

An Interlaced Extended Information Filter for Self-Localization in Sensor Networks

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Abstract—Wireless Sensor Networks (WSNs) are at the forefront of emerging technologies due to the recent advances in Microelectromechanical Systems (MEMSs). The inherent multidisciplinary nature of WSN attracted scientists coming from different areas stemming from networking to robotics. WSNs are considered to be unattended systems with applications ranging from environmental sensing, structural monitoring, and industrial process control to emergency response and mobile target tracking. Most of these applications require basic services such as self-localization or time synchronization. The distributed nature and the limited hardware capabilities of WSN challenge the development of effective applications. In this paper, the self-localization problem for sensor networks is addressed. A distributed formulation based on the Information version of the Kalman Filter is provided. Distribution is achieved by neglecting any coupling factor in the system and assuming an independent reduced-order filter running onboard each node. The formulation is extended by an interlacement technique. It aims to alleviate the error introduced by neglecting the cross-correlation terms by “suitably” increasing the noise covariance matrices. Real experiments involving MICAz Mote platforms produced by Crossbows along with simulations have been carried out to validate the effectiveness of the proposed self-localization technique.

Index Terms—Sensor networks, distributed applications, distributed network.

1 THE SELF-LOCALIZATION PROBLEM IN SENSOR NETWORKS

A sensor network consists of a collection of nodes deployed in an environment that cooperate to perform a task. Each node, which is equipped with a radio transceiver, a microcontroller, and a set of sensors, shares data to reach the common objective. Sensor networks provide a framework in which, exploiting the collaborative processing capabilities, several problems can be faced and solved in a new way. However, it comes along with several challenges such as limited processing, storage and communication capabilities, as well as limited energy supply and bandwidth. Performing a partial computation locally on each node, and exploiting internode cooperation, is the ideal way to use sensor networks. Unfortunately, this modus operandi is highly constrained by the reduced hardware capabilities as well as by the limited energy resources that makes communication very expensive in terms of lifetime for a node. As a consequence, these constraints must be taken into account when developing algorithms able to operate in a distributed fashion.

Sensor networks can be of interest to different areas of application, ranging from environmental monitoring [9], [41], civil infrastructures [23], [27], medical care [38], [32] to home and office applications [39], [25]. In each field, the deployment of a sensor network has provided interesting advantages. For instance, in the context of environmental monitor, the introduction of a sensor network made it possible to keep environments intrinsically threatening for

human beings [41] under surveillance, or in the context of medical care, it made it possible to remotely monitor the health condition of patients by continuously extracting clinical relevant information [32].

However, in order to build these applications, some basic services, such as time synchronization or nodes localization, are generally required. In fact, basic middle ware services, such as routing, often rely on location information, e.g., geographic routing [5], [40], [24]. Specifically, the localization problem in sensor networks consists of finding out the locations of nodes with regard to any topology or metric of interest. This problem turns out to be difficult to solve. In fact, in [21], [14], it was proved that a sufficient condition for a sensor network to be localizable cannot be easily identified. This holds even when considering the availability of perfect measurements. Further, several analyses showed that having reliable ranging information is fairly practical [42], [44], [2], especially when using the received signal strength indication (RSSI).

In this paper, a distributed formulation based on the Information version of the Kalman Filter is provided to deal with the self-localization problem in sensor networks. Distribution is achieved by neglecting any coupling factor in the system and assuming an independent reduced-order filter running onboard each node. The error introduced by this assumption is then mitigated by increasing the noise covariance matrices. This formulation is particularly convenient in all those scenarios where the dimension of the state space is lower than the dimension of the observations. Indeed, this is the case of the proposed sensor network scenario, where the dimension of the state space for each node is equal to 2, while the number of observations is strictly related to the number of nodes deployed into the environment.

The rest of the paper is organized as follows: In Section 2, the state of the art for the localization problem in sensor

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networks is given. In Section 3, some theoretical insights about the estimation problem in a probabilistic framework are provided. In Section 4, the interlacement technique is described. In Section 5, the sensor network scenario exploited in this work is detailed. In Section 6, the formulation of the information filter for the adopted scenario is proposed. In Section 7, the performance analysis is depicted, while in Section 8, the experimental results are described. In Section 9, the analysis of the computational complexity is detailed. Finally, in Section 10, we conclude the paper.

2 STATE OF THE ART

A taxonomy of localization algorithms for sensor networks can be drawn according to the computational organization, i.e., centralized and distributed, to the mechanism adopted for estimating location, i.e., range-based or range-free, and finally, with regard to the availability of anchors nodes, i.e., anchor-based or anchor-free.

Centralized algorithms exploit a central computer to perform all the complex computations using information gathered by nodes [12], [36], [6]. Distributed algorithms dispense the computation over the network, allowing each node to perform locally and compensating for the lack of global knowledge through an intensive collaborative processing [28], [11], [10]. Both schemes offer advantages and drawbacks. Centralized algorithms provide interesting performance but they lack in scalability and robustness. Distributed algorithms provide high robustness and scalability but the development of effective collaborative processing algorithms is challenging.

Range-based algorithms exploit point-to-point distances or angle estimates in order to perform the localization task [33], [37], [30]. Range-free algorithms do not make any assumption about the availability or reliability of this information [20], [26], [44]. Although range-free approaches are appealing as a cost-effective alternative to more expensive range-based approaches, their performance may lack in accuracy.

Anchor-based algorithms rely on the availability of location information for some special nodes in order to localize the network [15], [35]. Anchor-free methods determine the geometry of the network simply by exploiting local interaction among nodes [34], [45]. Anchor-based algorithms have the advantage of directly localizing nodes within a global reference frame, but their accuracy is affected by the number of anchor nodes and their distribution in the sensor field [7]. Conversely, anchor-free methods scale better and do not require expensive hardware, although only relative location estimates can be provided.

Centralized algorithms represent the first attempt to solve the localization problem in sensor networks. In [12], the authors propose the semidefinite programming approach (SDP) to solve the localization problem. The key idea is to model geometric constraints between nodes as linear matrix inequalities (LMIs) and then use the semidefinite programming theory to solve it. The result is a bounding region for each node, representing feasible locations where nodes are supposed to be. Although using a set of convex constraints in order to estimate the position of a node is

very elegant, it turns out to be inaccurate as constraints do not use precise data range. Moreover, the algorithm provides a good estimation only when having anchors densely deployed on the boundary of the sensor network, a condition that cannot always be guaranteed. The SDP approach is extended to deal with noisy distance measurements by taking advantage of an additional technique to mitigate inaccuracies [3]. In fact, the solution provided by the SDP, though not accurate, represents by the authors a good starting point for a gradient descent method. Furthermore, numerical results show that by means of this improvement, it is possible to obtain a solution very close to the optimal one. However, the distributed formulation is the result of a clusterization and a local execution of the algorithm within each subset. Therefore, the computational complexity is merely mitigated reducing the number of nodes but the approach still remains almost centralized. In [36], the authors propose an algorithm that uses connectivity information, i.e., which nodes are within the communication range of which others, to derive the locations of the nodes in the network. This algorithm is based on multi-dimensional scaling (MDS), a set of data analysis techniques that displays the structure of distance-like data as a geometrical picture [4]. It can be broken down into three steps. Starting with the given network connectivity information, an all-pairs shortest path algorithm is run to roughly estimate the distance between each possible pair of nodes. Then, the multidimensional scaling is applied over these data to derive node locations. Finally, location estimates are normalized with respect to nodes whose position is known.

Distributed algorithms better fit the inherent collaborative nature of sensor networks. In [28], the authors developed an algorithm focused on providing more robust local maps. The idea is to split the problem into a subset of smaller regions in which the localization is performed taking advantage of the probabilistic notion of *robust quadrilaterals*. A robust quad is a set of four nodes fully connected by distance measurements and well-spaced in such a way that no ambiguity can arise, even when in the presence of noise. The algorithm merges the subregions using a coordinate system registration procedure. Such a procedure maps local reference systems into a global one providing the best fitting matrix when in presence of a set of common nodes. An optional optimization step can be executed in order to refine the local map first. The weakness of this approach, as pointed out by the same authors, is that under conditions of low node connectivity or high measurement noise, the algorithm may be able to localize only a reduced number of nodes. In [10], the authors propose an approach, where localization is performed by exploiting clustering information. Starting from locally aware anchors, an initial set of calibrated nodes is built. This set is then expanded to include iteratively all the cluster heads, i.e., the representative node for the cluster. Due to the iterative nature of this approach, a refining step is required in order to provide reliable location estimates. Once the cluster heads have been fully localized, the remaining follower nodes, i.e., noncluster head nodes, can be localized.

Range-free algorithms instead may offer an alternative that anytime distance information is not available, due to the

stringent hardware limitations. In [20], a range-free localization algorithm called APIT is proposed. In this work, the environment is first isolated into triangular regions defined by beacons: localization is achieved by checking whether a node is inside or outside of these regions. Combinations of anchor positions can be used to reduce the diameter of the estimated area. Although an interesting insight on how localization error affects a variety of location-dependent applications such as geographical routing or target tracking is provided, an impractical number of beacons might be required to achieve satisfactory performances. In [44], a sequence-based RF localization algorithm called Ecolocation is proposed. The key idea is to determine the location of unknown nodes by examining the ordered sequence of received signal strength (RSS) measurements taken at multiple reference nodes. The authors propose a constraint-based approach that provides for robust location decoding even in the presence of random RSS fluctuations due to multipath fading and shadowing. However, the algorithm performance is heavily conditioned by the number of available reference nodes. In [8], the authors propose an RF-based distributed localization method, where location is estimated by simply averaging the positions of all the anchors it is connected to. Obviously, the accuracy of the estimation is strictly related to the density of anchors deployed in the environment and the density required to obtain an acceptable estimation is fairly practical.

