Building the compacted colored de Bruijn Graph



Construction of the compacted colored De Bruijn Graph from reference sequence

Bioinformatics, 33(24), 2017, 4024–4032 doi: 10.1093/bioinformatics/btw609 Advance Access Publication Date: 21 September 2016 Original Paper



Sequence analysis

TwoPaCo: an efficient algorithm to build the compacted de Bruijn graph from many complete genomes

Ilia Minkin¹, Son Pham² and Paul Medvedev^{1,3,4,*}

¹Department of Computer Science and Engineering, The Pennsylvania State University, University Park, PA 16802, USA, ²BioTuring Inc., San Diego, CA 92121, USA, ³Department of Biochemistry and Molecular Biology and ⁴Genomic Sciences Institute of the Huck, The Pennsylvania State University, University Park, PA 16802, USA

*To whom correspondence should be addressed.

Associate Editor: Alfonso Valencia

Received on April 3, 2016; revised on September 1, 2016; accepted on September 16, 2016



TwoPaCo: An efficient algorithm to build the compacted de Bruijn graph from many complete genomes

Ilia Minkin¹, Son Pham², Paul Medvedev¹

Pennsylvania State University¹
Salk Institute for Biological Studies²

8th July 2016

Motivation

- More and more complete genomes
- Pan-genome: analysis within same species
- Mammalian-sized genomes are coming soon

Motivation

- More and more complete genomes
- Pan-genome: analysis within same species
- Mammalian-sized genomes are coming soon

Key question: what is a handy data structure to represent genomes?

Motivation

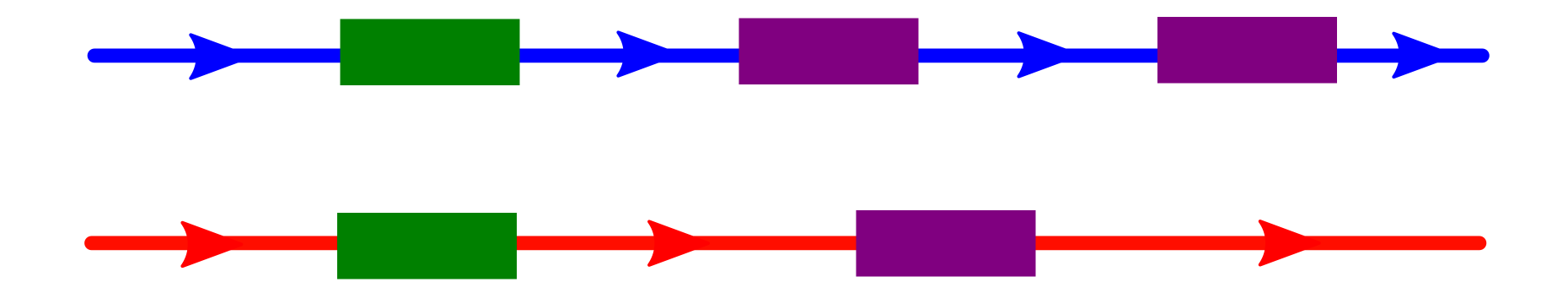
- More and more complete genomes
- Pan-genome: analysis within same species
- Mammalian-sized genomes are coming soon

Key question: what is a handy data structure to represent genomes?

The simplest way: string(s) of characters.

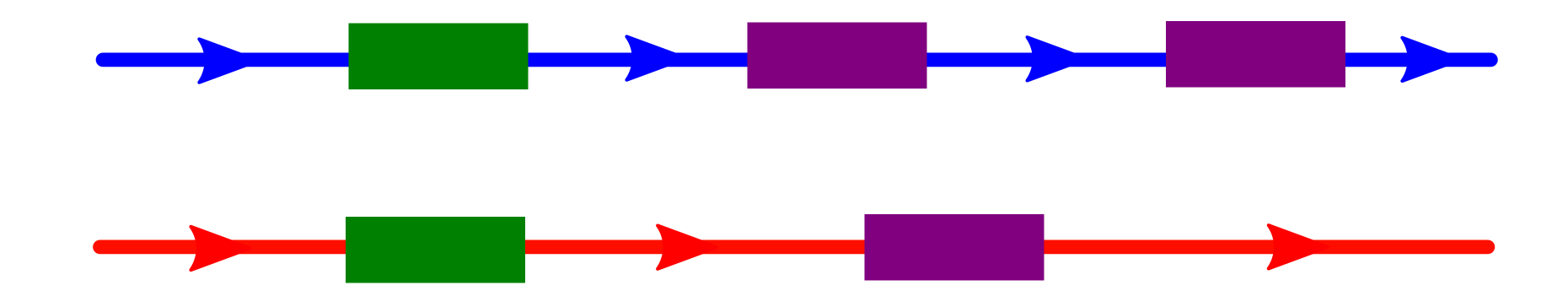
The Linear Representation

Two genomes:



The Linear Representation

Two genomes:

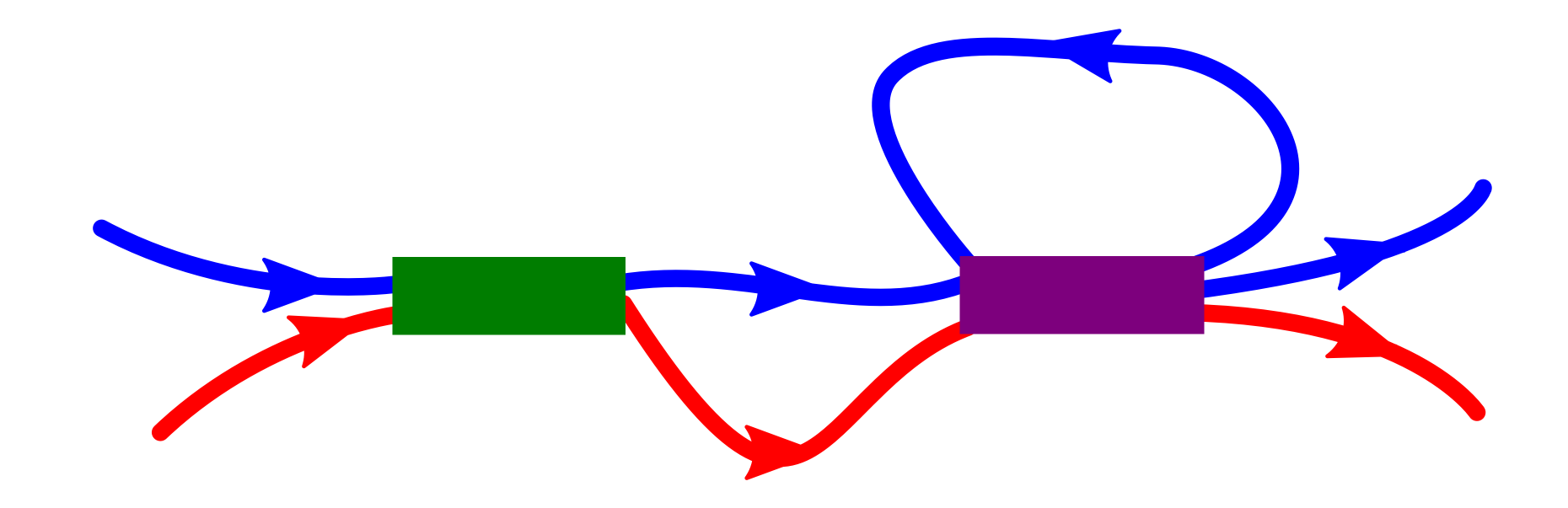


Issues:

- Homology between genomes?
- Duplications?
- Rearrangements?

Solution: a Graph Representation

What we want to see:



Why de Bruijn graph?

A simple object.

