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Finding the Polygon Hull of a Network Without Conditions on the Starting Vertex

Ahcène Bounceur · Madani Bezoui · Mohammad Hammoudeh · Loïc Lagadec · Reinhardt Euler

Abstract Many real-life problems arising within the fields of wireless communication, image processing, combinatorial optimisation, etc., can be modeled by means of Euclidean graphs. In the case of Wireless Sensor Networks (WSN), the overall topology of the graph is not known, because sensor nodes are often randomly deployed. One of the significant problems in this field is the search for boundary nodes. This problem is important in cases such as the surveillance of an area of interest, image contour reconstruction, graph matching problems, routing or clustering data, etc. In the literature, many algorithms are proposed to solve this problem, a recent one of which is the LPCN algorithm and its distributed version D-LPCN which are both based on the concept of a polar angle visit. An inconvenience of these algorithms is the determination of the starting vertex. In effect, the point with the minimum x-coordinate is a possible starting point but it has to be known at the beginning which considerably increases the algorithms’ complexity. In this article, we propose a new method called RRLPCN (Reset and Restart with Least Polar-angle Connected Node) which is based on the LPCN algorithm to find the boundary vertices of a Euclidean graph. The main idea is to start the LPCN algorithm from an arbitrary vertex and, whenever it finds a vertex with an x-coordinate smaller than that of the starting one, LPCN is reset and restarted from this new vertex. The algorithm stops as soon as it visits the
same edge for the second time in the same direction. In addition to finding the boundary vertices, RRLPCN also finds the vertex with minimum $x$-coordinate which is the last starting point of our algorithm. The distributed version of the proposed algorithm, called D-RRLPCN, is then applied to boundary node detection in Wireless Sensor Networks. It has been implemented using real sensor nodes (Arduino/XBee and TelosB). The simulation results have shown our algorithm to be very performant in comparison to other algorithms\footnote{This work is part of the research project PERSEPTEUR supported by the French Agence Nationale de la Recherche ANR.}.

**Keywords** Polygon Hull · Wireless Sensor Network · Boundary Node Detection · Polar angle

1 Introduction

Since the 1970s, the algorithmic search for a “border” (or “boundary”) circumscribing the elements of a set of points in the plane has attracted many researchers. One prominent example for such a border is the “convex hull” which could be illustrated by using a minimum size string to surround a set of nails planted in a plane. In 1971, Graham [28] was the first to propose an algorithm to find the convex hull of a set of points, which is based on the scan of the point set. One year later, Jarvis [31] proposed his version, based on the use of polar angles. After the work of these two pioneers, many other approaches have emerged: the Quickhull algorithm [18] in 1977, Andrew’s algorithm [3] in 1979, Kallay’s algorithm [32] in 1984, Chan’s algorithm [11] in 1996, etc.

In some situations, the convex hull does not sufficiently reflect the geometric properties of a data set. The “concave hull”, called here “polygon hull”, is more accurate for geometric analysis. This polygon hull approach is a more advanced method designed to capture the precise shape of the data set’s surface. However, the construction of such a hull is not straightforward. Most algorithms require the knowledge of a starting point, supposed to belong to the border.

The problem of finding the polygon hull of a connected Euclidean graph can be formulated as follows: Given an undirected Euclidean graph $G = (V, E)$, where $V = \{v_0, v_1, ..., v_{n-1}\}$ is the set of vertices of $G$ and $E$ its set of edges, we are looking for a closed cycle of minimum length in $G$ such that all vertices are either on or surrounded by that cycle. This cycle defines an area which is also called a Polygon Hull, and it can be described by a sequence of vertices $\mathbb{B}_V$ and a set of edges $\mathbb{B}_E$ as follows:

\begin{align*}
\mathbb{B}_V &= (v_0, v_1, ..., v_{i-1}, v_i, v_{i+1}, ..., v_h) \quad (1) \\
\mathbb{B}_E &= \{\{v_0, v_1\}, \{v_1, v_2\}, \cdots, \{v_{h-1}, v_h\}\} \quad (2)
\end{align*}

such that

$\quad v_0 = v_h.$
The search for a polygon hull of a Euclidean graph arises as an essential question in many fields such as Wireless Sensor Networks (WSN), statistics, data analysis, image processing, fingerprint matching tests, biological networks, etc. Unfortunately, there exist only a few methods to solve such a problem. In recent papers [36], [51], we have proposed an algorithm called LPCN and its distributed version called D-LPCN [51]. The principle of these algorithms is to start from the point with minimum $x$-coordinate and look for a neighbor which forms the minimum polar angle. This process is repeated until a polar angle is traversed for the second time. Despite the good performance of this algorithm and its optimality, demonstrated in [36], it requires an a priori knowledge of the starting vertex, which considerably increases its complexity.

In this paper, we propose a new method called Reset and Restart. It is an extension of the LPCN algorithm giving a new algorithm called RRLPCN, which allows to find the polygon hull of a Euclidean graph without any condition on the starting vertex. We also present a distributed version of this algorithm, called D-RRLPCN, together with an application in the field of WSNs.

A WSN consists of autonomous sensor nodes distributed in space to cooperatively monitor physical or environmental conditions such as temperature, sound, vibration, pressure, motion, etc. Typical WSN applications may require the random deployment of sensor nodes over a large target area. Besides, military surveillance systems may also require the detection of activities around the boundaries of the area under surveillance. Thus, the system should be able to detect and identify any object entering or leaving the monitored area. Therefore, developing mechanisms to identify network boundary nodes is a significant and challenging problem.

