An Optimal Filtering Algorithm for Non-Parametric Observation Models in Robot Localization

Jose-Luis Blanco, Javier González, and Juan-Antonio Fernández-Madrigal

Abstract—The lack of a parameterized observation model in robot localization using occupancy grids requires the application of sampling-based methods, or particle filters. This work addresses the problem of optimal Bayesian filtering for dynamic systems with observation models that cannot be approximated properly as any parameterized distribution, which includes localization and SLAM with occupancy grids. By integrating ideas from previous works on adaptive sample size, auxiliary particle filters, and rejection sampling, we derive a new particle filter algorithm that enables the usage of the optimal proposal distribution to estimate the true posterior density of a non-parametric dynamic system. Our solution avoids approximations adopted in previous approaches at the cost of a higher computational burden. We present simulations and experimental results for a real robot showing the suitability of the method for localization.

I. INTRODUCTION

Two prominent applications of Bayesian sequential estimation have received a huge attention by the robotics research community in the last decade, namely localization and simultaneous localization and map building (SLAM) [22], [23]. The former consists of estimating the pose of a mobile robot within a known environment, whereas in SLAM the map is also estimated while performing self-localization.

In both problems the choice for the representation of the environment determines the probabilistic estimation method that can be applied. In the case of landmarks, the map can be modeled by multivariate Gaussian distributions with Gaussian observation models, obtained by solving the problem of association [3], [4]. Thus, SLAM with landmark maps can be approached well through Gaussian filters such as the EKF [12]. However, these methods are not applicable to other types of map representations, as occupancy grid-maps [17], forcing a sample-based representation of the joint probability density. In this case, sequential estimation is carried-out by Monte-Carlo simulations, or particle filters [5].

In this paper we focus on the problem of localization using occupancy grids, although the proposed method can be also applied to other maps compatible with a sample-based representation of distributions, e.g. topological maps. Some advantages of occupancy grids are the precise dense information they provide and the direct relation of the map with the sensory data, which avoids the problem of data association. Their main drawback is that the probabilistic observation model for grid maps can be evaluated only pointwise and lacks a parametric form [23], in contrast to analytical models available for landmark maps [3], [4]. Standard particle filter algorithms like the Sequential Importance Sampling (SIS) filter [20] and the SIS with resampling (SIR) filter [10], [21] allow us to perform sequential filtering provided only the ability to draw samples according to the system transition model (the robot motion model) and to pointwise evaluate the observation model. However, the efficiency of these algorithms is greatly compromised by peaky sensor models and outliers, which make most of the particles to be discarded in a resampling step and lead to particle impoverishment or even to the divergence of the filter. For mobile robots this issue typically arises in robots equipped with low-noise sensors such as laser range finders.

A theoretical solution that enables the efficient representation of probability densities through perfectly distributed particles was proposed by Doucet et al. [7], consisting of an optimal proposal distribution from which to draw samples at each time step. However, a direct application of this approach requires an observation model with a parametric distribution from which to draw random samples (as in [15]), whereas for grid maps we can evaluate it only pointwise [23].

The contribution of this work is a new particle filter algorithm that, given the same requirements as the original SIS and SIR algorithms, dynamically generates the minimum number of particles that best represent the true distribution within a given bounded error, thus providing optimal sampling. Our method is grounded on previous works related to optimal sampling [6], [7], auxiliary particle filters (APF) [19], rejection sampling [14], and adaptive sample size for robot localization [8]. In the context of mobile robots, the proposed algorithm represents an important contribution since no Gaussian approximations are assumed while generating new particles, which is the case of previous works [11], [15]. Moreover, our method is based on the formulation of a general particle filter, thus it does not depend on the reliability of scan matching as previous works and can be applied to a wider variety of problems.

The rest of this paper is structured as follows. In section II we review previous particle filter algorithms used in robotics and highlight the differences with our proposal, which is introduced in section III. We provide simulations and experimental results with real data in section IV, and finally we remark some conclusions.

