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► **To cite this version:**

Antoine Thomas, Jean-Stéphane Varré, Aïda Ouangraoua. Genome dedoubling by DCJ and reversal. BMC Bioinformatics, BioMed Central, 2011, 12 (Suppl 9), pp.S20. <10.1186/1471-2105-12-S9-S20>. <hal-00784416>

HAL Id: hal-00784416

<https://hal.inria.fr/hal-00784416>

Submitted on 4 Feb 2013

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PROCEEDINGS

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Genome dedoubling by DCJ and reversal

Antoine Thomas, Jean-Stéphane Varré*, Aïda Ouangraoua*

From Ninth Annual Research in Computational Molecular Biology (RECOMB) Satellite Workshop on Comparative Genomics
Galway, Ireland. 8-10 October 2011

Abstract

Background: Segmental duplications in genomes have been studied for many years. Recently, several studies have highlighted a biological phenomenon called *breakpoint-duplication* that apparently associates a significant proportion of segmental duplications in Mammals, and the *Drosophila* species group, to breakpoints in rearrangement events.

Results: In this paper, we introduce and study a combinatorial problem, inspired from the breakpoint-duplication phenomenon, called the *Genome Dedoubling Problem*. It consists of finding a minimum length rearrangement scenario required to transform a genome with duplicated segments into a non-duplicated genome such that duplications are caused by rearrangement breakpoints. We show that the problem, in the Double-Cut-and-Join (DCJ) and the reversal rearrangement models, can be reduced to an APX-complete problem, and we provide algorithms for the Genome Dedoubling Problem with 2-approximable parts. We apply the methods for the reconstruction of a non-duplicated ancestor of *Drosophila yakuba*.

Conclusions: We present the *Genome Dedoubling Problem*, and describe two algorithms solving the problem in the DCJ model, and the reversal model. The usefulness of the problems and the methods are showed through an application to real *Drosophila* data.

Introduction

Gene duplication is an important source of variations in genomes. Recently, several studies have highlighted biological evidence for abundant segmental duplications that occur around breakpoints of rearrangement events in the evolution of eukaryotes.

In mammals, an evidence for a strong association between duplications, genomic instability and large-scale chromosomal rearrangements in primate evolution was first reported in [1]. Later in [2], a study of all evolutionary rearrangement breakpoints between human and mouse genomes reported that 53% of the breakpoints were associated with segmental duplications, as compared to 18% expected in a random assignment of breaks. In [3], a first study of the human-mouse rearrangement breakpoints, considering only synteny blocks of

length larger than 100Kb and duplicated sequences of length larger than 10Kb, showed that 25% (122/461) of the breakpoints contained duplicated sequences of length greater than 10kb.

The association between segmental duplications and regions of breaks of synteny was also reported in the *Drosophila* species group. In [4], an analysis of the breakpoints of *Drosophila yakuba* compared to two related species, *Drosophila simulans* and *Drosophila melanogaster*, revealed that the breakpoint regions of 59% of the reversals (17/29) were associated with inverted duplications of genes or other nonrepetitive sequences. Further evidences of the recurrent presence of repetitive sequences near breakpoints of rearrangement in the evolution of *Drosophila* were also reported in [5-7].

A rearrangement event is an operation that modifies the organization of a given genome by cutting the genome at some points called *breakpoints* to glue the exposed extremities in a different way. The biological

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phenomenon called *breakpoint-duplication* results in the presence of the same genomic segment on both extremities of a breakpoint in a rearrangement. Several biological models have been presented to explain the presence of duplicated sequences at rearrangement breakpoint regions. These models are based on DNA breaks repairs that produce duplicated segments because of staggered Single-Strand-Breaks [3,4], or non-reciprocal genetic exchange in Double-Strand-Breaks [5]. Most of these biological models support a nonrandom model of chromosomal evolution that implicates a predominance of recurrent small-scale duplications and large-scale evolutionary rearrangements within specific *fragile* regions. Moreover, the genetic instability of such regions is often suggested to be the cause rather than the consequence of duplicated genomic architecture [3,8]. Interestingly, a growing number of the breakpoint-duplications detected in Supra-primates evolution have also been linked to recurrent chromosomal rearrangements associated with common diseases in the human population [1-3,8,9]. In [10], breakpoint-duplications were also identified in human sex chromosomes, allowing to order rearrangement events in time, based on the degree of divergence of the breakpoint-duplicated sequences.

In this paper, we are interested in using the segmental duplications of a given present-day genome that has undergone breakpoint-duplication rearrangements, in order to reconstruct a non-duplicated ancestral genome. We formally define the breakpoint-duplication phenomenon, and introduce a combinatorial problem called the *Genome Dedoubling problem*. Given a genome that has undergone breakpoint-duplication rearrangements, possibly with other rearrangements events, the problem asks to find an ancestral genome such that the number of rearrangement events needed to transform the ancestor into the given genome is minimal. Note that the Genome Dedoubling problem asks to find a non-duplicated ancestor of a given duplicated genomes, as the *Genome Halving problem* introduced in [11] that consists of, given a genome that has undergone a whole-genome duplication followed by rearrangement events, finding the ancestral genome that was present right before the whole-genome duplication event. However, the two problems and their solutions are different as they aim at recovering different types of duplication events, breakpoint-duplications and whole-genome duplications. As the Genome Halving problem is motivated by the whole-genome duplication events in molecular evolution, the Genome Dedoubling problem is motivated by breakpoint-duplication events in molecular evolution. Both problems are useful for the comparison of genomes with duplicated segments.

In the following, we study the Genome Dedoubling problem under the Double-Cut-and-Join (DCJ) and the

reversal rearrangement models. In Section **Methods**, we formally present breakpoint-duplication (BD) rearrangements and the Genome Dedoubling Problem. We show that the problem can always be regarded as a Dedoubling Problem on totally duplicated genomes. In Section **Genome dedoubling by DCJ**, we study the problem under the DCJ model, on multichromosomal then unichromosomal genomes. We prove the NP-completeness of the problems by reduction to an APX-complete problem, and provide algorithms with a linear time complexity, except for an APX-complete part that is 2-approximable. In Section **Genome dedoubling by reversal**, we study the problem under the reversal model on oriented genomes, making use of some results of the Hannenhalli-Pevzner (HP) theory [12] on sorting by reversal described in [13,14]. We provide an algorithm with a quadratic time complexity, except for an APX-complete part that is 2-approximable. In Section **Application**, an application for the reconstruction of a non-duplicated ancestor of *Drosophila yakuba*, using data from [4], is presented.

