On Accurate and Efficient Statistical Counting in Sensor-Based Surveillance Systems

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Abstract

Sensor networks have been used in many surveillance systems, providing statistical information about monitored areas. Accurate counting information (e.g., the distribution of the total number of targets) is often important for decision making. As a complementary solution to double-counting in communication, this paper presents the first work that deals with double-counting in sensing for wireless sensor networks. The probability mass function (pmf) of target counts is derived first. This, however, is shown to be computationally prohibitive when a network becomes large. A partitioning algorithm is then designed to significantly reduce computation complexity with a certain loss in counting accuracy. Finally, two methods are proposed to compensate for the loss. To evaluate the design, we compare the derived probability mass function with ground truth obtained through exhaustive enumeration in small-scale networks. In large-scale networks, where pmf ground truth is not available, we compare the expected count with true target counts. We demonstrate that accurate counting within 1 ~ 3% relative error can be achieved with orders of magnitude reduction in computation, compared with an exhaustive enumeration-based approach.

1 Introduction

Wireless sensor networks have been widely used to monitor many types of environments such as battlefields [1], buildings [16] and habitats [19, 21]. One of the key design objectives of these monitoring systems is to acquire and verify information about the number of targets/events within the system at any given point of time. For example, (i) in a battlefield, a commander needs to estimate enemy capability by counting different types of targets in an area to issue a counter-force attack strategically; (ii) in a building, a manager might want to turn off some facilities if the number of people in a certain area is less than a certain threshold; (iii) in geyser fields monitoring, the number of eruptions indicates the activity pattern underneath. In all these cases, although it is not necessary to have precise counting information, it is important to obtain reasonable total count values through a sensor network.

In general, there are two types of errors that would lead to inaccurate counts: miss-detection and double-counting. Miss-detection is normally addressed by using reliable sensing hardware [4] and/or robust detection algorithms [13, 8, 22], while double-counting is a more challenging problem, because it involves duplicates in both communication and sensing. Several excellent projects have investigated how to avoid the double-counting problem in communication. For example, synopsis diffusion [14] uses energy-efficient multi-path routing schemes to transmit order-and duplicate-insensitive (ODI) data aggregates. Recently, CountTor-
2 Related Work

To obtain accurate target counts, a monitoring sensor system shall prevent miss-detection as well as double-counting. Miss-detection can be reduced by introducing reliable hardware design. For example, XSM notes [4] incorporate a band-pass filter to enhance the detection of acoustic emission, a digital potentiometer to detect a wide range of signals, and a polyethylene film to reduce the effect of sunlight. Besides hardware enhancements, advanced detection algorithms [13, 8, 22] have also been proposed to avoid mis-detection with minimal energy consumption. VigilNet [8] utilizes a multi-level detection algorithm with in-situ adaptive thresholds to avoid both false positive and false negative detections in changing weather conditions. Feng et al. [22] propose a collaborative tracking algorithm with distributed Bayesian estimation to improve reliability based on current and previous estimation (beliefs) from sets of sensors.

Even with reliable detection at individual nodes, accurate total counts would not be obtained if a target is counted multiple times (double-counting). Double-counting problem has been investigated in the context of communication. The summarization of total counts without duplicates could be achieved by building a spanning tree rooted at the base. Individual counts are aggregated along a tree from leaves to the root as suggested in TAG [12]. However, this spanning-tree-based approach could suffer loss of counts severely, due to node or communication failures. For example, a single node failure could lose the count of a whole subtree beneath it. To address this limitation, synopsis diffusion [14] utilizes multi-path routing to deliver count information. The authors prove that duplicate-insensitive (ODI) count aggregation can be achieved by using Flajolet and Martin’s algorithm (FM) [6], which counts distinct elements in a multiset. Recently, CountTorrent [10] investigates in the context of communication. The summarization of total counts without duplicates could be achieved by building a spanning tree rooted at the base. Individual counts are aggregated along a tree from leaves to the root as suggested in TAG [12]. However, this spanning-tree-based approach could suffer loss of counts severely, due to node or communication failures. For example, a single node failure could lose the count of a whole subtree beneath it. To address this limitation, synopsis diffusion [14] utilizes multi-path routing to deliver count information. The authors prove that duplicate-insensitive (ODI) count aggregation can be achieved by using Flajolet and Martin’s algorithm (FM) [6], which counts distinct elements in a multiset. Recently, CountTorrent [10] allocates binary labels to individual nodes using an Abstract Prefix Tree and disseminates the (label,count) pairs through multi-path routing. Count values are aggregated only when two binary labels differ only in their last bit. Labeled aggregates ensure all values are counted only once during communication.

Although double-counting can be eliminated in communication, the final aggregated count could still be incorrect, if the targets within overlapping regions are counted more than once. According to [7], the percentage of overlapping region in sensor networks under random deployment is indeed significant. For example, with an average node density of 5, the overlapping percentage is 86% and with an average node density of 14, the overlapping percentage would be as high as 99.9%! Therefore, we argue double-counting is common in sensing and hence needs to be addressed accordingly.

3 Problem Definition and Assumptions

We consider a network model where counting sensor nodes are randomly deployed in a region (e.g., an open area or a room in a building) with known locations [18, 11]. They are used to monitor different types of targets, such as vehicles on the road, people in the room, or any other objects of interests. Counting capability is supported, using photoelectric-based sensors such as the one in [17]. The count values at individual sensors are reported to a base node, where the probability mass function (pmf) of the total number of distinct targets is calculated. Since the system-wide total count is the objective, a centralized solution at the base is a natural approach for sensor networks, which is also compatible with counting communication methods in TAG [12], Synopsis Diffusion [14], and CountTorrent [10].

To simplify the description, the sensing range of these nodes are treated as circles. It should be noted that the accuracy of our method only depends on the size of the area, not the shape of the area. In case of irregular sensing areas, methods proposed in [9] shall be used to obtain the size of sensing areas.