Demonstrated utility in:

- Assembly
- Read mapping
- Synteny identification

$$k = 2$$

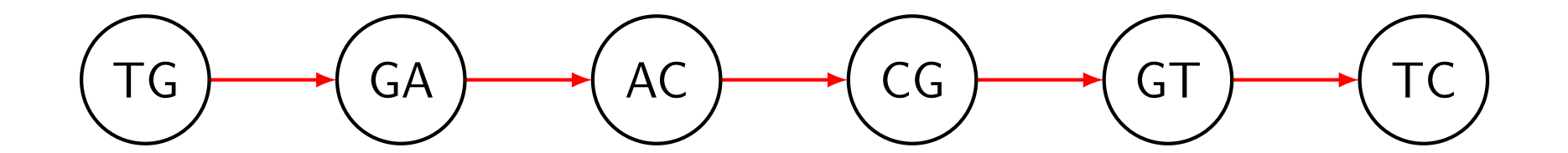
TGACGTC

TGACTTC

$$k = 2$$

TGACGTC

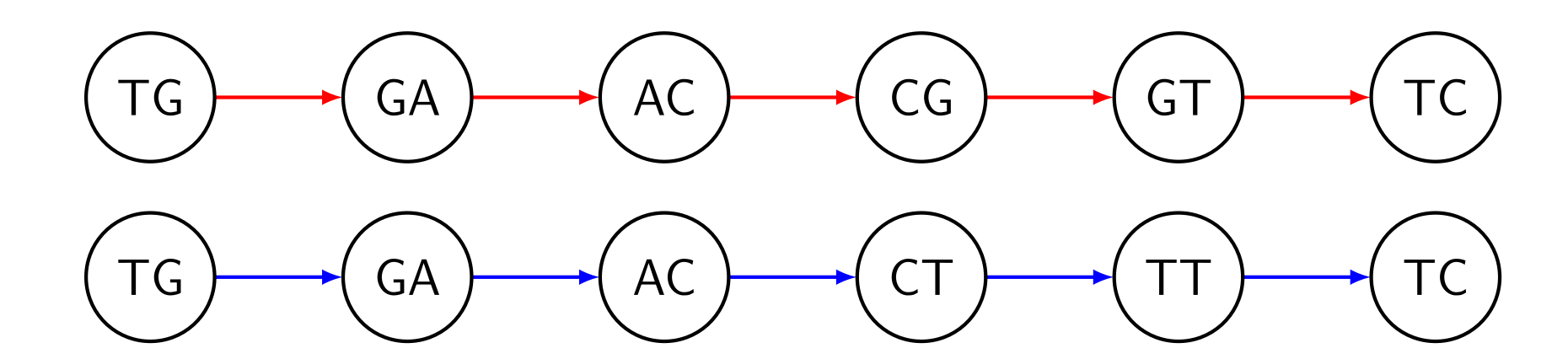
TGACTTC

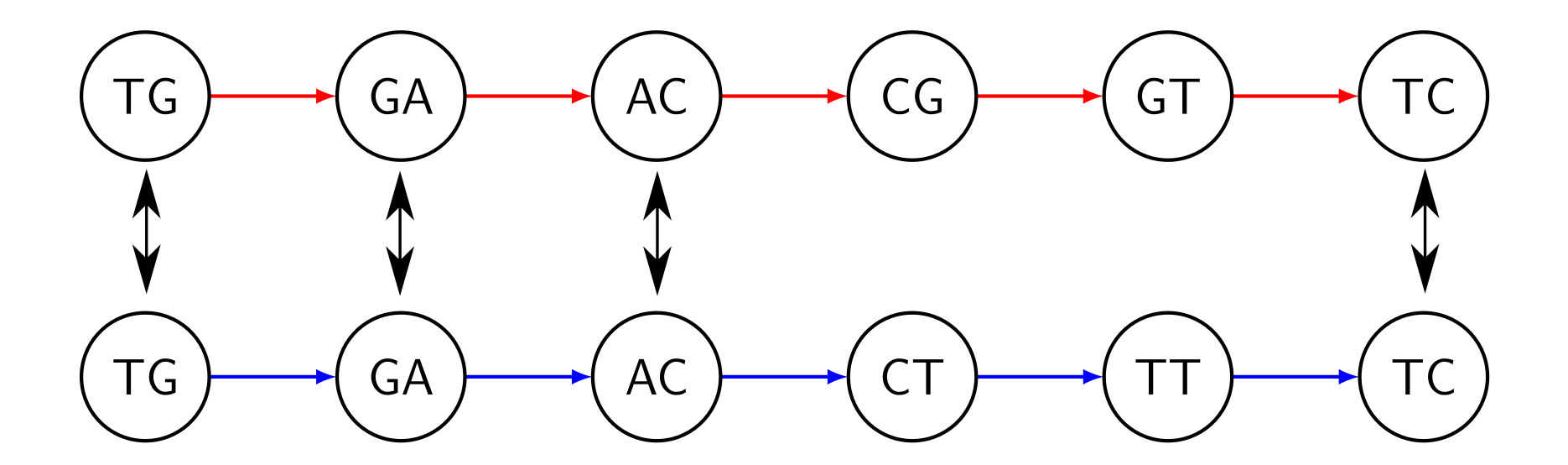


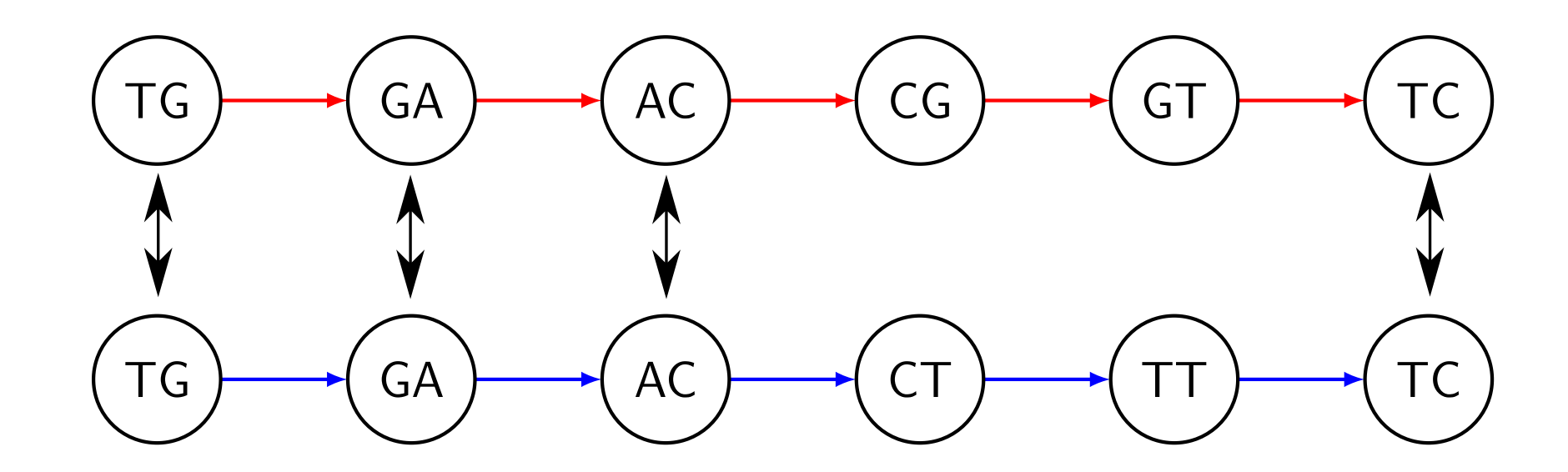
$$k = 2$$

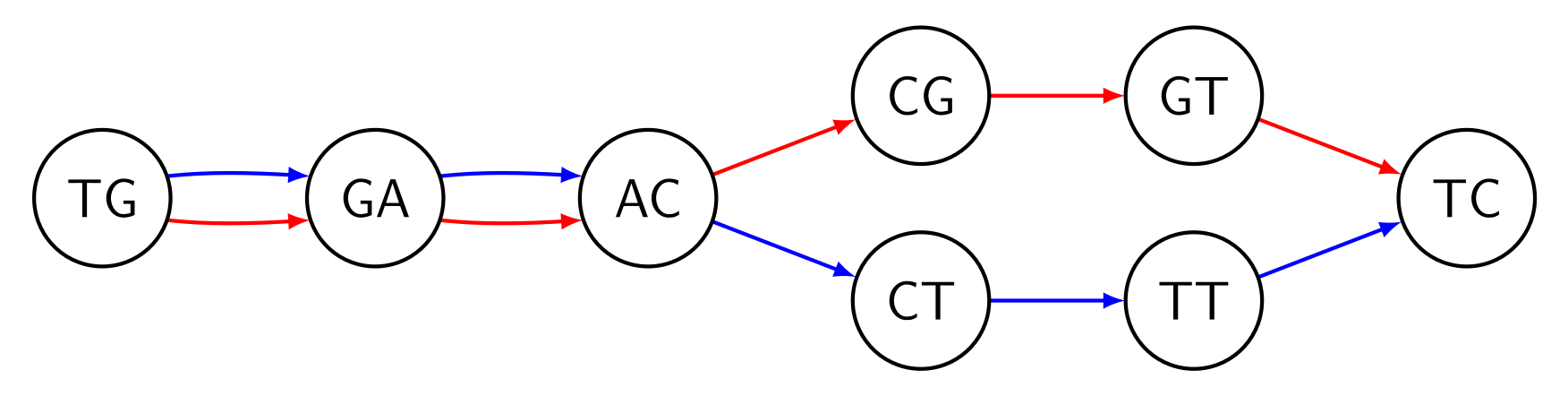
TGACGTC

TGACTTC

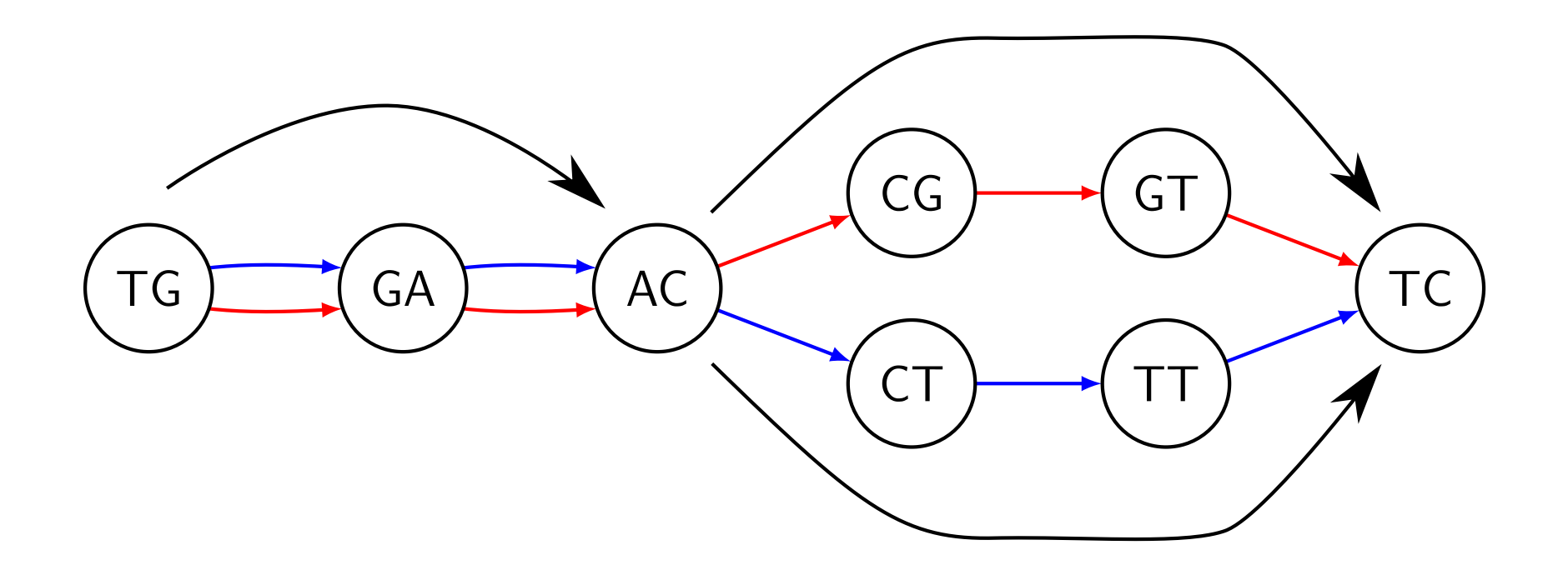




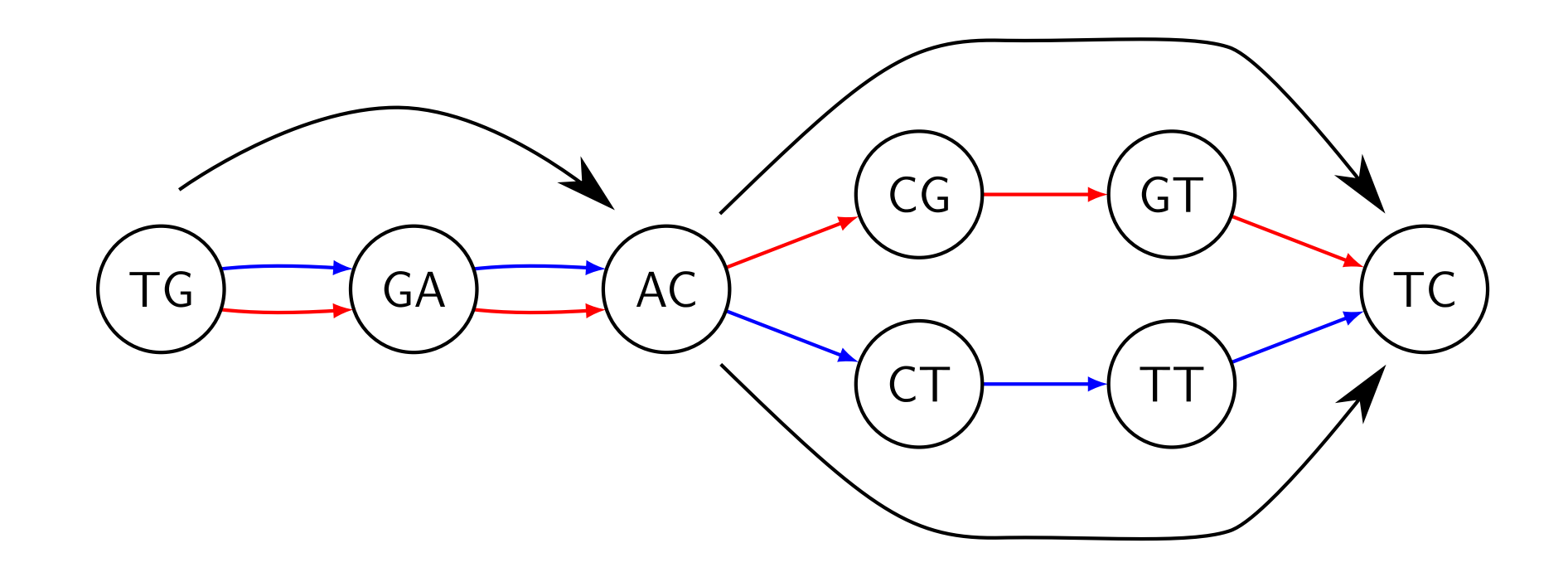




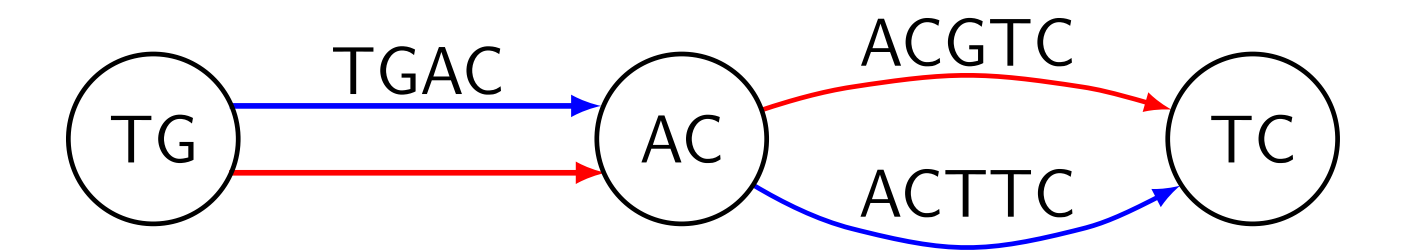
Compaction



Compaction



After compaction:



The Challenge

Construct the compacted graph from many large genomes **bypassing** the ordinary graph traverse.