Methods

In this section we give the main definitions and notations of duplicated genomes and rearrangements. Next, we generalize the definitions of rearrangements in order to introduce a formal definition of *breakpoint-duplication rearrangements*, and the *Genome Dedoubling Problem* studied in the paper.

Duplicated genomes

A genome consists of linear or circular chromosomes that are composed of genomic markers. Markers are represented by signed integers such that the sign indicates the orientations of markers in chromosomes. By convention, $-x = x$. A linear chromosome is represented by an ordered sequence of signed integers surrounded by the unsigned marker \circ at each end indicating the telomeres. A circular chromosome is represented by a circularly ordered sequence of signed integers. For example, $(1\ 2\ -3)\ (\circ\ 4\ -5\ \circ)$ is a genome composed of one circular and one linear chromosome.

Each genome contains at most two occurrences of each marker. Two copies of a same marker in a genome are called paralogs. If a marker x is present twice, one of the paralogs is represented by \bar{x} . By convention, $\bar{\bar{x}} = x$. Here, such markers represent segments duplicated by a breakpoint-duplication rearrangement.

Definition 1 A duplicated genome is a genome in which a subset of the markers are duplicated.

For example, $(1\ 2\ -3\ -\bar{2})\ (\circ\ 4\ -5\ \bar{1}\ \bar{5}\ \circ)$ is a duplicated genome where markers 1, 2, and 5 are duplicated. A *non-duplicated genome* is a genome in which no marker is duplicated. A *totally duplicated*

genome is a duplicated genome in which all markers are duplicated. For example, $(1 \ 2 \ \bar{2}) (\circ \ -3 \ \bar{1} \ \bar{3} \ \circ)$ is a totally duplicated genome.

An adjacency in a genome is a pair of consecutive markers. Since a genome can be read in two directions, the adjacencies $(x \ y)$ and $(-y \ -x)$ are equivalent. For example, the genome $(1 \ 2 \ \bar{2}) (\circ \ -3 \ \bar{1} \ \bar{3} \ \circ)$ has seven adjacencies, $(1 \ 2), (2 \ \bar{2}), (\bar{2} \ 1), (\circ \ -3), (-3 \ \bar{1}), (\bar{1} \ \bar{3}),$ and $(\bar{3} \ \circ)$.

Definition 2 A dedoubled genome is a duplicated genome G such that for any duplicated marker x in G , either $(x \ \bar{x})$, or $(\bar{x} \ x)$ is an adjacency of G .

For example, $(1 \ \bar{1} \ -2 \ \bar{2}) (\circ \ -3 \ 4 \ -5 \ \bar{5} \ \circ)$ is a dedoubled genome. The reduction of a dedoubled genome G , denoted by G^R , is the genome obtained from G by replacing every pair $(x \ \bar{x})$, or $(\bar{x} \ x)$ by a single marker x . For example the reduction of $G = (1 \ \bar{1} \ -2 \ \bar{2}) (\circ \ -3 \ 4 \ -5 \ \bar{5} \ \circ)$ is $G^R = (1 \ -2) (\circ \ -3 \ 4 \ -5 \ \circ)$.

Rearrangement

A rearrangement operation on a given genome cuts a set of adjacencies of the genome called *breakpoints* and forms new adjacencies with the exposed extremities, while altering no other adjacency. In this paper, we consider two types of rearrangement operation called *double-cut-and-join (DCJ)* and *reversal*. In the sequel, the breakpoints of a rearrangement operation are indicated in the genome by the symbol \blacktriangle , and the new adjacencies are indicated in the genome by dots.

A DCJ operation on a genome G cuts two different adjacencies in G and glues pairs of the four exposed extremities to form two new adjacencies. For example, the following DCJ cuts adjacencies $(1 \ 2)$ and $(-5 \ \bar{1})$ to produce $(1 \ \bar{1})$ and $(-5 \ 2)$.

$$(1 \ \blacktriangle \ 2 \ -3 \ \bar{2}) (\circ \ 4 \ -5 \ \blacktriangle \ \bar{1} \ -3 \ \bar{5} \ \circ) \rightarrow (\circ \ 4 \ -5 \ \cdot \ 2 \ -3 \ \bar{2} \ 1 \ \cdot \ \bar{1} \ -3 \ \bar{5} \ \circ)$$

A reversal on a genome G is a DCJ operation that cuts two adjacencies $(a \ b)$ and $(c \ d)$ in a chromosome of G of the form $(\dots a \ b \ \dots c \ d \ \dots)$ to form two new adjacencies $(a \ -c)$ and $(-b \ d)$, thus reversing the orientation of the segment of G beginning with marker b and ending with marker c . For example, the following reversal cuts adjacencies $(-5 \ \bar{1})$ and $(\bar{5} \ \circ)$ and reverses the segment $\bar{1} \ -3 \ \bar{5}$.

$$(1 \ 2 \ -3 \ \bar{2}) (\circ \ 4 \ -5 \ \blacktriangle \ \bar{1} \ -3 \ \bar{5} \ \blacktriangle \ \circ) \rightarrow (1 \ 2 \ -3 \ \bar{2}) (\circ \ 4 \ -5 \ \cdot \ \bar{5} \ \bar{3} \ -\bar{1} \ \cdot \ \circ)$$

A DCJ (resp. reversal) scenario between two genomes A and B is a sequence of DCJ (resp. reversal) operations allowing to transform A into B . The length of a scenario is the number of rearrangement operations composing the scenario.

The DCJ (resp. reversal) distance between two genomes A and B is the minimum length of a DCJ (resp. reversal) scenario between A and B .

Breakpoint-duplication rearrangements

We now generalize the definitions of rearrangement operations to account for possible duplications at their breakpoints.