This work assumes spatial distribution of targets within the area is known (e.g., complete spatial randomness, spatial aggregation or spatial inhibition). Without loss of generality, we use Poisson distribution [20, 15, 2] as a concrete exemplary distribution to present our methodology through the paper. We expect our high-level idea can be applied to non-Poisson distributions, although mathematical derivation would be quite different.

Under the Poisson distribution, targets are uniformly distributed in the area of interest with intensity of \( \lambda \). The \( \lambda \) value can be either known a prior or estimated online (as we explain later). The probability that there are \( k \) targets in the region \( s \) of size \( S \) can be computed as follows:

$$P(N(s) = k) = \frac{e^{-\lambda S} (\lambda S)^k}{k!}. \quad (1)$$

Suppose there are in total \( N \) sensor nodes. The \( i_{th} \) sensor node \( v_i \) whose sensing area is circle \( C_i \) has detected \( n_i \) targets in its sensing range, where \( 1 \leq i \leq N \). Suppose the \( N \) sensing circles of these nodes divide the whole area into \( M \) non-overlapping subareas. Each subarea, \( M_k \), where \( 1 \leq k \leq M \), may belong to one or more circles. As a result, each circle is the union of a subset of all these subareas. We say \( M_k \subset C_i \) if \( M_k \) is within the subset of the \( i_{th} \) circle \( C_i \). If we further use \( N(M_k) \) to denote the number of distinct targets in subarea \( M_k \), we will have the following equation:

$$n_i = N(C_i) = \sum_{M_k \subset C_i} N(M_k). \quad (2)$$

Since the subareas are non-overlapping, the total number of distinct targets detected by the \( N \) sensor nodes (denoted as \( T \)) will be equal to the sum of the number of targets in each subarea, which can be computed by the following equation:

$$T = N(\bigcup_{i=1}^{N} C_i) = \sum_{k=1}^{M} N(M_k). \quad (3)$$

The objective of this work is to find the probability mass function (pmf) of the total target count \( T \) in Equation (3), given the individual counts \( n_i \) from all nodes, with low computation complexity. Particularly, the probability that the total number of distinct targets equals to \( t \) given the count information can be expressed as

$$P(T = t | N(C_1) = n_1, N(C_2) = n_2, \cdots, N(C_N) = n_N).$$

![Figure 1. A simple example: Two-Circle Case](image-url)
calculate the probability that there are in total two (or one) distinct targets in this area, respectively. For simplicity, we define notation $<m_1,m_2,m_3>$ as the joint probability that $M_1$ has $m_1$ targets, $M_2$ has $m_2$ targets and $M_3$ has $m_3$ targets. An example is shown as follows:

$$<1,0,1> = P(N(M_1) = 1)P(N(M_2) = 0)(N(M_3) = 1) = 1.$$ (5)

Using the definition of conditional probability, from (2), (3) and (4), we get,

$$P(T = 2|N(C_1) = 1,N(C_2) = 1) = \frac{P(T = 2,N(C_1) = 1,N(C_2) = 1)}{P(N(M_1) = 1,N(C_2) = 1)} = \frac{P(N(M_1) = 1,N(M_2) = 0,N(M_3) = 1)}{\sum_{k=0}^{1}P(N(M_1) = 1 - k,N(M_2) = k,N(M_3) = 1 - k) < 1,0,1>} = \frac{\sum_{k=0}^{1} <1 - k,k,1 - k> < 1,0,1> = <1,0,1> + <0,1,0> = \frac{1}{2}.}$$ (6)

The penultimate equality holds because $M_1$, $M_2$ and $M_3$ are non-overlapping subareas and the numbers of targets in these subareas are independent random variables. Since each term in Equation (6) can be computed using Equation (1), we can finally compute the probability for any given m and thus compute the conditional pmf of the total number of distinct targets.

Noting that all the terms in the denominator of Equation (6) ($<1,0,1>$ and $<0,1,0>$) are the probability of possible solutions that satisfy the count condition of all circles ($N(C_1) = 1,N(C_2) = 1$) while the numerator ($<1,0,1>$) is the probability of the only solution that satisfies both the count condition and the total number of distinct targets condition ($N(C_1) = 1,N(C_2) = 1, T = 2$). Similarly, we can derive the equation for a more general case, i.e., for a region that has been divided into $M$ subareas by $N$ sensor nodes, Equation (4) can be computed as

$$P(T = t|N(C_1) = n_1,N(C_2) = n_2,\cdots,N(C_N) = n_N) = \frac{P(T = t,N(C_1) = n_1,N(C_2) = n_2,\cdots,N(C_N) = n_N)}{P(N(C_1) = n_1,N(C_2) = n_2,\cdots,N(C_N) = n_N)} = \frac{\sum_{\{m_k\} \in A(A|B) <m_1,m_2,\cdots,m_M>} \sum_{\{m_k\} \in A} <m_1,m_2,\cdots,m_M>}{<1,m_1,\cdots,m_M>}$$ (7)

where $<m_1,m_2,\cdots,m_M>$ is defined similarly as before, but extended to a more general case, $\{m_k\}$ denotes the set of $\{m_1,m_2,\cdots,m_M\}$, $\{m_k\} \in A$ means for each term in the denominator, the corresponding $\{m_k\}$ satisfies the count condition of the $N$ circles so that it is a solution to a set of equations $A$ defined as follows:

$$A: \begin{cases} \sum_{M_k \subset C_1} m_k = n_1 \\
\sum_{M_k \subset C_2} m_k = n_2 \
\vdots \\
\sum_{M_k \subset C_N} m_k = n_N \\
N(M_k) = m_k \geq 0, \forall 1 \leq k \leq M
\end{cases}$$ (8)

