# Accelerating Genome Analysis Using New Algorithms and Hardware Designs

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**Preferred Networks** 

16 December 2019





## SAFARI Research Group

31 © = 1 Professor, 2 Lecturers & Senior Researchers, 3 Senior Researchers, 12 PhD Students, 3 Masters, 8 Interns, 2 Admins



Think BIG, Aim HIGH!

## Switzerland, Zurich, ETH Zurich, CS



#### Professor Mutlu's Bio

#### Onur Mutlu

- Professor @ ETH Zurich CS, since September'15, started May'16
- □ Strecker Professor @ Carnegie Mellon University ECE (CS), 2009-2016, 2016-...
- PhD from UT-Austin, worked @ Google, VMware, Microsoft Research, Intel, AMD
- https://people.inf.ethz.ch/omutlu/
- omutlu@gmail.com (Best way to reach me)
- Publications: <a href="https://people.inf.ethz.ch/omutlu/projects.htm">https://people.inf.ethz.ch/omutlu/projects.htm</a>

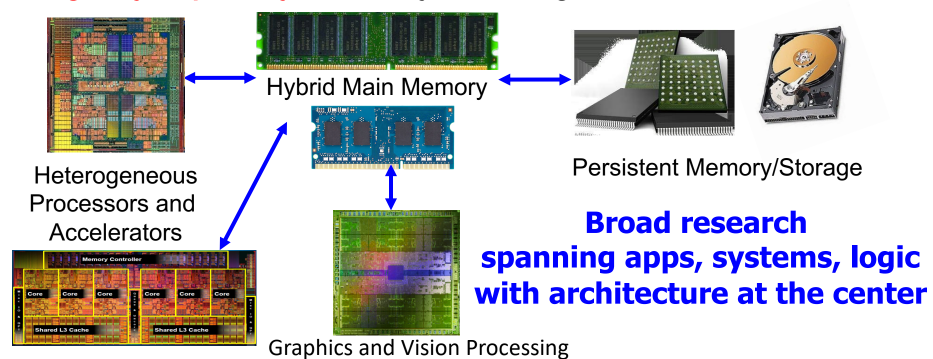
#### Research, Education, Consulting in

- Computer architecture and systems, bioinformatics
- Memory and storage systems, emerging technologies
- Many-core systems, heterogeneous systems, core design
- Interconnects
- Hardware/software interaction and co-design (PL, OS, Architecture)
- Predictable and QoS-aware systems
- Hardware fault tolerance and security
- Algorithms and architectures for genome analysis
- ...

#### Current Research Focus Areas

#### Research Focus: Computer architecture, HW/SW, security, bioinformatics

- Memory and storage (DRAM, flash, emerging), interconnects, security
- Heterogeneous & parallel systems, GPUs, systems for data analytics
- System/architecture interaction, new execution models, new interfaces
- Energy efficiency, fault tolerance, hardware security, performance
- Genome sequence analysis & assembly algorithms and architectures
- Biologically inspired systems & system design for bio/medicine



## Openings @ SAFARI

 We are hiring enthusiastic and motivated students and researchers at all levels.

Join us now:

safari.ethz.ch/apply





This talk is NOT about how to analyze biological data using available tools.

- Why Genome Analysis?
- What is Genome Analysis?
- How we Map Reads?
- What Makes Read Mapper Slow?
- Algorithmic & Hardware Acceleration
  - Seed Filtering Technique
  - Pre-alignment Filtering Technique
  - Read Alignment Acceleration
- Where is Read Mapping Going Next?

- Why Genome Analysis?
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## Why Genome Analysis? Why Bother?

- Personalized Medicine.
- Genome-Wide Association Studies (GWASs).
- City-Scale Microbiome Profiling.

**...** 

#### 1-Personalized Medicine

Nan-Byo Difficult + Illness

Coined in 1972 by the Japanese Ministry of Labor, Health, and Welfare.

https://www.nanbyo-research.jp/nanbyo



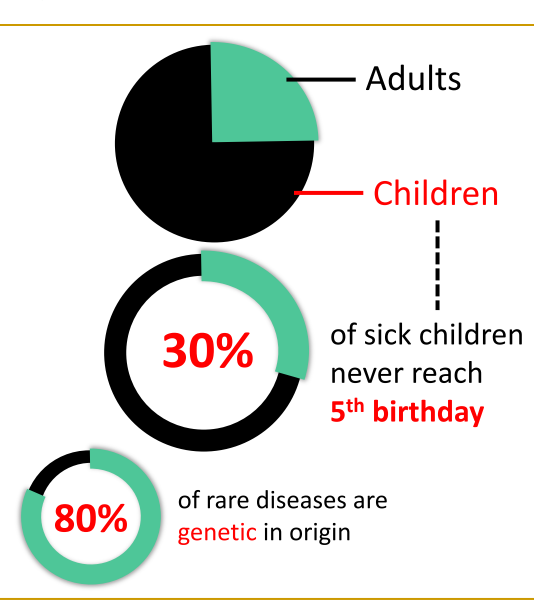
#### 1-Personalized Medicine



1 in 17 people

in the world have a rare disease

That's 350 Million



## Rare Diseases in Japan

"We don't know exactly how many people in Japan have a rare disease, which is why we want to design the rare disease platform to be as comprehensive as possible. There are thousands of rare diseases. So even though the number of patients with each disease is very small, there are many people who have one. Out of 20 of your friends, for example, one will have a rare disease," explains Matsuda.



