Distributed, simple and stable network localization

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Abstract—We propose a simple, stable and distributed algorithm which directly optimizes the nonconvex maximum likelihood criterion for sensor network localization, with no need to tune any free parameter. We reformulate the problem to obtain a gradient Lipschitz cost; by shifting to this cost function we enable a Majorization-Minimization (MM) approach based on quadratic upper bounds that decouple across nodes; the resulting algorithm happens to be distributed, with all nodes working in parallel. Our method inherits the MM stability: each communication cuts down the cost function. Numerical simulations indicate that the proposed approach tops the performance of the state of the art algorithm, both in accuracy and communication cost.

Index Terms—Distributed algorithms, non-convex optimization, distributed iterative sensor localization, sensor networks, maximum-likelihood estimation.

I. PROBLEM STATEMENT

The sensor network is represented as an undirected connected graph \( G = (V,E) \). The node set \( V = \{1,2,\ldots,n\} \) denotes the sensors with unknown positions. There is an edge \( i \sim j \in E \) between sensors \( i \) and \( j \) if a noisy range measurement between nodes \( i \) and \( j \) is available at both, and if \( i \) and \( j \) can communicate with each other. The set of sensors with known positions, named anchors, is denoted by \( A = \{1,\ldots,m\} \). For each \( i \in V \), we let \( A_i \subset A \) be the subset of anchors (if any) relative to which node \( i \) also possesses a noisy range measurement.

Let \( \mathbb{R}^p \) be the space of interest (\( p = 2 \) for planar networks, and \( p = 3 \) otherwise), \( x_i \in \mathbb{R}^p \) the position of sensor \( i \), and \( d_{ij} \) the noisy range measurement between sensors \( i \) and \( j \), known by both \( i \) and \( j \). Without loss of generality, we assume \( d_{ij} = d_{ji} \). Anchor positions are denoted by \( a_k \in \mathbb{R}^p \). Similarly, \( r_{ik} \) is the noisy range measurement between sensor \( i \) and anchor \( k \), available at sensor \( i \).

The distributed network localization problem addressed in this work consists in estimating the sensors’ positions \( x = \{x_i : i \in V\} \), from the available measurements \( \{d_{ij} : i \sim j\} \cup \{r_{ik} : i \in V, k \in A_i\} \), through collaborative message passing between neighboring sensors in the communication graph \( G \).

Under the assumption of zero-mean, independent and identically-distributed, additive Gaussian measurement noise, the maximum likelihood estimator for the sensor positions is the solution of the optimization problem

\[
\minimize f(x), \tag{1}
\]

where

\[
f(x) = \sum_{i \sim j} \frac{1}{2}(\|x_i-x_j-d_{ij}\|^2) + \sum_i \sum_{k \in A_i} \frac{1}{2}(\|x_i-a_k-r_{ik}\|^2).
\]

Problem (1) is nonconvex and difficult to solve. Even in the centralized setting (i.e., all measurements are available at a central node) currently available iterative techniques don’t claim convergence to the global optimum. Also, even with noiseless measurements, multiple solutions might exist due to ambiguities in the network topology itself (1).

