

## MIT Open Access Articles

### *Computing Signed Permutations of Polygon*

The MIT Faculty has made this article openly available. **Please share** how this access benefits you. Your story matters.

**Citation:** Aloupis, Greg et al. "Computing Signed Permutations of Polygon." International Journal of Computational Geometry & Applications 21.01 (2011): 87-100.

**As Published:** <http://dx.doi.org/10.1142/s0218195911003561>

**Publisher:** World Scientific

**Persistent URL:** <http://hdl.handle.net/1721.1/73973>

**Version:** Author's final manuscript: final author's manuscript post peer review, without publisher's formatting or copy editing

**Terms of use:** Creative Commons Attribution-Noncommercial-Share Alike 3.0



**Massachusetts Institute of Technology**

# Computing Signed Permutations of Polygons \*

Greg Aloupis <sup>†</sup>    Prosenjit Bose<sup>‡</sup>    Erik D. Demaine<sup>§</sup>

Stefan Langerman<sup>†</sup>    Henk Meijer<sup>¶</sup>    Mark Overmars<sup>||</sup>  
Godfried T. Toussaint<sup>†</sup>

## Abstract

Given a planar polygon (or chain) with a list of edges  $\{e_1, e_2, e_3, \dots, e_{n-1}, e_n\}$ , we examine the effect of several operations that *permute* this edge list, resulting in the formation of a new polygon. The main operations that we consider are: *reversals* which involve inverting the order of a sublist, *transpositions* which involve interchanging subchains (sublists), and *edge-swaps* which are a special case and involve interchanging two consecutive edges. Using these permuting operations, we explore the complexity of performing certain actions, such as convexifying a given polygon or obtaining its mirror image. When each edge of the given polygon has also been assigned a *direction* we say that the polygon is *signed*. In this case any edge involved in a reversal changes direction. The complexity of some problems varies depending on whether a polygon is signed or unsigned. An additional restriction in many cases is that polygons remain simple after every permutation.

## 1 Introduction

Much focus has been placed recently on the problem of sorting a permutation of  $n$  integers by *reversals* [HP98, BP95]. As one might guess, a single reversal

---

\*This research was partly funded by NSERC and FCAR.

<sup>†</sup>School of Computer Science, McGill University. [{athens,sl,godfried}@uni.cs.mcgill.ca](mailto:{athens,sl,godfried}@uni.cs.mcgill.ca)

<sup>‡</sup>Carleton University. [jit@scs.carleton.ca](mailto:jit@scs.carleton.ca)

<sup>§</sup>MIT Laboratory for Computer Science, 200 Technology Square, Cambridge, MA 02139, USA, [edemaine@mit.edu](mailto:edemaine@mit.edu)

<sup>¶</sup>Queens University. [henk@cs.queensu.ca](mailto:henk@cs.queensu.ca)

<sup>||</sup>University of Utrecht. [markov@cs.uu.nl](mailto:markov@cs.uu.nl)

is applied to a consecutive set of these integers and the result is that their order is inverted. The key problem that arises is determining the minimum number of reversals necessary to sort a given permutation. This number is called the *reversal distance* of the permutation. A variation of this problem involves *signed* permutations [BMY01]. In this case any integer affected by a given reversal also changes *parity*.

Each of these interesting combinatorial problems has its roots in bioinformatics and molecular biology [HP98, HP96, BP95, CJM<sup>+</sup>00, BMY01]. Specifically, genomes have been modeled as linear or cyclic sequences, where each element in a sequence is a *block* of smaller elements that are never separated. A popular model for mutation involves inverting parts of these sequences. In order to determine the number of such mutations needed to transform one genome to another, one may compute the reversal distance of the associated permutations. An extension of this model is to consider the direction of each *block*. This leads to the study of signed permutations. We illustrate signed inversions in Figure 1 which has been modified from [BP95].

