx)))
                        new_state{old_choice} = new_state{old_choice}.
                    end
                end
            end
        end
    end
end
% occluded in the last frame of old_state
occ_start = find(isnan(old_state(:,targ_idx(1))),1);
save_state(:,targ_idx) = optimizeOcclusion(...
old_state(1:occ_start-1,targ_idx),...
i-occ_start,cur_state(1,targ_idx));
end
end
test_state = save_state;
for j=1:num_targ
targ_idx = (dim*(j-1)+1):(dim*j);
% note that the "end" could be a "1" since cur_state has
% one row
if any(isnan(cur_state(end,targ_idx)))
% occluded in the last frame of old_state
occ_start = find(isnan(test_state(:,targ_idx(1))),1);
prepare_state(:,targ_idx) = optimizeOcclusion(...
prepare_state(1:occ_start-1,targ_idx),...
i-occ_start+1,zeros(0,dim));
end
end

new_state{old_choice} = save_state;
e = totalPosterior(test_state,allObs(1:i),tstamps(1:i),1);
err(old_choice) = sum(e.*e);
end
[min_val min_idx] = min(err);
new_opt{new_choice} = new_state{min_idx};
new_logic_opt{new_choice} = [logic_opt{min_idx};new_obs_idx];
end
opt = new_opt;
logic_opt = new_logic_opt;
end

[b] = size(opt);
init = cell(1,b);
for i=1:b
    tmp = opt{i};
    for j=1:num_targ
        targ_idx = (dim*(j-1)+1):(dim*j);
        if any(isnan(tmp(end,targ_idx)))
            % occluded in the last frame of old_state
            occ_start = find(isnan(tmp(:,targ_idx(1))),1);
            tmp(:,targ_idx) = optimizeOcclusion(...
                tmp(1:occ_start-1,targ_idx),N-occ_start+1,zeros(0,dim));
        end
    end
    init{i} = tmp;
end

bootstrapXDistribution.m

function X = bootstrapXDistribution(num_targ, mean, obs, timestamps, mode_dynmodel, init0)
    [N] = size(obs);
    num_samples = 10;

    % The following parameters control how the multi-dimensional histogram is
    % created:
    % grain - the size of a bin (in a single dimension)
% window - the total distance covered by the histogram in a single
% dimension
% num_steps - the number of discrete values possible in a given dimension
grain = .1;
window = 1;
um_steps = window/grain;

[junk total_dim] = size(mean);

offset = mean(end,:)-ones(1,total_dim)*(window/2);
almost_zero = 0.0001;
 % ensure that each location has some small probability so that the
 % divergences can be compared.
X = almost_zero*ones(num_steps*ones(1,total_dim));
% create a multiplication matrix for finding the index
mul_mat = exp(log(num_steps)*(0:total_dim-1));

for i=1:num_samples
    sample = obs;
    for j=1:N
        sample{j} = sample{j}+0.01*randn(size(sample{j}));
    end
    init = batchPredict(num_targ, sample, timestamps, mode_dynmodel, init0);
y = find_map(num_targ, sample, timestamps, mode_dynmodel, init);
dataPoint = y(end,:);
    % including grain/2 does not affect bucketing but ensures that
    % rounding does not make idx too large
    delta = min(max(dataPoint-offset,grain/2),window-grain/2);
    idx = ceil(delta/grain);
    idx = round(sum(mul_mat.*idx-mul_mat)+1);
X(idx) = X(idx)+1;
end

%normalize
X = X/N;

calcKLDivergence.m

function kl = calcKLDivergence(P,Q)
kl = 0;

num_elts = numel(Q);
% we use the fact that a matrix can be indexed as a vector to allow us to
% index a matrix of arbitrary dimension.
for i=1:num_elts
p = P(i);
q = Q(i);
kl = kl + p*log(p/q)/log(2);
end
end

calcMeanCvKLDivergence.m

estimateBatch.m

function Y = estimateBatch(obs, tstamps, init0, criticalObsMode)