The Challenge

Construct the compacted graph from many large genomes **bypassing** the ordinary graph traverse.

Earlier work: based on suffix arrays/trees Sibelia & SplitMEM handled > 60 E.Coli genomes.

The Challenge

Construct the compacted graph from many large genomes **bypassing** the ordinary graph traverse.

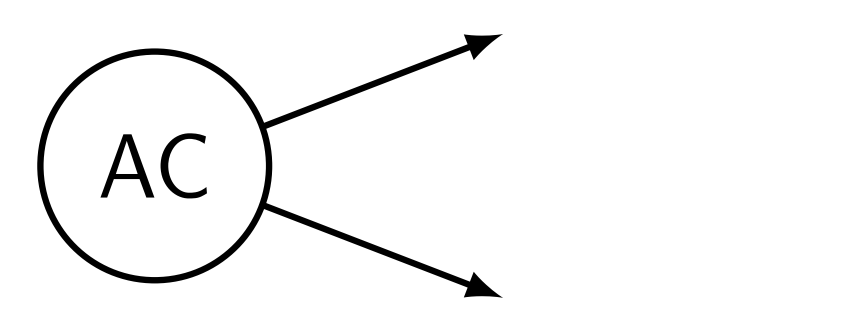
Earlier work: based on suffix arrays/trees Sibelia & SplitMEM handled > 60 E.Coli genomes.

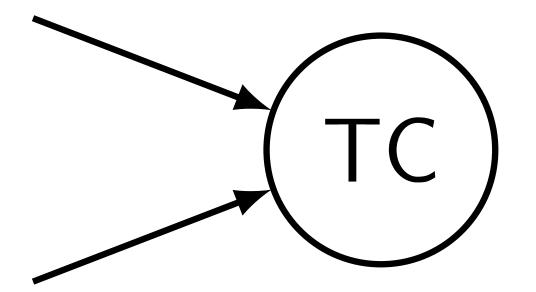
A recent advance: 7 Humans in 15 hours using 100 GB of RAM using a BWT-based algorithm by Baier et al., 2015, Beller et al., 2014.