RRLPCN is developed and validated theoretically using the CupCarbon and Tossim simulators. The distributed version of our proposed algorithm (D-RRLPCN) is applied to the boundary node detection problem in a wireless sensor network. The simulation results show that D-RRLPCN consumes less energy compared to some existing algorithms. After validation by simulation, a real implementation is conducted using two types of wireless sensor nodes, TelosB and Arduino Xbee.

The remainder of this paper is organized as follows. Section 2 presents related work from the fields of polygon hull determination and boundary node detection in wireless sensor networks. A mathematical formulation of the problem of polygon hull finding is proposed in Section 3. In Section 4, the LPCN algorithm and the Reset and Restart method are presented. Moreover, the integration of Reset and Restart into LPCN is described. In Section 4.3 the distributed version of RRLPCN is presented. The convergence of our method is demonstrated in Section 5 in which simulation results and an implementation into real sensor nodes are shown. Section 6 briefly describes the use of our approach for two other applications, and Section 7 concludes the paper.
2 Related work

In this section, we present a state of the art on algorithms to find convex, concave or polygon hulls of a set of points in the plane and of Euclidean graphs as well as on algorithms to detect boundary nodes of wireless sensor networks. In this paper, we do not deal with security and mobility problems. For this, it is suggested to refer to existing methods [4] [5] [25] [30] [2].

2.1 Finding hulls of a set of points in the plane

Chaudhuri et al. [12] have proposed an approach to find the concave hull of a plane point set \( S \). This technique is called perceptual boundary extraction, and it is based on a new definition, called \( s - \text{shape} \), which is simply the union of lattice cells containing all the points of \( S \). To obtain a polygon boundary, another \( r - \text{shape} \) is defined, where \( r \) is obtained from \( s \). For points \( p, q \in S \), the edge \( \{p, q\} \) is selected if and only if the boundaries of the disks centered around \( p \) and \( q \) intersect at a point which is on the boundary of the union of all the disks. The \( r - \text{shape} \) of \( S \) is then the union of the selected edges.

Garai et al. [24] have proposed an algorithm that starts with a convex hull of the given points and reaches the final limit in two stages: division and fusion. By splitting, one or more sides of the convex hull will be removed, and new sides are added to maintain the inherent convexity. To obtain a smooth polygon border, two or more sides are merged into one.

Adriano and Santos [48] have suggested an algorithm based on nearest neighbors. It assumes that the currently selected point is related to its \( k \) nearest neighbors. Then it selects the point forming the least polar angle with the current point, just as in Jarvis’ algorithm [31]. A polygon hull is not obtained for any set of points, and the value of \( k \) must be adapted to each case.

Gheibi et al. [26] have presented a shape reconstruction algorithm that finds a polygon hull. The algorithm starts from the convex hull of the initial pattern and gradually “concaves” it to obtain a polygon output. Thus, at any step, each selected edge will be replaced by two new edges constructed so that the shape remains a polygon. This process is repeated until the stop condition is met.

Park and Se-Jong [49] have proposed a four-step algorithm. In the first step, a set of convex hull edges is selected and a threshold value \( \eta \) is determined. In the second step, a decision distance between these closest internal points and the edge points is calculated. Third, if \( \frac{\text{length of edge}}{\text{decision distance}} > \eta \), the search process is executed. Finally, the second and third steps are repeated until there is no more internal point to find.

The Regularized Geometric Hull (RGH) algorithm proposed by Korner et al. [34] is used principally for biomedical image segmentation. It converts the points of a data set into a set of triangles. Each triangle having its maximal edge length greater than a given parameter \( \zeta \) will be excluded. The value of \( \zeta \) regularizes the convexity or concavity of the geometric hull.
The algorithm proposed by Methirumangalath et al. [47] starts by constructing the Delaunay graph $G$ [56]. Afterwards, it creates a priority queue of the outside edges of $G$ in decreasing order of edge lengths. Then it removes, repeatedly, the outside edge of the external triangle from the head of the priority queue and the current graph $G$, but only if it satisfies the defined circle constraint and if $G$ without the removed edge is regular. Once the outside edge is removed from $G$, the adjacent sides of the external triangle are added to the priority queue by maintaining the decreasing order of the edge lengths. This process is repeated until there is no possibility of removing any outside edges from the graph $G$.

The main idea of Braune et al.’s algorithm [10] is to start from the convex hull of the entire data set. The convex hull is iteratively shrunk by replacing edges that are too long by new edges that fit the data more accurately, and then, to recursively split the convex hull path into two separate closed paths as soon as this process converges to one point.

2.2 Finding boundary vertices of a Euclidean graph

To the best of our knowledge, LPCN introduced by Bounceur et al. [8,9,36] is the only algorithm that can find the polygon hull of a Euclidean graph. This algorithm is inspired by Jarvis’ algorithm designed to find the convex hull of a set of points. The main idea of this algorithm is to start from the point with minimum $x$-coordinate and to select a neighbor that forms the minimum polar angle. This step will be repeated by every selected point with respect to its neighbors, and the process stops as soon as a selected point is chosen a second time. LPCN is presented in Section 3.2. A distributed version has been published in [51].

2.3 Boundary node detection in Wireless Sensor Networks

Several solutions have been proposed to detect the boundary nodes in a wireless sensor network. These solutions can be classified into three categories:

**Geometrical approach:** The algorithms of this category assume that the sensor nodes know their exact location within the network. Martincic and Schwiebert [45] have presented an algorithm that requires each node to know the positions and communication links of its 2-hop neighborhood. Using this information, the node decides whether it is enveloped by a circle of other nodes. If this occurs, the node is located in the interior of the network.

The algorithm of Deogun et al. [15] only requires a node to be able to determine the distances to its direct neighbors. The node selects four of its closest neighbors which are far from each other. Then it checks whether three of these neighbors are surrounded or not.
The approach proposed by Fang et al. [19] needs to know the position of the nodes. Using a Delaunay graph and local searches, holes are identified. Zhang et al. [59] apply localized Voronoi polygons for which each node has to collect the positions of its direct neighbors.