[one total_steps] = size(obs);
[junk dim] = size(obs{1});
[one b] = size(init0);
num_targ = b/dim;
total_obs = {obs{1}};
total_time = tstamps(1);
Y = init0;
for step=2:total_steps,
    %step
    % --- new observation...
    newobs = obs{step};
    newtime = tstamps(step);
    % --- list of obervations to use
    total_obs = {total_obs{:} newobs};
    total_time = [total_time; newtime];
    % --- prediction
    init0 = batchPredict(init0);
    % --- OPTIMIZATION
    %[Y,hh,Htot] = find_map(2, total_obs, total_time, 1, init0);
    y = find_map(num_targ, total_obs, total_time, 1, init0);
    Y = [Y;y(end,:)];
    init0 = Y;
end

drawTrajectory(y(:,1:dim),'g-');
drawTrajectory(y(:,dim+1:2*dim),'g-');

estimateIntelBatch.m

function Y = estimateIntelBatch(obs, tstamps, init0, criticalObsMode)

active_count = 6;

[one total_steps] = size(obs);
[junk dim] = size(obs{1});
[one data] = size(init0);
num_targ = data/dim;

active_obs = {obs{1}};
active_time = tstamps(1);
Y = init0;
old_y = init0;
init = init0;

tmp_style = 'gmy';

for step=2:total_steps,
    disp(step);
    % --- new observation...
    newobs = obs{step};
    newtime = tstamps(step);
    % --- list of observations to use
    active_obs = {active_obs{:} newobs};

    active_time = [active_time; newtime];
    % --- prediction
    tmp_init = init0;%= reshape(init0',dim,num_targ)';
    init = batchPredict(num_targ, active_obs, active_time, 1, tmp_init);
    %init = {[init0;rand(step-1,num_targ*dim)]};
    % --- OPTIMIZATION
    %[Y, hh, Htot] = find_map(2, total_obs, total_time, 1, init);
    [y J init_idx] = find_map(num_targ, active_obs, active_time, 1, init);
    %y = find_map2(num_targ, active_obs, active_time);
y = matchTrajectories(num_targ,old_y,y);
Y = [Y;y(end,:)];

color = tmp_style(mod(step,3)+1);
drawTrajectory(y(:,1:dim),[color ' - ']);
drawTrajectory(y(:,dim+1:2*dim),[color ' - ']);

[one num_obs] = size(active_obs);
if num_obs>active_count
    switch criticalObsMode
        case 0
            % do nothing. This is normal batching.
            idx = 0;
        case 1
            % remove the earliest observation
            idx = 1;
        case 2
            % Compare Euclidian distances
            dist = zeros(1,active_count);
            for i=1:num_obs-1,
                [obs_red, time_red, y_red] = ...
                    removeObservation(i,active_obs,active_time,y);
                Yred = ...;
                find_map(num_targ, obs_red, time_red, 1, {y_red});
                delta = Yred(end,:)-y(end,:);
                delta = delta.*delta;
                delta = add_cols(delta,2);
                dist(i) = sum(sqrt(delta));
        end
    end
end
end
[d idx] = min(dist);

case 3
%
% Compare KL Divergence...this is the first try

dist = zeros(1,active_count);
Q = bootstrapXDistribution(num_targ, y, active_obs,...
   active_time, 1, init0);
for i=1:num_obs-1,
    [obs_red, time_red, y_red] = ...% removeObservation(i,active_obs,active_time,y);
P = bootstrapXDistribution(num_targ, y_red,obs_red,...
   time_red,1, init0);
dist(i) = calcKLDivergence(P,Q);
end
[d idx] = min(dist);

end

case 4
%
% Use mean/covariance for KL divergence

dist = zeros(1,active_count);
[Q_mean Q_Cv] = jacobianXDistribution(y,J);
for i=1:num_obs-1
    [obs_red, time_red, init_red] = ...
        removeObservation(i,active_obs,active_time,y);
    [y_red J_red] = ...
        find_map(num_targ, obs_red, time_red, 1, init_red);
    [P_mean P_Cv] = jacobianXDistribution(y_red,J_red);
dist(i) = ...
        calcMeanCvKLDivergence(P_mean,P_Cv,Q_mean,Q_Cv);
end
[d idx] = min(dist);
end
[active_obs, active_time, y] = ...
    removeObservation(idx,active_obs,active_time,y);

active_time
end

old_y = y;
pause(0.01);
end

find_map.m

% find synchrony between x1 and x2
% Ntarg: number of targets
% returns H : cov of y ??? Maybe not..
% Hlast: cov of last estimate ??? Maybe not..
function [y J init_idx] = ...
    find_map(Ntarg, allObs, timestamps, mode_dynmodel, init)

[one N] = size(allObs);
[junk dim] = size(allObs{1});
[one num_init] = size(init);