II. RELATED RESEARCH

In this section we briefly review the applications of particle filters to robot localization and SLAM. A more compre-
hensive review of particle filter techniques can be found elsewhere [1], [5].

The probabilistic approach to localization and SLAM includes the estimation of the posterior distribution of the robot poses up to the current instant of time given the whole history of available data. In the case of localization (the focus of this work), let \( x_t \) denote the robot pose at time step \( t \) and \( z^t \) and \( u^t \) represent the sequences of observations and actions up to \( t \), respectively. Then, the posterior of the robot pose can be computed sequentially by applying the Bayes rule:

\[
p(x_t | z^t, u^t) \propto p(z_t | x_t) \prod_{i=1}^{M_i} p(x_t | z_i^t, u^t)
\]

Under the assumptions of linearity and Gaussianity, the Kalman filter [13] represents a closed-form, optimal solution to (1). Some improvements have been proposed to overcome the assumption of linearity, where the most widely known is the Extended Kalman Filter (EKF) [12]. The EKF approach to localization and SLAM has been the predominant one for several years [4], but the limitations of this Gaussian filter led to the popularization of particle filters for global localization [9], and, more recently, also for mapping [11], [16].

As opposed to parametric probability distributions (e.g. Gaussians and sum of Gaussians), the distributions estimated by a particle filter are represented by a finite set of hypotheses, or particles, which are weighted according to importance sampling. The simplest particle filter algorithm is the SIS filter [20], described next in the context of localization. Concretely, let \( \{ x_i^t \}_{i=1}^{M_i} \) denote a set of robot pose hypotheses for the time step \( t \), approximately distributed according to the posterior, that is, \( x_i^t \sim p(x_t | z^t, u^t) \) for \( i = 1, ..., M_i \). Note that virtually all previous particle filter techniques rely on \( M_i \) representing a constant number of particles for all time steps \( t \) (an exception in the work by Fox in [8]).

In general, the particles will be not distributed exactly according to the true posterior, hence they are assigned importance weights \( \omega_i^t \) to obtain an unbiased estimation of the density. The SIS algorithm consists of simulating the Bayes update in (1) by drawing samples for the new robot pose from some proposal distribution, that is, \( x_i^{t+1} \sim q(x_t | x_{t-1}^i, z^t, u^t) \), and then updating the weights by [6]:

\[
\omega_i^t \propto \omega_i^{t-1} \frac{p(z_t | x_t, x_{t-1}^{i-1}, z^t, u^t)}{q(x_t | x_{t-1}^{i-1}, z^t, u^t)}
\]

The simplest choice for the proposal distribution \( q(\cdot) \) is the robot motion model – the prior in (1). In this paper we will refer to this choice as the standard proposal. In this case, widely employed in robotics [8], [9], [16], the weight update in (2) simplifies to the product of the previous weights with the evaluation, at each particle, of the observation model \( p(z_t | x_t^{i-1}, z^t, u^t) \). Note how the SIS filter requires only the ability of drawing samples from the robot motion model and evaluating the observation likelihood pointwise. In spite of its simplicity, the SIS filter is not a practical solution, since it has been demonstrated that the variance of the weights increases over time [6], which eventually leads to the degeneracy of the filter. This is the reason for the introduction of the SIS with resampling (SIR) algorithm [10], where a resampling step replaces those particles with low weights by copies of more likely particles. For the case of SLAM, Rao-Blackwellized Particle Filters (RBPF) are a practical solution [18] that has been applied to landmark maps (FastSLAM [16]) and occupancy grids [11].

However, the efficiency of all the above particle filter algorithms is strongly influenced by the choice of the proposal distribution \( q(\cdot) \): the larger the mismatch between the proposal and the observation likelihood, the more particles are wasted in non-relevant areas of the state space. In particular, this is the case of mobile robots equipped with accurate sensors like laser scanners [11]. This is partly overcome with the Auxiliary Particle Filter (APF) [19], through a one-step look-ahead resampling. In general, an APF reduces the number of wasted particles, but it is still a sub-optimal solution since particles are propagated using the standard proposal distribution.