The work introduced by Shirsat and Bhargava [54] only requires a node to know an anti-clockwise order of its neighbors. Each node checks for empty cones and cordless paths in the connectivity graph of its 2-hop neighborhood.

Luthy et al. [43] have presented an algorithm that draws an exact image to decide whether the boundary of the nodes’ communication range is entirely covered by the communication ranges of its neighbors.

Sahoo et al. [50] have proposed the sequential boundary node selection algorithm SBNS and the distributed boundary node selection algorithm DBNS. The first algorithm starts from the sink. Then it uses the right-hand rule to select boundary nodes sequentially. The second algorithm defines extreme nodes as boundary nodes and then connects them to form cycles enclosing boundaries. An extreme node is defined as a node that has either maximum or minimum values in its coordinates compared to those of its one-hop neighbors.

The algorithm of Lara-Alvarez et al. [38] is based on $Ircod$ sensors. These sensors determine a cyclic order neighbor and choose the first neighbor in this order. Then the Right Hand without Crossings (RHWoC) rule is used to determine the complete boundary.

Zhao et al. [61] have proposed two distributed algorithms based on computational geometry, Distributed Sector Cover Scanning (DSCS) and Directional Walk (DW). The first is used to identify the nodes on the hole borders and the outer boundary. The second is used to locate the coverage holes based on the boundary nodes identified with DSCS and which they enclose.

Saoudi et al. [51] have presented a distributed version of the LPCN (Least Polar-angle Connected Node) algorithm introduced in [36]. In each iteration, the current boundary node chooses the nearest polar angle given by the previously chosen node and its neighbors, and the first node has to be a boundary node. This node can be automatically determined using the Minimum Finding algorithm. The algorithm works with any connected network, given as planar or not, and it can determine for a disconnected network all the boundaries of the different connected components.

**Statistical Approach:** This type of algorithm tries to exploit statistical properties such as node degrees to detect boundary nodes. As long as nodes are evenly distributed, this approach works quite well since boundary nodes usually have fewer neighbors than interior nodes. However, as soon as node degrees fluctuate noticeably, most statistical approaches produce misclassifications. Besides, these algorithms often require unrealistically high average node degrees.

Prominent statistical approaches are Fekete et al. [20,21], and Bi et al. [7]. Fekete et al. first analyze the node degree distribution in a theoretical study. Their implementation in a second work requires data being collected over the entire network to compute a histogram of node degrees. Using the histogram, they determine a threshold value by which each node can classify itself as an
inner node or a boundary node. In the approach proposed by Bi et al., nodes need information only of their neighborhood localization. Each node compares its node degree with the average node degrees of its 2-hop neighbors to decide whether it is on the network boundary.

**Topological Approach:** Algorithms using this approach concentrate on information given by the connectivity graph and try to infer boundaries from its topological structure. They often require nodes to gather information about a large neighborhood or entail sophisticated algorithms with high computational cost.

Funke [22] and Funke and Klein [23] have described algorithms that construct iso-contours and check whether their contours are broken. If a node detects that a contour is broken, it classifies the corresponding contour endpoints as boundary nodes. The first algorithm requires that the whole network is flooded, starting from some seed nodes. The second algorithm is distributed, based on 6-hop neighborhoods.

The methods proposed by Ghrist and Muhammad [27] and de Silva and Ghrist [14] detect holes by utilizing algebraic homology theory. They are centralized and rely on restrictive assumptions on the communication model.

The algorithm of Kroller et al. [35] identifies complex combinatorial structures called flowers. Such flowers exist with high probability under some hypotheses on the communication model. The algorithm requires that every node knows its 8-hop neighborhood.

Wang et al. [57] have introduced an algorithm that works well even in networks with low average node degree. It involves multiple steps, some of which require that the whole network is flooded.

Dong et al. [17] have described a distributed algorithm based on topological transformations of the connectivity graph. It is, in particular, aimed at locating small holes.

The approach presented by Li and Hunter [41] needs to determine all pairs of nodes with overlapping sensing ranges (e.g., by comparing sensing results). The graph induced by this information is used to detect holes in the network. Boundary nodes are determined by finding circles in their 1-hop neighborhood. Using knowledge of their k-hop neighborhood, holes of up to $4k+2$ ($k = 1, 2, \ldots$) hops in perimeter are found.

Dinh [16], Schieferdecker et al. [53], and Chu and Ssu [13] have introduced algorithms that exploit the same basic information. Each node constructs a graph induced by its neighbors at the exact 2-hop distance. It is checked whether these graphs form closed circles. This is done by verifying the connectivity of sub-graphs in Dinh’s case, by tree construction and analysis in the approach of Chu and Ssu, who further provide a correctness proof for their method.

Saukh et al. [52] have introduced an algorithm that tries to identify distinct patterns in the neighborhood of a node. Under specific conditions, they can guarantee that all nodes that are classified as interior nodes lie inside the network. The algorithm is distributed and every node needs information only
on its $k$-hop neighborhood. The radius $k$ depends on the node density. For low
density, $k = 6$ is used.

Sitaynah et al. [55] have presented a heuristic based on finding boundary
cycles inside a network. The principal idea of this approach is to start from a
first cycle which will be increased gradually by inserting other vertices until
it hits the boundary of the network. A vertex with the highest degree will be
chosen as the center of the cycle. In case that other cycles are found, they will
be merged. This approach can work with low-density wireless sensor networks.