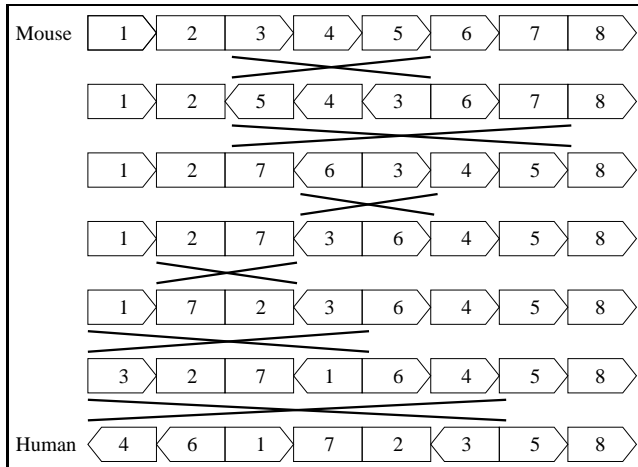


Figure 1: A most parsimonious evolutionary scenario for the transformation of human into mouse chromosome assuming that the X chromosome evolves solely by inversions [BP95]. Each block represents a conserved linkage group of genes. Reversal distance is equal to six.

The problem of computing *transposition distance* also stems from bioin-

formatics. In this case, a transposition involves exchanging two disjoint sets of consecutive integers in a permutation. Computing reversal distance has been shown to be NP-hard [Cap99] for unsigned permutations, but for the signed version a linear time algorithm exists [BMY01]. Computing transposition distance is of unknown complexity [CJM<sup>+</sup>00]. The reader may also be interested in [BHK01, BP98, EEK<sup>+</sup>01].

In this paper we extend the ideas mentioned above from one dimension to two. Instead of considering permutations of integers, we consider permutations of *edges* which form polygons or chains. We define operations such as *edge-swaps*, *reversals* and *transpositions*, in analogy to  $\mathbb{R}^1$ . We introduce the notions of *signed permutations of polygons and chains*. These concepts give rise to a wide range of problems to be solved.

## 2 Definitions

First we introduce the notion of a *signed polygon* or *signed permutation* of a polygon. Any polygon  $P$  may be described by a list of edges  $\{e_1, e_2, e_3, \dots, e_{n-1}, e_n\}$ . A signed polygon is no different, except that each edge is also assigned a direction. The same holds for chains. This is a generalization of the notion of parity that is used in  $\mathbb{R}^1$ . If the directions of all edges are consistent as we traverse a polygon or chain, then this polygon or chain is *oriented*. In Figure 2 we illustrate some signed polygons and chains.

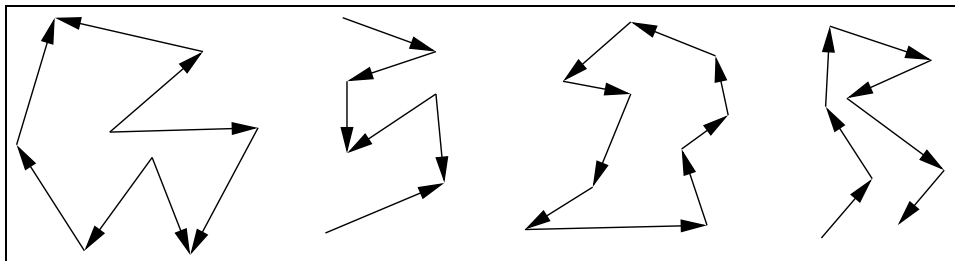


Figure 2: From left to right, a signed polygon, signed chain, oriented polygon, oriented chain.

Without loss of generality, suppose that we are dealing with an oriented polygon. A *transposition* of two edges  $A$  and  $B$  involves interchanging their positions so that the resulting polygon remains oriented. This is illustrated in

Figure 3. If  $A$  and  $B$  are consecutive, this operation is defined as an *edge-swap* or plainly *swap* (Figure 4a). It is not difficult to see that entire subchains may also be transposed. A *single-edge* transposition involves transposing an edge with an empty subchain. One may also think of this operation as a transposition between the single edge and one of its neighboring subchains (Figure 4b).

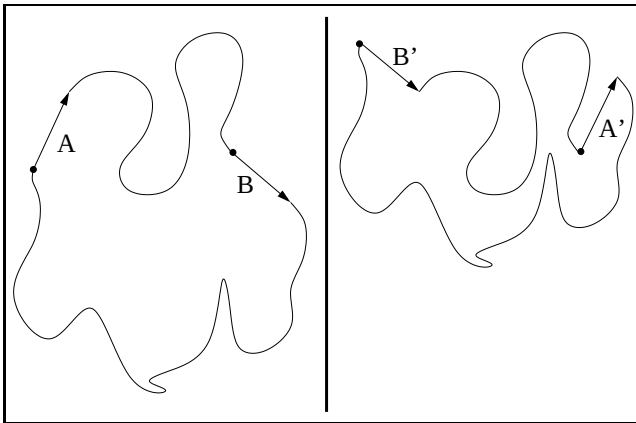


Figure 3: Transposing two edges  $A$  and  $B$ .