Anchor-free algorithms may finally represent an alternative solution in case prior knowledge about location is not available and an estimation with regard to a global reference frame is not required. In [34], the authors propose the Anchor-Free Localization (AFL) algorithm, an algorithm where all nodes concurrently calculate and refine their coordinate information. The key idea is the introduction of fold freedom: a fold-free embedding of a graph is an embedding where every cycle has the correct clockwise/counterclockwise orientation of nodes, modulo global reflection, with respect to the original graph. In detail, AFL is composed of two steps. During the first step, a folder-free graph embedding is computed starting from the original embedding and selecting five ad hoc reference nodes used to approximate the polar coordinate of any other node. Successively, a mass-spring-based optimization is performed in order to correct and balance localized errors. In [45], an anchor-free node localization protocol, which exploits clusterization to achieve scalability, is proposed. Such a protocol consists of three steps: network bootstrapping, local position discovery, and global localization. During the first step, clusters are identified and a “breadth-first spanning tree” rooted at the head of each cluster is performed. Since each node is able to measure distances from its neighbors (by exploiting some TOA technique) and a route exists from it to the cluster headset, all local distance informations are sent to the cluster heads. This information will be used during the second step to build a local map at each cluster head. Finally, in the third step, cluster heads collaborate in order to obtain a global map of the network. Such a global coordinate system can be built from the local maps by simply exploiting matrix rotations, translations, and mirroring.

In this paper, a novel distributed, range-based algorithm, namely the Interlaced Extended Information Filter (IEIF), is

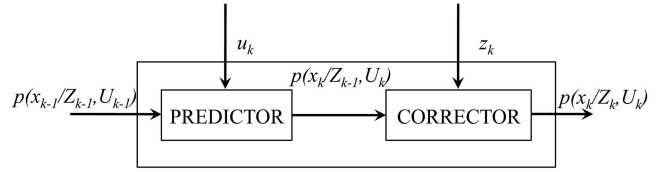


Fig. 1. Bayesian filter.

proposed. Starting from a centralized formulation, distribution is achieved by neglecting any coupling factor in the system and assuming an independent reduced-order filter running onboard each node. This formulation is successively extended by an interlacement technique aiming to alleviate the error introduced by neglecting the cross-correlation terms by “suitably” increasing the noise covariance matrices. The proposed algorithm can provide global localization by assuming anchors are available. In the same way, also relative localization among nodes can be achieved by relaxing the assumption of anchors availability. The effectiveness of this distributed approach has been thoroughly investigated by experiments carried out with MICAz Mote platforms produced by Crossbows, while its scalability has been analyzed by means of simulations.

3 THEORETICAL BACKGROUND

3.1 Bayesian Framework

The probability theory provides a suitable framework for modeling the self-localization problem in sensor networks. Let us consider a system described by the following set of equations:

$$\begin{aligned} x_k &= f(x_{k-1}, u_k, w_k), \\ z_k &= h(x_k, v_k), \end{aligned} \quad (1)$$

where x_k is a stochastic variable representing the locations of the nodes, u_k is the control input, w_k and v_k are noises that affect the system, while $f(\cdot)$ and $h(\cdot)$ are mathematical relations that characterize the state transition and the observation z_k , respectively.

In the probabilistic context, the localization problem consists of computing the probability distribution $p(x_k|Z_k, U_k)$ for all times k . This probability distribution describes the *joint posterior density* of the sensor locations (x_k) given the recorded observations (Z_k) and control inputs (U_k) up to time k . To apply this approach in a real context, it is often required to perform the above-mentioned computation online. Therefore, a recursive formulation should be provided in terms of Bayesian filter, graphically depicted in Fig. 1.

The idea is to provide at each time step k a new estimate by combining the available estimate of the *joint posterior distribution* $p(x_{k-1}|Z_{k-1}, U_{k-1})$ at time $k-1$, with the control u_k and the observation z_k . In this way, both the *state transition model* and the *observation model*, describing, respectively, the stochastic effects of the control input and observation, are required.

From a probabilistic point of view, the state transition model can be described in terms of the *joint prior density* $p(x_k|x_{k-1}, u_k)$. Such probability distribution exploits that the

state transition is assumed to be a Markov process in which the next state x_k depends only on the immediately preceding state x_{k-1} and the applied control u_k and is independent of the observations.

On the other hand, the observation model describes the probability of retrieving an observation z_k when the sensor locations are known, and is generally stated in the form $p(z_k|x_k)$.

The localization algorithm can be implemented in a standard two-step recursive prediction (*time update*)

$$p(x_k|Z_{k-1}, U_k) = \int_{\Xi} p(x_k|x_{k-1}, u_k)p(x_{k-1}|Z_{k-1}, U_{k-1})dx_{k-1} \quad (2)$$

and correction (*measurement update*) form

$$p(x_k|Z_k, U_k) = \frac{p(z_k|x_k)p(x_k|Z_{k-1}, U_k)}{p(z_k|Z_{k-1}, U_k)}. \quad (3)$$

Equations (2) and (3) provide a recursive procedure for calculating the joint posterior $p(x_k|Z_k, U_k)$; however, they cannot be implemented on a digital computer in their general form stated above, as the joint posterior over the state space is a density over a continuous space, hence, has infinitely many dimensions. Therefore, any effective localization algorithm has to resort to additional assumptions.

3.2 The Kalman Filter

A common approach is represented by the use of Kalman filter [22]. In this context, a linear or linearized system model is required:

$$\begin{aligned} x_k &= F_k x_{k-1} + B_k u_k + w_k, \\ z_k &= H_k x_k + v_k, \end{aligned} \quad (4)$$

where $w_k \sim \mathcal{N}(0, Q_k)$, $v_k \sim \mathcal{N}(0, R_k)$, and $x_0 \sim \mathcal{N}(\hat{x}_0, P_0)$ are mutually independent Gaussian variables for each pair of time instant (k, k') . The joint posterior $p(x_k|Z_k, U_k)$ is modeled by a unimodal Gaussian density. The mode of this density (\hat{x}_k) yields the current positions of the nodes, and the variance (P_k) represents the current uncertainty. As only these two parameters have to be computed to propagate uncertainty, there is no need to discretize the state space. In this way, the prediction becomes

$$\begin{aligned} \hat{x}_{k|k-1} &= F_k \hat{x}_{k-1|k-1} + B_k u_k, \\ P_{k|k-1} &= F_k P_{k-1|k-1} F_k^T + Q_k, \end{aligned} \quad (5)$$

while the correction requires the computation of the well-known Kalman gain matrix

$$K_k = P_{k|k-1} H_k^T [H_k P_{k|k-1} H_k^T + R_k]^{-1} \quad (6)$$

before update the estimate

$$\begin{aligned} \hat{x}_{k|k} &= \hat{x}_{k|k-1} + K_k (z_k - H_k \hat{x}_{k|k-1}), \\ P_{k|k} &= P_{k|k-1} - K_k [H_k P_{k|k-1} H_k^T + R_k] K_k^T. \end{aligned} \quad (7)$$

The advantage of Kalman filter lies in its efficiency and in the high accuracy that can be obtained; however, it is not able to cope with high nonlinear system and multimodal distributions. Therefore, in most practical situation, Kalman filter cannot be applied. Instead, one is forced to use

approximations or suboptimal solutions. Over the years, a large number of approximate nonlinear filters have been proposed in the literature [13]. Some are fairly general, while others are more tailored to a particular application.

Here, only analytic approximations have been considered: in this category, it is included the Extended Kalman Filter (EKF). The main feature of this filter is that it linearizes the nonlinear functions in the state transition and observation models. The EKF is derived for nonlinear systems with additive noise

$$\begin{aligned} x_k &= f(x_{k-1}, u_k) + w_k, \\ z_k &= h(x_k) + v_k, \end{aligned} \quad (8)$$

where w_k and v_k are mutually independent, zero-mean white Gaussian random sequences, having covariance matrices Q_k and R_k , respectively. The nonlinear functions $f(\cdot)$ and $h(\cdot)$ are approximated by the first term in their Taylor series expansion. The joint posterior density is approximated by a Gaussian density and computed recursively as follows:

- Prediction

$$\begin{aligned} \hat{x}_{k|k-1} &= f(\hat{x}_{k-1|k-1}, u_k), \\ P_{k|k-1} &= J_x^f P_{k-1|k-1} J_x^{fT} + Q_k. \end{aligned} \quad (9)$$

- Update

$$\begin{aligned} K_k &= P_{k|k-1} J_x^h [J_x^h P_{k|k-1} J_x^{hT} + R_k]^{-1}, \\ \hat{x}_{k|k} &= \hat{x}_{k|k-1} + K_k (z_k - h(\hat{x}_{k|k-1})), \\ P_{k|k} &= P_{k|k-1} - K_k [J_x^h P_{k|k-1} J_x^{hT} + R_k] K_k^T, \end{aligned} \quad (10)$$

where J_x^f and J_x^h are the Jacobians of the nonlinear functions $f(\cdot)$ and $h(\cdot)$, respectively.

As only Kalman filters are used in the sequel, only these techniques have been reported due the space limit; however, a complete review can be found in [13].

