The Open Source Toolbox for Genomic Assembly & Analysis

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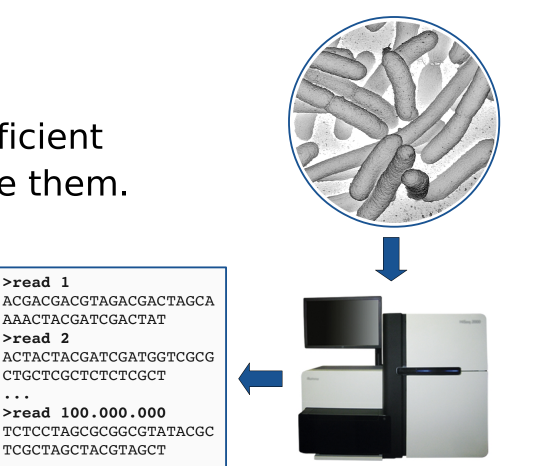
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1. What is GATB ?

Motivation

NGS technologies produce terabytes of data. Efficient and fast NGS algorithms are essential to analyze them.



2. Software Solution

The GATB philosophy proposes a 3-layer construction to analyze NGS datasets

1. GATB-CORE: a C++ library holding all the services needed for developing software dedicated to NGS data.

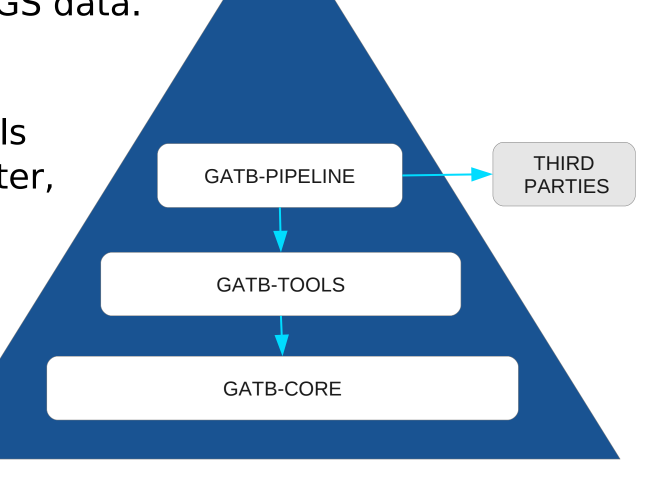
Objective

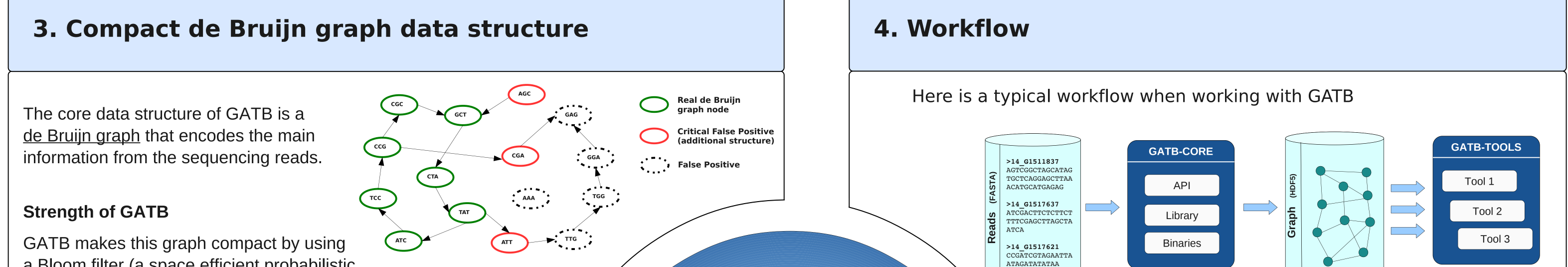
The Genome Assembly Tool Box (GATB)

- ► is an <u>open-source software</u>
- provides an easy way to develop efficient and fast NGS tools
- ► is based on data structure with <u>a very low memory footprint</u>
- allows complex genomes to be processed on desktop computers

2. **GATB-TOOLS**: a set of elementary NGS tools mainly built upon the GATB library (k-mer counter, contiger, scaffolder, variant detection, etc.).

3. **GATB-PIPELINE**: a set of NGS pipeline that links together tools from the previous layer.





a <u>Bloom filter</u> (a space efficient probabilistic data structure) and by using a CFP additional structure that avoids false positive answers from the Bloom filter due to its probabilistic nature.

5. GATB helps you as a NGS user

<u>GATB's de Bruijn graph: a basis for families of tools</u>

- ► Data error correction
- Assembly
- a whole human genome sequencing reads can be handled with 5 GBytes
- Biological motif detection _ of memory

Several tools based on GATB are already available

Bloocoo K-mer spectrum based read error corrector for large datasets

Short read assembler based on a de Bruijn graph. Results are Minia of similar contiguity and accuracy to other de Bruijn assemblers (e.g. Velvet)

DiscoSNP Discover Single Nucleotide Polymorphism (SNP) from nonassembled reads

How to Analyze Complex Genomes on a Simple Desktop Computer ?

GATB-CORE transforms the reads into a de Bruijn graph, saves it in a HDF5 file that can be opened by other tools developed with the GATB-CORE API.

6. GATB helps you as a NGS developer

The GATB C++ library gives you the opportunity to quickly develop new NGS tools that fit your needs.

<u>Major facts about the GATB C++ library</u>

- Object Oriented Design
- Simple and powerful graph API
- Simple and powerful multithreading model
- ► HDF5 usage for data storage
- ► Fully documented with numerous code samples

TakeABreak Detects inversion breakpoints without a reference genome by looking for fixed size topological patterns in the de Bruijn graph

Complete test suite

Publications

G. Rizk, D. Lavenier, R. Chikhi, **DSK: k-mer counting with very low memory usage**, Bioinformatics, 2013 Mar 1;29(5):652-3

R. Chikhi, G. Rizk. Space-efficient and exact de Bruijn graph representation based on a Bloom filter, Algorithms for Molecular Biology 2013, 8:22

G. Collet, G. Rizk, R. Chikhi, D. Lavenier, Minia on Raspberry Pi, assembling a 100 Mbp genome on a Credit Card Sized Computer, Poster at the JOBIM conference, 2013 Jul 1-4 (Toulouse) Best poster award.



K.I Salikhov, G. Sacomoto, G. Kucherov, Using Cascading Bloom Filters to Improve the Memory Usage for de Brujin Graphs, Algorithms in Bioinformatics, Lecture Notes in Computer Science, Volume 8126, 2013, pp 364-376

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Proprietary licencing for software editors or services providers is currently being studied.

For more details on GATB:

http://gatb.inria.fr