Related work: The literature on sensor network localization can be divided on centralized and distributed approaches; the first category encompasses methods that use a central processing node, which collects all measurements and estimates the sensor positions. Distributed approaches require that each node computes its own position, working only with data collected locally and exchanged with neighbors. The body of work on centralized approaches to the problem is vast (e.g., [2]–[6]). At a smaller scale, distributed techniques based on convex relaxations of the problem are also abundant (for example, [7]–[11]). But distributed, and maximum likelihood (thus nonconvex) approaches to the sensor network localization problem are much less common. The algorithm presented in [12] is a nonlinear Gauss-Seidel approach: only one node works at a time and solves a source localization problem with neighbors playing the role of anchors. The nodes activate sequentially in a round-robin scheme. Thus, the time to complete just one cycle becomes proportional to the network size. Parallel algorithms — the ones we are interested in this paper — avoid altogether this issue, as all nodes operate simultaneously; moreover, adding or deleting a node raises no special synchronization concern. The work presented in [13] puts forward a two-stage algorithm which is parallel: in a first consensus phase, a Barzilai-Borwein (BB) step size is calculated, followed by a local gradient computation phase. It is known that BB steps do not necessarily decrease the objective function; as discussed in [14], an outer globalization scheme involving line searches is needed to ensure its stability. However, line searches are cumbersome to implement in a distributed setting and are, in fact, absent in [13]. Further, the algorithm requires the step size to be computed via consensus, and thus the number of consensus rounds needed is a parameter to tune. We will present an algorithm with simple implementation which is both parallel and stable, with no free parameters. We will compare experimentally the performance.
of our method with the distributed, parallel, state of the art method in [13] in Sec. [16].

Contributions: We tackle directly the nonconvex problem in (1), with a simple and efficient algorithm which:
1) is parallel;
2) does not involve any free parameter definition;
3) is proven not to increase the value of the cost function at each iteration;
4) has better performance in positioning error and cost value than the state of the art method, while expending less in communications.

The first and second claims are addressed in Sec. III and the last one in Sec. V, dedicated to numerical experiments.

II. PROBLEM REFORMULATION

We can reformulate Problem (1) as

\[ \min_{x_i, y_{ij}, w_{ik}} \frac{1}{2} \sum_{i \neq j} \|x_i - x_j - y_{ij}\|^2 + \sum_{i} \frac{1}{2} \sum_{j \in A_i} \|x_i - a_k - w_{ik}\|^2 \]

subject to \( \|y_{ij}\| = d_{ij}, \|w_{ik}\| = r_{ik}, \)

and rewrite (2) as

\[ \min_{x_i, y_{ij}, w_{ik}} \frac{1}{2} \|Ax - y\|^2 + \sum_{i} \frac{1}{2} \|x_i \otimes 1 - a_i - w_i\|^2 \]

subject to \( \|y_{ij}\| = d_{ij}, \|w_{ik}\| = r_{ik}, \)

with concatenated vectors \( x = (x_i)_{i \in V}, y = (y_{ij})_{i \sim j}, a_i = (a_{ik})_{k \in A_i}, \) and \( w_i = (w_{ik})_{k \in A_i}. \) In [3], the symbol 1 stands for the vector of ones. Matrix \( A \) is the result of the Kronecker product of the arc-node incidence matrix \( G \) with the identity matrix \( I_p: A = C \otimes I_p. \) Problem (3) is equivalent to

\[ \min_{x_i, y_{ij}, w_{ik}} \frac{1}{2} \| [A - I] \begin{bmatrix} x \\ y \\ w \end{bmatrix} \|^2 + \frac{1}{2} \| Ex - \alpha - w \|^2 \]

subject to \( \|y_{ij}\| = d_{ij}, \|w_{ik}\| = r_{ik}, \)

where \( \alpha = (a_i)_{i \in V}, w = (w_i)_{i \in V}, \) and \( E \) is a matrix with zeros and ones, selecting the entries in \( \alpha \) and \( w \) corresponding to each sensor node. We now collect all the optimization variables in \( z = (x, y, w), \) and rewrite our problem as

\[ \min_{z} \frac{1}{2} \| [A - I] 0 \| z \|^2 + \frac{1}{2} \| E 0 - I \| z - \alpha \|^2 \]

subject to \( z \in Z, \)

where \( Z = \{ z = (x, y, w) : \|y_{ij}\| = d_{ij}, i \sim j, w_{ik} = r_{ik}, i \in V, k \in A_i \}. \) Problem (4) can be written as

\[ \min_{z} f(z) = \frac{1}{2} z^T M z - b^T z \]

subject to \( z \in Z, \)

for \( M \) and \( b \) defined as

\[ M = M_1 + M_2, \quad b = \begin{bmatrix} E^T \\ 0 \end{bmatrix} \alpha, \]

\[ M_1 = \begin{bmatrix} A^T \\ -I \end{bmatrix} [A - I 0], \quad M_2 = \begin{bmatrix} E^T \\ 0 \end{bmatrix} \begin{bmatrix} E & 0 \end{bmatrix} - I. \]