Yan et al. [58] focus on detecting small coverage holes. They assume the
communication range of each node to be two times its sensing range and the
nodes know or can easily determine whether they are located on the outer
boundary of the network. Their theoretical reasoning implies the topologi-
cal Cech and Rips complexes. In their distributed implementation, each node
needs to find a Hamilton cycle in its 2-hop neighborhood.

In another work, Dong et al. [17] also focus on discovering small coverage
holes. They make the same assumptions as Yan et al. [58] regarding commu-
nication ranges and periphery awareness.

Khan et al. [33] devise a topology based method for hole detection, in which
the connectivity between nodes and their $x$-hop neighbors is used without the
location of nodes.

In this article, we propose an algorithm to find the polygon hull in a con-
nected Euclidean graph based on LPCN (Least Polar-angle Connected Node),
which allows to start from any point of the graph. A distributed version of the
algorithm is proposed in view of its application to boundary node detection in
wireless sensor networks.

3 Basic concepts

Before we begin the presentation of our proposed algorithm, let us give some
definitions that will help the reader to better understand our method.

Let $G = (V, E)$ be a connected Euclidean graph in the plane, where $V$
and $E$ are, respectively, the set of vertices and edges of $G$. Unless otherwise
specified, these definitions apply throughout this document.

3.1 Definitions

Definition 1
A boundary vertex of $G$ is a vertex whose deletion will modify the geometric
area formed by the vertices of $G$.

Definition 2
If $v$ is a boundary vertex of $G$, then there is no subset of $V$ forming a polygon
and containing $v$ in its interior.
We can illustrate Definition 1 in Figure 1(a) which shows a graph with two kinds of vertices. The boundary vertices are those that belong to the orange edges, and the other ones are non-boundary vertices. The orange area of Figure 1(b) shows the geometrical area formed by the graph. Figure 1(c) shows a case where one boundary vertex is removed. As we can see, the new shape of the graph is different from the original one. This means that the deleted vertex is a boundary vertex. However, if we remove a non-boundary vertex, as shown by Figure 1(d), the new geometrical area formed by the vertices of the graph is the same as the original one, which means that the deleted vertex is a non-boundary one.

3.2 Least Polar-angle Connected Node Algorithm (LPCN)

In this section, we review the LPCN algorithm used to find the polygon hull of a Euclidean graph $G = (V, E)$. The algorithm starts from a vertex of $G$ with minimum $x$-coordinate and determines the vertices $v_0, v_1, ..., v_{h-1}$ of the polygon hull. The main step of the algorithm is the selection of the neighbors of the current vertex which forms the least polar angle. The process stops when the angle corresponding to the starting vertex is visited twice. The LPCN pseudo-code is described in Algorithm 1, where $\mathbb{B}_V$ and $\mathbb{B}_E$ are the sets of, respectively, boundary vertices and boundary edges, defined by Equation 1.
and Equation 2. In this algorithm $N(p)$ represents the set of the neighbours of the point $p$.

**Algorithm 1** Least Polar-angle Connected Node (LPCN).

1. procedure LPCN($V, E$)
2. $P_0 ← P_c ←$ a point having the minimum $x$-coordinate
3. $P_p ←$ a fictitious point situated to the left of $P(c)$
4. $B_V ← \{P_c\}$
5. $B_E ← \emptyset$
6. once $←$ true
7. repeat
8. $A ← \{P ∈ N(P_c) / B_E \cap \{P_c, P\} = \emptyset\}$
9. $P_{min} ← \arg\min_{P ∈ A} \{\varphi(P_p, P_c, P)\}$
10. $B_V ← B_V \cup \{P_{min}\}$
11. $B_E ← B_E \cup \{\{P_c, P_{min}\}\}$
12. $P_p ← P_c$
13. $P_c ← P_{min}$
14. if (once = true) then
15. once $←$ false
16. $P_{first} ← P_{min}$
17. end if
18. until ($P_c = P_0$ and ($P_{min} = P_{first}$))
19. return $B_V, B_E$
20. end procedure

4 Boundary detection without condition on the starting vertex

4.1 Reset and Restart (R&R) technique

In this section, we describe a new method called Reset and Restart which will be combined with the centralized algorithm LPCN in order to find the boundary vertices of a connected Euclidean graph by starting from any vertex. As a by-product, this method also allows to find the vertex with the minimum $x$-coordinate in the graph. However, if the considered system is distributed then the Reset and Restart concept must be combined with the D-LPCN algorithm. In this paper we assume that the nodes/vertices of the graph are not mobile. However, the mobility is considered as future work.

To illustrate the Reset and Restart technique let us take the graph shown in Figure 2, where the vertex with minimum $x$-coordinate is linked by a chain to the starting vertex. Suppose that the algorithm starts from vertex 1 as shown by Figure 2(a). First, we will mark it and set the value of $x_{min} = x_1$, the $x$-coordinate of vertex 1 and go to vertex 2, as shown by Figure 2(b), and select it. Since the $x$-coordinate of vertex 2 is greater than $x_{min}$, we select vertex 3 and compare its $x$-coordinate to $x_{min}$ (cf. Figure 2(c)). Again, the $x$-coordinate of vertex 3 is greater than $x_{min}$. Now vertex 4 is selected (cf. Figure 2(d)). Since the $x$-coordinate of vertex 4 is less than $x_{min}$, ‘Reset’
is launched, which consists first in unselecting all the selected vertices (Figure 2(e)). After that ‘Restart’ executes LPCN with the last selected vertex 4 and updates $x_{min}$ to the value $x_4$, the $x$-coordinate of vertex 4 (Figure 2(f)). Then by executing LPCN, we select vertex 5 (Figure 2(g)). Again, since the $x$-coordinate of vertex 5 is less than $x_{min}$ (Figure 2(h)), we will execute the Reset and Restart process (Figure 2(i)). Now, if we run LPCN from vertex 5, we will find the polygon hull of the graph because the algorithm will be run until it selects vertex 5 for the second time, which is the stop condition of the algorithm.