A 1-breakpoint-duplication DCJ (1-BD-DCJ) operation on a genome G is a rearrangement operation that alters two different adjacencies $(a \ b)$ and $(c \ d)$ of G , by:

- first adding marker \bar{a} at the appropriate position to produce segment $(a \ \bar{a} \ b)$,
- then applying a DCJ operation that cuts adjacencies $(a \ \bar{a})$ and $(c \ d)$ to produce either $(a \ d)$ and $(c \ \bar{a})$, or $(a \ -c)$ and $(-\bar{a} \ d)$.

A 2-breakpoint-duplication DCJ (2-BD-DCJ) operation on a genome G is a rearrangement operation that alters two different adjacencies $(a \ b)$ and $(c \ d)$ of G , by:

- first adding markers \bar{a} and \bar{c} at the appropriate positions to produce segments $(a \ \bar{a} \ b)$ and $(c \ \bar{c} \ d)$,
- then applying a DCJ operation that cuts adjacencies $(a \ \bar{a})$ and $(c \ \bar{c})$ to produce either $(a \ \bar{c})$ and $(c \ \bar{a})$, or $(a \ -c)$ and $(-\bar{a} \ \bar{c})$.

Definition 3 A breakpoint-duplication DCJ (BD-DCJ) operation on a genome G is either a 1-BD-DCJ operation, or a 2-BD-DCJ operation.

In the sequel, if some markers are duplicated by a BD-DCJ operation, they are indicated in bold font in the initial genome. For example, the following rearrangement is a 2-BD-DCJ operation that acts on adjacencies $(-2 \ -1)$ and $(4 \ -3)$, and duplicates markers 2 and 4. The intermediate step resulting in the duplication of markers 2 and 4 is shown above the arrow.

$$(1 \ \blacktriangle \ 2) (\circ \ 3 \ \blacktriangle \ -4 \ \circ) \xrightarrow{(1 \ \bar{2} \ \blacktriangle \ 2) (\circ \ 3 \ \bar{4} \ \blacktriangle \ -4 \ \circ)} (\circ \ 3 \ \cdot \ \bar{4} \ \cdot \ 2 \ 1 \ \bar{2} \ \cdot \ -4 \ \circ)$$

To summarize, a BD-DCJ operation consists of a *first step* in which one or two markers are duplicated, followed by a *second step* where a DCJ operation is applied. Similarly, we now define a *breakpoint-duplication reversal (BD-reversal)* operation.

Definition 4 A breakpoint-duplication reversal (BD-reversal) operation on a genome G is a BD-DCJ operation such that the DCJ operation applied in the second step of the BD-DCJ operation is a reversal.

For example, the following rearrangement is a BD-reversal that is a 1-BD-DCJ operation that acts on adjacencies $(2 \ -1)$ and $(-3 \ 4)$, and duplicates marker 2.

$$(\circ \ 1 \ \blacktriangle \ -2 \ -3 \ \blacktriangle \ 4 \ \circ) \xrightarrow{(\circ \ 1 \ \bar{2} \ \blacktriangle \ -2 \ -3 \ \blacktriangle \ 4 \ \circ)} (\circ \ 1 \ \cdot \ \bar{2} \ \cdot \ 3 \ 2 \ \cdot \ 4 \ \circ)$$

A BD-DCJ scenario (resp. BD-reversal scenario) between a non-duplicated genome A and a duplicated

genome B is a sequence composed of BD-DCJ (resp. BD-reversal) operations and possibly DCJ (resp. reversal) operations allowing to transform A into B .

Definition 5 Given a non-duplicated genome A and a duplicated genome B , the BD-DCJ distance (resp. BD-reversal distance) between A and B is the minimal length of a BD-DCJ (resp. BD-reversal) scenario between A and B .

We now give an obvious, but useful property allowing to reduce a BD-DCJ scenario to a DCJ scenario.

Proposition 1 Given a non-duplicated genome A and a duplicated genome B , for any a BD-DCJ (resp. BD-reversal) scenario between A and B , there exists a DCJ (resp. reversal) scenario of same length between a dedoubled genome D and B such that the reduction of D is A ($D^R = A$).

Proof. Let S be a BD-DCJ (resp. BD-reversal) scenario between A and B . D is the genome obtained from A , by adding, for any marker x duplicated by a BD-DCJ operation in S , the marker \bar{x} in a way to produce either adjacency $(\bar{x} \ x)$, or $(x \ \bar{x})$ as done in S . Thus, $D^R = A$. The DCJ (resp. reversal) scenario between D^R and B having the same length as S , is the sequence of DCJ (resp. reversal) contained in S or in BD-DCJ (resp. BD-reversal) operations of S , with the same order as in S . ■

For example, in the following, a BD-reversal scenario of length 4 between $A = (\circ \ 1 \ 2 \ 3 \ 4 \ 5 \ \circ)$ and $B = (\circ \ 1 \ \bar{4} \ 2 \ \bar{3} \ \bar{5} \ \bar{2} \ \bar{1} \ 4 \ -3 \ 5 \ \circ)$ induces a reversal scenario of length 4 between $D = (\circ \ 1 \ \bar{1} \ \bar{2} \ 2 \ \bar{3} \ 3 \ \bar{4} \ 4 \ \bar{5} \ 5 \ \circ)$ and B .