III. MAJORIZATION-MINIMIZATION

To solve Problem (4) in a distributed way we must deal with the complicating off-diagonal entries of \( M \) that couple the sensors’ variables. We emphasize a simple, but key fact:

Remark 1. The function optimized in Problem (4) is quadratic in \( z \) and, thus, has a Lipschitz continuous gradient [15], i.e.,

\[ \| \nabla f(x) - \nabla f(y) \| \leq L \| x - y \|, \]

for some \( L \) and all \( x, y. \)

From this property of function \( f \) we can obtain the upper bound (also found in [15]) \( f(z') \leq f(z^t) + \langle \nabla f(z^t), z - z' \rangle + \frac{L}{2} \| z - z' \|^2, \) for any point \( z' \) and use it as a majorizer in the Majorization-Minimization framework [16]. This majorizer decouples the variables and allows for a distributed solution. Our algorithm is simply:

\[ z^{t+1} = \arg \min_{z \in Z} f(z') + \langle \nabla f(z^t), z - z' \rangle + \frac{L}{2} \| z - z' \|^2. \] (7)

The solution of (7) is the projected gradient iteration [15]

\[ z^{t+1} = P_Z \left( z^t - \frac{1}{L} \nabla f(z^t) \right), \] (8)

where \( P_Z(p) \) is the projection of point \( p \) onto \( Z. \) The gradient in (8) can be easily computed as the affine function \( \nabla f(z) = M z - b. \) See the recent work [17] for interesting convergence properties of the recursion (8). Particularly, we emphasize that the cost function is non increasing per iteration.

We now compute a Lipschitz constant \( L \) for the gradient of the quadratic function in Problem (4), such that it is easy to estimate in a distributed way.

\[ L = \lambda_{\text{max}}(M) \]

\[ \leq \lambda_{\text{max}}(M_1) + \lambda_{\text{max}}(M_2) \]

\[ = \lambda_{\text{max}}(AA^T + I) + \lambda_{\text{max}}(EE^T + I) \]

\[ \leq \lambda_{\text{max}}(A^T A) + \lambda_{\text{max}}(EE^T) + 2 \]

\[ \leq 2\delta_{\text{max}} + \max_{i \in V} |A_i| + 2, \]

where \( \lambda_{\text{max}} \) denotes the largest eigenvalue, \( |A| \) is the cardinality of set \( A, \) and \( \delta_{\text{max}} \) is the maximum node degree of the network. We note that \( \lambda_{\text{max}}(A^T A) \) is the maximum eigenvalue of graph \( G \) laplacian matrix; the proof that it is upper-bounded by \( 2\delta_{\text{max}} \) can be found in [18]. This Lipschitz constant can be computed in a distributed way by, e.g., a diffusion algorithm (c.f. [19] Ch. 9)).
Algorithm 1 Distributed nonconvex localization algorithm

Input: $x_0; L; \{d_{ij} : j \in N_i\}; \{r_{ik} : k \in A_i\};$
Output: $\hat{x}$