Also, for each term in the numerator, the corresponding $\{m_k\}$ satisfies both the count condition and the total number condition. As a result, each $\{m_k\}$ is a solution to both $A$ and $B$ where $A$ is defined in Equation (8) and $B$ is defined as follows:

$$B: m_1 + m_2 + \cdots + m_M = t$$ (9)

### 4 Complexity Analysis

In order to determine the conditional pmf of the total number of distinct targets $T$, we need to compute the probability in Equation (7) for every possible value of $t$. Several interesting observations can be captured from Equation (7). First, the numerator is actually a subset of the denominator for a particular value of $t$. All these subsets are disjoint and sum to the denominator, which is consistent with the fact that all the values of the probability mass function sum up to 1. Second, in order to compute the denominator, we need to solve the equations of $A$ and try to find all the solutions. As a result, the complexity of computing the probability for a single value of $t$ is exactly the same as computing the whole probability mass function since we need to find all the solutions to $A$ anyway. Third, since all the variables in $A$ are non-negative integers, we have to exhaustively list all the solutions of $A$. Based on these observations, we develop an algorithm using an exhaustive enumeration-based method to compute Equation (7) as shown in Algorithm (1).

The complexity of finding the conditional pmf using Algorithm (1) can be computed as

$$f_1 = O\left(\prod_{k=1}^{M} UB(m_k)\right)$$ (10)

where $UB(m_k)$ is defined as the maximum possible number of targets in subarea $M_k$ and can be computed as

$$UB(m_k) = \max\{n_i\} \text{ where } M_k \subset C_i.$$ (11)

Obviously, $f_1$ is an exponential function of $M$. Remember $M$ is the number of non-overlapping subareas divided by the $N$ circles. As a result, $M$ would be much greater than $N$. For example, even in a linear network where sensor nodes are uniformly deployed, $M$ is almost twice of $N$. Exponential complexity makes it prohibitive for Algorithm (1) to obtain an accurate count in large-scale sensor networks, using a reasonable amount of time.

### 5 Partitioning Design

In the previous section, we have concluded that a large $M$ value makes the computation time intolerable, which also indicates that reducing $M$ can significantly reduce computation complexity. Figure (2) shows that $N$ circles belong to two disjoint groups $G_1$ and $G_2$ at initial deployment time. We note this deployment rarely happens in a dense network, however, we use this example to show the power of partitioning in reducing complexity. Suppose the numbers of subareas in $G_1$ and $G_2$ are $M_{G_1}$ and $M_{G_2}$, respectively. Since $G_1$...
and \( G_2 \) are disjoint, the total numbers of distinct targets in \( G_1 \) and \( G_2 \) (denoted by \( T_1 \) and \( T_2 \)) are independent random variables. As a result, we can compute the pmf of \( T_1 \) and \( T_2 \) separately and then combine the two functions to compute the pmf of \( T \) which is equal to the sum of these two independent random variables: \( T = T_1 + T_2 \). The method used to combine two independent distributions can be found in textbooks [3] and will not be discussed. Here we are interested in how much complexity can be reduced by partitioning. It’s clear that the complexity of the combination process is the product of the sizes of the sample space of \( G_1 \) and \( G_2 \). This value is negligible compared to the complexity of the enumeration process. Thus, the total complexity of this method can be computed in Equation (12).

\[
f_2 = O\left(\prod_{k=1}^{M_1} (UB(m_k)) + \prod_{k=M_1+1}^{M_1+M_2} (UB(m_k))\right). 
\] (12)

If the values of \( M_1 \) and \( M_2 \) are similar in \( G_1 \) and \( G_2 \), \( f_2 \) is much less than \( f_1 \), especially when \( M \) is large. Let’s compare \( f_2 \) with \( f_1 \) using the example shown in Figure (2). Suppose all sensors detect \( n \) targets for simplicity. If we compute the conditional pmf using Equation (7) directly, the cost is \( O(n^{22}) \). If we compute the pmf for \( G_1 \) and \( G_2 \) separately and then combine them to get the total, the complexity according to Equation (12) is \( O(n^{11}) \). Generically, if multiple disjoint groups exist in the area and the maximum size of each group \( G_i \) is bounded, the computation complexity is:

\[
f_3 = O(M \prod_{k=1}^{MAX} UB(m_k))
\] (13)

where MAX is the maximum number of subareas of each group. It’s obvious that the \( f_3 \) is a polynomial function of \( M \).

### 5.1 Deleting Zero Count Circles

In the previous section, we have shown that disjoint groups reduce the complexity significantly. However, given a space covered by sensor nodes, it’s not always the case that the circles are disjoint. Therefore, it is necessary to partition the nodes into groups as well as compensate for the loss of accuracy caused by partitioning.

Recall that \( <m_1,m_2,m_3> \) is defined as the probability that \( M_1 \) has \( m_1 \) targets, \( M_2 \) has \( m_2 \) targets and \( M_3 \) has \( m_3 \) targets. Since \( N(M_k) \) is all independent due to the fact that \( M_k \)s are non-overlapping, the decomposability of \( <m_1,m_2,m_3> \) can be easily derived from Equation (5) as follows:

\[
< m_1,m_2,m_3 > = < m_1,m_2,* > < *,*,m_3 > = < m_1,*,* > < *,m_2,* > < *,*,m_3 >
\] (14)

where the symbol "*" means the number within the corresponding subarea can be any value. Based on this property, the effect of eliminating a zero-count node can be studied.