Fig. 2: Reset and Restart technique.
The major disadvantage of LPCN is the need for an a priori knowledge of the vertex of $G$ with minimum $x$-coordinate. Unfortunately, this information is often unknown, and its search increases the complexity of the algorithm. This is the main motivation for the proposed improvement of LPCN using the Reset and Restart technique. The complexity of the proposed algorithm is the same as that of the LPCN algorithm repeated each time the starting process is executed. This means that $O(RRLPCN) = w \times O(LPCN) = w \times O(dn_b + n)$, where $n$ is the number of nodes, $n_b$ is the number of the boundary nodes and $d$ is the maximum degree of the network/graph.

The main steps of the new algorithm are described below:

1. In the Restart step, the starting vertex will select that vertex, which forms the minimum polar angle between the starting vertex, a fictitious vertex, which is an imaginary vertex situated on its left and its neighbors, as shown by Figure 3. In this figure, the edge between the starting vertex and the selected one is colored green.

![Fig. 3: Minimum polar angle.](image)

2. After the selection of the next vertex, we have two possible cases: Either the $x$-coordinate of this vertex is greater or equal to $x_{\text{min}}$ or it is less than $x_{\text{min}}$.
   (a) In the first case, we continue the execution of LPCN and select a subsequent vertex.
   (b) In the second case, we run Reset and set $x_{\text{min}}$ equal to the current vertex $x$-coordinate and we Restart the algorithm from this vertex.

Figure 4 shows an example where all the $x$-coordinates of the selected vertices are greater than $x_{\text{min}}$.

We can conclude that finding the boundary vertices matches with finding a vertex with minimum $x$-coordinate. Thus, Algorithm 2 shows the pseudo-code of RRLPCN to find the boundary vertices by starting from any vertex of a
network. Additionally, a flowchart is presented in Figure 5, which resumes the main steps of the introduced method.

Fig. 4: An example for polygon hull finding.

![Flowchart](image)

**Fig. 5: A flowchart of RRLPCN.**

Figure 6 shows the execution of Algorithm 2 on a graph of 10 vertices.
Algorithm 2 Reset and Restart Least Polar-angle Connected Node (RRLPCN).

1: **procedure** RRLPCN($V, E$)
2: $P_c \leftarrow$ an arbitrary vertex chosen from $V$.
3: $B_V \leftarrow \{P_c\}$
4: $P_{first} \leftarrow P_c$
5: $P_p \leftarrow$ a fictitious vertex situated to the left of $P_f$
6: $X_{first} \leftarrow$ the $x$-coordinate of $P_{first}$
7: repeat
8:   $\mathcal{A} \leftarrow \emptyset$
9:   $P_v \leftarrow \arg\min_{P_j \in N(P_c) \& P_j \notin \mathcal{A}} \{\varphi(P_p, P_c, P_j)\}$
10:   $X_v \leftarrow$ the $x$-coordinate of $P_v$
11:   if $X_v < X_{first}$ then
12:     Reset: $B_V \leftarrow \emptyset$; $B_E \leftarrow \emptyset$
13:     Restart: $P_c \leftarrow P_v$; go to step 3;
14:   end if
15:   if $B_E \cap \{P_c, P_v\} \neq \emptyset$ then
16:     $\mathcal{A} \leftarrow \mathcal{A} \cup \{P_v\}$
17:     Go to 9
18: end if
19: $B_V \leftarrow B_V \cup \{P_v\}$
20: $B_E \leftarrow B_E \cup \{P_c, P_v\}$
21: $P_p \leftarrow P_c$
22: $P_c \leftarrow P_v$
23: until $P_v = P_f$
24: return $B_V, B_E$
25: **end procedure**

The algorithm starts from the red vertex (see part (a)), which is set as a starting vertex and with an $x$-coordinate equal to $x_{min}$. Then it computes the next vertex among its neighbors, which is the neighbor forming the minimum polar angle between a fictitious vertex, itself (the red vertex) and its neighbors. In our example, the next vertex is the green vertex (see part (b)). Since the $x$-coordinate of the green vertex is less than the $x$-coordinate of the red vertex, the red vertex is Reset, the new starting vertex is the green vertex whose $x$-coordinate is set to the new $x_{min}$ (see part (c)). In the same way, the new starting vertex computes the next vertex (see part (d)). Since the $x$-coordinate of the next vertex is less than $x_{min}$, the next vertex is the new starting vertex, its $x$-coordinate is set as $x_{min}$ and the previously selected vertex is Reset (part (f)). Using the same principle, the new starting vertex Restart the process and computes the next vertex, since this time the $x$-coordinate of the next vertex is greater than $x_{min}$. This process continues without Restart, until reaching the most left green vertex (part (g)). Since the $x$-coordinate of this vertex is less than $x_{min}$, all previously selected vertices (part (h)) are Reset, and the process is Restart from this vertex. The algorithm selects all boundary vertices, until arriving at the starting vertex, which computes the next vertex as the same vertex selected in the previous round, and the stop condition is satisfied, i.e., all the boundary vertices are selected (part (i)).
4.3 Reset and Restart combined with the Distributed Least Polar-angle Connected Node algorithm (D-RRLPCN)

Boundary node detection is an essential problem in wireless sensor network applications. It is necessary for keeping track of the coverage range and events entering or leaving the region and for tracking communication with the external environment. It can be used for extracting further information about the structure and robustness of the network which is employed for routing, controlling and management purposes.