BD-reversal scenario	Reversal scenario
$A = (\circ \ 1 \ \blacktriangle \ 2 \ 3 \ \blacktriangle \ 4 \ 5 \ \circ)$	$D = (\circ \ 1 \ \blacktriangle \ \bar{1} \ \bar{2} \ 2 \ \bar{3} \ 3 \ \bar{4} \ \blacktriangle \ 4 \ 5 \ \bar{5} \ \circ)$
$(\circ \ 1 \ \bar{4} \ \blacktriangle \ -3 \ \bar{2} \ \blacktriangle \ \bar{1} \ 4 \ 5 \ \circ)$	$(\circ \ 1 \ \bar{4} \ \blacktriangle \ -3 \ \bar{3} \ \bar{2} \ \blacktriangle \ \bar{2} \ \bar{1} \ 4 \ 5 \ \bar{5} \ \circ)$
$(\circ \ 1 \ \bar{4} \ 2 \ 3 \ \blacktriangle \ \bar{2} \ \bar{1} \ 4 \ \blacktriangle \ 5 \ \circ)$	$(\circ \ 1 \ \bar{4} \ 2 \ \bar{3} \ 3 \ \blacktriangle \ \bar{2} \ \bar{1} \ 4 \ \blacktriangle \ 5 \ \bar{5} \ \circ)$
$(\circ \ 1 \ \bar{4} \ 2 \ \blacktriangle \ 3 \ \bar{4} \ \bar{1} \ \bar{2} \ 5 \ \blacktriangle \ \circ)$	$(\circ \ 1 \ \bar{4} \ 2 \ \bar{3} \ \blacktriangle \ 3 \ \bar{4} \ \bar{1} \ \bar{2} \ 5 \ \blacktriangle \ \bar{5} \ \circ)$
$(\circ \ 1 \ \bar{4} \ 2 \ \bar{3} \ \bar{5} \ \bar{2} \ \bar{1} \ 4 \ -3 \ \bar{5} \ \circ)$	$(\circ \ 1 \ \bar{4} \ 2 \ \bar{3} \ \bar{5} \ \bar{2} \ \bar{1} \ 4 \ -3 \ \bar{5} \ \circ)$

Genome dedoubling problem

We now state the genome dedoubling problems considered in this paper.

Genome dedoubling problem: Given a duplicated genome G , the DCJ (resp. reversal) genome dedoubling problem consists of finding a non-duplicated genome H such that the BD-DCJ (resp. BD-reversal) distance between H and G is minimal.

Given a duplicated genome G , we denote by $d_{dcj}(G)$ (resp. $d_{rev}(G)$), the minimum BD-DCJ (resp. BD-reversal) distance between any non-duplicated genome and G . From Proposition 1, the following proposition is straightforward.

Proposition 2 Given a duplicated genome G , the DCJ (resp. reversal) genome dedoubling problem on G is equivalent to finding a dedoubled genome D such that

the DCJ (resp. reversal) distance between D and G is minimal.

The next proposition describes a further reduction of the genome dedoubling problem on a duplicated genome G .

Proposition 3 Given a duplicated genome G , the DCJ (resp. reversal) genome dedoubling problem on G is equivalent to the DCJ (resp. reversal) genome dedoubling problem on the totally duplicated genome G^T obtained from G by replacing every maximal subsequence of non-duplicated markers beginning with a marker x by the pair $x \ \bar{x}$.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

For example, solving the DCJ (resp. reversal) genome dedoubling problem on $G = (\circ \ 1 \ 4 \ \bar{7} \ \bar{1} \ \bar{5} \ 10 \ \bar{8} \ \bar{4} \ 2 \ 6 \ \bar{9} \ 3 \ \bar{8} \ \circ)$ is equivalent to solving it on $G^T = (\circ \ 1 \ 4 \ \bar{7} \ \bar{7} \ \bar{1} \ \bar{5} \ \bar{5} \ \bar{8} \ \bar{4} \ 2 \ \bar{2} \ \bar{8} \ \circ)$. The transformations applied on G to obtain G^T are indicated in bold font.

In the sequel, G will always denote a totally duplicated genome, and we focus in Sections **Genome dedoubling by DCJ** and **Genome dedoubling by reversal** on the problem of finding a dedoubled genome D such that the DCJ (resp. reversal) distance between D and G is minimal.

Results

In this section, we first study the Genome Dedoubling Problem under the DCJ model. Next, we study the problem under the reversal model on oriented genomes described in the Hannenhalli-Pevzner (HP) theory on sorting by reversal [12-14].

Genome dedoubling by DCJ

In this section, G denotes a totally duplicated genome. In order to give a formula for the DCJ dedoubling distance of G , $d_{dcj}(G)$, we use a graph called the *dedoubled adjacency graph* of G .

Dedoubled adjacency graph

Definition 6 The dedoubled adjacency graph of G , denoted by $\mathcal{A}(G)$, is the graph whose vertices are the adjacencies of G , and for any marker x there is one edge between the vertices $(x \ u)$ and $(v \ \bar{x})$, and one edge between the vertices $(y \ x)$ and $(\bar{x} \ z)$.

An example of dedoubled adjacency graph is depicted in Fig. 1. In the following, we will simply refer to dedoubled adjacency graphs as adjacency graphs.

Note that all vertices in $\mathcal{A}(G)$ have degree one or two. Thus, the connected components of $\mathcal{A}(G)$ are only paths and cycles. These paths and cycles are called *elements* of $\mathcal{A}(G)$.

Given a couple of paralogous markers (x, \bar{x}) , an element of the graph $\mathcal{A}(G)$ is said to *contain* the couple

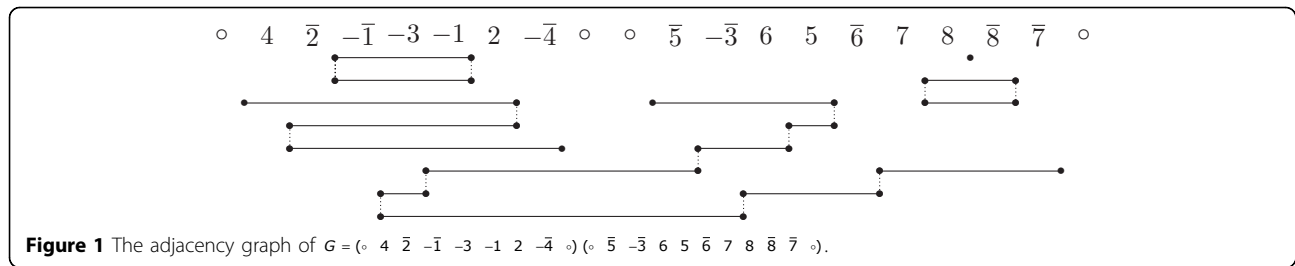


Figure 1 The adjacency graph of $G = (\circ 4 \bar{2} \bar{-1} -3 -1 2 \bar{-4} \circ \circ 5 \bar{-3} 6 5 \bar{6} 7 8 \bar{8} \bar{7} \circ)$.