It has been demonstrated by Doucet et al. [7] that the variance of the particle weights is minimized by choosing an optimal proposal distribution, which incorporates the information of the most recent observation while propagating particles. It has been applied to landmark maps (FastSLAM 2.0 [15]), but it is not directly applicable to map representations without parametric observation models, like occupancy grids.

Recent work by Grisetti et al. [11] overcomes this by approximating the sensor model with a Gaussian whose mean value is obtained by scan matching over the grid map. This approximation has demonstrated its practical utility allowing the efficient mapping of large environments. However, the observation likelihood may not be appropriately approximated by a Gaussian in many situations, thus the posterior distribution would be severely distorted. Even in those cases where the observation likelihood could resemble a Gaussian, it can not be proven that the mean of the real posterior

<table>
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<th>Proposal distribution</th>
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<td>Standard</td>
<td>Non-Gaussian</td>
<td>SIR [10], APF [19], RBPF [18], FastSLAM [16]</td>
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<tr>
<td>Optimal</td>
<td>Non-Gaussian</td>
<td>FastSLAM 2.0 [15], Grisetti et al. [11]</td>
</tr>
<tr>
<td>Optimal</td>
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Fig. 1. The theoretic model of our optimal particle filter. An initial set of $M_{t-1}$ particles is first replicated into a set of auxiliary particles, which are then propagated according to the optimal proposal distribution (simulated by rejection sampling). Then, a resampling stage (with an adaptive sample size) chooses the final set of $M_t$ samples from the updated auxiliary particles, taking each one of them with a probability proportional to its weight. As a result, all the final particles have equal importance weights (omitted in the graph by this reason).

coincides with the result of scan matching. Actually, there are some practical situations where scan matching techniques fail. It has been proposed to discard the information of the corresponding observations [11], but observe that even in those cases we could obtain a more precise posterior by integrating all the available information, which is lost otherwise, and which would provide valuable information when facing ambiguous sensor measurements.

To summarize the above discussion we have represented the previous methods in Table I, where our method also appears for comparison.

III. THE OPTIMAL PARTICLE FILTER

A. Preliminary definitions

It has been shown that the optimal proposal distribution that minimizes the variance of the next weights for any generic particle filter is given by [7]:

$$x_t[i] \sim q(x_t|x_{t-1}[i], z^t, u^t) = \frac{p(x_t|x_{t-1}[i], z^t, u^t)}{p(z_t|x_t, x_{t-1}[i], z^{t-1}, u^t)p(x_t|x_{t-1}[i], z^{t-1}, u^t)}$$

(3)

For mobile robots this proposal requires drawing samples from the product of the transition (robot motion) and observation models, which are the terms that appear in the numerator of (3), respectively. Since the system state for the last time step $(x_t)$ does not appear in the denominator, this is a constant value $\mu_i$ for each particle $i$. Therefore, to draw samples from the optimal proposal is equivalent to draw:

$$\omega_t[i] \propto \omega_t[i-1]p(z_t|x_{t-1}[i], z^{t-1}, u^t)$$

(5)

At this point, we state that the purpose of our optimal particle filter algorithm is to generate samples exactly distributed according to the density in (3), while dynamically adapting the number of samples to assure a good representation of the true posterior at each moment.

To avoid the problem of particle depletion we have found two different approaches in other works. The first one is to resample particles at every time step as required to assure that they represent well the true posterior. Another solution consists of resampling only when a measure of the representativeness of the samples is below a given threshold [21]. We will employ the first approach for the derivation of our optimal algorithm. As discussed later on, this generic optimal filter fits perfectly to the problem of mobile robot localization. A variation using selective resampling can be devised for SLAM, but this will be not addressed here due to space limitation.

B. Derivation of the optimal filter algorithm

In the following we derive the algorithm for generating a dynamically-sized set of samples according to the exact posterior being estimated. To clarify the exposition we have summarized the process graphically in Fig. 1.