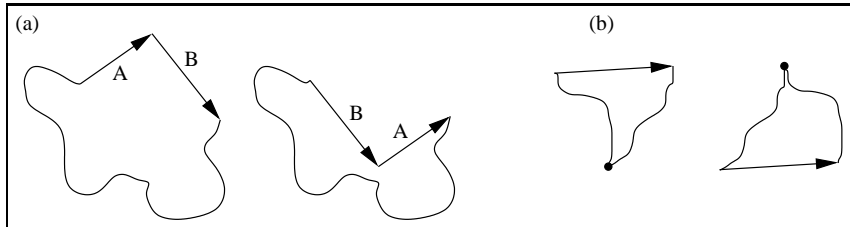


Figure 4: (a) An edge swap. (b) A single-edge transposition.

A *reversal* of a subchain belonging to a polygon involves inverting the order of the edges in the subchain. Geometrically this rotates the subchain rigidly in the plane by an angle of  $\pi$  so that its endpoints are placed exactly at each other's original location. For unsigned polygons this operation appears identical to the *flipturn* operation introduced by Joss and Shannon [GZ01]. However, here we allow reversals to take place on any subchain, not only

on pockets. For signed polygons the direction of each edge involved in the reversal is switched, as is done for parity in  $\mathbb{R}^1$ . In Figure 5 we illustrate a reversal of subchain  $\{e_i, \dots, e_j\}$  for a signed (initially oriented) polygon. One

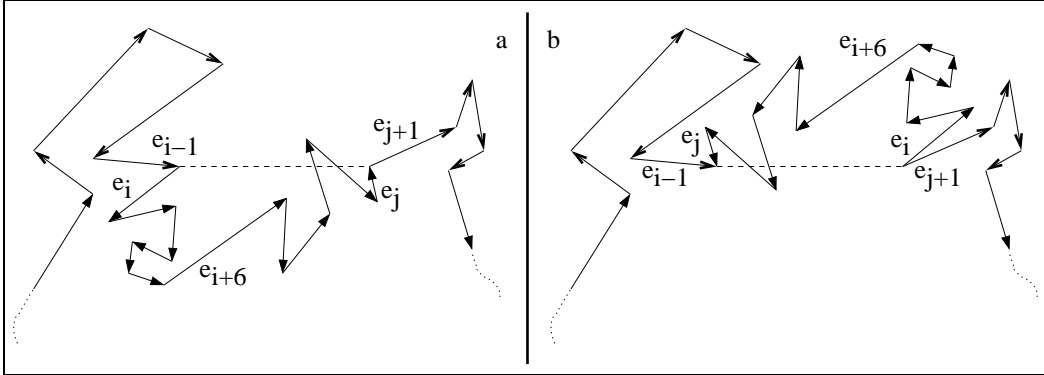


Figure 5: Reversing a subchain of a signed polygon.

can see that for unsigned polygons, an edge-swap is merely a transposition *or* a reversal of two consecutive edges. For signed polygons there is a difference in the resulting direction of each edge.

Each of the operations above results in the same shape when used on a polygon, regardless of the directions of its edges. In other words, to compute how a polygon changes shape, one can imagine that it is oriented. However, for chains alternate definitions exist. For example consider the oriented chain in Figure 6. We may choose to perform a reversal on edges  $(A, B, C)$  in at least two ways. One way (shown on top) is identical to what is done for polygons. This is convenient but also means that the endpoints of the chain will never move. A second way (shown at the bottom) is to preserve orientation. This may allow the chain to form more interesting configurations. We use the latter definition in Theorem 3.6 in the next section.

### 3 Permuting Polygons

Scott [Sco82] has shown that precisely two permutations of an edge list form oriented convex polygons, and these have maximal area. It is also known that if the longest edge of a polygon has unit length, this polygon may be permuted to fit into a circle of radius  $\sqrt{5}$  [GY79].