Junctions

A vertex v is a **junction** if:

• v has ≥ 2 distinct outgoing or incoming edges:





Junctions

A vertex v is a **junction** if:

 \triangleright v has ≥ 2 distinct outgoing or incoming edges:



v is the first or the last k-mer of an input string

Junctions

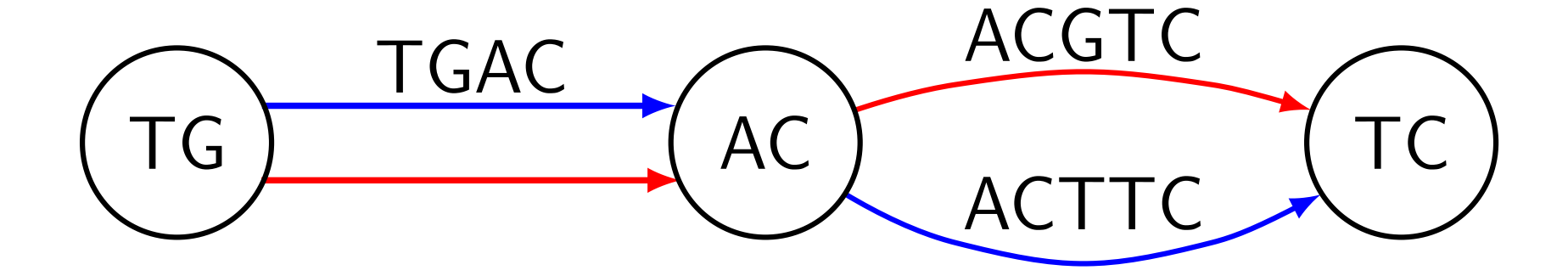
A vertex v is a **junction** if:

 \triangleright v has ≥ 2 distinct outgoing or incoming edges:

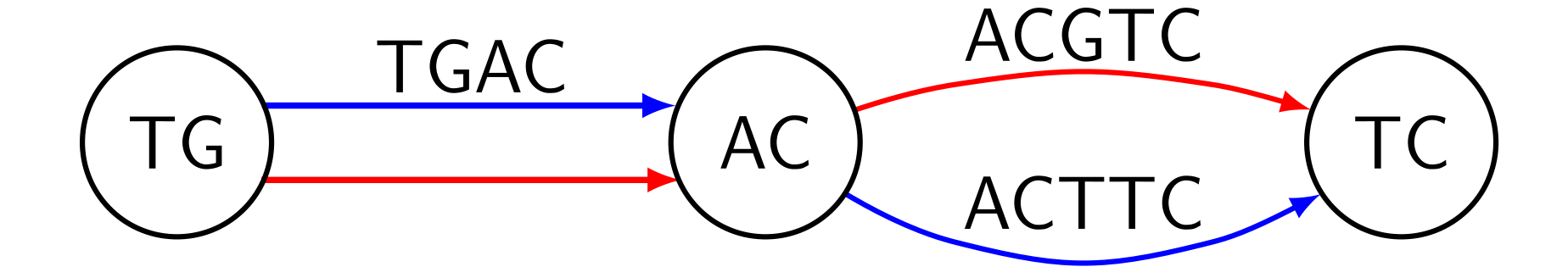


- ► *v* is the first or the last *k*-mer of an input string Facts:
 - Junctions = vertices of the compacted graph
 - Compaction = finding positions of junctions

Observations

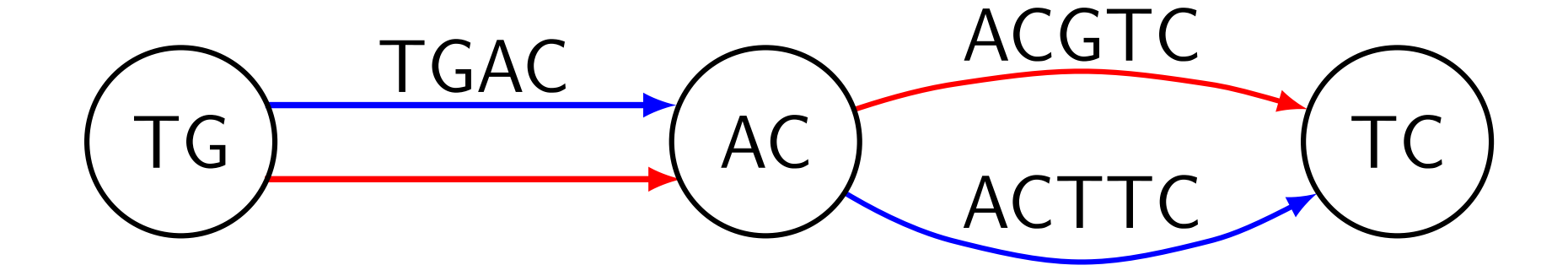


Observations



TG GA AC CG GT TC

Observations



TG GA AC CG GT TC $TG \rightarrow AC \rightarrow TC$

The Observation

The observation only works when we have complete genomes.

Once we know junctions, construction of the edges is simple.

We can simply traverse input strings and record junctions in the order they appear.

How to identify junctions?

The Naive Algorithm

A naive way:

- Store all (k + 1)-mers (edges) in a hash table
- Consider each vertex one by one
- Query all possible edges from the table
- \triangleright If found > 1 edge, mark vertex as a junction

Simple algorithm in more detail

Algorithm 1. Filter-Junctions

```
Input: strings S = \{s_1, \ldots, s_n\}, integer k, and an empty set data structure E. A candidate set of marked junction positions C \supseteq J(S, k)
is also given. When the algorithm is run naively, all the positions would be marked.
```

Output: a reduced candidate set of junction positions.

```
1: for s \in S do
    for 1 \leq i < |s| - k do
     if C[s, i] = marked then
                                                                             \triangleright Insert the two (k+1)-mers containing the k-mer at i into E.
     Insert s[i..i+k] into E.
     Insert s[i-1..i-1+k] into E.
6: for s \in S do
    for 1 \leq i < |s| - k do
        if C[s, i] = \text{marked and } s[i..i + k - 1] is not a sentinel then
       in \leftarrow 0
                                                                                                                   Number of entering edges
      out \leftarrow 0
                                                                                                                    Number of leaving edges
                                                                              > Consider possible edges and count how many of them exist
           for c \in \{A, C, G, T\} do
                if v \cdot c \in E then

    ▷ The symbol · depicts string concatenation

                   out \leftarrow out + 1
                if c \cdot v \in E then
                   in \leftarrow in + 1
             if in = 1 and out = 1 then
                                                                                                          \triangleright If the k-mer at i is not a junction.
                C[s, i] \leftarrow Unmarked
18: return C
```

The Naive Algorithm

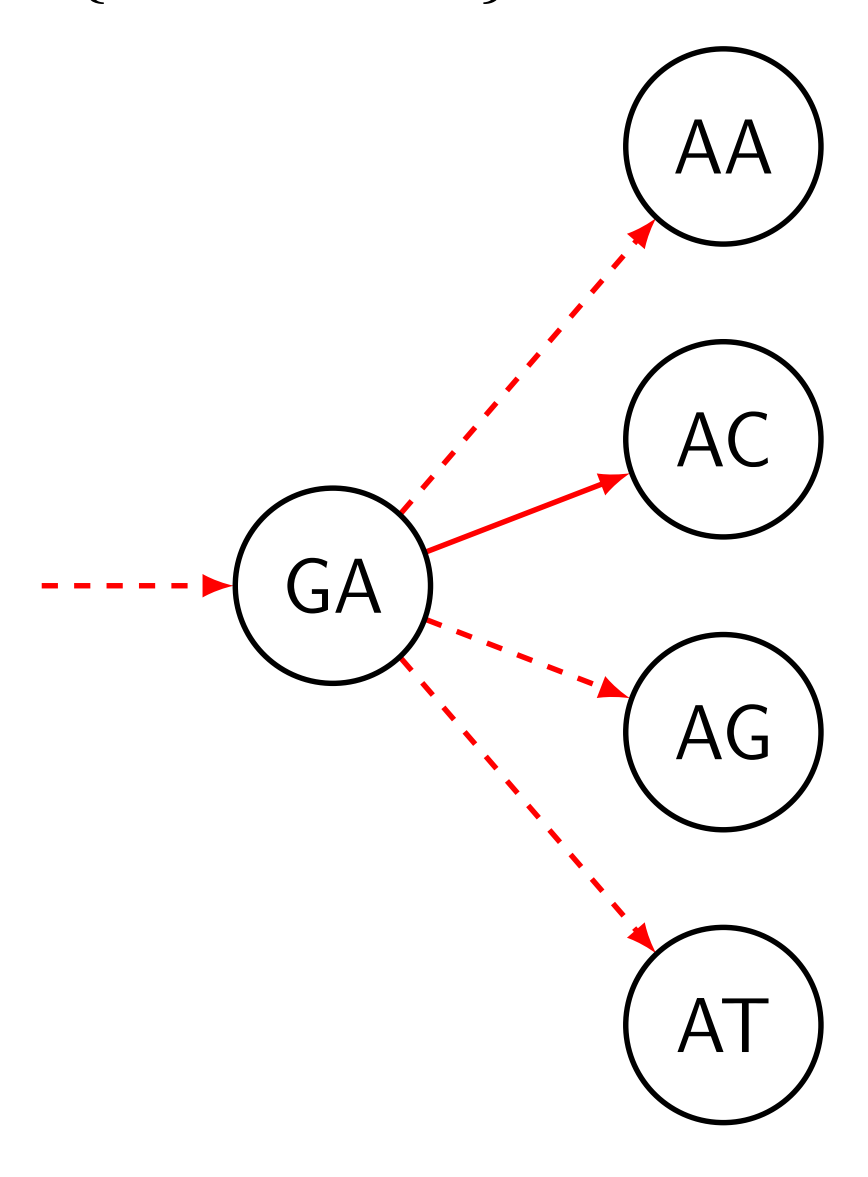
A naive way:

- Store all (k + 1)-mers (edges) in a hash table
- Consider each vertex one by one
- Query all possible edges from the table
- \blacktriangleright If found > 1 edge, mark vertex as a junction

Problem: the hash table can be too large.