Suppose node \( v_1 \), whose sensing circle is \( C_1 \) as shown in Figure (3), detects zero targets, which means \( n_1 = 0 \). It’s possible that there still exist other zero-count circles besides \( v_1 \) but they would not affect the discussion here. From Equation (2) we have

\[
n_1 = \sum_{M_k \subset C_1} m_k = 0.
\] (15)

Suppose there are \( z \) subareas in \( C_1 \). (\( z = 5 \) in the example shown in Figure (3)). For simplicity, these subareas are named as \( m_1, m_2, \cdots, m_z \). Equation (15) then becomes

\[
n_1 = \sum_{k=1}^{z} m_k = 0.
\] (16)

Note that Equation (16) is also an equation in \( A \). Since all the \( m_k \)s are nonnegative, all the solutions to \( A \) satisfy the condition that \( m_k = 0, \forall 1 \leq k \leq z \), which can be easily interpreted as the number of targets in any subarea within circle \( C_1 \) should be zero. Based on this, Equation (7) can be further rewritten as

\[
P(T = t | N(C_1) = n_1, \cdots, N(C_N) = n_N) = \frac{\sum_{m_k' \in \{A\cap B\}} < m_1', m_2', \cdots, m_M' >}{\sum_{m_k \in A} < m_1, m_2, \cdots, m_M >} = \frac{\sum_{m_k' \in \{A\cap B\}} < m_1', \cdots, m_z', *, *, \cdots, m_M' >}{\sum_{m_k \in A} < m_1, \cdots, m_z, *, *, \cdots, m_M >} = \frac{< 0, \cdots, 0 > \sum_{m_k' \in \{A\cap B\}} < *, *, *, m_2', \cdots, m_M' >}{< 0, \cdots, 0 > \sum_{m_k \in A} < *, *, *, m_2, \cdots, m_M >} = \frac{\sum_{m_k' \in \{A\cap B\}} < m_2', \cdots, m_M' >}{\sum_{m_k \in A} < m_2, \cdots, m_M >}.
\] (17)

The last equality holds because \( < 0, \cdots, 0 > \) is a constant value which can be computed using Equation (1) and hence is canceled out in the function.

From Equation (17) we can conclude that deleting a zero-count node does not cause any loss of accuracy since the result in Equation (17) is the same as the computation in a similar network where \( C_1 \) is excluded. As a result, given a number of sensor nodes, the zero-count circles can be deleted first before computation. There are two major benefits from doing this. Firstly, by deleting the zero-count circles the number of subareas is reduced. Reducing \( M \) further reduces the complexity as we have discussed before. Secondly, the whole graph can sometimes be partitioned into groups by deleting these circles, especially when there are several zero-count circles. If the circles can be divided into groups that are not overlapping with each other as \( G_1 \) and \( G_2 \) shown in Figure (3), the complexity can be significantly reduced.
5.2 Partition with Balanced Minimal Cuts

In the previous section, we have shown that deleting a zero-count node simplifies the computation without losing accuracy. In this subsection we describe how to divide a sensor network into several balanced groups, each with bounded number of subareas, while incurring minimal loss of accuracy. Our solution is based on observing that we have less uncertainty in number of counts, if 1) the size of the overlapping area between different groups is small, and/or 2) the number of targets in the overlapping area is small.

With the consideration of the complexity of cutting and future compensation algorithm, our partition algorithm is recursive and pairwise optimal. The network is firstly divided into two groups, one of which has a bounded number of subareas. If the size of the other group is still out of range, the partitioning algorithm is applied again until all groups have bounded number of subareas and their pmfs can be computed separately. The algorithm is described in more detail in the next few subsections.

5.2.1 Optimization Objectives

Based on the layout of overlapping areas, a sensor network can be modeled into a topology \( G(V,E) \), where \( V \) is the set of the \( N \) sensor nodes \( v_1, v_2, \cdots, v_N \) and edge \( e_{ij} \) exists between node \( v_i \) and \( v_j \) if and only if the sensing areas of the two nodes \( v_i \) and \( v_j \) overlap with each other. The weight of the edge \( e_{ij} \) is decided by both the size of overlapping area and the count values of nodes \( v_i \) and \( v_j \). Formally,

\[
e_{ij} = e_{ji} = \begin{cases} W(rs_{ij}, n_i, n_j) & \text{if } C_i \cap C_j \neq \emptyset \\ 0 & \text{if } C_i \cap C_j = \emptyset \end{cases}
\]

where \( rs_{ij} \) is the percentage that the overlapping area between circles \( i \) and \( j \) out of the total area of circle \( i \) and circle \( j \). \( W \) is an increasing function of \( rs_{ij} \), \( n_i \) and \( n_j \), respectively. A good example of \( W \) is \( W(rs_{ij}, n_i, n_j) = rs_{ij} \times (n_i + n_j) \). Figure (4) shows the corresponding \( G(V,E) \) of the sensor network in Figure (3).

If we partition \( G \) into two subgroups \( G_1 \) and \( G_2 \), we can define the objective function \( f_{obj} \) as the sum of the weights of all the edges cut by the partition. More precisely, \( f_{obj} \) can be expressed by the following equation:

\[
f_{obj} = \sum_{v_i \in G_1, v_j \in G_2} e_{ij}
\]

The objective is to find a partition that minimizes \( f_{obj} \).

5.2.2 Partition Algorithm

We develop a partition algorithm based on the Fiduccia-Mattheyses (FM) Algorithm [5]. For a given graph, the goal is to find a partition that divides the circles into two groups and minimizes \( f_{obj} \) as described before. We bound the size of the first group \( G_1 \) so that the pmf of \( T_1 \) can be computed directly. We apply the partitioning algorithm to \( G_2 \) recursively, until the size of \( G_2 \) is small enough and we can compute the pmf of \( T_2 \) directly as well.

We name the objective function \( f_{obj} \) as the cutting size since it denotes the total weight of cutting edges by a partition. The size of \( G_1 \) should always be smaller than the maximum size such that the pmf of \( T_1 \) can be computed directly. The size of each group should also be greater than a minimum size in order to maintain the accuracy of counting. These requirements on the size of the two groups are termed as the balance requirements (BR). The goal of our algorithm is to find a partition that has the minimum cutting size while satisfying the balance requirements.