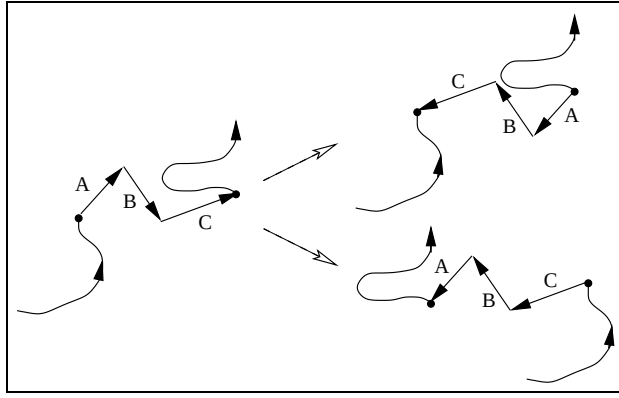


Figure 6: Two ways that a reversal may be defined on a chain.

For the remainder of this section we present our results concerning permutations of polygons or chains. We impose the restriction that simplicity must be maintained at all times, unless mentioned otherwise.

In Figure 7a we show a polygon which does not admit any edge-swaps. Examples such as this one may be extended easily to create any  $n$ -gon which will not admit edge-swaps. In Figure 7b we show a polygon which does not admit single-edge transpositions, with the exception of a few edge-swaps for some edges that are almost collinear. These transpositions cannot change the basic shape. Thus we see that sometimes local permutations will not be sufficient to achieve desired reconfigurations.

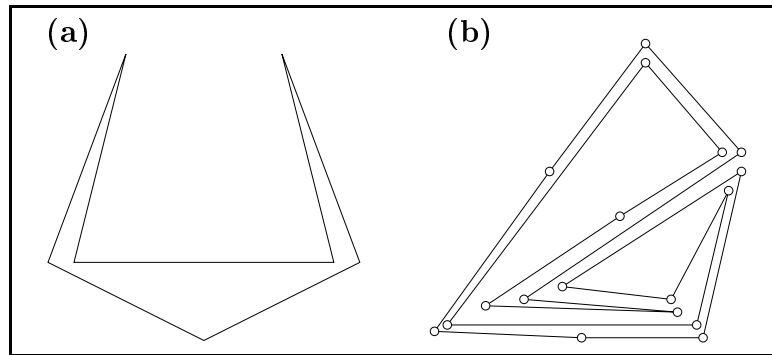


Figure 7: Polygon (a) does not admit edge-swaps. Polygon (b) does not admit single-edge transpositions.

**Theorem 3.1** A simple polygon may be convexified with  $O(n^2)$  reversals while maintaining simplicity after each reversal.

*Proof:* This result holds for the more restricted reversal operation of flipturns  $[ABC^{+00}]$ .  $\square$

**Theorem 3.2** A star-shaped polygon can be convexified with  $O(n^2)$  edge-swaps while maintaining star-shapedness after each edge-swap, and this bound is tight in the worst case.

*Proof:* Let  $k$  be a point in the kernel and without loss of generality suppose that the polygon is oriented clockwise. If the polygon is not convex, there must exist two successive edges  $\overrightarrow{ab}$  and  $\overrightarrow{bc}$  which form a left hand turn (see Figure 8a). Since the polygon is star-shaped,  $b$  is the only vertex in

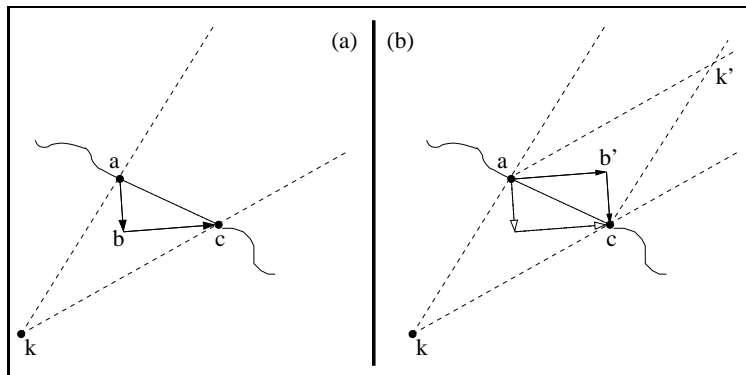


Figure 8: Using edge-swaps to convexify a star-shaped polygon.

the cone formed by the half-lines  $ka$  and  $kc$ . If we edge-swap  $\overrightarrow{ab}$  and  $\overrightarrow{bc}$ , we obtain the configuration shown in Figure 8b. The new position of  $b$  (shown as  $b'$ ) must be somewhere in the triangle  $(a, c, k')$ . The swapped edges are still visible from  $k$ , and they do not interfere with the other edges of the polygon. Thus the polygon remains star-shaped. Furthermore any point in the kernel remains in the kernel and any point in the polygon remains in the polygon.