$(v \bar{x})$ if it contains the edge linking vertices $(x u)$ and $(v \bar{x})$, or the edge linking vertices $(y x)$ and $(\bar{x} z)$. By definition, a couple (x, \bar{x}) can possibly be contained in only one element A of $\mathcal{A}(G)$ if element A contains both edges $((x u), (v \bar{x}))$ and $((y x), (\bar{x} z))$. In this case, A is said to *contain twice* the couple (x, \bar{x}) , and A is called a *duplicated* element of $\mathcal{A}(G)$. If an element A contains no couple (x, \bar{x}) twice, then it is called a *non-duplicated* element of $\mathcal{A}(G)$. If the two edges $((x u), (v \bar{x}))$ and $((y x), (\bar{x} z))$ belong to two different elements A and B of $\mathcal{A}(G)$, then A and B will both contain (x, \bar{x}) . In this case, we say that A and B *intersect*. If two elements A and B do not intersect, then we say that A and B are *independent*. For example in Fig. 1 the two paths of the adjacency graph are duplicated, while the three cycles are non-duplicated. The leftmost path and the leftmost cycle intersect because they both contain the couple $(2, \bar{2})$, while the two paths are independent. Given an element A of $\mathcal{A}(G)$, the *set induced by A* is the set of couples (x, \bar{x}) contained in A .

General sorting

In this section, we prove the following theorem:

Theorem 1 *Let n be the number of couples of paralogous markers in G . Let C_i be the maximum size of a subset of non-duplicated pairwise independent cycles in $\mathcal{A}(G)$. The DCJ dedoubling distance of G is $d_{dcj}(G) = n - C_i$.*

For example, in Fig. 1, the maximum size of a subset of non-duplicated pairwise independent cycles is 2 as there are three cycles, and the two rightmost cycles intersect. The distance would then be $d_{dcj}(G) = 8 - 2 = 6$. To prove Theorem 1, we use the following properties:

Property 1 *Let n be the number of couples of paralogous markers in G .*

1. *The maximum size C_i of a set of non-duplicated pairwise independent cycles in the graph $\mathcal{A}(G)$ is n .*

2. *If G is dedoubled genome, then $\mathcal{A}(G)$ contains n non-duplicated pairwise independent cycles, each containing a single couple of paralogous markers, plus possibly other cycles. In this case, $C_i = n$.*

3. *A DCJ operation can only alter the maximum size C_i of a set of non-duplicated pairwise independent cycles by $-1, 0$ or $+1$.*

Proof. See proof in Additional file 1 (Supplemental proofs). ■

Algorithm 1 is an algorithm that provides a $n - C_i$ length DCJ scenario transforming G into a dedoubled genome.

Algorithm 1 Finding a shortest DCJ scenario transforming a totally duplicated genome G into a dedoubled genome

```

1: Construct  $\mathcal{A}(G)$ .
2: Choose a maximum size set  $S_i$  of non-duplicated pairwise independent cycles.
3: For any couple  $(x, \bar{x})$  of paralogous markers do
4:   If  $(x, \bar{x})$  is contained in a cycle  $c$  of  $S_i$  containing more than one couple then
5:     Perform the DCJ that creates adjacency  $(x \bar{x})$  or  $(\bar{x} x)$  by splitting  $c$ 
       into two cycles, one of the cycles containing only the couple  $(x, \bar{x})$ .
6:     Replace  $c$  in  $S_i$  by the two new cycles.
7:   Else
8:     Perform any DCJ that creates adjacency  $(x \bar{x})$  or  $(\bar{x} x)$ .
9:   End if
10: End for
    
```

We now have all the pre-requisites to give the proof of Theorem 1. The proof can be found in Additional file 1 (Supplemental proofs).

Lemma 1 *Choosing a maximum size set of non-duplicated pairwise independent cycles in $\mathcal{A}(G)$ is an APX-complete problem, approximable with an approximation ratio of 2.*

Proof. See proof in Additional file 1 (Supplemental proofs). ■

From Lemma 1, the complexity of the Genome Dedoubling problem by DCJ follows immediately.

Corollary 1 *The Genome Dedoubling problem by DCJ is NP-complete. Algorithm 1 solves the problem in linear time complexity, except for the computation of the set of cycles S_i that is 2-approximable.*

Sorting between linear unichromosomal genomes

In this section, we search for a minimum length DCJ scenario that transforms a duplicated genome consisting of a single linear chromosome into a dedoubled genome consisting of a single linear chromosome. The results of this section will then be used in the next section for the study of the Genome Dedoubling problem under the reversal model.

In this section and the sequel, G denotes a totally duplicated genome consisting of a single linear chromosome. In this case, the graph $\mathcal{A}(G)$ contains exactly one path, and possibly several cycles.

Definition 7 *The path in $\mathcal{A}(G)$ is said to be valid if it contains every couple (x, \bar{x}) of paralogous markers in G .*

A DCJ operation that merges a cycle c of $\mathcal{A}(G)$ in the path p is a DCJ operation that acts on an adjacency of c and an adjacency of p , thus gathering c and p into a longer path.

Note that if G is a dedoubled genome, then the path in $\mathcal{A}(G)$ is necessarily valid. We call such a genome a *dedoubled linear genome*. So, if the path in $\mathcal{A}(G)$ is not valid, then any DCJ scenario transforming G into a dedoubled linear genome will merge, in the path, cycles containing the couples (x, \bar{x}) that are not contained in the path.

In the following, we always denote by m the minimum number of cycles required to make the path of $\mathcal{A}(G)$ valid. We also denote by C_i the maximum size of a subset of non-duplicated pairwise independent cycles. First, we have the following property:

Property 2 Let C be the number of cycles in $\mathcal{A}(G)$. We have $C_i = C - m$.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

From Property 2, we then have the following lemma.

Lemma 2 Let n be the number of couples of paralogous markers in G . Let C be the number of cycles in $\mathcal{A}(G)$. The minimum length d of a DCJ scenario transforming G into a dedoubled genome consisting of a single linear chromosome equals $d = n - C + 2m$.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

From Property 2 and Lemma 2, we immediately have the following complexity.