3.3 The Information Filter

An Information Filter (IF) is essentially a Kalman Filter (KF) expressed in terms of measures of *information* about the parameters (state) of interest rather than direct state estimates and their associated covariances [19]. The two key information-analytic variables are the *information matrix* and the *information state vector*, where the term “information” is used according to the Fisher definition.

The Fisher information matrix Ψ_k is the amount of information that an observable random variable z carries about an unobservable parameter x upon which the likelihood function of z , $L(x) = p(z|x)$, depends. It can be derived as the covariance of the *score function*, that is, the partial derivative, with respect to some parameter x , of the logarithm (commonly the natural logarithm) of the likelihood function. If the observation is z and its likelihood is $L(x) = p(z|x)$, then the score $S_k(x)$ can be described as follows:

$$S_k(x) = \nabla_x \ln p(z_k | x_k) \quad (11)$$

$$= \frac{\nabla_x p(z_k | x_k)}{p(z_k | x_k)}. \quad (12)$$

Moreover, being the expectation of the score:

$$E[S_k(x)] = \int \frac{\nabla_x p(z_k | x_k)}{p(z_k | x_k)} p(z_k | x_k) dz_k \quad (13)$$

$$= \nabla_x \int p(z_k | x_k) dz_k \quad (14)$$

$$= \nabla_x 1 = 0. \quad (15)$$

The Information matrix Ψ_k is simply the second-order moment of the score function $S_k(x)$, as follows:

$$\Psi_k = E[S_k(x)S_k(x)^T] \quad (16)$$

$$= E[\{\nabla_x \ln p(z_k | x_k)\}\{\nabla_x \ln p(z_k | x_k)\}^T]. \quad (17)$$

Furthermore, if the following regularity condition holds:

$$\int H_x(p(z_k | x_k)) = \nabla_x \nabla_x^T p(z_k | x_k) = 0, \quad (18)$$

where H_x is the square matrix of the second-order partial derivatives (i.e., Hessian Matrix), the Information matrix Ψ_k can be also written as

$$\Psi_k = -E[\nabla_x \nabla_x^T \ln p(z_k | x_k)]. \quad (19)$$

At this point, when the likelihood function $p(z | x)$ is a Gaussian distribution and the posterior conditional distribution is Gaussian as well, described as $p(x_k | z) \sim \mathcal{N}(\hat{x}_k, P_k)$, then it can be proved [29] that the Information Matrix is equal to the inverse of the covariance matrix P_k as follows:

$$\Psi_k = P_k^{-1}. \quad (20)$$

Likewise, the information state vector y_k can be easily derived as the product of the inverse of the information matrix and the state estimate as follows:

$$y_k = \Psi_k x_k \quad (21)$$

$$= P_k^{-1} x_k. \quad (22)$$

The information filter formulation can be easily derived from the Kalman Filter formulation under the assumption of Gaussianity previously stated. In particular, by performing the substitutions given in (22) and (20), the following set of equations is obtained:

- Prediction

$$\begin{aligned} \Psi_{k|k-1} &= [F_k(\Psi_{k-1|k-1})^{-1}F_k^T + Q_k]^{-1}, \\ L_{k|k-1} &= \Psi_{k|k-1}F_k\Psi_{k-1|k-1}^{-1}, \\ \hat{y}_{k|k-1} &= L_{k|k-1}\hat{y}_{k-1|k-1} + \Psi_{k|k-1}B_k u_k. \end{aligned} \quad (23)$$

- Estimation

$$\begin{aligned} \Psi_{k|k} &= \Psi_{k|k-1} + \Phi_k, \\ \hat{y}_{k|k} &= \hat{y}_{k|k-1} + i_k, \\ \Phi_k &= H_k^T R_k^{-1} H_k, \\ i_k &= H_k^T R_k^{-1} z_k. \end{aligned} \quad (24)$$

The information filter can be extended to a linearized estimation algorithm for nonlinear system, the Extended

Information Filter (EIF). The idea is to apply the analytic approximations used in EKF and the substitutions of IF to build up an estimation method for nonlinear systems. The EIF presents several interesting features, among the others an easy initialization of matrices and vectors, a reduced computational load, and aptitude to be distributed for parallel computation. The EIF equations can be found as follows:

- Prediction

$$\begin{aligned} \Psi_{k|k-1} &= [J_x^f(\Psi_{k-1|k-1})^{-1}J_x^{fT} + Q_k]^{-1}, \\ \hat{y}_{k|k-1} &= \Psi_{k|k-1}f(\hat{x}_{k-1|k-1}, u_k). \end{aligned}$$

- Estimation

$$\begin{aligned} \Psi_{k|k} &= \Psi_{k|k-1} + \Phi_k, \\ \hat{y}_{k|k} &= \hat{y}_{k|k-1} + i_k, \\ \Phi_k &= J_x^h{}^T R_k^{-1} J_x^h, \\ i_k &= J_x^h{}^T R_k^{-1} [z_k - h(\hat{x}_{k|k-1}) + J_x^h \hat{x}_{k|k-1}]. \end{aligned} \quad (25)$$

A more comprehensive description of the information filter derivation is given in [29].

4 ON THE INTERLACEMENT OF EKF AND EIF

The interlacement technique has been developed [18] to reduce the computational load of a nonlinear filter by means of splitting the estimation of the state variables into parallel subfilters. The key idea is derived from the multiplayer dynamic game theory, where each player chooses its own strategy as the optimal response to the strategy adopted by the other players. In the framework of estimation, players are represented by subfilters, strategy by estimate, whereas the optimal response depends on the estimation algorithm. The interlacement technique can be applied both to the EKF and the EIF, as detailed below.

4.1 Interlaced Extended Kalman Filter

The Interlaced Extended Kalman Filter (IEKF) has been introduced to distribute the estimate of an EKF over a network of processors, each one devoted to estimate a subspace of the state variables minimizing the loss of cross-correlation links. For the sake of clarity, let us consider a system whose model can be decomposed into two subsystems and rewritten as (for the first filter, $i = 1$ and $j = 2$, while for the second, $i = 2$ and $j = 1$):

$$\begin{aligned} x_k^{(i)} &= f^{(i)}(x_{k-1}^{(i)}, x_{k-1}^{(j)}, u_k) + w_k^{(i)}, \\ z_k^{(i)} &= h^{(i)}(x_k^{(i)}, x_k^{(j)}) + v_k^{(i)}. \end{aligned} \quad (26)$$

The IEKF equations proceed from the EKF filter equations (see Fig. 2). At the k th step, each subfilter forms a prediction exploiting both its own estimation and the one of the other filter, according to the following equation:

$$\hat{x}_{k|k-1}^{(i)} = f^{(i)}(\hat{x}_{k-1|k-1}^{(i)}, \hat{x}_{k-1|k-1}^{(j)}, u_{k-1}), \quad (27)$$

$$P_{k|k-1}^{(i)} = J_{x,i}^{f,i} P_{k-1|k-1}^{(i)} J_{x,i}^{f,iT} + \tilde{Q}_k^{(i)}, \quad (28)$$

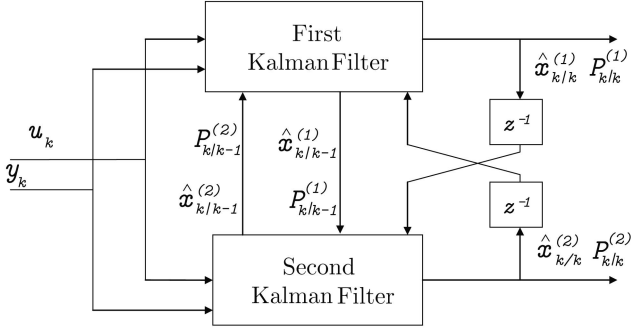


Fig. 2. Interlaced Kalman filter.

where

$$\tilde{Q}_k^{(i)} = Q_k^{(i)} + J_{x,j}^{f,i} P_{k-1|k-1}^{(j)} J_{x,j}^{f,iT}, \quad (29)$$

being $J_{x,i}^{f,i}$ and $J_{x,j}^{f,i}$ the Jacobians of the relation $f^{(i)}(\cdot)$ with respect to $x_k^{(i)}$ and $x_k^{(j)}$.

After the prediction step, the estimates elaborated by the two subfilters are exchanged and used during the update step.

In this step, the observation prediction is formed and compared with the measure z_k provided by the system:

$$\hat{x}_k^{(i)} = \hat{x}_{k|k-1}^{(i)} + K_k^{(i)} [z_k - h^{(i)}(\hat{x}_{k|k-1}^{(i)}, \hat{x}_{k|k-1}^{(j)})], \quad (30)$$

$$P_{k|k}^{(i)} = P_{k|k-1}^{(i)} - K_k^{(i)} J_{x,i}^{h,i} P_{k|k-1}^{(i)}, \quad (31)$$

where the Kalman gain is computed applying the relation

$$K_k^{(i)} = P_{k|k-1}^{(i)} J_{x,i}^{h,iT} [J_{x,i}^{h,i} P_{k|k-1}^{(i)} J_{x,i}^{h,iT} + \tilde{R}_k^{(i)}]^{-1}$$

in which

$$\tilde{R}_k^{(i)} = R_k + J_{x,j}^{h,i} P_{k|k-1}^{(j)} J_{x,j}^{h,iT}, \quad (32)$$

where $J_{x,j}^{h,i}$ and $J_{x,j}^{h,i}$ are the Jacobians of $h^{(i)}(\cdot)$ with respect to $x_k^{(i)}$ and $x_k^{(j)}$.