Every edge  $e$  may be found only within a halfplane determined by a line parallel to  $e$  that passes through  $k$ . Now suppose that two edges,  $\overrightarrow{ab}$  and  $\overrightarrow{cd}$  form a right hand turn. This means that  $b$  and  $c$  coincide as shown

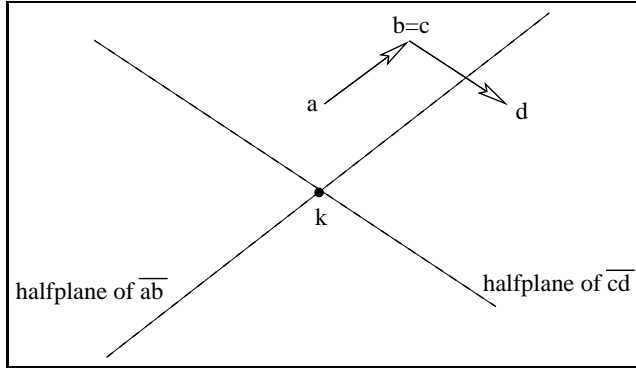


Figure 9: Any pair of edges may be swapped at most once.

in Figure 9. It is impossible to move these edges within their respective halfplanes and into a left hand turn without obstructing visibility from  $k$  to either  $b$  or  $c$ . Thus once a pair of edges forming a left hand turn are swapped, they will never form a left hand turn again. The polygon will become convex only when there are no swaps to be made on left hand turns. Since any pair of edges may be swapped at most once,  $O(n^2)$  swaps suffice to convexify a star-shaped polygon. In Figure 10 we show that this bound is tight. Every edge  $e_i$  ( $2 \leq i \leq n-2$ ) must be swapped with edges  $e_1, \dots, e_{i-1}$  for the polygon to become convex.  $\square$

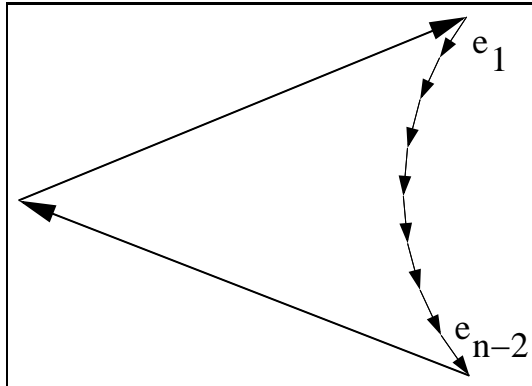


Figure 10: A star-shaped polygon which requires  $\Omega(n^2)$  edge-swaps to become convex.

For the following two theorems we do not enforce simplicity.

**Theorem 3.3** *Determining whether a signed polygon may be permuted using transpositions so that its shape is rotated by an angle of  $\pi$  takes  $\Theta(n \log n)$  time in the algebraic decision tree model of computation.*

*Proof:* Since only transpositions are allowed, each edge of the polygon must have its *opposite* also present in the polygon. This property also suffices if we do not impose the restriction of maintaining simplicity at all times. By “opposite” we mean an edge with the same angle, but opposite direction. For example, in Figure 11 edges  $a, b, c, d, e$  of the polygon on the left are matched by edges  $\bar{a}, \bar{b}, \bar{c}, \bar{d}, \bar{e}$ . This means that the shape of this polygon may be rotated by an angle of  $\pi$  (shown on right) with appropriate transpositions.

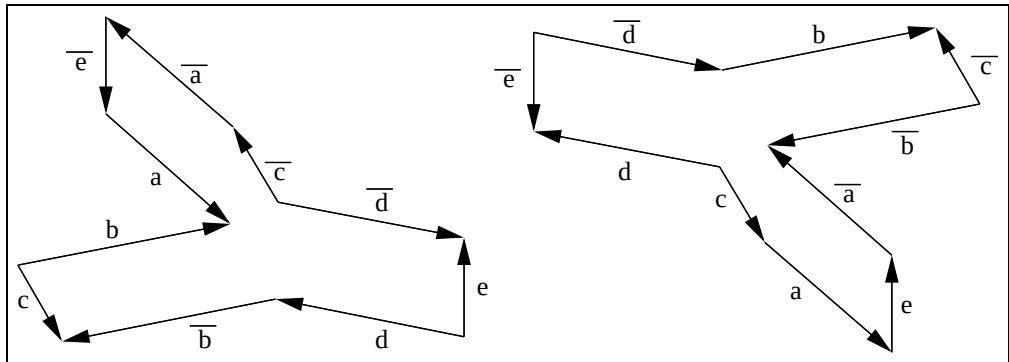


Figure 11: Left: a signed polygon for which every edge is matched by an “opposite”. Right: a permutation of the polygon with the same shape rotated by an angle of  $\pi$ .