% USING LSQNONLIN and Trying to avert local minima...
y_candidates = cell(size(init));
J_candidates = cell(size(init));
err=zeros(size(init));
options = optimset('Display', 'off', 'LargeScale', 'off', ...
    'LargeScale', 'on', 'TolX', 0.0001); % Turn off Display
for i=1:num_init
    allInit = mat2vec(init{i});
    [y_vec resnorm residual exitflag output lambda J] = ...
        lsqnonlin(@ppfunc, allInit, [], [], options);

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y = vec2mat(y_vec,Ntarg*dim);
y_candidates{i} = y;
J_candidates{i} = J;
p = totalPosterior(y, allObs, timestamps, mode_dynmodel);
p = sum(p.*p);
err(i) = p;
end

[min_err min_idx] = min(err);
y = y_candidates{min_idx};
J = J_candidates{min_idx};
init_idx = min_idx;

function y = ppfunc(st_vec)
st_mat = vec2mat(st_vec,Ntarg*dim);
y = totalPosterior(st_mat, allObs, timestamps, mode_dynmodel);
end
end

matchTrajectories.m

% This function matches the trajectories in new_y to the trajectories in
% old_y.
function y = matchTrajectories(num_targ, old_y, new_y)
[N total_dim] = size(old_y);
dim = total_dim/num_targ;

new_perms = perms(1:num_targ);
[num_perms junk] = size(new_perms);
err = zeros(1,num_perms);
for i=1:num_perms
    idx_vec = new_perms(i,:);
    idx_vec = ...  
        repmat(dim*(idx_vec-ones(1,num_targ)),dim,1)+repmat((1:dim)',1,num_targ);
    idx_vec = reshape(idx_vec,1,total_dim);

    cur_y = new_y(1:end-1,idx_vec);
    dist = cur_y-old_y;
    dist = dist.*dist;
    dist = add_cols(dist,dim);
    dist = sqrt(dist);
    err(i) = sum(sum(dist));
end

[min_err min_idx] = min(err);
idx_vec = new_perms(min_idx,:);
idx_vec = ... 
        repmat(dim*(idx_vec-ones(1,num_targ)),dim,1)+repmat((1:dim)',1,num_targ);
idx_vec = reshape(idx_vec,1,total_dim);
y = new_y(:,idx_vec);
end

optimizeOcclusion.m

function y = optimizeOcclusion(pre_y, num_steps_occ, post_y)

    [num_pre total_dim] = size(pre_y);
    [num_post total_dim] = size(post_y);

if num_pre>0 && num_post>0
    start_loc = pre_y(end,:);
end_loc = post_y(1,:);

delta = end_loc-start_loc;
step = delta/(num_steps_occ+1);
else if num_pre>1
    start_loc = pre_y(end,:);
    step = start_loc-pre_y(end-1,:);
else if num_pre>0
    start_loc = pre_y(end,:);
    step = zeros(1,total_dim);
else if num_post>1
    % this should never be reached
    disp('ERROR: unreachable code reached!');
else if num_post>0
    start_loc = post_y(1,:);
    step = zeros(1,total_dim);
else
    start_loc = zeros(1,total_dim);
    step = zeros(1,total_dim);
end
end
end

occ_y = repmat(step,num_steps_occ,1);
occ_y(1,:) = occ_y(1,:)+start_loc;
occ_y = cumsum(occ_y);
\[ y = \text{[pre}_y; \text{occ}_y; \text{post}_y]; \]
end

**runBatch.m**

function \( Yb = \text{runBatch}(\text{obs}, \text{tstamps}, \text{init0}); \)
styles = ['b-';'r-';'g-'];

% Fix init0 so it is in the proper form
[num_traj dim] = size(init0);
batch_init0 = reshape(init0',1, num_traj*dim);

Yb = \text{estimateBatch}(\text{obs}, \text{tstamps}, \text{batch_init0}, 0);
for i=0:(num_traj-1)
    drawTrajectory(Yb(:,(dim*i+1):(dim*(i+1))), styles(i+1,:));
end

end

**runIntelBatch.m**

function \( Yib = \text{runIntelBatch}(\text{obs}, \text{tstamps}, \text{init0}, \text{criticalObsMode}); \)
\text{color} = 'brg';
styless = {'-' '--' '--' '-.' ':'};

% Fix init0 so it is in the proper form
[num_traj dim] = size(init0);
batch_init0 = reshape(init0',1, num_traj*dim);