From (29) and (32), it can be noticed that the process and measurement noise covariance matrices $Q_k^{(i)}$ and $R_k^{(i)}$ are suitable increased by addition of positive semidefinite quantities that take into account the error introduced by the decoupling operation. It is easy to recognize that the term added to $R_k^{(i)}$ in (32) represents the cross correlation between the filters due to innovation process, while the term added to $Q_k^{(i)}$ in (29) is related to the cross correlation induced by the propagation process.

4.2 Interlaced Extended Information Filter

The IEIF is the counterpart of the IEKF in the information space. As already mentioned, the IEF is suitable for distributing the estimation process over parallel computation units, due to the loose correlation between the elements of the information vector. In the information space, indeed, the correlation between information variables, which are not explicitly connected or directly involved in a measurement, is not represented, whereas the covariance matrix explicitly stores this relation in the corresponding off-diagonal entries. Apart from this rough correlation between information variables, there is still a coupling factor that has

to be taken into account even in the presence of distributed implementation of EIF to prevent the divergence of the filter itself. In order to consider this coupling factor, the IEIF is introduced in this work. Let us consider again a system whose model can be decomposed into two subsystems having model equations expressed by (26). The IEIF schema is represented in Fig. 2, after substituting IEKF with IEIF while recalling the relation given by (20). At each time, the filter computes a prediction and estimation step exploiting the equations below:

- Prediction

$$\begin{aligned} \Psi_{k|k-1}^{(i)} &= [J_{x,i}^{f,i} (\Psi_{k-1|k-1}^{(i)})^{-1} J_{x,i}^{f,iT} + \tilde{Q}_k]^{-1}, \\ \hat{y}_{k|k-1}^{(i)} &= \Psi_{k|k-1}^{(i)} f(\hat{x}_{k-1|k-1}^{(i)}, \hat{x}_{k-1|k-1}^{(j)}, u_k), \\ \tilde{Q}_k^{(i)} &= Q_k^{(i)} + J_{x,j}^{f,i} (\Psi_{k-1|k-1}^{(j)})^{-1} J_{x,j}^{f,iT}. \end{aligned} \quad (33)$$

- Estimation

$$\begin{aligned} \Psi_{k|k}^{(i)} &= \Psi_{k|k-1}^{(i)} + \Phi_k^{(i)}, \\ \hat{y}_{k|k}^{(i)} &= \hat{y}_{k|k-1}^{(i)} + i_k^{(i)}, \\ \Phi_k^{(i)} &= J_{x,i}^{h,iT} \tilde{R}_k^{(i)-1} J_{x,i}^{h,i}, \\ i_k^{(i)} &= J_{x,i}^{h,iT} \tilde{R}_k^{(i)-1} z_k^{(i)}, \\ \tilde{R}_k^{(i)} &= R_k^{(i)} + J_{x,j}^{h,i} (\Psi_{k|k-1}^{(j)})^{-1} J_{x,j}^{h,iT}, \\ z_k^{(i)} &= \nu_k^{(i)} + J_{x,i}^{h,i} \hat{x}_{k|k-1}^{(i)} + J_{x,j}^{h,i} \hat{x}_{k|k-1}^{(j)}, \\ \nu_k^{(i)} &= z_k^{(i)} - h^{(i)}(\hat{x}_{k|k-1}^{(i)}, \hat{x}_{k|k-1}^{(j)}). \end{aligned} \quad (34)$$

After every single step, the subfilters exchange their results in terms of best estimate and the associate covariance. The estimate is used to compute the expected measurement, whereas the covariance matrix is involved in the computation of the matrices \tilde{Q}_k and \tilde{R}_k . These matrices have the same meaning introduced for IEKF and convey the coupling factor between information variables in subsystems i and j .

5 SENSOR NETWORK SCENARIO

In this paper, a group of Ω nodes deployed on a planar environment is considered. A typical sensor network node's hardware consists of a microprocessor with reduced computational capability, a radio component, several sensor devices, a minimal data storage unit, and a battery with limited life. Furthermore, a few nodes are equipped with an absolute position system device so that localization with regard to a global frame can be obtained for the whole network. Finally, nodes are assumed to be motionless.

The state of the node i at time k is described by its location with respect to a global frame as follows:

$$x_k^{(i)} = [p_{x,k}^{(i)} p_{y,k}^{(i)}]^T. \quad (35)$$

Thus, the state of the whole system is the vector obtained by collecting the locations of all nodes:

$$x_k = [x_k^{(1)T}, \dots, x_k^{(\Omega)T}]^T. \quad (36)$$

5.1 System Model

Since nodes are assumed to be still, the model of the i th node is simply given by

$$x_k^{(i)} = x_{k-1}^{(i)} + w_k^{(i)}, \quad (37)$$

where $w_k^{(i)} \in \mathbb{R}^2$ is a zero-mean white noise vector with covariance matrix $Q_k^{(i)}$.

Note that the system is naturally fully decoupled as the state transition of a node does not depend upon other nodes. This property turns out to be very useful for the distributed formulation of the filter. Furthermore, the framework allows to mix static nodes with mobile ones simply by changing the state transition model according to the kinematics of each sensor node [31].

5.2 Observation Model

Nodes are equipped with several sensor devices. In particular, a way to measure internode distances is assumed to be available. The related observation model can be obtained considering the euclidean distance as follows:

$$z_k^{(i,j)} = h^{(i,j)}(x_k^{(i)}, x_k^{(j)}) + v_k^{(i)} \quad (38)$$

$$= \|x_k^{(i)} - x_k^{(j)}\| + v_k^{(i)} \quad (39)$$

$$= \sqrt{(p_{x,k}^{(i)} - p_{x,k}^{(j)})^2 + (p_{y,k}^{(i)} - p_{y,k}^{(j)})^2} + v_k^{(i)}, \quad (40)$$

where $v_k^{(i)} \in \mathbb{R}$ is a zero-mean white noise vector with covariance $R_k^{(i)}$.

6 THE INFORMATION FILTER FOR SENSOR NETWORKS

Due to the nonlinear nature of the observation model, the linear information filter previously introduced cannot be applied as it is. An extension to deal with the nonlinearity of the observation model is required. Note that having a linear prediction model results in a ‘‘hybrid’’ information filter: with the prediction equation of a linear IF and the estimation equation of an EIF. In the following, a centralized formulation of the filter is proposed. Then, a distributed one based on simplifying assumptions is devised. However, both filters can be summarized by the same two-stage formulation:

- Prediction

$$\begin{aligned} \Psi_{k|k-1} &= [\Psi_{k-1|k-1}^{-1} + Q_k]^{-1}, \\ L_{k|k-1} &= \Psi_{k|k-1} \Psi_{k-1|k-1}^{-1}, \\ \hat{y}_{k|k-1} &= L_{k|k-1} \hat{y}_{k-1|k-1}. \end{aligned} \quad (41)$$

- Estimation

$$\begin{aligned} \Psi_{k|k} &= \Psi_{k|k-1} + \Phi_k, \\ \hat{y}_{k|k} &= \hat{y}_{k|k-1} + \hat{i}_k, \\ \Phi_k &= J_x^T R_k^{-1} J_x^h, \\ \hat{i}_k &= J_x^T R_k^{-1} z_k', \\ z_k' &= \nu_k + J_x^h \hat{x}_{k|k-1}, \\ \nu_k &= z_k - h(\hat{x}_{k|k-1}). \end{aligned} \quad (42)$$

Differences between the centralized formulation and the distributed formulation are merely related to the state space dimension and to the construction of the Jacobian matrix J_x^h .

6.1 Centralized EIF

In the case of the centralized formulation, the whole state of the system as given in (36) is considered. Therefore, the computation of the complete Jacobian matrix involves all the interdistance measurements available over the network. In particular, given a generic observation $z_k^{(i,j)}$, representing the distance from the node i to the node j measured by the node i , the related Jacobian row is

$$J^{h(i,j)} = [\mathbf{0} \quad J_{x,i}^{h(i,j)} \quad \mathbf{0} \quad J_{x,j}^{h(i,j)} \quad \mathbf{0}], \quad (43)$$

where

$$J_{x,i}^{h(i,j)} = \begin{bmatrix} \frac{p_x^{(i)} - p_x^{(j)}}{d} & \frac{p_y^{(i)} - p_y^{(j)}}{d} \end{bmatrix} = -J_{x,j}^{h(i,j)} \quad (44)$$

with

$$d = \sqrt{(p_x^{(i)} - p_x^{(j)})^2 + (p_y^{(i)} - p_y^{(j)})^2}. \quad (45)$$

According to this notation, given the following set of observations $\mathbf{z}_k = [z_k^{(i,j)}, z_k^{(l,j)}, z_k^{(i,l)}, z_k^{(l,i)}]^T$ among three nodes $\{x_i, x_j, x_l\}$, the resulting Jacobian matrix J_x^h is

$$\begin{aligned} J_x^h &= [J^{h(i,j)T} \quad J^{h(l,j)T} \quad J^{h(i,l)T} \quad J^{h(l,i)T}]^T \\ &= \begin{bmatrix} \mathbf{0} & J_{x,i}^{h(i,j)} & \mathbf{0} & J_{x,j}^{h(i,j)} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & J_{x,l}^{h(l,j)} & J_{x,j}^{h(l,j)} & \mathbf{0} \\ \mathbf{0} & J_{x,i}^{h(i,l)} & J_{x,l}^{h(i,l)} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & J_{x,i}^{h(l,i)} & J_{x,l}^{h(l,i)} & \mathbf{0} & \mathbf{0} \end{bmatrix}. \end{aligned} \quad (46)$$