An Example

 $\mathsf{Hash}\;\mathsf{table} = \{\;\mathsf{GA} \to \mathsf{AC}\;\}$



What is the Bloom filter

A probabilistic data structure representing a set

Properties:

- Occupies fixed space
- May generate false positives on queries
- False positive rate is low

What is the Bloom filter

A probabilistic data structure representing a set

Properties:

- Occupies fixed space
- May generate false positives on queries
- False positive rate is low

```
Example: Bloom Filter = \{GA \rightarrow AC\}
```

Is $GA \rightarrow AC$ in the set? Yes.

What is the Bloom filter

A probabilistic data structure representing a set

Properties:

- Occupies fixed space
- May generate false positives on queries
- False positive rate is low

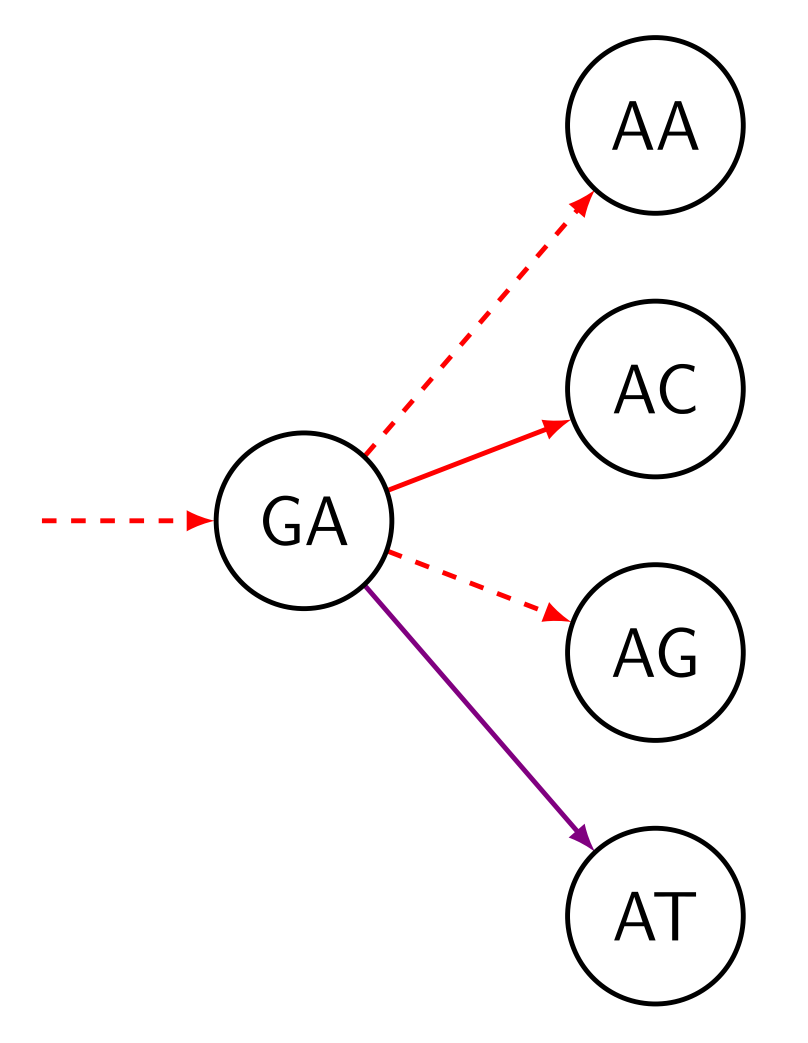
```
Example: Bloom Filter = \{GA \rightarrow AC\}
```

Is $GA \rightarrow AC$ in the set? Yes.

Is $GA \rightarrow AT$ in the set? **Maybe** no.

An Example

Bloom Filter = $\{ GA \rightarrow AC, GA \rightarrow AT \}$



The purple edge is a false positive.

The Two Pass Algorithm

How to eliminate false positives?

The Two Pass Algorithm

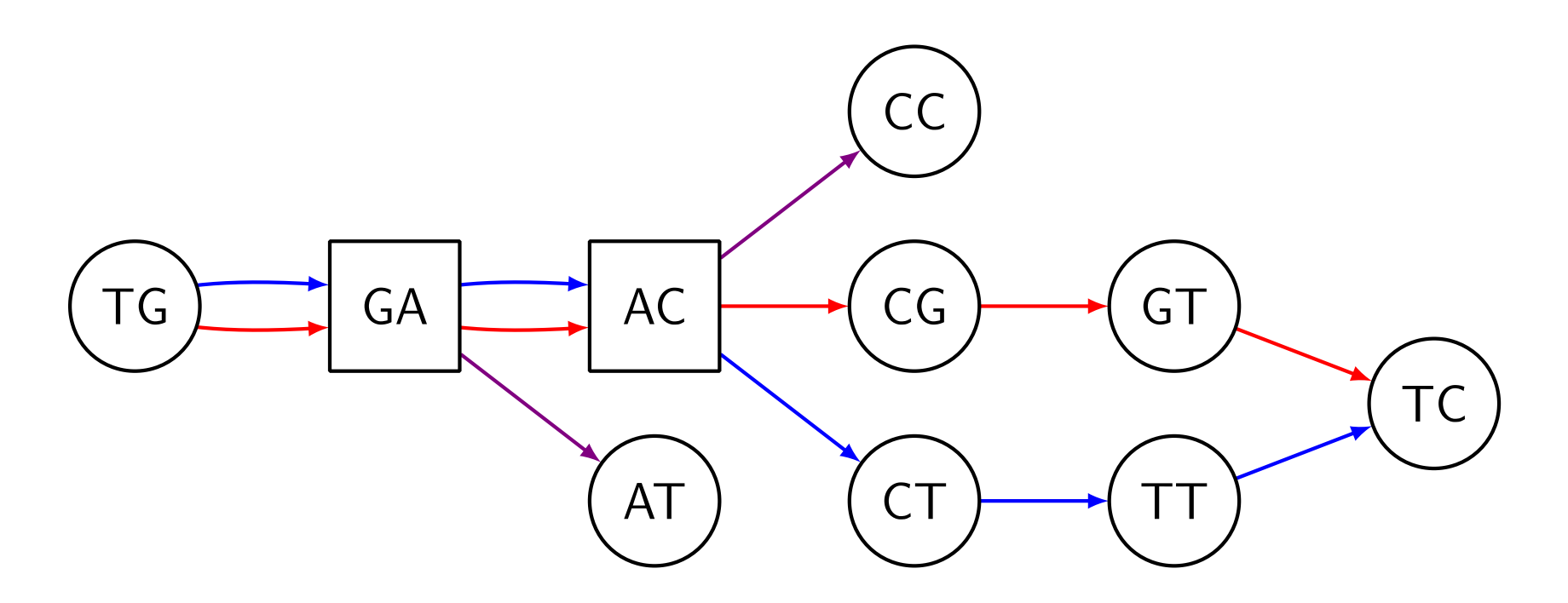
How to eliminate false positives?