Yib = \text{estimateIntelBatch}(\text{obs}, \text{tstamps}, \text{batch_init0}, \text{criticalObsMode});
for i=0:(num_traj-1)
    drawTrajectory(Yib(:,(dim*i+1):(dim*(i+1))),...
A.5 Condensation and Condensation DKF

calcCondLoc.m

function x = calcCondLoc(samples, sample_pi, dim)
[one k] = size(samples);
[num_targ x_dim] = size(samples{1});

mean = zeros(num_targ,dim);
data = zeros(k,num_targ*dim);
for i=1:k
    sample = samples{i};
data(i,:) = reshape(sample(:,x_dim-2*dim+1:x_dim-dim)',1,num_targ*dim);
    mean = mean+sample_pi(i)*sample(:,x_dim-2*dim+1:x_dim-dim);
% note that the weights are normalized, so the above gives the average
end
x = reshape(mean',1,num_targ*dim);

return;

% If we want to use mean-shift to find a peak...
dist_thresh = 0.001; % target error below which we stop
% err = 1;
% h = 1; %window radius
while err>dist_thresh
    x_rep = repmat(x,k,1);
    dist = (x_rep-data)/h;
    dist = sum(dist.*dist,2);
    g_vec = g(dist);
    g_vec = g_vec.*(sample_pi')*k;
    g_vec_rep = repmat(g_vec,1,num_targ*dim);
    m = sum(g_vec_rep.*data,1)/sum(g_vec);
    err = m-x;
    err = sum(err.*err);
    %disp(m-x);
    x = m;%x = .8*m+.2*x;
    %disp(m);
    %disp(x);
    %disp(err);
    %pause();
end

x = meanShift(data,sample_pi',dist_thresh,x);

function [x weight] = meanShift(data,weights,err_thresh,init)
    x = init;
    err = err_thresh+1;
    while err>err_thresh
        x_rep = repmat(x,k,1);
        dist = (x_rep-data);
        dist = sum(dist.*dist,2);
        g_vec = g(dist);
        g_vec = g_vec.*weights*k;
        g_vec_rep = repmat(g_vec,1,num_targ*dim);
x = sum(g_vec_rep.*data,1)/sum(g_vec);
end
end

function y = g(s)
y = 1/2*exp(-s/2);
end
end

cluster.m

function prob = cluster(dist_points, dist_weights, point)

[junk dim] = size(point);
[num_samples num_dist] = size(dist_weights);

% compute the probability that point came from each distribution

point_rep = ones(num_samples,1)*point;
point_rep = repmat(point_rep,1,num_dist);

coord_error = abs(point_rep-dist_points);
coord_error = coord_error .* coord_error;

sum_mat = [];
for i=1:num_dist
    sum_mat = blkdiag(sum_mat,ones(dim,1));
end
error = coord_error*sum_mat;

prob = dist_weights./(1+error);
prob = 1/num_samples*ones(1,num_samples)*prob;

end

condKLDivergence.m

function dKL = condKLDivergence(pi1,pi2)
dKL = sum(pi1.*log(pi1./pi2));

condPostProb.m

function W = condPostProb(samples, obs_set, tstamps, dim)
[num_targ x_dim] = size(samples{1});
%[num_obs data] = size(obs_set);
[one k] = size(samples);

W = zeros(1,k);
tmp = [];
for sample=1:k
    % Get the current sample and remove the velocity information
    state = samples{sample};
    state = state(:,1:end-dim);
    % Format for targetLogLikelihood
    x_state = state(:,1:2:end);
    y_state = state(:,2:2:end);
    state = [x_state(1,:)’ y_state(1,:)’ x_state(2,:)’ y_state(2,:)’];
    %state = reshape(state’, (x_dim-dim)/(dim), num_targ*dim);

    %disp(state)
    %w = totalPosterior(state, obs_set, tstamps’, 1);% For |AS|>=2
[junk w] = targetLogLikelihood(state,obs_set{1});% FOR |AS|=1
w = w.*w;
%disp(w)
%pause()
t2 = sum(w.*w);
tmp = [tmp;w t2 exp(1/t2)];
W(sample)=1/sum(w);%1/sum(w.*w);%exp(-1*sum(w));%1/w;
end