6.2 Distributed EIF

A distributed formulation can be introduced by means of some simplifying assumptions. The system model is linear and fully decoupled, thus, suitable for a distributed implementation, while the Jacobian matrix J_x^h features some couplings. In particular, for each node i , the following Jacobian block $J_x^{h(i)}$ can be considered:

$$J_x^{h(i)} = \begin{bmatrix} \mathbf{0} & J_{x,i}^{h(i,j)} & \mathbf{0} & J_{x,j}^{h(i,j)} & \mathbf{0} \\ \mathbf{0} & J_{x,i}^{h(i,l)} & J_{x,l}^{h(i,l)} & \mathbf{0} & \mathbf{0} \end{bmatrix}. \quad (47)$$

Furthermore, according to (47), it can be noticed that if a node i considers its neighbors as anchors at each time step, the partial derivatives of node j are always naughts for a generic Jacobian row $J_x^{h(i)}$. Therefore, the related Jacobian block $J_x^{h(i)}$ becomes

$$\begin{aligned} J_x^{h(i)} &= \begin{bmatrix} \mathbf{0} & J_{x,i}^{h(i,j)} & \mathbf{0} & J_{x,j}^{h(i,j)} & \mathbf{0} \\ \mathbf{0} & J_{x,i}^{h(i,l)} & J_{x,l}^{h(i,l)} & \mathbf{0} & \mathbf{0} \end{bmatrix} \\ &= \begin{bmatrix} \mathbf{0} & J_{x,i}^{h(i,j)} & \mathbf{0} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & J_{x,i}^{h(i,l)} & \mathbf{0} & \mathbf{0} & \mathbf{0} \end{bmatrix} \\ &= \begin{bmatrix} \mathbf{0} & |J_{x,i}^{h(i,i)}| & \mathbf{0} & \mathbf{0} & \mathbf{0} \end{bmatrix}. \end{aligned}$$

In this way, the complete Jacobian matrix J_x^h , described in (46), turns out to be a block matrix. Therefore, the centralized formulation can be easily decomposed into a set of Ω reduced-order filters, each one run by a single node with the aim of estimating its location with respect to information (in terms of observations and latest estimates) coming from the other nodes.

Furthermore, the capability of the algorithm to perform the localization process with or without anchors can be explained by the fact that for how the algorithm is conceived, neighbors are always considered as anchors. Thus, the availability of real anchors does not affect the formulation, but the accuracy of the localization process.

6.3 Interlaced EIF

Considering the neighborhood of each node as a set of anchors helps to distribute the formulation of the EIF. However, at the same time, an error is introduced into the estimation process as a consequence of this approximation. The Interlacement technique introduced in Section 4.1 turns out to be an effective solution to mitigate the error introduced by this simplifying assumption. The resulting formulation for the sensor network scenario is as follows:

- Prediction

$$\begin{aligned} \Psi_{k|k-1}^{(i)} &= [(\Psi_{k-1|k-1}^{(i)})^{-1} + Q_k^{(i)}]^{-1}, \\ L_{k|k-1}^{(i)} &= \Psi_{k|k-1}^{(i)} (\Psi_{k-1|k-1}^{(i)})^{-1}, \\ \hat{y}_{k|k-1}^{(i)} &= L_{k|k-1}^{(i)} \hat{y}_{k-1|k-1}^{(i)}. \end{aligned} \quad (48)$$

- Estimation

$$\begin{aligned} \Psi_{k|k}^{(i)} &= \Psi_{k|k-1}^{(i)} + \Phi_k^{(i)}, \\ \hat{y}_{k|k}^{(i)} &= \hat{y}_{k|k-1}^{(i)} + i_k^{(i)}, \\ \Phi_k^{(i)} &= J_{x,i}^{h,iT} \tilde{R}_k^{(i)-1} J_{x,i}^{h,i}, \\ i_k^{(i)} &= J_{x,i}^{h,iT} \tilde{R}_k^{(i)-1} z_k^{(i)}, \\ \tilde{R}_k^{(i)} &= R_k^{(i)} + \sum_{j \in \mathcal{N}(i)} J_{x,j}^{h,i} (\Psi_{k|k-1}^{(j)})^{-1} J_{x,j}^{h,iT}, \\ z_k^{(i')} &= \nu_k + J_{x,i}^{h,i} \hat{x}_{k|k-1}^{(i)}, \\ \nu_k^{(i)} &= z_k^{(i)} - h^{(i)}(\hat{x}_{k|k-1}^{(i)}, \xi_k^{(i)}), \end{aligned} \quad (49)$$

where $\xi_k^{(i)} = \{\hat{x}_{k|k-1}^{(j)} : j \in \mathcal{N}(i)\}$. Note that, the interlacement contribution does not add any significant complexity to the estimation process as the Jacobian term $J_{x,j}^{h,i}$ is simply obtained by negation of the term $J_{x,i}^{h,j}$ and the term $(\Psi_{k|k-1}^{(j)})^{-1}$ is broadcasted by the neighbors.

6.4 Algorithmic Derivation

From an algorithmic point of view, a possible implementation of the distributed EIF running onboard each node is given in Algorithm 1. In detail, at each iteration k , a given node i performs the following four steps: it listens for a prefixed τ amount of time waiting for new data $\{z_k^{(i)}, \Psi_k^{(i)}\}$ broadcasted by each other node within its range of visibility; successively, it updates its estimate $x_k^{(i)}$ by executing the set of equations given in (48) and (49), where the Jacobian

matrix $Jh^{(i)}$ is built, as previously described in (48), according to the collected data $\{z_k^{(i)}, \Psi_k^{(i)}\}$; finally, it notifies to the network its latest estimate $(\hat{y}_{k|k}^{(i)}, \Psi_{k|k}^{(i)})$. Note that, no clock synchronization is required for the sensor network; indeed, the temporal index k for the data coming from neighboring nodes is simply meant as the most recent available so far.

Algorithm 1. Reduced-Order Filter

Data: $\{\hat{y}_{k-1|k-1}^{(i)}, \Psi_{k-1|k-1}^{(i)}\}$
Result: $\{\hat{y}_{k|k}^{(i)}, \Psi_{k|k}^{(i)}\}$

```

/* Data Collecting */
{z_k^{(i)}, \Psi_k^{(i)}} ← listening_procedure(\tau)
where
z_k^{(i)} = {z_k^{(i,j_1)}, \dots, z_k^{(i,j_{M_i})}},
\Psi_k^{(i)} = {\hat{y}_{k-1|k-1}^{(j_1)}, (\Psi_{k-1|k-1}^{(j_1)})^{-1}, \dots, \hat{y}_{k-1|k-1}^{(j_{M_i})}, (\Psi_{k-1|k-1}^{(j_{M_i})})^{-1}}
/* Updating Step */
{\hat{y}_{k|k-1}^{(i)}, \Psi_{k|k-1}^{(i)}} ←
prediction_proc(\hat{y}_{k-1|k-1}^{(i)}, \Psi_{k-1|k-1}^{(i)})
/* Estimation Step */
{\hat{y}_{k|k}^{(i)}, \Psi_{k|k}^{(i)}} ← estimation_procedure(z_k^{(i)}, \Psi_k^{(i)})
/* Notification Step */
notification_procedure(\hat{y}_{k|k}^{(i)}, \Psi_{k|k}^{(i)})

```

7 PERFORMANCE ANALYSIS

Several computer simulations have been performed in order to investigate the effectiveness of the proposed distributed interlaced extended information filter on a large scale. Moreover, a comparison with an interlaced extended Kalman filter has been carried out as well. Note that in this work, the attention is focused on the design of an interlacement technique within the information filtering. For this reason only, a comparison between distributed versions of the algorithms is provided. The reader is referred to [16], [17] for a detailed analysis concerning a comparison with the centralized versions of the algorithms.

In particular, the following aspects of interest have been considered:

- level of noise of observations,
- scalability of the algorithm,

while the following two indexes of quality have been used:

- estimation accuracy and
- convergence velocity.

The former is given in terms of distance between the estimated and the real location of a node. The euclidean distance is adopted as metric. Maximum, minimum, and average errors computed over the whole network are considered. The latter is given in terms of number of steps required by the algorithm to settle around the best estimation. This index provides an evaluation of the “reactivity” of the algorithm.

Table 1 describes how the performance indexes vary with respect to different levels of noise. Convergence is

TABLE 1
Statistical Analysis: Indexes of Quality

| Conf | Noise Std 0.05 [m] | | Noise Std 0.2 [m] | | Noise Std 0.5 [m] | |
|-----------------|-----------------------|------|----------------------|------|----------------------|-------|
| | IEIF | IEKF | IEIF | IEKF | IEIF | IEKF |
| Max Error [cm] | 2.88 | 1.79 | 11.64 | 6.97 | 28.52 | 18.12 |
| Min Error [cm] | 0.14 | 0.06 | 0.56 | 0.24 | 1.47 | 0.62 |
| Mean Error [cm] | 1.09 | 0.57 | 4.38 | 2.30 | 11.09 | 5.92 |
| Std. Dev. [cm] | 0.60 | 0.36 | 2.40 | 1.44 | 6.00 | 3.77 |
| Conv. Step | 21 | 39 | 21 | 42 | 25 | 45 |
| % Failure | 3 | 10 | 4 | 3 | 4 | 4 |

assumed to be reached when the fluctuation of the estimate was bounded within a predefined interval, ± 0.5 cm. A configuration involving 90 nodes with 30 anchors deployed on a 30 m \times 30 m environment is considered. According to the obtained results, the accuracy of the estimation is considerably influenced by the level of noise, while it does not seem to significantly affect neither the convergence time nor the percentage of failures, i.e., the number of unsuccessful trials, of the algorithm.