Suppose we have a wireless sensor network with all nodes deployed randomly. There can be two kinds of nodes: those which are on the boundary and the interior ones. We assume that each node can get its localization within the deployment zone. Many existing methods can be used for this information [37, 44, 60].

In order to define a wireless sensor network formally, we model it as a Euclidean graph $G = (V, E)$. We use the letter $n$ to denote $|V|$, the number...
of nodes in the network. For each node $i$ of $G$, we use the notation $\text{nbrs}_i$ to denote the ”number of neighbors” of $i$, that is those nodes located in the communication range of $i$.

We also define a message alphabet, described in Table 1, and the functions used in the algorithm in Table 2.

<table>
<thead>
<tr>
<th>Message</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>AC</td>
<td>Request coordinates</td>
</tr>
<tr>
<td>CS</td>
<td>Send coordinates</td>
</tr>
<tr>
<td>SN</td>
<td>Select a node</td>
</tr>
<tr>
<td>DN</td>
<td>Reset node</td>
</tr>
</tbody>
</table>

Table 1: Different messages and their description.

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>getId()</td>
<td>Returns the node identifier</td>
</tr>
<tr>
<td>getCoord()</td>
<td>Returns the node coordinates $(x, y)$</td>
</tr>
<tr>
<td>getNumberOfNeighbors()</td>
<td>Returns $\text{out} - \text{nbrs}_i$, the number of neighbors of the node</td>
</tr>
<tr>
<td>send(m, adr)</td>
<td>Sends a message $m$ to the node having address $adr$,</td>
</tr>
<tr>
<td></td>
<td>if $adr = \ast$ it sends a message to all outgoing neighbors</td>
</tr>
<tr>
<td>reset()</td>
<td>Reset node</td>
</tr>
</tbody>
</table>

Table 2: Functions used by D-RRLPCN.

Algorithm 3 exhibits the main steps performed by each sensor node for the identification of the boundary nodes of a wireless sensor network. These steps can be explained by the flowchart of Figure 7. In this case, the nodes are completely independent and the x-coordinate of the starting node is sent to the next boundary node in each iteration. If a node receives an x-coordinate which is smaller than its own x-coordinate, this node will send to the previous found boundary nodes a message to reset them as non-boundary nodes. Then, it will restart the process of finding the boundary nodes.

5 Validation of the proposed algorithms

5.1 Analytical validation

In order to validate the proposed algorithm regarding optimality and convergence, we will present, in the following, some mathematical concepts that will allow us to define the final version of the proposed algorithm.

**Proposition 1**

*After the application of LPCN starting from a non-boundary vertex $v_1 \in V$, with $X_{v_1}$ its x-coordinate and the degree of $v_1$ being equal or greater than one,*
Algorithm 3 Distributed Reset and Restart Least Polar-angle Connected Node (D-RRLPCN).

1: boundary = false;
2: once = true;
3: cid = getId();
4: cx = getX();
5: cy = getY();
6: i = 0;
7: n = getNumberOfNeighbors();
8: nid = 0;
9: nid_ref = -1;
10: phi_min = 10;
11: first = id of the starting node;
12: prevList=();
13: repeat
14:   if (cid==first and once) then
15:     boundary = true;
16:     once = false;
17:   end if
18:   id = read();
19:   type = read();
20:   if (type=="AC") then
21:     data = cid+"|"+"AC";
22:     send(data, *);
23:   end if
24:   if (type=="CS") then
25:     id = read();
26:     nx = read();
27:     ny = read();
28:     phi = angle(px, py, cx, cy, nx, ny);
29:     if (phi<phi_min) then
30:       phi_min = phi;
31:     end if
32:     nid = id;
33:     i = i + 1;
34:     if (i==n) then
35:       if (nid==nid_ref and cid==first) then
36:         STOP
37:       else
38:         if (nid_ref<0) then
39:           nid_ref = nid
40:         end if
41:       end if
42:     end if
43:     data = cid+"|"+"SN"+"|"+cx+"|"+cy+"|"+xmin;
44:     send(data, nid);
45:   end if
46: end if
47: i = 0;
48: data = cid+"|"+"SN"+"|"+cx+"|"+cy+"|"+xmin;
49: send(data, nid);
50: end if
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52: if (type=="SN") then
53:    boundary = true;
54:    n = getNumberOfNeighbors();
55:    nid = 0;
56:    phi_min = 10;
57:    px = read();
58:    py = read();
59:    xmin = read();
60:    if (cx<xmin) then
61:       delay(t)
62:       once = true;
63:       fi
64:       first = cid;
65:       data = cid+"|"+"DN"+"|"+first;
66:       send(data, id);
67:    else
68:       prevList.add(id);
69:       data = cid+"|"+"AC";
70:       send(data, *);
71:    end if
72:    end if
73: if (type=="DN") then
74:    boundary = false;
75:    first = read();
76:    if (prevList.size()>0) then
77:       data = cid+"|"+"DN"+"|"+first;
78:       previous = prevList.getLast();
79:       prevList.removeLast();
80:       if previous.hasPrevious() then
81:          send(data, previous);
82:       end if
83:    end if
84:    end if
85: until false

The sub-graph visited by LPCN is a cycle containing at least one vertex \( v_0 \in V \) with:

\[ X_{v_0} < X_{v_1} \]  

(3)

where \( X_{v_0} \) is the \( x \)-coordinate of \( v_0 \).

Proof

Suppose the opposite, i.e., \( v_1 \) is a non-boundary vertex and there exists no vertex visited by LPCN that has a smaller \( x \)-coordinate. This implies that \( X_{v_1} \) is the minimal \( x \)-coordinate of the resulting subgraph \( G' \). But \( G' \) is a boundary of \( G \) and \( v_1 \in G' \) which means that \( v_1 \) is a boundary vertex. This is a contradiction to the fact that \( v_1 \) is a non-boundary vertex.