% Normalize weights in W
%disp(W);
W_sum = sum(W);
W = W./W_sum;
%disp(tmp);
%disp(W);

condPredict.m

function [samples tStamps] = condPredict(new_samples,tStamps,dim,new_t)
% [k, d] = size(samples);
% A=[1, 0, 1, 0; 0, 1, 0, 1; 0, 0, 1, 0; 0, 0, 0, 1];
% B=[0.05, 0, 0.1, 0; 0, 0.05, 0, 0.1; 0, 0, 0.1, 0; 0, 0, 0, 0.1];
% new_samples = A*(samples')+B*randn(4,k);
% new_samples = new_samples';

[num_targ x_dim] = size(new_samples{1});
[one k] = size(new_samples);
samples = cell(1,k);

tStamps = [tStamps new_t];
dynMat = blkdiag(eye(x_dim-2*dim),...
            [eye(dim) eye(dim) zeros(dim); zeros(dim) eye(dim) eye(dim)]);

for i=1:k
    sample = new_samples{i};
    %disp(sample);
    sample = sample*dynMat;
    %sample = sample(:,dim+1:end);%pulls off firs
    noise = 0.05*randn(num_targ,dim);
    sample = sample+[zeros(num_targ,x_dim-dim) noise noise];%[noise noise]
    %sample = sample(:,dim+1:end);%pulls off firs
    samples{i} = sample;
    %disp(sample);
end

estimateIntelCondensation.m

function Y = estimateIntelCondensation(obs, tstamps, init0)
    [num_targ dim] = size(init0);
    [one total_steps] = size(obs);

    active_count = 1;
    k = 150; % number of samples in condensation algorithm

    % initialize samples
    samples = cell(1,k);
    new_samples = cell(1,k);
sample_pi = ones(1,k)/k; % the likelihood associated with each sample

for i=1:k
    samples{i}=[[init0(:,1)';init0(:,2)'] 0.03*randn(num_targ,dim)];
end

sample_t = [1];
obs_set = obs(1);
Y = reshape(init0,1,num_targ*dim);

%figure(2);

for step=2:total_steps
    % compute the sum of the weights up to each index
    disp(step);

    obs_set = [obs_set obs{step}];
    new_t = tstamps(step);

    if mod(step,1)==0

        c = cumsum(sample_pi);
        weight_sum = sum(sample_pi);
        for sample=1:k
            % pick a random sample from the list
            %i = ceil(k*rand());
            r=weight_sum*rand();
            i=1;
            % search through until the sum of the weights is less than r
            while i <= k && c(i)<r
                i = i+1;
            end
        end
    end
end
end

new_samples{sample} = samples{i};
end

samples = new_samples;

end

[samples sample_t] = condPredict(samples, sample_t, dim, new_t);
[samples obs_set sample_t sample_pi] = filterSamples(samples, ...)
    sample_pi, obs_set, sample_t, dim, active_count);

% Compute the current location as the weighted average of the samples
x = calcCondLoc(samples, sample_pi, dim);
Y = [Y;x];

end

end

filterSamples.m

function [filtered_samples filtered_obs_set filtered_tStamps filtered_pi] = ... filterSamples(samples, sample_pi, obs_set, sample_t, dim, active_count)

DKF_mode = 0;

% First pass, just discard the last sample. This should work as before.

[a b] = size(samples{1});
active_set_size = b/dim-1;
if (active_set_size <= active_count)
    filtered_samples = samples;
    filtered_tStamps = sample_t;
    filtered_obs_set = obs_set;

    filtered_pi = condPostProb(samples, obs_set, sample_t, dim);
    return % if we haven’t stored enough frames yet, don’t discard any.
end

if DKF_mode == 0
    [filtered_samples filtered_obs_set filtered_tStamps] = ...
       sparsify(samples, obs_set, sample_t, dim, 1);
    filtered_pi = condPostProb(filtered_samples, ...
       filtered_obs_set, filtered_tStamps, dim);

    return
end

if DKF_mode == 1
    opt_x = calcCondLoc(samples, sample_pi, dim);
end

opt_score = 0;