Table 2 investigates the scalability of the proposed algorithm. In this analysis, the level of noise is fixed to 0.1 for all the configurations. Furthermore, anchors are supposed to be uniformly distributed and both the environmental size and the ration between the number of nodes and number of anchors are kept constant. According to the obtained results, the higher is the number of nodes, the quicker is the convergence of the algorithm. In the same way, the number of failures decreases with an increasing number of nodes. On the other hand, the accuracy of the estimation is not significantly influenced by the dimension of the sensor network, being this related to the hardware sensing capabilities.

The algorithm might converge to alternative admissible solutions. Indeed, given Ω nodes with Θ anchors, “symmetrical” solutions may exist with regard to the deployment of the anchors. In the case of perfect communication, i.e., no packet is lost, and fully connectivity among nodes, placing anchors on the boundary is a sufficient condition to have a unique solution. However, in a real system, some nodes may not be able to communicate with other nodes; therefore, in practice, alternative plausible solutions with respect to the available data may always exist.

8 EXPERIMENTAL RESULTS

Experimental results have been performed to validate the proposed distributed interlaced extended information filter in a real context. In particular, apart from the IEKF, a comparison against a third algorithm, i.e., the ESDP algorithm, has been considered. In detail, the ESDP algorithm is an SDP relaxation approach proposed in [43] for which the code is freely available at <http://www.stanford.edu/~yyye>.

The network is composed of MICAz (MPR2400) platform, a generation of Motes from Crossbow Technology. The MPR2400 (2,400-2,483.5 MHz band) uses the Chipcon CC2420, the IEEE 802.15.4 compliant, ZigBee ready radio frequency transceiver integrated with an Atmega128L

TABLE 2
Statistical Analysis: Indexes of Quality

| Conf | N:120 A:30 | | N:180 A:45 | | N:360 A:90 | |
|-----------------|------------|------|------------|------|------------|------|
| | IEIF | IEKF | IEIF | IEKF | IEIF | IEKF |
| Max Error [cm] | 5.59 | 3.53 | 4.23 | 3.18 | 4.11 | 3.01 |
| Min Error [cm] | 0.27 | 0.11 | 0.13 | 0.07 | 0.11 | 0.05 |
| Mean Error [cm] | 2.17 | 1.16 | 1.52 | 0.96 | 1.30 | 0.8 |
| Conv. Step | 20 | 45 | 17 | 31 | 15 | 27 |
| % Failure | 4 | 6 | 1 | 0 | 0 | 0 |

microcontroller. It provides also a flash serial memory, as well as a 51 pin I/O connector that allows several sensor and data acquiring boards to be connected to it. MICAz platform comes along with TinyOS, an open-source event-driven operating system designed for wireless embedded sensor networks. It features a component-based architecture, which enables rapid innovation and implementation while minimizing code size as required by the severe memory constraints inherent in sensor networks. TinyOS component library includes network protocols, distributed services, sensor drivers, and data acquisition tools, all of which can be used as is or be further refined for a custom application.

8.1 Ranging Technique

A ranging technique based on the Time of Arrival (ToA) principle is exploited to compute internode distances. The implementation consists of a node sending first an RF packet and emitting an acoustic signal right after. For the receiving node, the RF signal, whose propagation can be assumed instantaneous, is used to trigger a timer, while the acoustic signal, whose propagation delay is measurable, is used to stop it. By the measurement of such a propagation delay and by knowing the propagation velocity of the acoustic signal, the distance is then computed.

Regarding the MICAz platform, the MTS300 and MTS310 boards, both providing a sounder and a microphone, have been exploited. The sounder is a simple 4 kHz fixed frequency piezoelectric resonator, while the microphone can be used either for acoustic ranging or for general acoustic recording and measurement. Therefore, the RF and acoustic (sounder) signals are exploited for the implementation of the proposed ranging technique.

The proposed ranging technique for MICAz platforms has been thoroughly investigated in order to determine the achievable performance. A significant amount of internode distances (more than 200 measurements) was collected and a statistical analysis was performed. A precision of 3 ~ 8 cm with a standard deviation of 8 ~ 14 cm was experienced considering distances ranging from 20 cm to 2.5 m.

In addition, experiments have been carried out to verify whether the mutual orientation of nodes might influence the measured distance. For such a reason, two nodes have been arranged on the floor at the distance of 54 cm from each other. Such a distance has been manually measured from the sounder of the emitter to the microphone of the receiver. Successively, data have been collected considering different orientations of nodes, in order to simulate a realistic random deployment on the ground. Table 3 shows the statistic results using again more than 200 measurements for each configuration. According to this analysis,

TABLE 3
Internode Ranging Technique: Experimental Results

| Exp. | Mean value [m] | Std dev [m] | node 1 orientation [rad] | node 2 orientation [rad] |
|------|----------------|-------------|--------------------------|--------------------------|
| 1 | 0.5781 | 0.1229 | $\pi/2$ | $3\pi/2$ |
| 2 | 0.5734 | 0.1331 | $3\pi/2$ | 0 |
| 3 | 0.5888 | 0.1146 | $3\pi/2$ | $3\pi/2$ |
| 4 | 0.5696 | 0.1052 | $3\pi/2$ | π |
| 5 | 0.5933 | 0.1098 | $3\pi/2$ | $\pi/2$ |
| 6 | 0.6008 | 0.1230 | $5\pi/4$ | $3\pi/4$ |
| 7 | 0.5972 | 0.1217 | $5\pi/4$ | $\pi/4$ |
| 8 | 0.5853 | 0.1136 | $5\pi/4$ | $5\pi/4$ |
| 9 | 0.5683 | 0.1181 | $5\pi/4$ | $5\pi/2$ |
| 10 | 0.5892 | 0.1186 | $5\pi/4$ | π |
| 11 | 0.5786 | 0.1239 | $5\pi/4$ | $7\pi/4$ |
| 12 | 0.5668 | 0.1299 | 0 | 0 |

differential mutual orientations do not significantly influence the measure of distances. However, as mentioned above, data present a bias as well as a considerable standard deviation that makes their use challenging.

The bias and the standard deviation describe the uncertainty in the observing process. There are several sources of such uncertainty. First of all, the parameters used to characterize the propagation velocity of an acoustic wave in the air have been considered fixed, while they change according with humidity and temperature. Second, the transmission protocol introduces a delay, which cannot be taken into account, as it is not directly observable.

8.2 Deployment and Evaluation

Several network deployments have been considered for the algorithm evaluation. Each deployment has been obtained by taking advantage of the regularity of the flooring grid and real locations were manually measured exploiting such a regularity. Note that the extent of the deployment region has been constrained by the hardware capability of the MICAz nodes. Indeed, experiments reveal measurements to be sufficiently reliable only within a range of approximately 2 meters. Here, results related only to two configurations are reported.

Fig. 3 shows the first deployment, where 10 nodes are considered. Each node is ideally within the communication

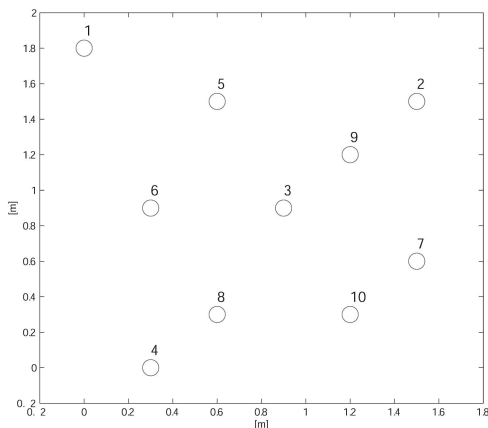


Fig. 3. First deployment: three (variable) anchors—seven nodes.

TABLE 4
First Deployment: IEIF versus IEKF versus ESDP

| Interlaced Extended Information Filter | | | | |
|--|----------------|----------------|-----------------|---------------|
| Anchors | Max Error [cm] | Min Error [cm] | Mean Error [cm] | Conv. [Steps] |
| {1,2,3} | 12.0 | 2.9 | 6.4 | 8 |
| {1,6,7} | 11.8 | 4.1 | 6.7 | 14 |
| {5,9,10} | 12.8 | 1.3 | 5.2 | 12 |
| Interlaced Extended Kalman Filter | | | | |
| Anchors | Max Error [cm] | Min Error [cm] | Mean Error [cm] | Conv. [Steps] |
| {1,2,3} | 11.1 | 4.1 | 7.9 | 25 |
| {1,6,7} | 11.0 | 2.5 | 6.5 | 26 |
| {5,9,10} | 12.7 | 1.9 | 5.6 | 24 |
| Edge-based SDP relaxation (ESDP) | | | | |
| Anchors | Max Error [cm] | Min Error [cm] | Mean Error [cm] | Conv. [Steps] |
| {1,2,3} | 7.3 | 1.6 | 5.5 | - |
| {1,6,7} | 11.7 | 2.6 | 6.5 | - |
| {5,9,10} | 11.6 | 2.5 | 6.8 | - |

range of each other. This way, a full connected graph is achieved.