We start by assuming that a set of $M_{t-1}$ particles $x_{t-1}[i]$ is available which are exactly distributed according to the posterior of our system for the time step $t-1$, that is:

$$x_{t-1}[i] \sim p(x_{t-1}|z^{t-1}, u^{t-1})$$

(6)

Since these samples are optimally distributed, all of them will have equal importance weights, and so they can be omitted. The assumption of perfectly distributed particles for the previous time step is not a problem but for the first iteration of the filter. Typical assumptions for the initial belief
include uniform or Gaussian distributions, depending on the available information and the specific problem.

Now we introduce a set of auxiliary particles \( \tilde{x}^{[i,j]} \) with associated importance weights \( \tilde{\omega}^{[i,j]} \), such that \( \tilde{x}^{[i,j]} = x^{[i]}_{t-1} \) for \( j = 1, ..., N \) and \( \tilde{\omega}^{[i,j]} = 1/(NM_{t-1}) \). That is, we replicate \( N \) times each particle \( x^{[i]}_{t-1} \), assigning equal weights to all of them. Notice that this process does not modify the sample-based estimation of the posterior, since each particle \( i \) is replicated the same number of times. We will use these auxiliary particles just as a computation artifact: in practice only a few of them need to be generated, as will become clear below. Therefore, the value \( N \) is left undefined here, although it is convenient to think of it as a large value, ideally the infinity.

The auxiliary particles are propagated according to the optimal proposal, in (4), in order to obtain a large amount (ideally infinity) of optimally distributed particles \( \tilde{x}^{[i,j]} \), from which we will finally keep only the required ones for providing a good representation of the posterior. This is achieved by generating the new set of particles \( x_t^{[k]} \) by resampling the set of auxiliary samples \( \tilde{x}^{[i,j]} \).

The key point that allows us to directly generate the optimally distributed particles without computing all the auxiliary ones is that all the auxiliary particles \( \tilde{x}^{[i,j]} \) coming from a given particle \( x^{[i]}_{t-1} \) will have equal weights. This property follows from the fact that the concrete value of the particle at time step \( t \) does not appear in the computation of the new weights, as described by (5). These groups of equally-weighted samples are sketched in Fig. 1.

Thus, the optimal particles \( x_t^{[k]} \) are generated by resampling the auxiliary set at time step \( t \). Similarly to auxiliary particle filters [19], we perform this by drawing indexes \( i \) of particles for the previous time step, in our case with a probability proportional to the weights \( \tilde{\omega}^{[i,j]} \), which are given by:

\[
\tilde{\omega}^{[i,j]} = \tilde{\omega}^{[i,j]}_t p(z_t | x_t^{-1}[i], z_t^{-1}, u_t)
\]  

(7)

Here the a priori likelihood of the observation \( z_t \) can be expanded using the law of total probability:

\[
p(z_t | x_t^{-1}[i], z_t^{-1}, u_t) = \int p(x_t | x_{t-1}[i], u_t) p(z_t | x_t, x_{t-1}[i], z_t^{-1}) dx_t
\]  

(8)

The terms that appear inside the integral above are the system transition and observation models, respectively. Since we are assuming in this work that we can only draw samples from the system transition model and evaluate pointwise the observation model, a Monte-Carlo approximation of the integral \( \hat{p}(z_t | \cdot) \approx p(z_t | \cdot) \) can be obtained by means of:

\[
\hat{p}(z_t | x_t^{-1}[i], z_t^{-1}, u_t) = \frac{1}{B} \sum_{n=1}^{B} p(z_t | x_t^{[n]}_t, x_t^{-1}[i], z_t^{-1})
\]  

(9)

with the \( B \) samples \( x_t^{[n]}_t \) generated according to the system transition model, e.g., the robot motion model for localization and SLAM. The number \( B \) is a parameter of our algorithm, and will be typically in the range 10 to 200 depending on the specific problem addressed by the filter.