Two-pass algorithm:

- 1. Use the Bloom filter to identify **junction** candidates
- 2. Use the hash table, but store **only edges that touch candidates**

An Example: the First Step

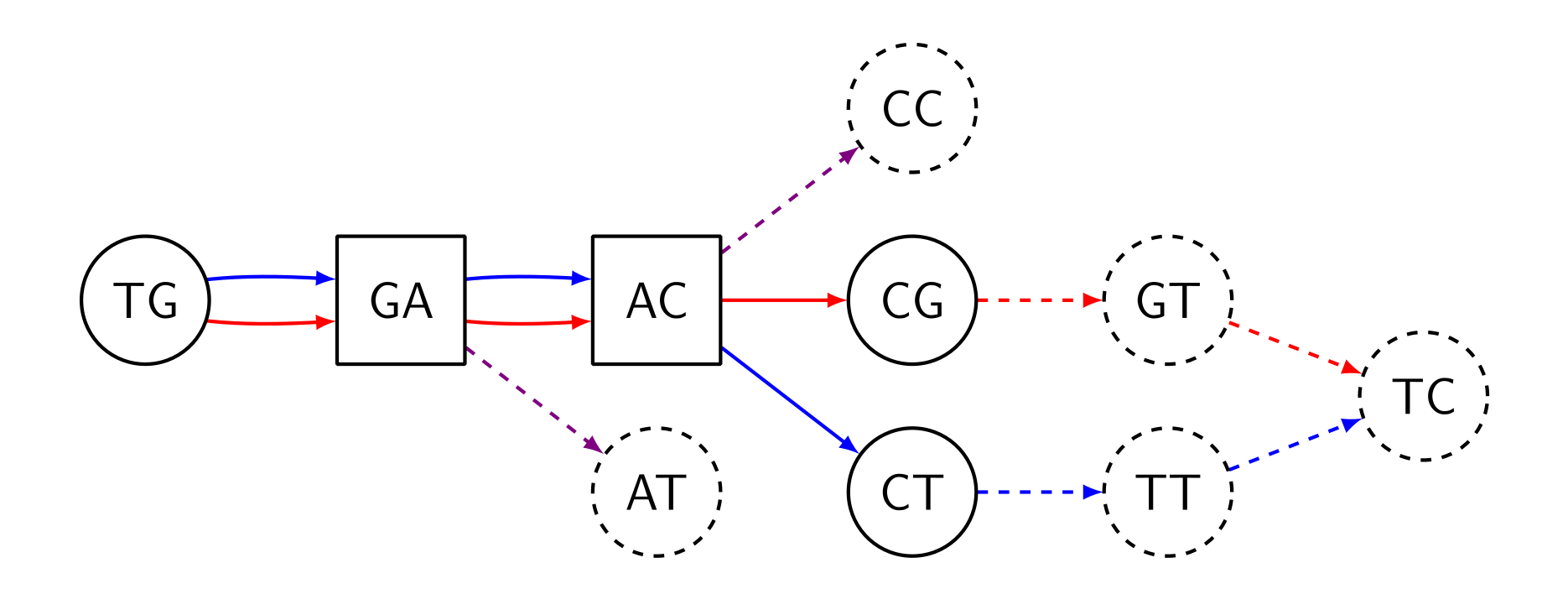
Here edges stored in the Bloom filter, purple ones are false positives:



Junction candidates: GA & AC

An Example: the Second Step

Edges stored in the hash table. We kept only edges touching junction candidates:



Junction: AC

The TwoPass Algorithm

Algorithm 2. Filter-Junctions-Two-Pass

```
Input: strings S = \{s_1, \dots, s_n\}, integer k, a candidate set of junction positions C_{\text{in}}, integer b
```

Output: a candidate set of junction positions C_{out}

- 1: $F \leftarrow$ an empty Bloom filter of size b
- 2: $C_{\text{temp}} \leftarrow Filter Junctions(S, k, F, C_{\text{in}})$ \triangleright The first pass
- 3: $H \leftarrow$ an empty hash table
- 4: $C_{\text{out}} \leftarrow Filter Junctions(S, k, H, C_{\text{temp}}) \triangleright \text{The second pass}$
- 5: return C_{out}

The TwoPaCo algorithm

Algorithm 3. TwoPaCo

```
Input: strings S = \{s_1, \ldots, s_n\}, integer k, integer \ell, integer b
Output: the compacted de Bruijn graph G_c(S, k)
1: Initialize counters c_0, \ldots, c_{q-1} to zeroes
2: F \leftarrow an empty Bloom filter of size b
3: for s \in S do
4: for 1 \le i \le |s| - k + 1 do
5: b \leftarrow s[i..i+k-1]
    if h not in F then
      Insert h into F
     c_{f(b)} \leftarrow c_{f(b)} + 1
                                            \triangleright Mean number of k-mers per partition
12: p_i \leftarrow \text{biggest integer larger than } p_{i-1} \text{ such that } (\sum_{p_{i-1} \leq j < p_i} c_j) \leq T, \text{ or } \min\{\ell, p_{i-1} + 1\} \text{ if it does not exist.}
13: C<sub>init</sub> ← Boolean array with every position unmarked
14: for 1 \leq i \leq \ell do
15: C_i \leftarrow \text{mark every position of } C_{\text{init}} that starts a k-mer h with hash value p_{i-1} \leq f(h) < p_i
16: C'_i \leftarrow \text{Filter} - \text{Junctions} - \text{Two} - \text{Pass}(S, k, b, C_i)
17: C_{\text{final}} = \bigcup C'_{i}
18: return Graph implied by C_{\text{final}}, as described in Section 3.
```

Results

Datasets:

- ▶ 7 humans: 5 versions of the reference +
 2 haplotypes of NA12878 from 1000 Genomes
- ▶ 93 simulated humans (FIGG)
- ► 8 primates available in UCSC genome browser

Results

Format: minutes (GB)

Table 2. Benchmarking comparisons

	DSK+BCALM	Minia	Sibelia	SplitMem	bwt-based from	Baier et al. (2015)	TwoPaCo	
				Single strand	Single strand	Both strands	1 thread	15 threads
62 E.coli (k = 25)	6 (1.57)	151 (0.9)	10 (12.2)	70 (178.0)	8 (0.85)	12 (1.7)	4 (0.16)	2 (0.39)
62 E.coli (k = 100)	13 (2.50)	114 (1.9)	8 (7.6)	67 (178.0)	8 (0.50)	12 (1.0)	4 (0.19)	2 (0.39)
7 humans ($k = 25$)	444 (22.44)	968 (48.09)	_	_	867 (100.30)	1605 (209.88)	436 (4.40)	63 (4.84)
7 humans ($k = 100$)	1347 (221.65)	1857 (222.0)	_	_	807 (46.02)	1080 (92.26)	317 (8.42)	57 (8.75)
8 primates ($k = 25$)	2088 (85.62)	_	_	_	_	_	914 (34.36)	111 (34.36)
8 primates ($k = 100$)	_	_	_	_	_	_	756 (56.06)	101 (61.68)
(43+7) humans $(k=25)$	_	_	_	_	_	_		705 (69.77)
(43 + 7) humans $(k = 100)$	_	_	_	_	_	_		927 (70.21)
(93 + 7) humans $(k = 25)$	_	_	-	-	-	_		1383 (77.42)

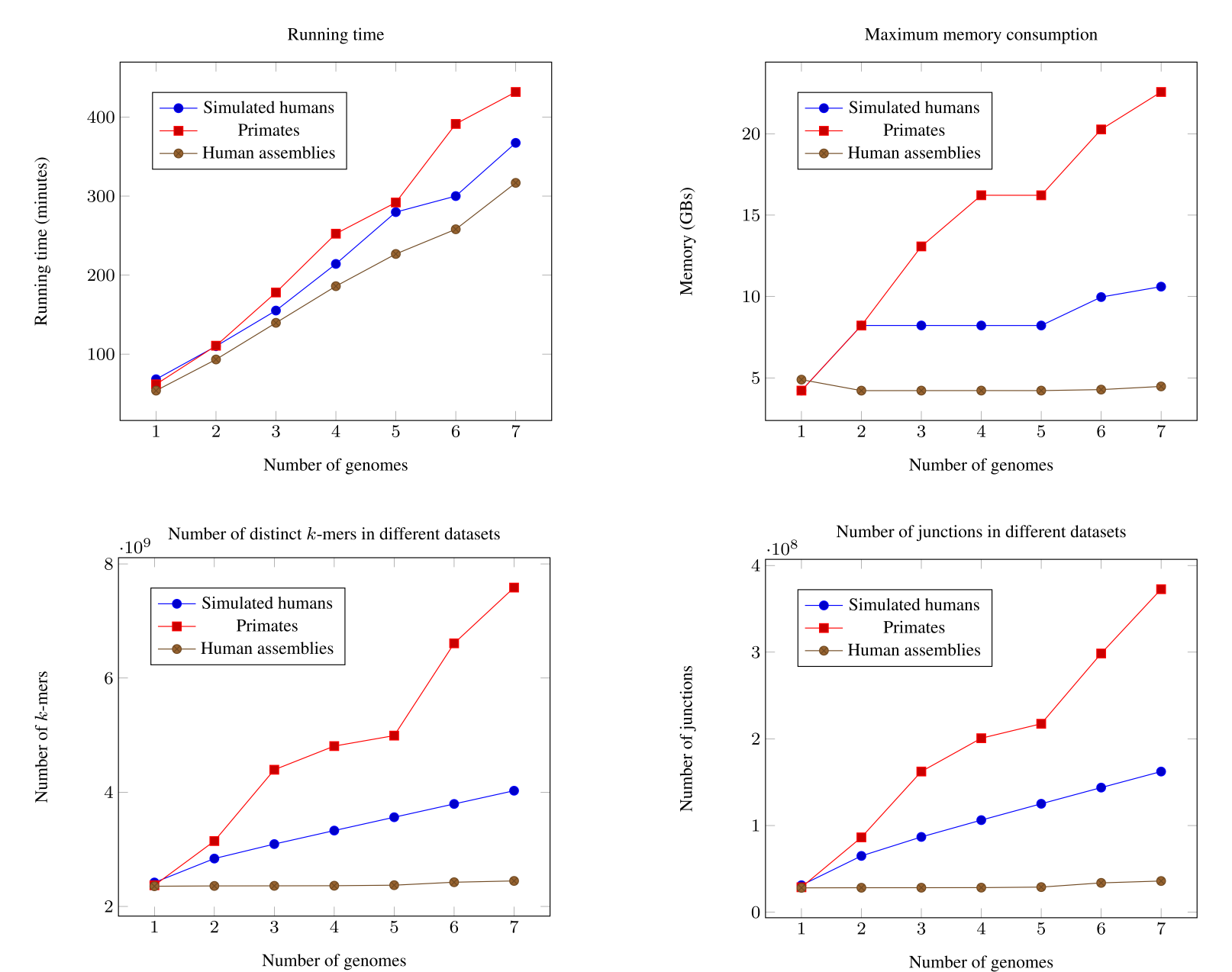
Note: Each cell shows the running time in minutes and the memory usage in parenthesis in gigabytes. TwoPaCo was run using just one round, with a Bloom filter size b = 0.13 GB for E.coli, 4.3 GB for 7 humans with k = 25, b = 8.6 GB with k = 100, b = 34 GB for primates, and b = 69 GB for (43 + 7) and larger human dataset. A dash in the SplitMem and bwt-based columns indicates that they ran out of memory, a dash in the Sibelia column indicates that it could not be run on such large inputs, a dash in the minia column indicates that it did not finish in 48 h, a dash in the BCALM column indicates that it ran out of disk space (4 TB). A double dash indicates that the software had a segmentation fault. An empty slot indicates that the experiment was not done.