If we translate every edge to the origin (so that they are directed away from the origin), we obtain a set of  $n$  points. The shape of the given polygon can be rotated if and only if every such point has a reflection through the origin. This can be determined in  $O(n \log n)$  time with a radial sort, and the matching lower bound is obtained by a reduction from *Set Equality* (see [Ead88]).  $\square$

**Theorem 3.4** *Determining whether we can obtain the mirror image of a signed polygon using transpositions takes  $\Theta(n \log n)$  time in the algebraic decision tree model of computation.*

*Proof:* In order to be able to obtain a mirror image, there must exist an axis through which every edge has its reflection present (allowing translation). For example consider the polygon on the left in Figure 12. If we take a vertical line as an axis of symmetry, then edges  $d$  and  $j$  are reflections of each other. The same holds for pairs  $(b, h)$  and  $(f, k)$ . Vertical edges do not need a matching edge. If such an axis exists, then a mirror image of the polygon can be obtained using transpositions. As in Theorem 3.3 we can place every edge at the origin to obtain a set of  $n$  points. The symmetries of this point set may be found using the Knuth-Morris-Pratt string matching algorithm [KMP77]. The overall time complexity is  $O(n \log n)$ . This is pointed out by Eades [Ead88] who also mentions that such reflection tests have  $\Omega(n \log n)$  lower bounds on fixed degree decision tree machines.  $\square$

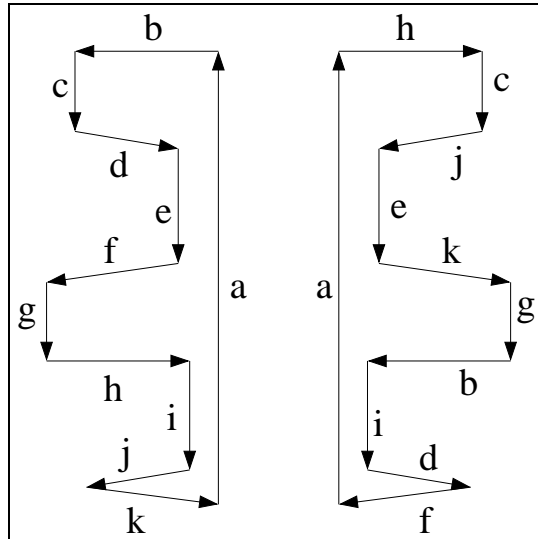


Figure 12: Two polygons that are mirror images and have different permutations of the same edge list.

**Theorem 3.5** *Given an oriented polygon  $P$  and a rectangle  $R$ , deciding whether  $P$  can be permuted by transpositions into an oriented polygon  $P'$  that can be drawn inside  $R$  is (weakly) NP-complete.<sup>1</sup>*

---

<sup>1</sup>This result also holds if  $P'$  is to be placed inside a strip or circle, instead of a rectangle.

*Proof:* Consider an integer partition problem with  $S = \{a_0, a_1, \dots, a_{n-1}\}$  and  $a_i > 0$  for all  $i$ . Let  $A = \sum_{a \in S} a/2$ . Deciding whether there is a subset  $S'$  of  $S$  with  $\sum_{a \in S'} a = A$  is (weakly) NP-hard. Consider the following polygon  $P$ . Denote the edges of  $P$  in counter-clockwise order by  $\{e_0, e_1, \dots, e_{2n+3}\}$ . The edges with even indices are parallel to the  $x$ -axis; the edges with odd indices are parallel to the  $y$ -axis. Let  $\epsilon$  be a positive number less than one. The edge  $e_0$  has length  $a_0 + \epsilon$ . Edges  $e_i$  for  $i = 2, 4, 6, \dots, 2n - 2$  have length  $a_{i/2}$ . Edge  $e_{2n}$  has length  $A$ . Edge  $e_{2n+2}$  has length  $A + \epsilon$ . Edges  $e_i$  for  $i = 1, 3, 5, \dots, 2n + 1$  have length 1. Edge  $e_{2n+3}$  has length  $n + 1$ .