As shown in Figure 5, the partition algorithm consists of several iterations, called passes. Each pass has several steps. In each step, a vertex that has the best gain is selected and moved to the other group. The gain of a vertex represents how much the cutting size can be reduced by moving this vertex to the other group. A vertex can be only moved once in a single pass and BR should always be satisfied. A single pass process stops when all vertices have been moved once or moving any unmoved vertex violates BR. Then the best partition (the one has the minimum cutting size) during the whole pass is selected as the starting partition of the next pass process.

An example of how the first pass works in a simple network is shown in Figure (5).

- **Step 1:** Initially, the vertices are divided into two groups randomly as shown in Figure (5) where \( v_1, v_2 \) belongs to one group and \( v_3, v_4 \) and \( v_5 \) belong to the other. This is the starting partition. Before performing any moving, the gain of each vertex is computed. Suppose all the edges in the figure have an equal weight of 1. Then for vertex \( v_3 \), the gain of moving it to the other group is 1 since the cutting size changes from 2 to 1. The gain of all the other vertices can also be computed in this way. We use \((0, 0, 1, -2, -1)\) to denote the gain of the vertices \( v_1, v_2, v_3, v_4, v_5 \). Suppose the BR in this example requires...
Algorithm 2 Partitioning Algorithm

Input: \( G \) and \( C_i \)

Output: Partition \( P_{\text{new}} : \{G_1, G_2\} \)

1: \( P_{\text{old}} \leftarrow \{G_1, G_2\} \) %random initial partition
2: \( \{V\} \leftarrow \) all vertices in \( G_1 \) or \( G_2 \)
3: repeat
4: \( \text{index} \leftarrow 0; P_{\text{old}} = P_{\text{new}} \)
5: repeat
6: Compute the gain for all the vertices in \( V \)
7: \( v \leftarrow \) An unlocked vertex satisfies BR and has the maximum gain
8: if \( v \in G_1 \) then
9: \( G_1 \leftarrow G_1 - v; G_2 \leftarrow G_2 + v \)
10: else
11: \( G_1 \leftarrow G_1 + v; G_2 \leftarrow G_2 - v \)
12: end if (record partition for each step)
13: \( P_{\text{save}}[\text{index}++] \leftarrow \{G_1, G_2\} \)
14: until no vertices in \( V \) can be moved
15: \( P_{\text{new}} \leftarrow \) A minimum cutting partition in \( P_{\text{save}} \)
16: Unlock all vertices
17: until \( P_{\text{old}} == P_{\text{new}} \)

the size of each group should be no less than 2. Due to this requirement, \( v_1 \) and \( v_2 \) cannot be moved since it violates BR. Based on these observations, \( v_3 \) is selected to move to the other group since it has the best gain. It is also marked as locked after it is moved. A locked vertex can not be moved any more in the following steps during the current pass process.

- **Step 2:** In step 2, the gain of each vertex is updated. \( v_4, v_5 \) can not be moved in this step due to the balance requirement although \( v_4 \) has the best gain. \( v_3 \) can not be moved either since it has been locked. As a result, \( v_2 \) is selected to move to the other group although its gain is negative, i.e., moving \( v_2 \) makes the result worse.

- **Other Steps:** Similar process continues until Step 6 when all vertices are locked.

- **Selection and Unlock:** The partition in Step 2 is selected as the starting partition of next pass process since it has the smallest cutting size. All vertices are unlocked, ready for next pass.

A pass, which includes the above steps, repeats itself until there is no positive gain from moving any more, i.e., the partition selected at the end of a pass is the same as its starting partition at the beginning of this pass. Then this partition is the final partition of the algorithm.

In the example shown in Figure (5), the output of the second pass process is the same as its starting partition which is the partition in Step 2 in the figure. This partition is the final result. The whole process of the algorithm is shown in Algorithm (2). The complexity of this algorithm is \( O(n^2) \).

6 Accuracy Compensation

In Section 5, we described how we can partition using minimal cutting. We can reduce the computational complexity to a certain level by setting the maximal size of each subgraph to a threshold \( \text{MAX} \). However, partitioning leads to loss of accuracy. In this section, we propose two methods to compensate for the loss of accuracy caused by partitioning.

We start with an example shown in Figure (6). In this figure, there are 10 circles in total. Suppose there is no zero count circle

as in this example. Using the partitioning algorithm described in Section 5, we can identify the best place of the first partitioning should be between \( C_1 \) and \( C_2 \), where there is only one overlapping subarea denoted as \( M_1 \) as shown in the figure. Then the whole graph will be divided into two groups with the six circles on the left belonging to \( G_1 \) and the four circles on the right belonging to \( G_2 \).

We can compute the \( \text{pmf} \) of the total number of distinct targets for \( G_1 \) and \( G_2 \) (denoted as \( T_1 \) and \( T_2 \)) separately. If we estimate the final \( \text{pmf} \) by simply combining the \( \text{pmf} \) of \( T_1 \) and \( T_2 \) assuming that they are independent, there would be two main factors making the result inaccurate.

1. \( T_1 \) and \( T_2 \) are actually dependent since the two groups have an overlapping subarea \( M_1 \).
2. The targets in subarea \( M_1 \) are counted twice since they belong to both \( G_1 \) and \( G_2 \).

We propose two methods to compensate for the errors caused by these two factors. The first method compensates for errors by deducting the \( \text{pmf} \) counts in the overlapping area, called Partitioning Compensation Minus (P.C.Minus). The second method compensates by adding the \( \text{pmf} \) counts in the overlapping area, called Partitioning Compensation Plus (P.C.Plus). P.C.Minus is simple and efficient for complex topologies and extremely large-scale, while P.C.Plus achieves high accuracy with more overhead.