Conclusion & Future Work

Can potentially facilitate:

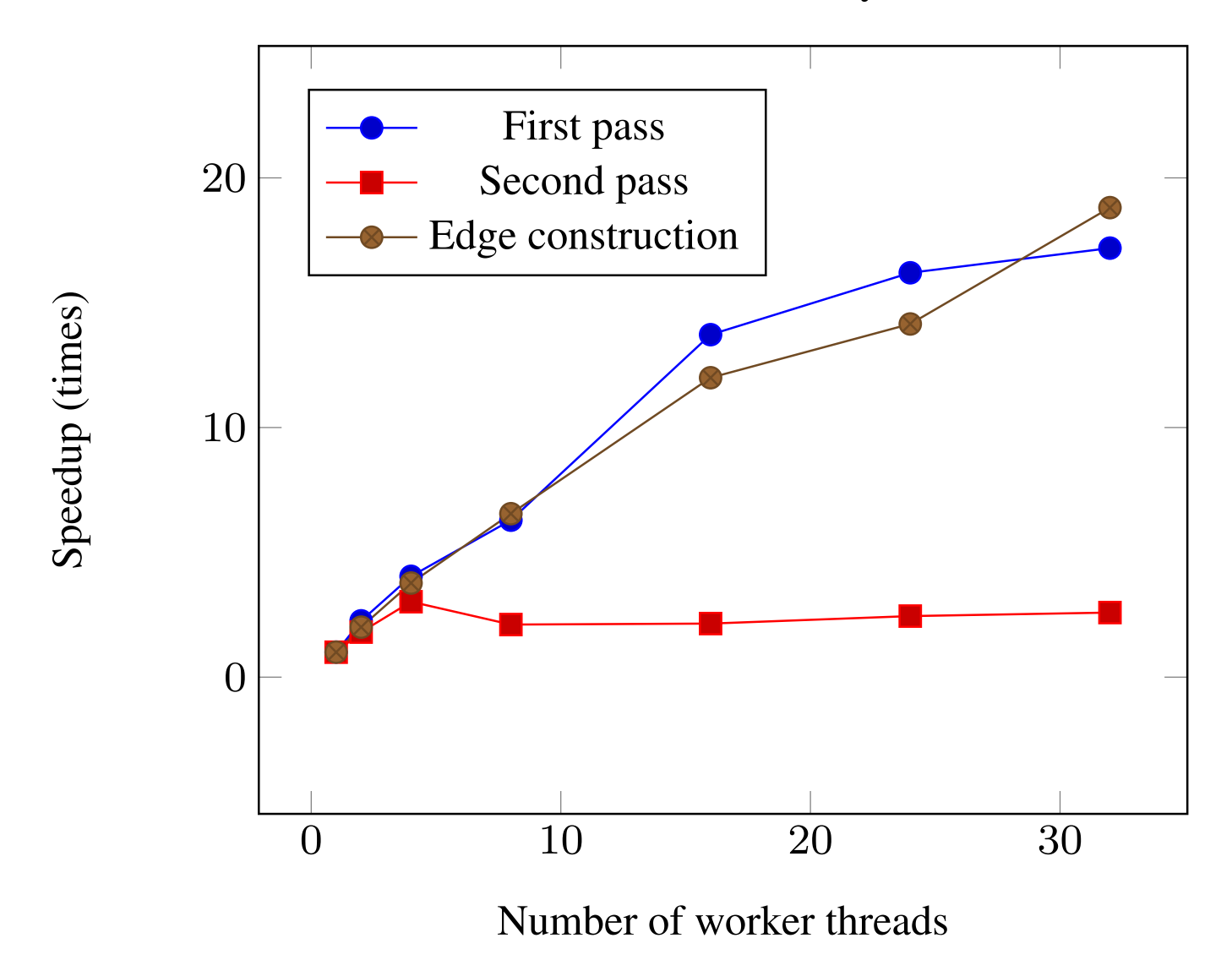
- Visualization
- Synteny mining (Sibelia)
- Structural variations analysis
- ...

Input Size vs. Performance



Parallel Scalability

Parallel scalability



Splitting

Table 1: The minimal number of rounds it takes to compress the graph without exceeding a given memory threshold.

Memory threshold	Used memory	Bloom filter size	Running time	Rounds
10	8.62	8.59	259	1
8	6.73	4.29	434	3
6	5.98	4.29	539	4
4	3.51	2.14	665	6

Can we do even better?

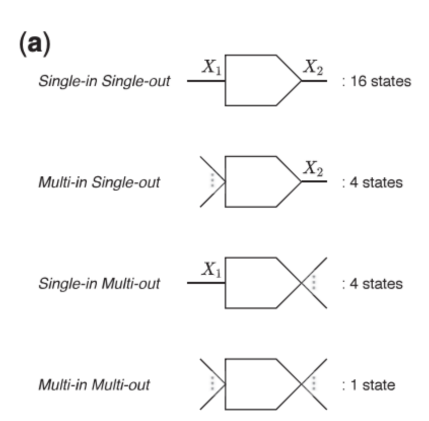
Bioinformatics, 37, 2021, i177–i186 doi: 10.1093/bioinformatics/btab309 ISMB/FCCB 2021

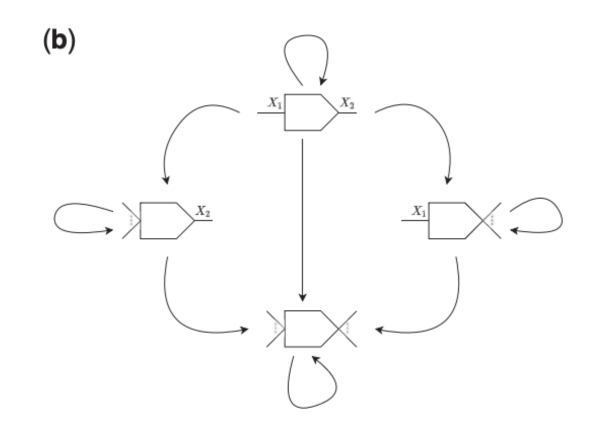


Cuttlefish: fast, parallel and low-memory compaction of de Bruijn graphs from large-scale genome collections

Jamshed Khan^{1,2} and Rob Patro^{1,2,*}

¹Department of Computer Science, University of Maryland, College Park, MD 20742, USA and ²Center for Bioinformatics and Computational Biology, University of Maryland, College Park, MD 20742, USA





Key ideas:

Just like TwoPaCo, make use of explicit traversal of references to identify junction nodes

Build a minimal perfect hash (BBhash here) over the set of k-mers

Associate each k-mer with a finite state automaton, denoting its topological status — 26 states requires 5-bits/k-mer

Walking the references and updating the states results in correct status for each k-mer, and unitigs can then be extracted as sequences between the junction nodes.