Going back to the resampling of the auxiliary particles, for each drawn index \( i \) we generate a new optimal particle by taking the value of any auxiliary particle in the \( i \)th group, since all of them have equal probability of being selected in the resampling. That is, the new optimal particle \( x_t^{[k]} \) is a copy of \( \tilde{x}^{[i,j]} \), where the value of \( j \) is irrelevant. The importance weights of the final particles given by our algorithm can be ignored, since particles obtained by resampling all have exactly the same weights.

We need to provide a method to compute the concrete value of the auxiliary particles \( \tilde{x}^{[i,j]} \) for some certain value of \( i \). We employ here the rejection sampling technique to draw from the product of the transition and observation densities – refer to (4). Basically, this technique consists of generating samples \( x_t^{[k]} \) following one of the terms of the product (the transition model), and accepting the sample with a probability \( \Delta \) proportional to the other term (the observation model) [14]:

\[
\Delta = \frac{p(z_t | x_t^{[k]}_t, x_{t-1}[i], z_t^{-1}, u_t)}{\hat{p}_{\text{max}} (z_t | x_t, x_{t-1}[i], z_t^{-1}, u_t)}
\]  

(10)

We must remark that this technique has a random execution time. The only quantity required to evaluate (10) is the maximum value of the observation model \( \hat{p}_{\text{max}}(z_t | \cdot) \), which can be estimated simultaneously to the Monte-Carlo approximation in (9) for the same set of samples \( x_t^{[n]}_t \), thus it does not imply further computational cost.

Up to this point we have shown how to generate one particle according to the true posterior given the set of particles for the previous time step. The above method can be repeated an arbitrary number of times to generate the required number of particles \( M_t \) for the new time step \( t \). To determine this dynamic sample size we propose to integrate here the method introduced by Fox in [8]. There it is used the concept of Kullback-Leibler distance (KLD) [2] to derive an expression for the minimum number of particles \( N_t \) such as the KLD between the estimated and the real distributions

<table>
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<td><strong>THE OPTIMAL PARTICLE FILTER ALGORITHM</strong></td>
</tr>
<tr>
<td><strong>algorithm</strong> OptimalParticleFilter : ( {x_t^{[i]}<em>{t-1}}</em>{i=1}^{N_t} \mapsto {x_t^{[k]}<em>t}</em>{k=1}^{N_t} )</td>
</tr>
<tr>
<td>1. For each particle ( x_t^{[i]}_t )</td>
</tr>
<tr>
<td>1.1. Generate a set of ( B ) samples ( x_t^{[n]}_t \sim p(x_t</td>
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<tr>
<td>1.2. Use them to compute ( \hat{p}(z_t</td>
</tr>
<tr>
<td>2. Generate ( N_t ) particles, with ( N_t ) determined by KLD-sampling [8].</td>
</tr>
<tr>
<td>2.1. Draw an index ( i ) with probability given by weights in (7).</td>
</tr>
<tr>
<td>2.2. Generate a new sample by rejection sampling:</td>
</tr>
<tr>
<td>2.2.1. Draw a candidate sample ( x_t^{[k]}_t \sim p(x_t</td>
</tr>
<tr>
<td>2.2.2. Compute ( \Delta ) through (10).</td>
</tr>
<tr>
<td>2.2.3. With a probability of ( 1 - \Delta ), go back to 2.2.1.</td>
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we have computed the KLD between the analytical and the histograms, shown in the middle row of Fig. 2 along with the analytical solution from the KF. To evaluate each algorithm, we have reconstructed the estimated densities by means of weighted histograms, shown in the middle row of Fig. 2 along with the analytical solution from the KF. To evaluate each algorithm, we have reconstructed the estimated densities by means of weighted histograms, shown in the middle row of Fig. 2 along with the analytical solution from the KF.