We also assign directions to the edges, so that the edges form a counter-clockwise traversal of  $P$ . All edges of length 1 go up. The edges  $e_i$  for  $i = 0, 2, 4, \dots, 2n - 2$  go from left to right. The few remaining edges go down and right to left, as illustrated in Figure 13(a) with  $n=7$ .

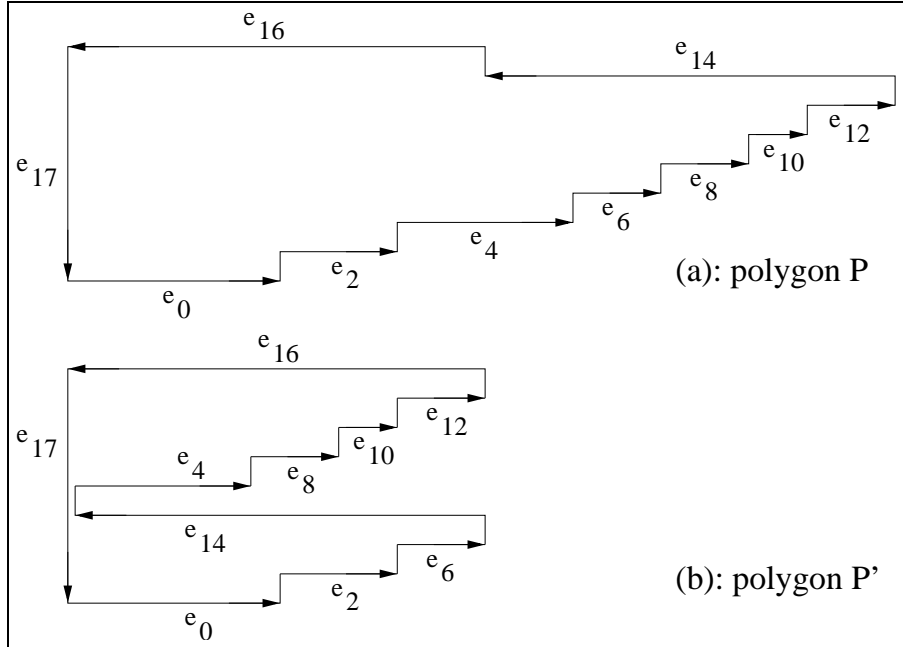


Figure 13: Polygons  $P$  and  $P'$  with 18 vertices.

Let  $R$  be a rectangle of size  $A + \epsilon$  by  $n + 1$ . W.l.o.g assume that  $R$  has  $(-\epsilon, 0)$  and  $(A, n + 1)$  as its left-bottom and right-top corner. Suppose  $P$  can be permuted into a polygon  $P'$  that can be drawn in the rectangle  $R$ . Again w.l.o.g assume that  $e_{2n+2}$  of  $P'$  lies along the top side of  $R$  and  $e_{2n+3}$

along the left side of  $R$ . This implies that the left and right endpoints of  $e_{2n}$  are  $(0, y)$  and  $(A, y)$  for some value of  $y$  with  $1 \leq y \leq n$ . Moreover the edge  $e_0$  lies below  $e_{2n}$ . The edges form a counter-clockwise traversal of  $P'$ . Since  $e_{2n}$  has a direction that goes from right to left, the horizontal edges above  $e_{2n}$  connect the left endpoint of  $e_{2n}$  with the right endpoint of  $e_{2n+2}$ , so their lengths must add up to  $A$ . Therefore the partition has a solution if and only if  $P$  can be permuted into a polygon  $P'$  that fits in  $R$ . Figure 13(b) shows a permutation of the polygon in Figure 13(a) that fits in rectangle  $R$ .  $\square$

**Theorem 3.6** *The maximum endpoint distance over all permutations of an oriented chain may be computed in  $O(n \log n)$  time.*