6.1 Partition Compensation Minus

A major factor that will cause the result to be inaccurate is that the targets in the overlapping area of the two groups have been counted twice. In order to eliminate such an error, we need to estimate the number of targets in the overlapping area and then subtract them from the final result.

As discussed in Section (4), the cost of computing the true \( \text{pmf} \) of the number of targets in the overlapping area (denoted as \( T_{ol} \)) is no less than the cost of computing the \( \text{pmf} \) of the total number of targets within the whole network. We only include a certain number of circles in the computation of \( T_{ol} \) in P.C.Minus. As shown in Figure (6), we can only include \( C_1 \) and \( C_2 \) in the computation of the \( \text{pmf} \) of \( T_{ol} \). We can also include \( C_3, C_8 \) and other circles in the computation. The more circles that are included, the more accurate the result is and the more computation overhead. However, if we include circles that are too far away from the overlapping area, the computation cost increases much faster than the accuracy we gain. This is because the further circles are away from each other, the less correlated they are. The number of subareas included in the computation of \( T_{ol} \) should also be no greater than \( \text{MAX} \) which is defined as the group size. In a small-scale network, this requirement can be satisfied since the size of overlapping subareas is usually small. However, in a large-scale complex network, more circles are clustered and more subareas will be included in a partition. The number of circles that are included in the computation of \( T_{ol} \) would be large. In this case, future partitioning would be needed and the
Algorithm 3 Partitioning Compensation Minus (P.C.Minus)

Input: $G_1, G_2, M_{ol}$
Output: $pmf$ of total target counts $T$

1: $\{C_i\} \leftarrow$ minimum set of circles cover all overlapping subareas of $G_1$ and $G_2$
2: $M_{ol} \leftarrow$ sizeof $\{C_i\}$
3: if $M_{ol} < MAX$ then
4: $C_q \leftarrow$ the nearest circle from overlapping subareas
5: $M_{ol} \leftarrow$ sizeof $\{\{C_i\} + C_q\}$
6: while $M_{ol} < MAX$ do
7: $\{C_i\} \leftarrow \{C_i\} + C_q$
8: end while
9: else
10: compute $T_{ol}$ by future partitioning
11: end if
12: Compute the $pmf$ of $T_1$ directly
13: Compute the $pmf$ of $T_2$ directly or by future partitioning
14: Compute the final $pmf$ of $T$
   where $T = T_1 + T_2 - T_{ol}$

$pmf$ of $T_{ol}$ will be computed recursively as the $pmf$ of $T_2$. Based on these analyses, we develop the algorithm of P.C.Minus as shown in Algorithm 3. In essence, it first obtains the $pmf$ of $T_1$, $T_2$, $T_{ol}$ of $G_1$, $G_2$ and $M_1$ respectively. The $pmf$ of total count $T$ is calculated as $T = T_1 + T_2 - T_{ol}$.

6.2 Partition Compensation Plus

The P.C.Minus algorithm reduces the duplicate count in the overlapping area and gives a certain level of compensation to the final result. However, it doesn’t solve the problem that $T_1$ and $T_2$ are not independent. Moreover, $T_{ol}$ is also not independent with $T_1$ and $T_2$. In order to improve accuracy further, we develop the P.C.Plus method in this section.

We start with a simple assumption that both of $C_1$ and $C_2$ have detected one target. The number of targets in the overlapping subarea $M_1$ will have only two possibilities: $N(M_1) = 0$ and $N(M_1) = 1$.

We define $C'_1 = C_1 - M_1$ and $C'_2 = C_2 - M_1$. $C_1'$ and $C_2'$ are both partial circles excluding the overlapping subarea $M_1$. For a fixed $N(M_1) = m_1$, the count information of $C_1'$ and $C_2'$ can be derived from the following equations:

$$
N(C'_1) = N(C_1) - N(M_1)
$$

$$
N(C'_2) = N(C_2) - N(M_1).
$$

(20)

![Figure 7. P.C.Plus Method](image)

We further define $G'_1 = G_1 - M_1$ and $G'_2 = G_2 - M_1$ where $G'_1$ and $G'_2$ are disjoint as shown in Figure (7) and the corresponding $T'_1$ and $T'_2$ are independent. We use $T'_1(m_1)$ and $T'_2(m_1)$ to denote the counts in $G'_1$ and $G'_2$, respectively, under the condition that there are $m_1$ targets within $M_1$. The $pmf$ of $T'_1(m_1)$ and $T'_2(m_1)$ are computed for each particular $m_1$, using the count and size information of $C'_1$, $C'_2$. Similarly, $T(m_1)$ denotes the total count under the condition that there are $m_1$ targets within $M_1$. Finally, $T$ can be expressed as follows:

$$
T = P(N(M_1) = 0)(T'_1(0) + T'_2(0)) + P(N(M_1) = 1)(T'_1(1) + T'_2(1) + 1)
$$

(21)

where $P(N(M_1) = m_1)$ denotes the probability that there are $m_1$ targets in subarea $M_1$. From Equation (21) we could see that if we find the $pmf$ of $N(M_1)$ we can compute the final $pmf$ of $T$ by enumerating all the possible $m_1$ values, computing the $pmf$ of $T'_1(m_1)$ and $T'_2(m_1)$ for each $m_1$ and then combining them with different weights $P(N(M_1) = m_1)$.

In the general case where there are $k$ overlapping subareas $M_1, M_2, \cdots, M_k$, the process is quite similar. The $pmf$ of $\{N(M_1), N(M_2), \cdots, N(M_k)\}$ is computed first. Then, for a particular value $\{N(M_1) = m_1, N(M_2) = m_2, \cdots, N(M_k) = m_k\}$, the count information of the overlapping circles are updated. $T(m_1, \cdots, m_k)$ can then be computed by simply combining the two independent random variables $T'_1(m_1, \cdots, m_k)$ and $T'_2(m_1, \cdots, m_k)$ plus the sum of $\{m_k\}$. The $pmf$ of $T$ can be computed as the sum of these $T(m_1, \cdots, m_k)$ values with different weight. The algorithm for the general case is shown as Algorithm 4:

It’s not difficult to find that the P.C.Plus method has eliminated error caused by both of the two factors. The only inaccuracy comes from the estimation of the $pmf$ of overlapping subareas. If we compute the $pmf$ accurately, the complexity will be no less than the complexity of computing the final distribution. Like in P.C.Minus, only a subset of circles are included in the computation of the $pmf$ of $\{N(M_1), \cdots, N(M_k)\}$.