To measure the accuracy of each particle filter we have compared the accuracy in the localization between our optimal PF and a standard PF. The most interesting conclusion from the results, in Fig. 3(b), is that our optimal PF has an excellent performance starting from just one particle (an average error of roughly 0.10m), whereas the standard proposal needs about 10 particles or more to avoid the filter to diverge (e.g. the average error of 6m for one particle means the filter has lost track of the localization).

On the other hand, our method requires more computation time than the standard approach. For example, for 100 particles, ours takes 50.56s for the whole experiment while the standard PF takes 9.91s only. Thus, one could argue that a standard PF with more particles would achieve a similar accuracy than our optimal PF for the same computation time. Actually, we can see in the graphs that our method always achieves a better accuracy than the standard approach, even

is kept below a certain threshold $\epsilon$ with a probability $1 - \delta$. Please, refer to [8] for further details.

To summarize the introduction of our algorithm we present an algorithmic description of the overall method in Table II.

IV. EXPERIMENTAL RESULTS

In this section we will first present simulations for comparing our filtering algorithm to others, and next we will show experiments where our approach is applied to robot localization (position tracking).

A. Simulations

We have considered here a one-dimensional linear system with Gaussian transition and observation models. Such a simple system allow us to contrast the output of the different filters with the analytical solution from a Kalman filter (KF) which provides us the exact posterior. The situation being simulated is that of an observation model much more peaked than the prior distribution obtained from the system transition model, a situation similar in mobile robotics to a poor motion model (such as odometry) and a very precise sensor (such as a laser scanner). The top graphs of Fig. 2 represent the location and weights of the obtained particles with three different algorithms: a SIR filter with a standard proposal distribution [21], an auxiliary particle filter (APF) [19], and our method. We can observe how the standard proposal leads to most of the particles being wasted in non relevant areas with negligible importance weights. The APF introduces a great improvement in this sense, since particles are more concentrated in the area of interest. However, the weights still contain a certain variance. In contrast, our optimal algorithm generates particles distributed exactly according to the true posterior, thus they all have the same weights. To measure the accuracy of each particle filter we have reconstructed the estimated densities by means of weighted histograms, shown in the middle row of Fig. 2 along with the analytical solution from the KF. To evaluate each algorithm, we have computed the KLD between the analytical and the estimated distributions for a range of sample sizes (we have disabled here the capability of automatically determining the sample size in our method for comparison purposes with the others). The average KLD for 1000 realizations, shown in the bottom row of Fig. 2, confirms that our approach gives estimations closer to the actual posterior (with less particles) than previous methods.

B. Localization

The following experiments consist of tracking the pose of a mobile robot equipped with a laser range finder while it is manually guided through an office environment. Concretely, the path and the (already built) map of the environment are shown in Fig. 3(a). The purpose of the first experiment is to compare the accuracy in the localization between our optimal sampling mechanism and the standard proposal distribution. The resolution of the grid is 0.04m, and the non-parametric observation model is the likelihood field described in [23].

The accuracy has been calculated by averaging the localization errors of all the particles at each time step, and using as the ground truth the robot poses estimated while the map was first built. Significant results are obtained by averaging over 100 executions for each sample size. The capability of adapting the sample size in our algorithm has been disabled in this first experiment to provide a fair comparison to a standard PF. The most interesting conclusion from the results, in Fig. 3(b), is that our optimal PF has an excellent performance starting from just one particle (an average error of roughly 0.10m), whereas the standard proposal needs about 10 particles or more to avoid the filter to diverge (e.g. the average error of 6m for one particle means the filter has lost track of the localization).

Fig. 2. A comparison of our method to other two particle filters for a linear, Gaussian system. The top row shows the particles and weights obtained for each algorithm. Below the weighted histograms of the samples is compared to the exact Gaussian density. This similarity is measured in the third row with the Kullback-Leibler distance between the real and the estimated distributions for different sample sizes. Observe how our method achieves a lower distance (a higher similarity) even for a few particles.
Additional work will be required to explore the interesting applications of this method to SLAM, where a dynamic number of optimally distributed particles will improve recent works on RBPF-based grid mapping.

**REFERENCES**