Table 4 describes the result of the experiment involving the first environment (Fig. 3). Three different arrangements of anchors are considered, each one involving three nodes. According to experimental results for this configuration, varying the set of anchors does not significantly influence the accuracy of estimation. In particular, the ESDP algorithm has proved to perform slightly better for all the anchors configurations. This can be explained by the fact that being the ESDP a centralized algorithm, it can take advantage of the full availability of information. Nevertheless, it is worthy to mention that a similar estimation accuracy is achieved also by the other two algorithms despite from the partial information availability. With regard to the convergence velocity, the EIF turns out to perform slightly better. This can be explained by the fact that while the Kalman filter requires an initialization for the covariance matrix P , which slows down the convergence, the information filter does not require any initialization for the information matrix [29].

Fig. 4 shows the second deployment, where 11 nodes are considered. Again, each node is ideally within the communication range of each other in order to have a full connected graph. However, it should be pointed out that at each iteration, only a portion of the network is able to collaborate, due to the high number of outliers occurring in the measurement process. This implies that only partial information is available to the nodes.

Table 5 describes the result of the experiment involving the second environment (Fig. 4). Also in this deployment, three different arrangements of anchors are considered, each one involving three nodes. According to experimental results for this configuration, varying the set of anchors still does not significantly influence the performance. However, a general deterioration of the estimation accuracy can be noticed, due to the large number of packets lost, which considerably reduces the amount of information available for localization. Furthermore, also in this scenario, the ESDP

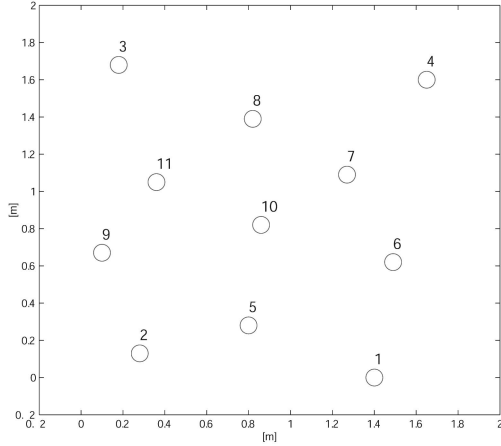


Fig. 4. Second deployment: three (variable) anchors—eight nodes.

algorithm has proved to perform slightly better. In particular, the advantages related to the availability of full information are more evident in this scenario, where a significant loss of packets was experienced.

9 COMPUTATIONAL COMPLEXITY ANALYSIS

Here, a comparative analysis regarding the computational complexity of both the IEIF previously introduced and the IEKF given in Section 4.1 is proposed. In order to achieve this, the asymptotic notation (a mathematical notation used to describe the asymptotic behavior of functions) is considered. Its purpose is to characterize a function behavior for very large (or very small) inputs in a simple but rigorous way that enables comparison to other functions [1].

Furthermore, in order to easily analyze the filter equations, a formalism has been introduced with the aim of describing the matrix operations and the related computational complexity:

TABLE 5
Second Deployment: IEIF versus IEKF versus ESDP

| Interlaced Extended Information Filter | | | | |
|--|----------------|----------------|-----------------|---------------|
| Anchors | Max Error [cm] | Min Error [cm] | Mean Error [cm] | Conv. [Steps] |
| {1,2,4} | 21.0 | 1.4 | 12.9 | 23 |
| {3,4,5} | 18.7 | 6.3 | 11.8 | 17 |
| {5,6,11} | 22.4 | 8.6 | 16.6 | 25 |

| Interlaced Extended Kalman Filter | | | | |
|-----------------------------------|----------------|----------------|-----------------|---------------|
| Anchors | Max Error [cm] | Min Error [cm] | Mean Error [cm] | Conv. [Steps] |
| {1,2,4} | 23.6 | 9.4 | 15.9 | 22 |
| {3,4,5} | 16.5 | 10.1 | 13.2 | 17 |
| {5,6,11} | 21.9 | 8.9 | 17.4 | 39 |

| Edge-based SDP relaxation (ESDP) | | | | |
|----------------------------------|----------------|----------------|-----------------|---------------|
| Anchors | Max Error [cm] | Min Error [cm] | Mean Error [cm] | Conv. [Steps] |
| {1,2,3} | 19.2 | 2.3 | 11.4 | - |
| {3,4,5} | 17.4 | 2.6 | 11.7 | - |
| {5,6,11} | 20.6 | 2.3 | 11.6 | - |

TABLE 6
Interlaced Extended Kalman Filter Computational Load

| | |
|--|--|
| $\hat{x}_{k k-1}^{(i)} = \hat{x}_{k-1 k-1}^{(i)}$ | - |
| $P_{k k-1}^{(i)} = [P_{k-1 k-1}^{(i)} + Q_k^{(i)}]$ | SUM (NxN, NxN) |
| $\hat{x}_{k k}^{(i)} = x_{k k-1}^{(i)} + K_k^{(i)} \nu_k$ | SUM (Nx1, Nx1) MUL (NxM, Mx1) |
| $\nu_k^{(i)} = z_k^{(i)} - h^{(i)}(\hat{x}_{k k-1}^{(i)}, \xi_k^{(i)})$ | SUB (Mx1, Mx1) |
| $K_k^{(i)} = P_{k k-1}^{(i)} J_x^{h,iT} (S_k^{(i)})^{-1}$ | MUL (NxN, NxM) MUL (NxM, MxM) INV (MxM) |
| $S_k^{(i)} = J_x^{h,i} P_{k k-1}^{(i)} J_x^{h,iT} + \tilde{R}_k^{(i)}$ | MUL (MxN, NxM) SUM (MxM, MxM) |
| $P_{k k}^{(i)} = (I - K_k^{(i)} J_x^{h,i}) P_{k k-1}^{(i)}$ | MUL (NxM, MxN) SUB (NxN, NxN) MUL (NxN, NxN) |
| $\tilde{R}_k^{(i)} = R_k^{(i)} + \sum_{j \in \mathcal{N}(i)} J_{x,j}^{h,i} P_{k k-1}^{(j)} J_{x,j}^{h,iT}$ | SUM (MxM, MxM) MUL (1xN, NxN) MUL (1xN, Nx1) |
| | M times |

- $\text{SUM}(NxM, NxM) = \mathcal{O}(N \cdot M)$.
- $\text{SUB}(NxM, NxM) = \mathcal{O}(N \cdot M)$.
- $\text{MUL}(NxM, MxP) = \mathcal{O}(N \cdot M \cdot P)$.
- $\text{INV}(NxN) = \mathcal{O}(N^3)$.

Note that, for the sake of simplicity, the asymptotic complexity assumed for these operations does not reflect the most efficient implementation available so far. However, it does not affect the validity of the analysis since the complexity of the most efficient implementations scale approximatively the same. Furthermore, all the elementary operations related to scalar values have been assumed with complexity $\mathcal{O}(1)$.

9.1 The IEKF

The complexity of the IEKF running onboard of a node can be summarized as in Table 6, where N is the dimension of each node state and M is the number of measurements (neighbors) for each node.

Three remarks are now in order:

- Only matrix operations have been taken into account.
- The complexity of the Jacobian construction has been neglected.
- The complexity of the observation evaluation has been neglected.

The first observation underlines that the asymptotic behavior of the algorithm is desired. The second observation comes from the consideration that the computational complexity of the Jacobian is always lighter compared to other operations. Thus, it will be omitted for the sake of clarity. The third observation follows the same reasoning as the second one.

9.2 The IEIF

The complexity of the IEIF running onboard of a node can be summarized as in Table 7.

The same considerations that have been done for the IEKF still hold here.

TABLE 7
Interlaced Extended Information Filter Computational Load

| | |
|---|---|
| $\Psi_{k k-1}^{(i)} = [(\Psi_{k-1 k-1}^{(i)})^{-1} + Q_k^{(i)}]^{-1}$ | INV (N×N) SUM (N×N, N×N) INV (N×N) |
| $L_{k k-1}^{(i)} = \Psi_{k k-1}^{(i)} (\Psi_{k-1 k-1}^{(i)})^{-1}$ | MUL (N×N, N×N) |
| $\hat{y}_{k k-1}^{(i)} = L_{k k-1}^{(i)} \hat{y}_{k-1 k-1}^{(i)}$ | MUL (N×N, N×1) |
| $\Psi_{k k}^{(i)} = \Psi_{k k-1}^{(i)} + \Phi_k^{(i)}$ | SUM (N×N, N×N) |
| $\Phi_k^{(i)} = J_x^{h,iT} (\tilde{R}_k^{(i)})^{-1} J_x^{h,i}$ | INV (M×M) (diag) MUL (N×M, M×M) (diag) MUL (N×M, M×N) |
| $\hat{y}_{k k}^{(i)} = \hat{y}_{k k-1}^{(i)} + i_k^{(i)}$ | SUM (N×1, N×1) |
| $i_k^{(i)} = J_x^{h,iT} (\tilde{R}_k^{(i)})^{-1} z_k^{(i)T}$ | MUL (N×M, M×1) |
| $z_k^{(i)T} = \nu_k^{(i)} + J_{x,i}^{h,i} x_{k k-1}^{(i)}$ | SUM (M×1, M×1) INV (N×N) MUL (N×N, N×1) |
| $\nu_k^{(i)} = z_k^{(i)} - h^{(i)}(\hat{x}_{k k-1}^{(i)}, \xi_k^{(i)})$ | SUB (M×1, M×1) |
| $\tilde{R}_k^{(i)} = R_k^{(i)} + \sum_{j \in \mathcal{N}^{(i)}} J_{x,j}^{h,i} (\Psi_{k k-1}^{(j)})^{-1} J_{x,j}^{h,iT}$ | SUM (M×M, M×M) MUL (1×N, N×N) MUL (1×N, N×1) M times |

9.3 IEKF versus IEIF

In order to find out the differences between the two algorithms, the matrix operations have been compared. Table 8, which summarizes the set of operations required by both algorithms at each iteration, can be simplified considering that from an asymptotical standpoint, some operations, such as sum, subtraction, or transposition, have a lower order than other ones, such as multiplication or inversion.