1: set $y_{ij} = P_{Y_{ij}}(x_0^i - x_0^j)$, $Y_{ij} = \{y : \|y\| = d_{ij}\}$ and $w_{ik}^0 = P_{W_{ik}}(x_0^i - a_k)$, $W_{ik} = \{w : \|w\| = r_{ik}\}$
2: $t = 0$
3: while some stopping criterion is not met, each node $i$ do
4: $x_{i}^{t+1} = b_i x_i^t + \frac{1}{L} \sum_{j \in N_i} \left( x_j^t + C_{(i \sim j,i)} y_{ij}^t \right) + \frac{1}{L} \sum_{k \in A_i} (w_{ik}^t + a_k)$
5: for all neighboring $j$, compute $y_{jk}^{t+1} = P_{Y_{ij}} \left( \frac{1}{L} y_{ij}^t + \frac{1}{L} C_{(i \sim j,i)} (x_i^t - x_j^t) \right)$
6: for each of the connected anchors $k \in A_i$, compute $w_{ik}^{t+1} = P_{W_{ik}} \left( \frac{L}{L-1} w_{ik}^t + \frac{1}{L}(x_i - a_{ik}) \right)$
7: broadcast $x_{i}^{t+1}$ to neighbors
8: $t = t + 1$
9: end while
10: return $\hat{x}_i = x_i^t$

IV. DISTRIBUTED SENSOR NETWORK LOCALIZATION

At this point, the recursion in Eq. (3) is already distributed, as detailed below. From Eq. (3) we will obtain the update rules for the variables $x$, $y$, and $w$. For this we write matrix $M$ as follows:

$$M = \begin{bmatrix} A^T A + E^T E & -A^T & -E^T \\ -A & I & 0 \\ -E & 0 & I \end{bmatrix}, \quad (10)$$

and denote $B = A^T A + E^T E$. Then, each block of $z$ is updated according to

$$x^{t+1} = \left( I - \frac{1}{L} B \right) x^t + \frac{1}{L} A^T y^t + \frac{1}{L} E^T (w^t + \alpha), \quad (11)$$

$$y^{t+1} = P_Y \left( \frac{L-1}{L} y^t + \frac{1}{L} A x^t \right), \quad (12)$$

$$w^{t+1} = P_W \left( \frac{L-1}{L} w^t + \frac{1}{L} E x^t - \frac{\alpha}{L} \right), \quad (13)$$

where $Y$ and $W$ are the constraint sets associated with the acquired measurements between sensors, and between anchors and sensors, respectively, and $N_i$ is the set of the neighbors of node $i$. We observe that each block of $z = (x, y, w)$ at iteration $t+1$ will only need local neighborhood information, as clarified in Algorithm 1. Each node $i$ will update the current estimate of its own position, each one of the $y_{ij}$ for all the incident edges $i \sim j$ and the anchor terms $w_{ik},$ if any. The symbol $C_{(i \sim j,i)}$ denotes the arc-node incidence matrix entry relative to edge $i \sim j$ (row index) and node $i$ (column index). The constant $b_i$ in step 4 of Algorithm 1 is defined as $\frac{L-1}{L} - |A_i|$. 

V. EXPERIMENTAL RESULTS

We present numerical experiments to ascertain the performance of the proposed Algorithm 1, both in accuracy and in communication cost. Accuracy will be measured in 1) mean positioning error per sensor, defined as

$$MPE = \frac{1}{n \cdot MC} \sum_{m=1}^{MC} \sum_{i=1}^{n} \| \hat{x}_i(m) - x_i^* \|, \quad (14)$$

where $MC$ is the total number of Monte Carlo trials, $\hat{x}_i(m)$ is the estimate generated by an algorithm at the Monte Carlo trial $mc$, and $x_i^*$ is the true position of node $i$, and 2) also by evaluating the cost function in (1), averaged by the Monte Carlo trials and number of sensors, as in (14). Communication cost will be measured taking into account that each iteration in Algorithm 1 involves communicating $pn$ real numbers. We will compare the performance of the proposed method with the Barzilai-Borwein algorithm in [13], whose communication cost per iteration is $n(2T + p)$, where $T$ is the number of consensus rounds needed to estimate the Barzilai-Borwein step size. We use $T = 20$ as in [13]. The setup for the experiments is a geometric network with 50 sensors randomly distributed in the two-dimensional square $[0, 1] \times [0, 1]$, with average node degree of about 6, and 4 anchors placed at the vertexes of this square. The network remains fixed during all the Monte Carlo trials. Both algorithms receive an initialization from a convex approximation method. The initialization will hopefully hand the nonconvex refinement algorithms a point near the basin of attraction of the true minimum. For this purpose we generate noisy range measurements according to