*Proof:* Fix one endpoint at the origin. Endpoint distance depends only on the direction of each edge. If we knew the direction in which to position the second endpoint, it would be a simple matter to select the direction of each edge in order to maximize the distance. Position two vectors at the origin for each edge, representing its possible directions. Sort the vectors radially and compute the sum of all vectors in one halfplane determined by a line  $\ell$  through the origin. This represents the maximum distance in a direction perpendicular to  $\ell$ . By rotating  $\ell$  and updating the vector sum whenever a vector enters or exits the rotating halfplane, we obtain the endpoint distance over all directions. The time complexity is dominated by the sorting step, so the entire procedure takes  $O(n \log n)$  time using  $O(n)$  space.  $\square$

## Acknowledgements

Much of this work was completed at the *Workshop on Geometric Aspects of Molecular Reconfiguration* organized by Godfried Toussaint at the Bellairs Research Institute of McGill University in Barbados, February 2002. We appreciate the helpful discussions with the other participants: David Bremner, Vida Dujmovic, Herbert Edelsbrunner, Jeff Erickson, Ferran Hurtado, Patrick Morin, Joseph O'Rourke, Suneeta Ramaswami, Ileana Streinu, Yusu Wang. We also appreciate the comments of an anonymous referee.

## References

- [ABC<sup>+</sup>00] Hee-Kap Ahn, Prosenjit Bose, Jurek Czyzowicz, Nicolas Hanusse, Evangelos Kranakis, and Pat Morin. Flipping your lid. *Geombinatorics*, X(2):57–63, 2000.
- [BHK01] Piotr Berman, Sridhar Hannenhalli, and Marek Karpinski. 1.375-approximation algorithm for sorting by reversals. Technical Report DIMACS TR:2001-41, 2001.
- [BMY01] David A. Bader, Bernard M.E. Moret, and Mi Yan. A linear time algorithm for computing inversion distance between signed permutations with an experimental study. *Journal of Computational Biology*, 8(5):483–491, 2001.
- [BP95] Vineet Bafna and Pavel A. Pevzner. Sorting by reversals: Genome rearrangements in plant organelles and evolutionary history of X chromosome. *Mol. Biol. Evol.*, 12(2):239–246, 1995.
- [BP98] Vineet Bafna and Pavel A. Pevzner. Sorting by transpositions. *SIAM J. Discrete Math.*, 11(2):224–240, May 1998.
- [Cap99] Alberto Caprara. Sorting permutations by reversals and Eulerian cycle decompositions. *SIAM J. Discrete Math.*, 12(1):91–110, 1999.
- [CJM<sup>+</sup>00] Mary E. Cosner, Robert K. Jansen, Bernard M.E. Moret, Linda A. Raubeson, Li-San Wang, Tandy Warnow, and Stacia Wyman. A new fast heuristic for computing the breakpoint phylogeny and experimental phylogenetic analyses of real and synthetic data. In *Proc. 8th Int'l Conf. on Intelligent Systems for Molecular Biology ISMB-2000, San Diego*, pages 104–115, 2000.
- [Ead88] Peter Eades. Symmetry finding algorithms. In *Computational Morphology: A Computational Geometric Approach to the Analysis of Form*, Godfried T. Toussaint (ed.), North-Holland. 1988.
- [EEK<sup>+</sup>01] Henrik Eriksson, Kimmo Eriksson, Johan Karlander, Lars Svensson, and Johan Wästlund. Sorting a bridge hand. *Discrete Math.*, 241(1–3):289–300, October 2001.
- [GY79] Lidija I. Golovina and Isaak M. Yaglom. *Induction in Geometry*. Mir Publishers, Moscow, 1979.

- [GZ01] Branko Grunbaum and Joseph Zaks. Convexification of polygons by flips and by flipturns. *Discrete Mathematics*, 241:333–342, 2001.
- [HP96] Sridhar Hannenhalli and Pavel A. Pevzner. To cut or not to cut (applications of comparative physical maps in molecular evolution). In *Seventh Annual ACM-SIAM Symposium on Discrete Algorithms*, pages 304–313, 1996.
- [HP98] Sridhar Hannenhalli and Pavel A. Pevzner. Transforming cabbage into turnip (polynomial algorithm for sorting signed permutations by reversals). In *Proc. 27th Annual ACM Symposium on the Theory of Computing*, pages 178–189, 1998.
- [KMP77] Donald E. Knuth, James H. Morris, and Vaughan R. Pratt. Fast pattern matching in strings. *SIAM Journal of Computing*, 6:322–350, 1977.
- [Sco82] Paul R. Scott. On a family of polygons. *Mathematics Magazine*, 55(2):104–105, March 1982.