Corollary 2 The problem of finding a DCJ scenario transforming G into a dedoubled genome consisting of a single linear chromosome is NP-complete. Algorithm 1, in which we add the line (2': Merge in the path all the cycles that are not in S_i) between lines 2 and 3, solves the problem in linear time complexity, except for the computation of the set of cycles S_i that is 2-approximable.

Genome dedoubling by reversal

We build and use a graph that behaves like the *arc overlap graph* used in [13] for the Hannenhalli-Pevzner theory of sorting by reversal [12]. The genome G consists of a single linear chromosome.

Dedoubled overlap graph

For any couple (x, \bar{x}) of paralogous markers in G , the segment of (x, \bar{x}) is the smallest segment of G containing both markers x and \bar{x} . For example, in genome $G = (\circ 1 3 \bar{1} -\bar{2} -4 -\bar{3} 2 -\bar{4} \circ)$, the segment of $(1, \bar{1})$ is $(1 3 \bar{1})$, and the segment of $(2, \bar{2})$ is $(-\bar{2} -4 -\bar{3} 2)$.

Definition 8 The dedoubled overlap graph of G , denoted by $\mathcal{O}(G)$, is the graph whose vertices are all the couples (x, \bar{x}) of paralogous markers of G , and there is an

edge between two vertices (u, \bar{u}) and (v, \bar{v}) if the segments of u and v overlap.

An example of dedoubled overlap graph is depicted in Fig. 4.a. In the following, we will simply refer to dedoubled overlap graphs as overlap graphs.

The vertex (x, \bar{x}) of the graph $\mathcal{O}(G)$ is *oriented* if x and \bar{x} have different signs in G , otherwise it is *unoriented*. If the vertex (x, \bar{x}) is oriented then there exists a reversal operation denoted by $\text{Rev}(x \bar{x})$ that produces the adjacency $(x \bar{x})$ and a reversal operation denoted by $\text{Rev}(\bar{x} x)$ that produces the adjacency $(\bar{x} x)$. For example, in genome $G = (\circ 1 3 \bar{1} -\bar{2} -4 -\bar{3} 2 -\bar{4} \circ)$, $(3, \bar{3})$ is an oriented vertex of $\mathcal{O}(G)$.

$$\begin{aligned} \text{Rev}(3 \bar{3}) &= (\circ 1 3 \bar{1} -\bar{2} -4 -\bar{3} 2 -\bar{4} \circ) \rightarrow (\circ 1 3 \bar{3} 4 \bar{2} -\bar{1} 2 -\bar{4} \circ) \\ \text{Rev}(\bar{3} 3) &= (\circ 1 3 \bar{1} -\bar{2} -4 -\bar{3} 2 -\bar{4} \circ) \rightarrow (\circ 1 4 \bar{2} -\bar{1} -3 -\bar{3} 2 -\bar{4} \circ) \end{aligned}$$

The overlap graph of G behaves like arc overlap graphs used in [13] for the Hannenhalli-Pevzner theory of sorting by reversal [12]. Indeed, given an oriented vertex (x, \bar{x}) of the graph $\mathcal{O}(G)$, performing the reversal $\text{Rev}(x \bar{x})$ or $\text{Rev}(\bar{x} x)$ complements the subgraph induced by (x, \bar{x}) and all its neighbouring vertices, and changes the orientation of all vertices in this subgraph (see Fig. 4.b).

A connected component of the graph $\mathcal{O}(G)$ is *oriented* if it contains at least one oriented vertex, otherwise it is *unoriented*. A genome is *oriented* if all connected components of the graph $\mathcal{O}(G)$ are oriented, otherwise it is *unoriented*.

Given an oriented vertex (x, \bar{x}) of the graph $\mathcal{O}(G)$, the score of (x, \bar{x}) is the number of oriented vertices in the overlap graph of the genome obtained after applying $\text{Rev}(x \bar{x})$ on G . Note that the same number of oriented vertices is obtained after applying $\text{Rev}(\bar{x} x)$ on G .

Property 3 Let (x, \bar{x}) be an oriented vertex of $\mathcal{O}(G)$ of maximum score. Performing $\text{Rev}(x \bar{x})$ or $\text{Rev}(\bar{x} x)$ does not create new unoriented connected components in the overlap graph of the genome obtained.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

In the sequel, we focus on sorting oriented genomes using reversal dedoubling scenarios. A totally duplicated genome G consisting of a single linear chromosome is called a *valid-path genome* if the single path in $\mathcal{A}(G)$ is valid. Otherwise, it is called a *non-valid-path genome*.

Sorting an oriented valid-path genome

In this section, we consider an oriented valid-path genome G . With n being the number of couples of paralogous markers in G , we have the following theorem:

Theorem 2 Let G be an oriented valid-path genome. Let C be the number of cycles in $\mathcal{A}(G)$. The reversal dedoubling distance of G is $d_{\text{rev}}(G) = n - C$.

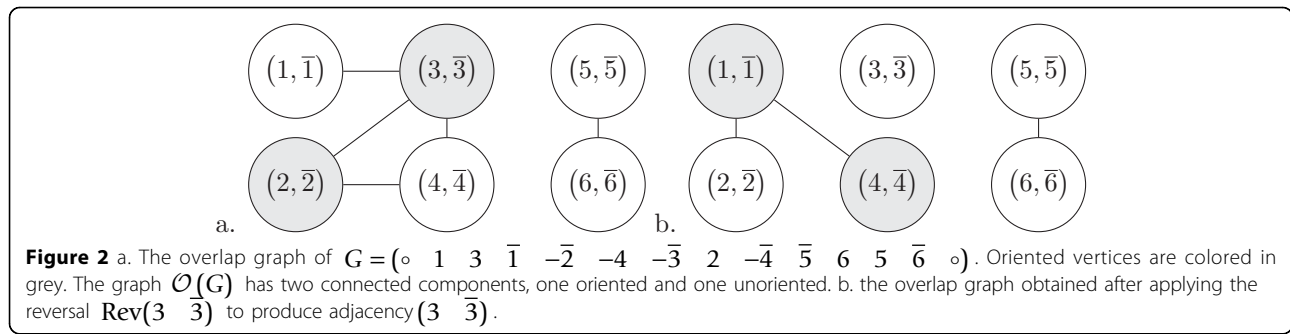


Figure 2 a. The overlap graph of $G = (o \ 1 \ 3 \ \bar{1} \ -\bar{2} \ -\bar{4} \ -\bar{3} \ 2 \ -\bar{4} \ \bar{5} \ 6 \ 5 \ \bar{6} \ o)$. Oriented vertices are colored in grey. The graph $\mathcal{O}(G)$ has two connected components, one oriented and one unoriented. b. the overlap graph obtained after applying the reversal $\text{Rev}(3 \ \bar{3})$ to produce adjacency $(3 \ \bar{3})$.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

Algorithm 2 Finding a shortest reversal scenario transforming an oriented genome G into a dedoubled genome.