$$d_{ij} = \|x_i^* - x_j^*\| + \nu_{ij}, \quad r_{ik} = \|x_i^* - a_k\| + \eta_{ik},$$

where $\{\nu_{ij} : i \sim j \in E\} \cup \{\eta_{ik} : i \in V, k \in A_i\}$ are independent gaussian random variables with zero mean and standard deviation $\sigma$. We conducted 100 Monte Carlo trials for each standard deviation $\sigma = (0.01, 0.05, 0.1)$. If we spread the sensors by a squared area with side of 1Km, this means measurements are affected by noise of 10m, 50m, and 100m. In terms of mean positioning error per sensor the proposed algorithm fares better than the benchmark: Table I shows the mean error defined in (14) after the algorithms have stabilized, or reached a maximum iteration number. In a square with 1Km sides, we improve the accuracy of the gradient descent with Barzilai-Borwein steps by about 1m per sensor, even for high power noise. Fig. 1 depicts the averaged evolution of the error per sensor of both algorithms as a function of the volume of accumulated communications, and also the evolution of the cost. The gradient descent with Barzilai-Borwein steps shows an irregular pattern for the error, only vaguely matching the variation in the corresponding cost (Fig. 1b), thus leaving some uncertainty on when to stop the algorithm and what.

| $\sigma$ | Proposed method | BB method |
|---------|----------------|-----------|
| 0.01    | 0.0053         | 0.0059    |
| 0.05    | 0.0143         | 0.0154    |
| 0.10    | 0.0210         | 0.0221    |
VI. CONCLUDING REMARKS

The monotonicity of the proposed method is a strong feature for applications of sensor network localization. Our method proves to be not only fast and resilient, but also simple to implement and deploy, with no free parameters to tune. The steady accuracy gain over the competing method also makes it usable in contexts with a wide range of measurement errors are expected. The presented method can be useful both as a refinement algorithm and as a tracking method, e.g., for mobile robot formations where position estimates computed on a given time step are used as initialization for the next one.

(a) The proposed method improves the comparing algorithm, both in accuracy and communication cost. Our proposed method improves the state of the art method in [13] by about 60 cm in mean positioning error per sensor, delivering a no surprises, stable progression of the error of the estimates.

(b) The final costs are, for the BB method, $1.7392 \times 10^{-4}$ and, for the proposed method $1.5698 \times 10^{-4}$. A small difference in cost that translates into a considerable distance in error, as depicted in Fig. 1(a) and Table 1.

Fig. 1. Noisy distance measurements with $\sigma = 0.01$, representing 10m for a square with 1Km sides. The proposed method shows a faster and smoother progression, while the comparing method bounces, always above the proposed method.

(a) For medium noise power the algorithms’ performance comparison follows the one under low noise power. The accuracy gain is more than 1m per sensor.

(b) Under medium noise the proposed method also reaches a smaller value for the average cost per sensor: 0.0031, against 0.0032 from the BB method.

Fig. 2. Distance measurements contaminated with noise, with $\sigma = 0.05$, representing 50m for a square with 1Km sides. The proposed method continues to outperform the comparing state of the art method, contrasting the instability of the BB method.

(a) The proposed algorithm tops the comparing method in error, under high noise power, by more than 1m, when considering a squared deployment area of 1Km sides.

(b) Under heavy noise the proposed method reaches a smaller value for the average cost per sensor: 0.0096, against 0.0099 from the BB method.

Fig. 3. Distance measurements contaminated with noise, with $\sigma = 0.1$, representing 100m for a square with 1Km sides.
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