Cuttlefish

Table 1. Time- and memory-performance benchmarking for compacting single input reference de Bruijn graphs

			Bifrost		deGSM		TwoPaCo		Cuttlefish	
	Thread- count	k	Build	Output	Build	Output	Build	Output	Build	Output
Human	1	31	04:54:50 (27.23)	15:18	01:54:41 (37.94)	25:06 (9.79)	01:13:19 (4.15)	39:38 (4.50)	32:59 (2.79)	19:23 (2.84
8		61	05:16:51 (50.19)	01:49	02:20:57 (84.16)	21:37 (8.77)	01:10:18 (6.02)	12:25 (4.35)	38:21 (3.06)	15:37 (3.08
	8	31	01:33:54 (27.23)	03:59	25:20 (37.94)	05:37 (9.80)	12:57 (5.04)	_	05:49 (2.79)	05:13 (2.92)
		61	01:20:28 (50.18)	00:40	47:52 (84.16)	03:55 (8.80)	11:28 (5.46)	_	07:45 (3.06)	03:20 (3.18)
	16	31	01:24:40 (27.24)	03:30	18:19 (37.94)	03:56 (9.80)	06:24 (5.57)	_	03:26 (2.79)	02:57 (2.93)
		61	01:12:33 (50.18)	00:52	46:34 (84.16)	02:35 (8.80)	07:12 (5.55)	_	04:23 (3.06)	01:54 (3.19)
Gorilla	1	31	05:44:10 (28.08)	16:30	01:34:29 (37.94)	24:26 (9.75)	01:00:15 (5.04)	43:25 (4.49)	31:46 (2.74)	17:07 (2.77
		61	05:31:06 (50.13)	02:05	02:11:33 (84.16)	22:03 (8.94)	01:11:29 (5.83)	17:52 (4.30)	38:15 (3.02)	15:59 (3.03)
	8	31	02:06:52 (28.08)	03:44	28:52 (37.94)	05:43 (9.76)	13:02 (5.82)	_	05:30 (2.74)	04:37 (2.87)
		61	01:24:21 (50.13)	00:54	47:45 (84.16)	03:59 (8.98)	10:03 (6.00)	_	07:58 (3.02)	02:54 (3.12)
	16	31	01:50:26 (28.08)	02:59	20:47 (37.94)	04:07 (9.76)	07:29 (5.52)	_	03:13 (2.74)	03:25 (2.87)
		61	01:10:06 (50.13)	04:04	38:45 (84.16)	02:40 (8.98)	06:24 (6.09)	_	04:29 (3.02)	02:06 (3.14)
Sugar	16	31	22:18:24 (229.17)	01:20:51	09:29:24 (145.23)	01:10:55 (119.18)	01:49:01 (61.93)	_	51:30 (14.24)	01:56:52
pine										(14.28)
		61	<i>X</i> (364.25)	_	<i>X</i> (166.54)	_	01:26:39 (64.86)	_	03:14:44 (20.88)	01:26:26
										(20.90)

Table 2 Time- and memory-performance benchmarking for compacting colored de Bruijn graphs (i.e. multiple input references) for k=31, using 16 threads

Dataset	Total genome-length (bp)	Distinct k-mers count	Bifrost	deGSM	TwoPaCo	Cuttlefish
62 E.coli	310 M	24 M	1 (0.47)	1 (3.34)	1 (0.80)	1 (0.96)
7 Humans	21 G	2.6 B	95 (29.06)	30 (37.94)	62 (6.14)	21 (2.88)
7 Apes	18 G	7.1 B	294 (100.25)	172 (145.23)	59 (28.87)	25 (7.42)
11 Conifers	204 G	82 B	_	_	981 (288.99)	525 (84.12)
100 Humans	322 G	28 B	_	_	1395 (126.25)	523 (28.75)

Cuttlefish 2

Scalable, ultra-fast, and low-memory construction of compacted de Bruijn graphs with Cuttlefish 2

Jamshed Khan^{1,2}, Marek Kokot^{3*}, Sebastian Deorowicz³ and Rob Patro^{1,2*}

Can generalize the cuttlefish algorithm to work on raw sequencing data in addition to reference genomes. Leads to a state-of-the-art compacted dBG construction algorithm.

Table 1 Time- and memory-performance results for constructing compacted de Bruijn graphs from short-read sets

			ABySS-BLOOM-DBG		BIFROST	DE GSM	BCALM 2	CUTTLEFISH 2		
Dataset	k	Thread-count	Small-memory	Large-memory				Default memory	Match second-best memory	Unrestricted memory
Human	27	8	22 h 18 min (39.3)	20 h 23 min (71.3)	11 h 43 min (48.5)	10 h 36 min (235.8)	04 h 23 min (6.7)	01 h 13 min (3.2)	01 h 10 min (6.2)	01 h (11.3)
		16	11 h 38 min (39.3)	11 h 02 min (71.3)	09 h 39 min (48.6)	07 h 08 min (235.8)	04 h 58 min (8.9)	56 min (3.3)	56 min (7.6)	51 min (11.3)
	55	8	16 h 32 min (34.0)	15 h 58 min (66.0)	05 h 43 min (43.8)	16 h 50 min (293.2)	04 h 01 min (7.4)	02 h 20 min (3.5)	01 h 08 min (7.1)	01 h 03 min (11.3)
		16	09 h 28 min (34.1)	08 h 37 min (66.1)	04 h 16 min (43.9)	15 h 54 min (293.3)	04 h 26 min (10.5)	02 h 02 min (3.7)	01 h 11 min (9.5)	51 min (11.3)
Human RNA-seq	27	8	11 h 47 min (33.7)	11 h 22 min (65.7)	06 h 04 min (7.2)	01 h 35 min (87.1)	02 h 58 min (3.8)	30 min (2.9)	-	18 min (80.1)
		16	11 h 38 min (39.3)	07 h 38 min (65.7)	07 h 24 min (7.2)	01 h 37 min (87.2)	02 h 46 min (3.9)	20 min (3.0)	_	12 min (80.1)
Gut microbiome	27	16	18 h 47 min (42.0)	20 h 12 min (74.0)	03 h 54 min (38.1)	02 h 28 min (157.2)	02 h 34 min (7.7)	26 min (3.5)	23 min (6.7)	20 min (26.8)
	55		1 day 17 h 43 min (35.9)	1 day 08 h 09 min (67.8)	02 h 44 min (46.7)	06 h 53 min (293.3)	03 h 02 min (12.5)	44 min (4.0)	25 min (11.3)	20 min (69.9)
Soil	27	16	1 d 18 h 35 min (150.4)	14 h 24 min (275.0)	15 h 28 min (274.1)	1 day 14 h 29 min (235.8)	19 h 39 min (52.0)	02 h 01 min (19.2))	02 h 18 min (40.9)	01 h 35 min (40.9)
	55		07 h 57 min (128.9)	06 h 36 min (256.8)	05 h 49 min (157.0)	1 day 11 h 05 min (293.3)	08 h 30 min (27.5)	03 h 02 min (11.1)	02 h 43 min (23.3)	01 h 38 min (23.3)
White spruce	27	16	*	X	Χ	†	2 days 06 h 12 min (36.8)	10 h 05 min (14.0)	07 h 47 min (35.2)	07 h 13 min (204.2)
	55		*	X	Χ	†	2 days 09 h 59 min (31.6)	10 h 12 min (23.8)	10 h 08 min (31.1)	07 h 24 min (279.3)

Table 2 Time- and memory-performance results for constructing compacted de Bruijn graphs from whole-genome reference collections

			B IFROST	DE GSM	BCALM 2	CUTTLEFISH 2		
Dataset (genome count)	k	Thread- count				Default memory	Unrestricted memory	
Human gut (30K)	27	8	06 h (155.1)	Δ	10 h 06 min (21.5)	01 h 39 min (15.2)	01 h 39 min (32.5)	
		16	05 h 30 min (155.1)		09 h 05 min (22.0)	01 h 01 min (15.5)	59 min (32.5)	
	55	8	08 h 47 min (279.2)		11 h 49 min (18.6)	04 h 14 min (20.6)	03 h 42 min (44.4)	
		16	08 h 20 min (279.2)		09 h 45 min (19.2)	03 h 50 min (20.9)	03 h 10 min (44.3)	
Human (100)	27	8	35 h 45 min (355.9)	19 h 23 min (235.8)	#	04 h 32 min (27.7)	04 h 09 min (59.7)	
		16	32 h 14 min (355.9)	14 h 07 min (235.8)	#	03 h 19 min (28.1)	02 h 49 min (59.7)	
	55	8	*	†	2 days 23 h 31 min (302.9)	15 h 08 min (56.0)	13 h 47 min (121.8)	
		16	*	†	*	12 h (56.2)	11 h 33 min (121.8)	
Bacterial archive (661K)	27	16	X	X	#	16 h 38 min (48.7)	16 h 24 min (104.9)	
	55				4 days 10 h 11 min (63.3)	22 h 44 min (59.9)	22 h 20 min (129.5)	