- 1: Construct $\mathcal{A}(G)$.
- 2: Construct $\mathcal{O}(G)$.
- 3: Choose a maximum size set S_c of non-duplicated pairwise independent cycles.
- 4: Merge in the path all the cycles that are not in S_c .
- 5: while G is not a dedoubled genome do
- 6: Pick a maximum score vertex (x, \bar{x}) in $\mathcal{O}(G)$.
- 7: if (x, \bar{x}) is contained in a cycle c of S_c then
- 8: Choose between $\text{Rev}(x \ \bar{x})$ and $\text{Rev}(\bar{x} \ x)$ the reversal that splits c into two cycles, and perform it.
- 9: Replace c in S_c by the two new cycles.
- 10: Apply the modification induced by the reversal on $\mathcal{O}(G)$.
- 11: else
- 12: Perform any reversal that creates adjacency $(x \ \bar{x})$ or $(\bar{x} \ x)$.
- 13: end if
- 14: end while

Sorting an oriented non-valid-path genome

In this section, G denotes an oriented non-valid path genome. At least m cycles of $\mathcal{A}(G)$ have to be merged in the path to make it valid.

An edge $((x \ u), (v \ \bar{x}))$ or $((y \ x), (\bar{x} \ z))$ of the adjacency graph $\mathcal{A}(G)$ is called oriented if markers x and \bar{x} have different signs. Note that extracting a cycle from any element of the graph $\mathcal{A}(G)$ requires this element to contain oriented edges. It is easy to see that given two adjacencies picked in a given element, a reversal acting on these adjacencies will extract a cycle if and only if the path linking these adjacencies contains an odd number of oriented edges. Thus, we have the following lemma:

Lemma 3 Let G be an oriented non-valid-path genome. Merging a cycle of $\mathcal{A}(G)$ in its path never creates unoriented connected components in the overlap graph of the genome obtained.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

Theorem 3 Let G be an oriented non-valid-path genome. Let C be the number of cycles in the graph $\mathcal{A}(G)$ and m be the minimum number of cycles to merge in the path to make it valid. The reversal dedoubling distance of G is $d_{\text{rev}}(G) = n - C + 2m$.

Proof. See proof in Additional file 1 (Supplemental proofs). ■

Prom Lemma 1 and Property 2, the complexity of the Genome Dedoubling problem by reversal on oriented genomes follows immediately.

Corollary 3 The Genome Dedoubling problem by reversal on oriented genomes is NP-complete. Algorithm 2 solves the problem in quadratic time complexity, except for the computation of S_c that is 2-approximable.

Application

We applied Algorithm 2 to reconstruct an ancestral chromosome for the chromosome 2 of *Drosophila yakuba* using a dataset obtained from [4] with *Drosophila melanogaster* used as the outgroup. The results obtained are in good agreement with the biological results explaining the evolution of the chromosome 2 from *Drosophila yakuba* to *Drosophila melanogaster* in the literature [4,15]. See Additional file 2 (Experimental results) for a description of the dataset and the results of the application.

Conclusion

In this paper, we introduced the genome dedoubling problem in the DCJ rearrangement model, NP-complete