Table 9 can be further simplified considering that from an asymptotical point of view, the number of occurrences, if not related to any of the parameters of interest, does not influence the complexity of the algorithm.

Table 10 describes the subset of operations characterizing the computational complexity of the two approaches. The

TABLE 8
Computational Complexity: Comparative Table 1

| IEKF | IEIF |
|-------------------|--|
| SUM (N×N, N×N) | INV (N×N) |
| SUM (N×1, N×1) | SUM (N×N, N×N) |
| MUL (N×M, M×1) | INV (N×N) |
| SUB (M×1, M×1) | MUL (N×N, N×N) |
| MUL (N×N, N×M) | MUL (N×N, N×1) |
| MUL (N×M, M×M) | SUM (N×N, N×N) |
| INV (M×M) | INV (M×M) (diag) |
| MUL (M×N, N×M) | MUL (N×M, M×M) (diag) |
| SUM (M×M, M×M) | MUL (N×M, M×N) |
| MUL (N×M, M×N) | SUM (N×1, N×1) |
| SUB (N×N, N×N) | MUL (N×M, M×1) |
| MUL (N×N, N×N) | MUL (M×N, N×1) |
| SUM (M×M, M×M) | SUM (M×1, M×1) |
| MUL (1×N, N×N) ×M | INV (N×N) |
| MUL (1×N, N×1) ×M | MUL (N×N, N×1) SUB (M×1, M×1) MUL (1×N, N×1) MUL (1×N, N×N) ×M MUL (1×N, N×1) ×M |

TABLE 9
Computational Complexity: Comparative Table 2

| IEKF | A.C.C. | IEIF | A.C.C. |
|---|--|---|--|
| MUL (N×M, M×1) | $\mathcal{O}(N \cdot M)$ | INV (N×N) INV (N×N) MUL (N×N, N×N) MUL (N×N, N×1) | $\mathcal{O}(N^3)$ $\mathcal{O}(N^3)$ $\mathcal{O}(N^3)$ $\mathcal{O}(N^2)$ |
| MUL (N×N, N×M) MUL (N×M, M×M) INV (M×M) MUL (M×N, N×M) | $\mathcal{O}(N^2 \cdot M)$ $\mathcal{O}(N \cdot M^2)$ $\mathcal{O}(M^3)$ $\mathcal{O}(N \cdot M^2)$ | INV (M×M) (diag) MUL (N×M, M×M) (diag) MUL (N×M, M×N) | $\mathcal{O}(M)$ $\mathcal{O}(N \cdot M)$ $\mathcal{O}(N^2 \cdot M)$ |
| MUL (N×M, M×N) | $\mathcal{O}(N^2 \cdot M)$ | MUL (N×M, M×1) MUL (M×N, N×1) | $\mathcal{O}(N \cdot M)$ $\mathcal{O}(M \cdot N)$ |
| MUL (N×N, N×N) MUL (1×N, N×N) ×M MUL (1×N, N×1) ×M | $\mathcal{O}(N^3)$ $\mathcal{O}(N^2 \cdot M)$ $\mathcal{O}(N \cdot M)$ | INV (N×N) MUL (N×N, N×1) MUL (1×N, N×N) ×M MUL (1×N, N×1) ×M | $\mathcal{O}(N^3)$ $\mathcal{O}(N^2)$ $\mathcal{O}(N^2 \cdot M)$ $\mathcal{O}(N \cdot M)$ |

dominant operation for the IEIF can be either the multiplication of a matrix $N \times M$ with a matrix $M \times N$ with complexity $\mathcal{O}(N^2 \cdot M)$ or the inversion of a matrix $N \times N$ with complexity $\mathcal{O}(N^3)$, where N is the dimension of the state space and M is the number of observations. Conversely, for the IEKF, the dominant operation can be either the inversion of a matrix $N \times N$ with complexity $\mathcal{O}(N^3)$ or the inversion of a matrix $M \times M$ with complexity $\mathcal{O}(M^3)$.

The use of one technique over the other depends upon the reciprocal dimension between the state space and the observations. If the dimension of the state space is lower than the dimension of the observations $N < M$, the IEIF turns out to be computationally more efficient than the IEKF. Conversely, if the dimension of the state space is higher than the dimension of the observations $N > M$, the IEKF performs better even though the complexity is the same from an asymptotical point of view. Indeed, this is due to the fact that several operations with cubic complexity in N are required by the IEIF at each iteration. Note that for the proposed sensor network scenario, the dimension of the state space for each node is fixed to $N = 2$, while the dimension of the observations is strictly related to the number of nodes Ω deployed into the environment. Therefore, the IEIF turns out to be more effective than the IEKF. The same considerations would apply even if a three-dimensional scenario ($N = 3$) for deployment were considered. Fig. 5 shows the computational load for the two algorithms with respect to an increasing number of nodes. Note that in this analysis, the dimension of the state space for each node was fixed to $N = 2$, the ratio between the number of nodes and the number of anchors was kept constant and so was the size of

TABLE 10
Computational Complexity: Comparative Table 3

| IEKF | A.C.C. | IEIF | A.C.C. |
|---|--|--|--|
| MUL (N×M, M×1) | $\mathcal{O}(N \cdot M)$ | INV (N×N) MUL (N×N, N×N) MUL (N×N, N×1) | $\mathcal{O}(N^3)$ $\mathcal{O}(N^3)$ $\mathcal{O}(N^2)$ |
| MUL (N×N, N×M) INV (M×M) MUL (M×N, N×M) | $\mathcal{O}(N^2 \cdot M)$ $\mathcal{O}(M^3)$ $\mathcal{O}(N \cdot M^2)$ | MUL (N×M, M×N) MUL (M×N, N×1) MUL (N×N, N×1) | $\mathcal{O}(N^2 \cdot M)$ $\mathcal{O}(M \cdot N)$ $\mathcal{O}(N^2)$ |

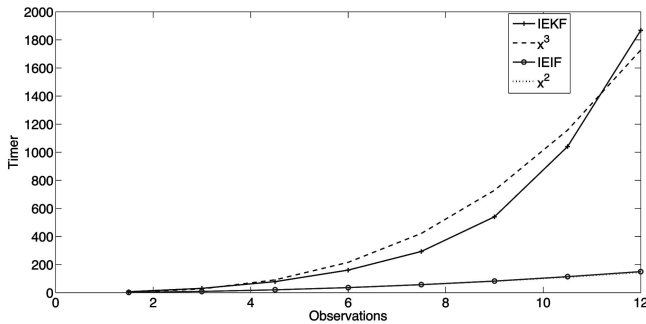


Fig. 5. Computational load: IEIF versus IEKF.

the environment. In this way, despite the random nature of the deployment, the number of observations available for each node was increasing with respect to an increasing number of nodes. According to the results given in Table 10, the computational load for the IEKF clearly shows a cubic trend while the computational load for the IEIF shows a quadratic trend.

9.3.1 Special Case: Single Observation Update

Thus far, an analysis where M observations were processed all together at each iteration has been provided. The computational complexity can be even further reduced if considering a single observation at a time. However, it should be noticed that this solution has two major drawbacks: the convergence time is significantly increased and the accuracy of the estimation can be highly affected by the order in which the measurement is processed, due to the nonlinear nature of the observations. In this case, for the IEKF, the inversion of the innovation is reduced to the inversion of a scalar. The dominant operation is given by the multiplication required for the computation of this scalar and its complexity becomes linear with the dimension of the state. However, since it has to be repeated M times, the real complexity becomes $\mathcal{O}(N \cdot M)$, which is indeed significantly lower compared to the previous one ($\mathcal{O}(M^3)$). Note that the situation for the IEIF is completely different. In fact, even if the computational load required for the construction of the Innovation matrix becomes linear with the dimension of the state, several inversions of matrices $N \times N$ are still required at each iteration. Therefore, in this case, any potential advantage simply vanishes.

10 CONCLUSIONS

In this paper, the IEIF for self-localization in sensor networks has been introduced. The centralized formulation has been distributed by neglecting any coupling factor in the system and assuming an independent reduced-order filter running onboard each node. The original formulation has been successively extended by an interlacement technique, which aims to alleviate the error introduced by neglecting the cross-correlation terms by "suitably" increasing the noise covariance matrices.

The effectiveness of the proposed formulation has been investigated via both computer simulations and real experiments involving the MICAz mote platforms produced by Crossbows. In addition, a comparison with an IEKF has been provided.

Computer simulations focused on investigating the efficacy of the proposed algorithm in a large scale. The obtained results evidence comparable performance underlying the algebraic equivalence of the two approaches. Experimental results focused on investigating the effectiveness of the proposed algorithm in a real scenario. Also in this case, the obtained results evidence comparable performance. However, according to the performed computational complexity analysis, the IEIF outperforms the IEKF anytime the dimension of the state space is lower than the dimension of the observations ($N < M$). Indeed, this is the case of the proposed sensor network scenario, where the dimension of the state space for each node is fixed to $N = 2$, while the dimension of the observations is strictly related to the number of nodes Ω deployed into the environment. Finally, the same considerations would apply even if a three-dimensional state space were considered.

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