Theorem 1

Except for the finally calculated angle of the starting point, the RRLPCN algorithm never calculates the same polar angle more than once.

Proof

We assume that the algorithm starts from vertex \( P_{\text{first}} \) and that at iteration
we can, without loss of generality, represent the current set of boundary nodes $B^i_V = \{P_{\text{first}}, \ldots, P_i\}$ by a vertex $PiBV$. On the other hand, we assume that angles are calculated only in anti-clockwise order. Hence, any angle, as given by two specific edges, will be calculated only once. Otherwise, if an angle is calculated a second time, then the algorithm must pass a second time through the vertex $PiBV$. This means that we pass a second time through the starting vertex $P_{\text{first}}$ which represents the stop condition of LPCN (cf. line 12 of Algorithm 2). This leads to a contradiction and therefore it is not possible to calculate an angle more than once.

**Theorem 2**

The RRLPCN algorithm never runs the Restart Step (step-14) from the same point.

**Proof**

Assume that at iteration $k$ we will start $RRLPCN$ with a vertex $P^k$ that has already been visited. Let $B^{k-1}_V$ be the set of the obtained vertices. Since the algorithm restarts from the same vertex, $X^k_P = \arg\min_{v \in B^{k-1}_V} X_v$, which means that $p_k$ is a boundary vertex according to Theorem 1. Which is the stop condition of LPCN.
Corollary 1
*The RRLPCN algorithm converges in a finite number of steps.*

Theorem 3 (See [36])
*The polygon hull found by LPCN is of minimum cardinality.*

Lemma 1
*The polygon hull found by RRLPCN is of minimum cardinality.*

Proof
When the algorithm executes the last iteration, it stops and finishes with a vertex \( v_f \) (stop condition satisfied). This means that there is no point \( v_i \in V \) such that \( x_{v_i} < x_{v_f} \). In other words, \( \forall v_i \in V : x_{v_i} < x_{v_f} \), consequently \( v_f \) is a vertex with minimum \( x \)-coordinate in the graph. Starting LPCN from this vertex will give us the polygon hull of the graph according to Theorem 3.

5.2 Application to Wireless Sensor Networks

5.2.1 Simulation results

To validate and test our distributed algorithm, we have used the two simulation environments CupCarbon [46] and TinyOS [40] Tossim [39]. The first one allowed us to develop our protocol and to make debugging, and the second one was used to test the final version and to analyze the performances.

CupCarbon [46] is a Smart City and Internet of Things simulator. Its objective is to design, visualize, debug and validate distributed algorithms for monitoring, environmental data collection, and to create ecological scenarios, generally within educational and scientific projects. CupCarbon proposes two simulation environments. The first is a multi-agent environment, which enables the creation of the mobility scenarios and the generation of events such as fires and gas as well as the simulation of mobiles such as vehicles and flying objects [42]. The second simulation environment uses the discrete event simulation of wireless sensor networks taking into account the scenario created by the first environment.

TOSSIM, the TinyOS Simulator, uses its hierarchical model by replacing lower level hardware components with software-emulated ones. This approach reduces the hole between the simulator and the real environment. By substituting low-level components, a high fidelity between reality and the simulation environment is achieved. One of the advantages of the simulation with TOSSIM is the reuse of the code since the code used for the simulation is the same that will be exported to the real sensor nodes.

Energy efficiency is one of the most critical factors for WSN applications. The lower the energy consumed by each node, the longer the network can perform its mission. For this reason, our simulations are based on this measure of performance. We calculate the average of energy consumed by each node in miliJule (mJ) according to simulation parameters.
Figure 8 represents the average consumption of nodes in mJ according to their position (boundary nodes, neighbors of boundary nodes and inner nodes). The purpose of this histogram is to compare the consumption of the nodes in the network according to their positions. As can be noted, the boundary nodes consume more energy, because they are the ones that participate most in the algorithm, followed by the neighbors of boundary nodes and finally the inner nodes.

![Energy consumption according to sensor node positions in D-RRLPCN.](image)

**Fig. 8:** Energy consumption according to sensor node positions in D-RRLPCN.

As we have seen in Figure 8, the boundary nodes consume more energy in comparison to other nodes. In Figure 9, the consumption of these nodes according to their number of neighbors is examined. There is a correlation between the energy consumption of these nodes and their number of neighbors. The more the number of neighbors increases the more they consume energy. One of the solutions to reduce this consumption is the reduction of the network density.

It is possible to not activate all network nodes at the same time, which will increase the lifetime of the network and reduce the consumption of the boundary nodes.
In order to prove the efficiency of D-RRLPCN regarding energy consumption, we have compared it with D-LPCN, and three other algorithms (D-LCPN, Min Finding, and D-LPCN+Min Finding).

Figure 10 shows a comparison histogram of the average of energy consumption of nodes depending on their position in the network (boundary nodes, neighbors of boundary nodes and inner nodes) for the two algorithms D-RRLPCN and D-LPCN. The graph allows us to deduce that whatever the position of the node in the network, D-RRLPCN consumes less energy than D-LPCN.
The D-LPCN algorithm consists of two phases. The first phase allows to find the starting node which is the node with minimum x-coordinate (in this paper we call this part “Min Finding”). The second phase allows to start from the starting sensor node found in the first phase to obtain a boundary of the network.

Figure 11 shows the energy consumption of the boundary nodes depending on their number of neighbors, for D-RRLPCN, the Min Finding, D-LPCN without the first part (Min Finding) and the last graph representing the sum of the two phases of D-LPCN. Considering the two phases of D-LPCN, D-RRLPCN is more efficient regardless of the number of neighbors.