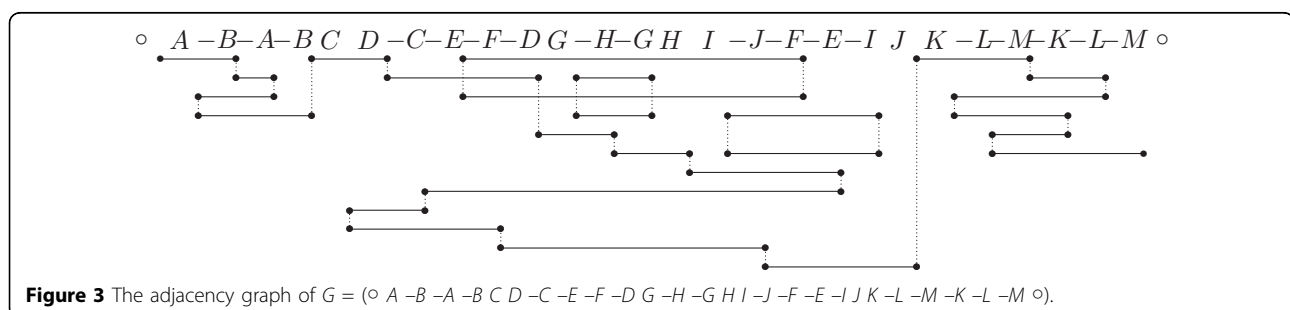
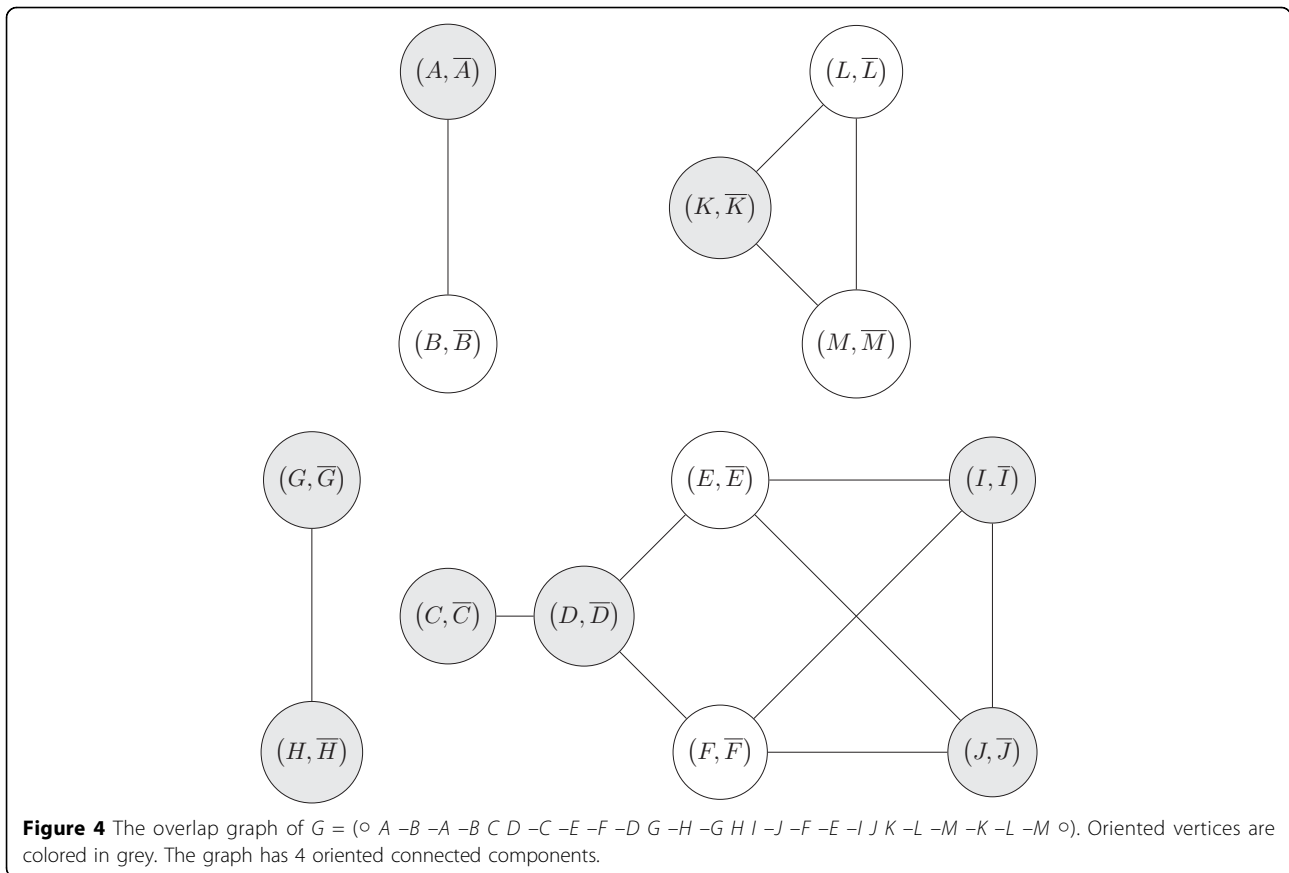


Figure 3 The adjacency graph of $G = (o \ A \ -B \ -A \ -B \ C \ D \ -C \ -E \ -F \ -D \ G \ -H \ -G \ H \ I \ -J \ -F \ -E \ -I \ J \ K \ -L \ -M \ -K \ -L \ -M \ o)$.



in both the multichromosomal and the linear unichromosomal case, by reduction to an APX-complete problem. For both cases, we described an algorithm solving the problems in linear time complexity, except for an APX-complete part that is 2-approximable. We also presented some results on the Genome Dedoubling problem by reversal, providing an algorithm solving the problem on oriented genomes in quadratic time complexity, except for an APX-complete part that is 2-approximable. The case of unoriented genomes in the reversal model will be treated in a future paper. Unsurprisingly, partial results obtained so far tend to show that the general distance formula can be written as $d_{rev}(G) = n - C + 2m + t$, with t corresponding to the cost of genome orientation. However, the cost t here differs from the orientation cost described in the classical reversal theory based on the unoriented component tree [14]. In our case, the structure of the graph $\mathcal{A}(G)$ allows to orient components while not decreasing the number of cycles, or even increasing it in some cases. This requires proper identification of different kinds of merging reversals and further extensions on the data structures presented in this paper.

The second obvious extension of the present work, as in the the Genome Halving problem theory [16], is to

generalize the Genome Dedoubling problem defined on a single genome, to the *Guided Genome Dedoubling problem*, that asks to find a non-duplicated genome that minimizes the breakpoint-duplication distance to a given duplicated genome, plus the distance to a given non-duplicated genome. A further extension of this work consists of taking account of the degree of divergence of the breakpoint-duplicated sequences to order the rearrangement operations in time as done in [10].

Additional material

Additional file 1: Supplemental proofs Additional file 1 is a PDF file containing the proofs of Proposition 3, Property 1, Lemma 1, and Property 3.

Additional file 2: Experimental results Additional file 2 is a PDF file containing a description of an application of the methods to real *Drosophila* data.

Acknowledgements

We would like to thank Anne Bergeron for her useful comments on the breakpoint-duplication phenomenon, and the anonymous reviewers of the paper for their useful comments on the first version of the document. This article has been published as part of *BMC Bioinformatics* Volume 12 Supplement 9, 2011: Proceedings of the Ninth Annual Research in Computational Molecular Biology (RECOMB) Satellite Workshop on

Comparative Genomics. The full contents of the supplement are available online at <http://www.biomedcentral.com/1471-2105/12?issue=S9>.

Authors' contributions

The work was divided in four steps: 1) Formal introduction and reduction of the Genome Dedoubling problem. 2) Design of the study. 3) Study of the Genome Dedoubling problem by DCJ. 4) Study of the Genome Dedoubling problem by reversal on oriented genomes. AT participated in 2) and 3), and carried out 4). JSV participated in 2), 3) and 4). AO carried out 1), 2) and 3), and participated in 4). All authors participated in writing the manuscript, and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Published: 5 October 2011

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doi:10.1186/1471-2105-12-S9-S20

Cite this article as: Thomas et al.: Genome dedoubling by DCJ and reversal. *BMC Bioinformatics* 2011 **12**(Suppl 9):S20.

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