Since the only error comes from the estimation of the $pmf$ of the overlapping area, we can adjust the level of accuracy by increasing the number of circles in the computation, regardless of the maximum size of each group. The complexity also increases as the required level of accuracy becomes higher and higher. In the extreme case, when all circles are included to compute the $pmf$, there would be no loss of accuracy, however also no savings of computation. In practice the maximum size is still fixed to $MAX$ in any $pmf$ computation.

The P.C.Plus algorithm works well when the topology is simple and the intensity $\lambda$ small, i.e., there are not too many overlapping subareas and the numbers of targets within most circles are
small. Otherwise the complexity will be too high because we need to compute $T'_1$ and $T'_2$ multiple times for every possible value of $m_1, m_2, \ldots, m_k$. The times we repeat on computing the pmf of the same block is exponential with respect to the number of overlapping subareas. It is also exponential with respect to the average of $\lambda$. As a result, the complexity is too high when the topology is complex and when $\lambda$ is large although we have already reduced the complexity to a polynomial function of $M$. However, P.C.Plus is a very good algorithm when used in a network with simple topology and moderate intensity as shown in the evaluation.

7 Evaluation

In this section we compare the performance of each algorithm in terms of both accuracy and computation overhead in different scales of networks consisting of 10 to 1000 nodes. If not specified, the nodes are randomly generated into a 2D unlimited space. Targets are also randomly generated with equal probability everywhere. In subsections (7.1), (7.2) and (7.3) we assume that the intensity $\lambda$ is already given in order to compare the accuracy of computation affected only by choosing a different algorithm. In (7.4) we show simulation results using an estimated $\lambda$ instead of the real one. In (7.5) we study the scenario when target distribution is not uniform.

7.1 On Small-Scale Networks

In this experiment, we compare the performance of different algorithms in small-scale networks, where ten sensor nodes are overlapped with each other with a total number of subareas $M = 20$. The size of each sensing circle is set to 9 units and the total coverage of these ten sensor nodes is 63 units. The true number of distinct targets in the region is 12 which corresponds to $\lambda \approx 0.2$. The reason that we study this small-scale scenario is to compare the pmf estimated by each algorithm with the true pmf directly, which is impossible in large-scale networks since the true pmf of Equation (7) is always too complex to compute.

Figure (8) shows a typical example of the pmfs with the ground-truth count of 12, using four different methods: Direct Computation (D.C.) which corresponds to (i) the True pmf in the figure, (ii) Partition Method with P.C. Plus, (iii) Partition Method with P.C. Minus and (iv) Partition Method without any compensation at all (P.O.). Compared with the True pmf in Figure (8), P.C. Plus gives the most accurate result that the pmf it computes is almost the same as the True pmf. P.C. Minus is less accurate than P.C. Plus, but is still a good estimation. The pmf computed by P.O. has a horizontal shift of about 2 due to the fact that the targets within the overlapping area have been counted twice. If we simply estimate the number of targets by adding the numbers reported by each node, we will get 17, which is worse than any of these methods.

We run similar simulations 200 times. In each simulation, $\lambda$ is fixed to 0.2 and targets are randomly regeneraged to make the count information different. The expected values of $T$ (denoted as $E(T)$) are calculated from the pmf computed by the four methods. The average absolute error of $E(T)$ is compared in Figure (9). Note that the true expected value of $T$ in each simulation is around 12, from which we can see the accuracy of P.C.Minus with a relative error of about 0.8% and P.C.Plus with a relative error of 0.3%.

Figure (10) shows the average runtime comparison (in log scale) between P.C.Plus, P.C.Minus and direct computation for various $\lambda$s from 0.07 to 0.21. We don’t include P.O. in the comparison here, because the complexity of P.O. is almost the same as that of P.C. Minus. From Figure (10), we can see when $\lambda \approx 0.2$, the runtime of P.C. Plus and P.C. Minus are less than 1% of the runtime of direct computation. Compared with the results shown in Figure (8) and Figure (9) we can conclude that P.C. Plus is the best choice in
small-scale network.

7.2 On Large-Scale Linear Networks

We place 100 nodes linearly. We study linear networks in order to compare the performance of both P.C. Plus and P.C. Minus, which is similar to the reason we study small-scale networks. The size of each sensing circle is still set to 9 and the total coverage is around 600. We vary $\lambda$ from 0.01 to 0.2 (The expected number of targets in the region varies from 10 to 130). Since this is a large-scale network where $M$ is around 200, the True pmf is no longer available. As a result, the average expected value of $T$ is compared in the evaluation.

For each $\lambda$, a fixed number of targets are generated randomly. Three partition methods (P.C. Plus, P.C. Minus and P.O.) are used to compute the pmf of $T$. Then $E(T)$ is calculated and recorded. We run similar simulations 200 times for each $\lambda$ and compare the average of $E(T)$ with the true number of targets generated in the simulation.

Figure (11) shows the comparison of average $E(T)$ computed by each algorithm. It also shows the estimation by the naive method of simply adding up all the count information which corresponds to the “UB” in the figure. From the figure we can see that in terms of average expected value of $T$, both P.C. Plus and P.C. Minus give a very good estimation while P.O. has a higher expected value as expected. However, P.O. is still much better than the naive method.