for i=1:active_set_size-1
    [cur_filtered_samples cur_filtered_obs_set cur_filtered_tStamps] = ...
       sparsify(samples, obs_set, sample_t, dim, i);
    cur_sample_pi = condPostProb(cur_filtered_samples, ...
       cur_filtered_obs_set, cur_filtered_tStamps, dim);
    if (DKF_mode == 1)
cur_x = calcCondLoc(cur_filtered_samples, cur_sample_pi, dim);
delta = cur_x - opt_x;
cur_score = sum(delta.*delta);
cur_score = 1/cur_score;
else
  kl = condKLDivergence(cur_sample_pi, sample_pi);
cur_score = 1/kl;
end
if (cur_score > opt_score)
opt_score = cur_score;
  filtered_samples = cur_filtered_samples;
  filtered_obs_set = cur_filtered_obs_set;
  filtered_tStamps = cur_filtered_tStamps;
  filtered_pi = cur_sample_pi;
end
end
end

function [sparse_samples sparse_obs_set sparse_tStamps] = ...
  sparsify(samples, obs_set, tStamps, dim, frame_to_remove)
sparsify = cell(size(samples));
for i=1:length(samples)
  sample = samples{i};
  sparse_samples{i} = sample(:,...
                          [1:dim*(frame_to_remove-1) dim*frame_to_remove+1:end]);
end
sparse_obs_set = obs_set([1:frame_to_remove-1 frame_to_remove+1:end]);
sparse_tStamps = tstamps([1:frame_to_remove-1 frame_to_remove+1:end]);
end

def getComNearData.m

% Finds the COM of all observations near to a state estimate, x
% "near" is defined as within the covariance

% x is the current state estimate
% cov is the covariance vector for that estimate
% obs is a list of current observations

% Note that x is a column vector with the same number of rows
% as each data point has entries.

function new_x = getComNearData(x, cov, obs)
[a b] = size(obs);

obs = obs';

% replicate x
rep_x = x*ones(1,a);
% extract the diagonal from the covariance matrix
cov = cov.*eye(size(cov));
% replicate the elements of the diagonal accross each row
rep_cov = cov*ones(b,a);
% calculate the distance between x and each observation
diff = abs(obs-rep_x);
% compare the distance to the covariance
comp = diff < rep_cov;
% sum "good" entries
good_sum = ones(1,b)*comp;
% if all entries in a row are good (i.e. the sum is b), then we keep it
comp2 = (good_sum == b*ones(1,a));
% calculate number of good observations
num_good = comp2*ones(a,1);

if num_good==0
  new_x = x;
else
  % replicate comp2 to make a mask
  comp2_rep = ones(b,1)*comp2;
  % mask obs to keep only the good ones
  obs = comp2_rep.*obs;
  % compute the average x and y values
  new_x = (obs*ones(a,1))/num_good;
end

end

runCondensation.m

function Yc = runCondensation(obs, tstamps, init0);
styles = ['y-';'y-';'y-'];

[num_traject dim] = size(init0);
Yc = estimateIntelCondensation(obs, tstamps, init0');
for i=0:num_traject-1
  %Y = estimateCondensation(obs, tstamps,init0(i,:));
  drawTrajectory(Yc(:,(dim*i+1):(dim*(i+1))), styles(i+1,:));
  %Yc = [Yc Y];
A.6 Kalman Filter

estimateKalman.m

function Y = estimateKalman(obs, tstamps, init0)

[dim num_track] = size(init0);

[one total_steps] = size(obs);

F = [eye(dim) eye(dim); zeros(dim) eye(dim)];
H = [eye(dim) zeros(dim)];
R = 0.5*eye(dim);
Q = 0.1*[zeros(dim,2*dim);zeros(dim,dim) (-.25*ones(dim)+1.25*eye(dim))];

x = [init0; zeros(dim, num_track)];

P=eye(2*dim);
Y = init0(:)’;

for step=2:total_steps
    x=F*x;
    P=F*P*F’+Q;
    G=(P*(H’))/((H*P*(H’))+R);
    cur_obs = obs{step}’;

end

end
\begin{verbatim}
y = multi_kalman_obs(Y, x, cur_obs);

x = x + G * (y - H * x);
P = (eye(2*dim) - G * H) * P;

est = x(1:dim,:);
est = est(:)';
Y = [Y; est];
end

end

multi_kalman_obs.m

function obs = multi_kalman_obs(Y, x, cur_obs)

thresh = 0.25;

[dim, num_obs] = size(cur_obs);
junk, num_filters = size(x);
dist_points = x(1:dim,:);
dist_points = dist_points(:)';
dist_weights = ones(1, num_filters);

prob = [];
points = [];