In Figure 12, a comparison with existing methods is given. We have compared the energy consumption of three algorithms, namely Hop-based [29], Distributed Boundary Detection (DBD) [58], and Located Voronoï Polygons (LVP) [1] with that of D-RRLPCN. The energy consumption of boundary nodes is calculated according to their number of neighbors.
5.2.2 Implementation in Real Sensor Nodes

Building a wireless sensor network system needs development and integration of many hardware and software components. Figure 13 shows the overall architecture of the wireless sensor network system that we have developed. The system includes a number of distributed wireless sensor nodes, where each sensor node is a combination of a microcontroller, an XBee radio transceiver, and a battery.

→ Design of Sensor Node
In this research, we have studied sensor nodes using Arduino and Digi XBee modules. Arduino is an open-source computer hardware and software company, project and user community that designs and produces devices for building digital applications and interactive objects that can sense and control the physical world. These devices are based on a family of microcontroller board designs produced primarily by Smart Projects in Italy, and also by numerous other vendors, using different 8-bit Atmel AVR microcontrollers or 32-bit Atmel ARM processors. In this work, we have used Arduino Uno R3 based on the Atmel Atmega328 microcontroller.

For wireless communication, we have used the XBee module from Digi. Xbee is basically used for this purpose. It offers the IEEE 802.15.4 connectivity in the 2.4GHz ISM band. There is a “pro” version which provides large communication range, denoted as XBP24, and a “normal” version, denoted as XB24. The term “series 1” is employed for the 802.15.4 version and “series 2” for the ZigBee version. The XBee module uses a UART (serial interface) to communicate with the main board. The advantage is simplicity and the possibility to re-use many serial tools. For our implementation, we have used the Xbee series 1. The indoor communication range of this module is 30 m
whereas the outdoor range is nearly 100 m. With low power consumption and data rates of up to 250 kbps, Xbee devices are particularly suitable for fast prototyping of wireless sensor network applications. It is possible to build a simple star-structured network or a complex mesh network using these devices. Table 3 summarizes the technical characteristics of the used sensor nodes.

<table>
<thead>
<tr>
<th>Devices</th>
<th>Specifications</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arduino UNO R3</td>
<td>Microcontroller</td>
<td>ATmega328P</td>
</tr>
<tr>
<td></td>
<td>Flash Memory</td>
<td>32 KB</td>
</tr>
<tr>
<td></td>
<td>Clock Speed</td>
<td>16 MHz</td>
</tr>
<tr>
<td></td>
<td>SRAM</td>
<td>2 KB</td>
</tr>
<tr>
<td></td>
<td>EEPROM</td>
<td>1 KB</td>
</tr>
<tr>
<td>Xbee XB24</td>
<td>Frequency Band</td>
<td>2.4 GHz</td>
</tr>
<tr>
<td></td>
<td>RF Data Rate</td>
<td>250 kbps</td>
</tr>
<tr>
<td></td>
<td>Indoor/Urban Range</td>
<td>30 m</td>
</tr>
<tr>
<td></td>
<td>Outdoor/RF Line-of-Sight Range</td>
<td>100 m</td>
</tr>
<tr>
<td></td>
<td>Transmit Current</td>
<td>45 mA</td>
</tr>
<tr>
<td></td>
<td>Receive Current</td>
<td>50 mA</td>
</tr>
<tr>
<td>Battery</td>
<td>Capacity</td>
<td>2200 mAh</td>
</tr>
</tbody>
</table>

Table 3: Technical characteristics of sensor nodes.

Figure 13 shows a wireless sensor network with 8 Arduino/Xbee sensor nodes deployed randomly. The sensor nodes which have a green led turned on are the boundary nodes detected by the D-RRLPCN algorithm.

Fig. 13: Execution of D-RRLPCN using real Arduino/XBee sensor nodes.
6 Other applications

There are several other problems and domains where we can apply the algorithms proposed in this paper (RRLPCN and D-RRLPCN). We can cite two examples:

Drawing object contours in images:
The algorithm combined with an algorithm to characterize the selected zone can be used to draw object contours in images. The user has to click on a pixel of the zone to study, and starting from this point, RRLPCN can detect the contour of the selected zone (object). This technique can be applied to identify the shape of a tumor, for example. RRLPCN can be very useful in the medical field. Indeed, detecting the boundary of a tumor with high precision is a very crucial task for the medical practitioner. With RRLPCN, this task can be done by selecting any pixel of the radio representing this tumor.

Defining the boundary of a two-dimensional Multi-Robot system:
a Multi-Robot system can be described as a set of robots operating in the same environment. In these systems, we can use a boundary as a formal practical definition of what is inside and outside the network. Knowing the boundary would allow us to estimate the perimeter of the configuration. For a surveillance application [6], robots on the boundary can specialize in target monitoring, and notify the network when a target has entered or left the tracking area. At each moment one of the robots can launch the D-RRLPCN algorithm, and the boundary robots will be detected.

7 Conclusion

In this article, we have proposed a new technique called Reset and Restart, which in combination with LPCN and D-LPCN allowed us to introduce two new algorithms: RRLPCN and D-RRLPCN. Both algorithms now find the polygon hull of a Euclidean graph without any assumption on the starting node where the second is the distributed version of the first and particularly suited for wireless sensor networks. We have also validated the proposed method theoretically and computationally.

The D-RRLPCN algorithm has been developed and simulated using the two simulation environments CupCarbon and TOSSIM. The results show that it consumes less energy than D-LPCN and Hop-based, DBD, and LVP. The algorithm has been implemented in two real sensor platforms, TelosB and Arduino Xbee. As future work, we suggest an extension of the RRLPCN algorithm to 3D space and a proposition of a distributed version for boundary node detection in 3D wireless sensor networks.
References