Figure (12) shows the comparison of average runtime of P.C. Plus and P.C. Minus. Again P.O. is not included in the figure since it almost has the same runtime as P.C. Minus. From the figure we can see that as the intensity $\lambda$ increases, P.C. Plus has a greater increase in runtime since it suffers too much from repeatedly computing the pmf of the same block. The runtime of P.C. Minus also increases when $\lambda$ increases, but much slowly than P.C. Plus. From these observations we conclude that in large-scale linear networks, P.C. Plus is a good choice when $\lambda$ is moderate while P.C. Minus is the best choice when $\lambda$ is large.

Figure (13) shows the runtime of P.C. Minus for networks consisting of different numbers of sensor nodes from 100 to 1000 when $\lambda$ is fixed to 0.1. From the figure we can see as the number of sensor nodes increases, the runtime of P.C. Minus increases linearly.

7.3 On 2-Dimensional Large-Scale Networks

We place 100 nodes randomly in a 2-Dimensional area. Since the topology this time is too complex, P.C. Plus is too complex to use since it needs too many iterations. As a result, only the performance of P.C. Minus and P.O. are compared here, using similar simulations as described in (7.2). Figure (14) shows the comparison of average $E(T)$ when $\lambda$ varies from 0.01 to 0.25. It also shows the estimation of the naive method UB. From the figure we can see that P.C. Minus gives a very good estimation in terms of expected value of $T$. P.O. gives a higher expected value as expected, but the estimation is still better than UB.

7.4 Impact of Estimated $\lambda$

In (7.1), (7.2) and (7.3) we use a given intensity $\lambda$ in the simulation in order to compare the performance of each algorithm. However, the intensity information is sometimes not available. As a result, $\lambda$ should be estimated using just the count information. One way to estimate $\lambda$ is to sum all the count information and then divide by the total area of the sensing circles, assuming that they are not overlapping. For example, in the small scale network, if the count information is $\{1,1,1,3,5,2,1,2,1,0\}$ and the size of each circle is 9, $\lambda$ can be estimated as $\lambda = \frac{17}{9}$. We repeat the simulation in (7.1) using estimated $\lambda$ in P.C. Plus, P.C. Minus and P.O. while keeping an actual $\lambda$ in the direct com-
putation to obtain the result of the True pmf. Figure (15) shows
the pmf computation result in the same scenario as in (7.1), where
the count information is \{1,1,1,3,5,2,1,2,1,3\} and the true number
of target is 12. Comparing Figure (15) with Figure (8) we can see the
pmf computed by the three algorithms are slightly changed, but
P.C.Plus and P.C.Minus are still good approximations.

Figure (16) shows the average absolute error of the expected
value of \( T \) in small-scale network. From the figure we can see the
average absolute error increases when estimated \( \lambda \) is not so accu-
rate. However, the difference is negligible due to the fact that \( E(T) \)
is around 12 and the relative error is extremely small especially for
P.C. Plus and P.C. Minus.

Figure (17) shows the absolute error of \( E(T) \) computed using an
estimated \( \lambda \) with \( E(T) \) computed using true \( \lambda \) in large-scale linear
network. From the figure we can see P.C.Plus is the most robust
when \( \lambda \) is inaccurate. The absolute error caused by the inaccuracy
of \( \lambda \) only causes a very small difference of \( E(T) \) which is negligible
due to the fact that \( E(T) \) is large according to Figure (11) and the
relative error is extremely small.

Based on all comparisons we conclude that our algorithm can be
easily applied to the real scenarios when the intensity information of
\( \lambda \) is not available.

7.5 Impact of Non-Uniform Distribution

In previous evaluation we have shown that the P.C. Plus and
P.C.Minus can approximate the pmf of \( T \) very well, under the as-
sumption that the targets are uniformly distributed. However, it’s
not always the case that the targets are uniformly distributed with
the same intensity \( \lambda \) everywhere. In this subsection we study the
case that targets are non-uniformly distributed, i.e., \( \lambda \) in each sub-
area can be different. Again an estimated \( \lambda \) is used in the computa-
tion of each algorithm.

We study large-scale linear networks to compare the perfor-
ance of all three partition methods. We divide the 100-node linear
network into 10 different sub-networks each of which has a random
intensity \( \lambda \) value when we distribute targets. In the simulation, \( \lambda \)
estimated using the same technique as in (7.4). Figure (18) shows the
simulation results of average \( E(T) \) in this scenario. From the
figure we can see in terms of average expected value of \( T \), P.C.Plus
and P.C.Minus are both robust to change of distribution models.

8 Conclusion

The double-counting problem has been sufficiently addressed in
the context of communication, however, to our knowledge, there is
no existing solution for double-counting problem in sensing. This
paper presents an efficient and accurate method to estimate the
number of targets within a monitored area with duplicate counts
among adjacent sensors.

Using a partition method, we significantly reduce the computa-
tion complexity of calculating the probability mass function of
total counts. Using several compensation methods, we improve the
accuracy with adjustable computational overhead. This work is
theoretical in nature, however, it can be practically applied since
most sensor systems do not require precise counts, but reasonable
accurate estimations. As a result, statistical counting is a viable
approach. To inspect whether our solution is practical with respect
to several assumptions we make, we evaluate the design with esti-
1mated \( \lambda \) values as well as non-uniform distributions. Results reveal
the accuracy of statistical counting degrades slightly when the true
\( \lambda \) is unknown and targets are non-uniformly distributed. Through
extensive simulation over various kinds of network settings, we

demonstrate that accurate statistical counting within 1 ~ 3% rel-

ative error can be obtained with orders of magnitude reduction in
computation, compared with the exhaustive enumeration-based ap-
proach. Without loss of generality, we use Poisson distribution as
a concrete example, we believe the ideas of partition with balanced
minimal cuts and accuracy compensation are applicable to other
target distributions as well.

9 References

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