% Determine the probability that a given observation belongs to a
% particular filter.
for i = 1:num_obs
    point = cur_obs(:,i)';
end
\end{verbatim}
points = [points;point];
dist_pr = cluster(dist_points,dist_weights,point);
prob = [prob; dist_pr];
end

% Now, make sure that each filter gets an observation
obs = zeros(dim,num_filters); % The sum of all observations given to each filter
obs_count = zeros(1,num_filters); % The number of observations given to each filter

for i=1:num_filters
    [p point_i] = max(prob(:,i));
    if p>thresh
        % If possible, give it a real observation
        point = points(point_i,:);
        prob = prob([1:(point_i-1) (point_i+1):end],:);
        points = points([1:(point_i-1) (point_i+1):end],:);
        obs(:,i)=obs(:,i)+point';
        obs_count(i)=obs_count(i)+1;
    else
        % Otherwise, give it back its prediction
        obs(:,i)=obs(:,i)+x(1:dim,i);
        obs_count(i)=obs_count(i)+1;
    end
end
% Divide up the rest of the points.
[a b] = size(prob);

for i=1:a
    pr = prob(a,:);
    [loc_pr c] = min(pr);
    if loc_pr > thresh
        point = points(a,:);
        obs(:,c)=obs(:,c)+point';
        obs_count(c)=obs_count(c)+1;
    end
end

for i=1:num_filters
    if obs_count(i)>0
        obs(:,i) = obs(:,i)/obs_count(i);
    end
end

runKalman.m

function Yk = runKalman(obs, tstamps, init0);
styles = ['b:';'r:';'g:'];

[num_trajectory dim] = size(init0);

% Fix init0 so it is in the proper form
kal_init0 = init0';
Yk = estimateKalman(obs, tstamps, kal_init0);

for i=0:(num_traject-1)
    drawTrajectory(Yk(:,(dim*i+1):(dim*(i+1))), styles(i+1,:));
end

end

A.7 Utility Functions

add_cols.m

% Adds the columns of A in groups of N
function B = add_cols(A,N)
[m n] = size(A);
post_n = n/N;
idx = 1:post_n;
idx = repmat(idx,N,1);
idx = reshape(idx,n,1);
C = eye(post_n);
C = C(idx,:);
B = A*C;
end

justify_obs.m

% Turns a cell list into a
function obs_mat = justify_obs(all_obs)
obs_mat=[];
[junk num_steps] = size(all_obs);
for i=1:num_steps
    cur_obs = all_obs{i};
[num_obs dim] = size(cur_obs);
[a b] = size(obs_mat);
if num_obs>a
    obs_mat = [obs_mat; NaN*ones(num_obs-a,b)];
else
    cur_obs = [cur_obs; NaN*ones(a-num_obs,dim)];
end
obs_mat = [obs_mat cur_obs];
end
end

mat2vec.m

function vec = mat2vec(mat)
vec = [];
[mat_nrows, N] = size(mat);
vec = reshape(mat,N*mat_nrows,1);

mergeObservations.m

% merge observations
function c = mergeObservations(a,b)
[m,n] = size(a);
for i=1:n,
    c{i} = [a{i};b{i}];
end

npermk.m

function num_perm = npermk(n,k)
num_perm = factorial(n)/factorial(n-k);
end
nquad.m

% Performs an n-integral on a function
%
% func = the function to be evaluated. It must be a function of a 1xn
% vector x, and be evaluated on m such vectors by giving it an mxn matrix.
% bounds = the bounds on the indices of the vectors
function int = nquad(func,bounds,tol)

[a b] = size(bounds);
if a == 1
    int = quad(@wrapFunc,bounds(1,1),bounds(1,2),tol);
else
    int = quad(@integrand,bounds(1,1),bounds(1,2),tol);
end

function out = integrand(X)
    [a b] = size(X);
    out = zeros(size(X));
    for i=1:b
        x = X(i);
        y = nquad(@subFunc,bounds(2:end,:),tol);
        out(i) = y;
    end
end

function subOut = subFunc(X)
    %X = X';
    [a b] = size(X);
    new_in = [x*ones(a,1) X];
    subOut = func(new_in);
end
function f = wrapFunc(X)
    f = func(X');
end

end

vec2mat.m

% Converts a column vector to a matrix with N columns
function mat = vec2mat(vec, N)
mat = [];
[vec_nrows junk] = size(vec);
mat_nrows = vec_nrows/N;
mat = reshape(vec,mat_nrows,N);
Bibliography