Prof. Fumihiko Matsuda, Director of the Center for Genomic Medicine, Kyoto University

https://www.nanbyo-research.jp/feature/43/japan%E2%80%99s-rare-disease-database-expedites-more-effective-research

## Personalized Medicine in Japan

## European Journal of Human Genetics

Policy Open Access | Published: 05 July 2017

## Japan's initiative on rare and undiagnosed diseases (IRUD): towards an end to the diagnostic odyssey

Takeya Adachi ⊡, Kazuo Kawamura, Yoshihiko Furusawa, Yuji Nishizaki, Noriaki Imanishi, Senkei Umehara ⊡, Kazuo Izumi ⊡ & Makoto Suematsu

European Journal of Human Genetics 25, 1025–1028(2017) | Cite this article





> 600 million

JPY annually



#### Personalized Medicine in UK

npj Genomic Medicine

www.nature.com/npjgenmed

NPJ Genom Med. 2018; 3: 10.

PMCID: PMC5884823

Published online 2018 Apr 4. doi: <u>10.1038/s41525-018-0049-4</u>

PMID: <u>29644095</u>

## Rapid whole-genome sequencing decreases infant morbidity and cost of hospitalization

Lauge Farnaes, #1,2 Amber Hildreth, #1,2 Nathaly M. Sweeney, #1,2 Michelle M. Clark, 1 Shimul Chowdhury, 1

Shareef Nahas, 1 Julie A. Cakici, 1 Wendy Benson, 1 Robert H. Kaplan, 3 Richard Kronick, 4 Matthew N. Bainbridge, 1

Jennifer Friedman, 1,2,5 Jeffrey J. Gold, 1,5 Yan Ding, 1 Narayanan Veeraraghavan, 1 David Dimmock, 1 and

Stephen F. Kingsmore 1



reduced inpatient cost by \$100,000-\$300,000

National Institute for Health Research

"From 2019, all seriously ill children in UK will be offered whole genome sequencing as part of their care"

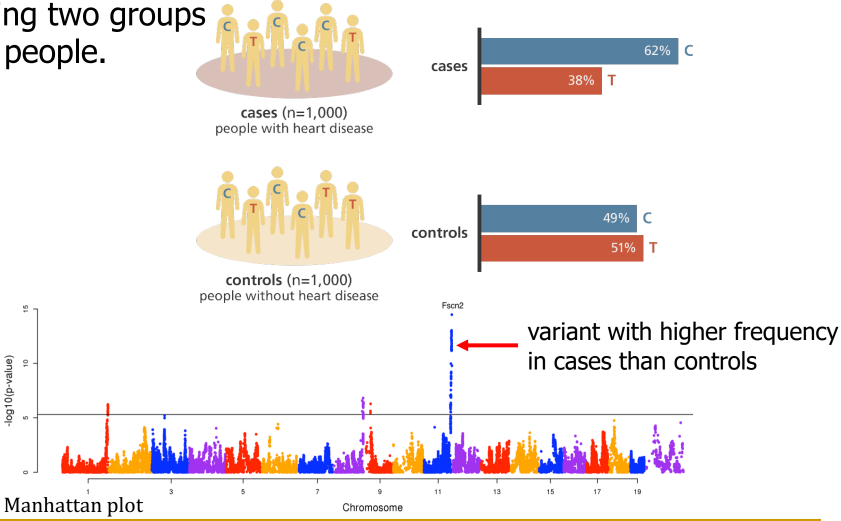
SAFARI

Farnaes+, "Rapid whole-genome sequencing decreases infant morbidity and cost of hospitalization", NPJ Genom Med. 2018

## 2-Genome-Wide Association Study (GWAS)

Detecting genetic variants associated with phenotypes

using two groups of people.



### Finding SNPs Associated with Complex Trait

SNP1	SNP2	<b>Blood Pressure</b>
ACATGCCGACATT	CATAGGCC	180
ACATGCCGACATT	CATAAGCC	175
ACATGCCGACATT	CATAGGCC	170
ACATGCCGACATT	CATAAGCC	165
ACATGCCGACATT	CATAGGCC	160
ACATGCCGACATT	CATAGGCC	145
ACATGCCGACATT	CATAAGCC	140
ACATGCCGACATT	CATAAGCC	130
ACATGTCGACATTT	CATAGGCC	120
ACATGTCGACATTT	CATAAGCC	120
ACATGTCGACATTT	CATAGGCC	115
ACATGTCGACATTT	CATAAGCC	110
ACATGTCGACATTT	CATAGGCC	110
ACATGTCGACATTT	CATAAGCC	110
ACATGTCGACATT1	CATAGGCC	105
ACATGTCGACATT1	CATA AGCC	100

Eleazar Eskin: Discovering the Causal Variants Involved in GWAS Studies, CGSI 2018, UCLA

Different

individuals

## Mirror Phenotypes of 593 Kb CNVs



**AUTISM** 

Weiss, *N Eng J Med* 2008 Deletion of 593 kb



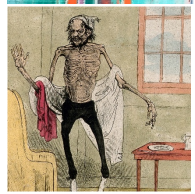
#### **SCHIZOPHRENIA**

McCarthy, *Nat Genet* 2009 Duplication of 593 kb



**OBESITY** 

Walters, *Nature* 2010 Deletion of 593 kb



#### **UNDERWEIGHT**

Jacquemont, *Nature* 2011 Duplication of 593 kb



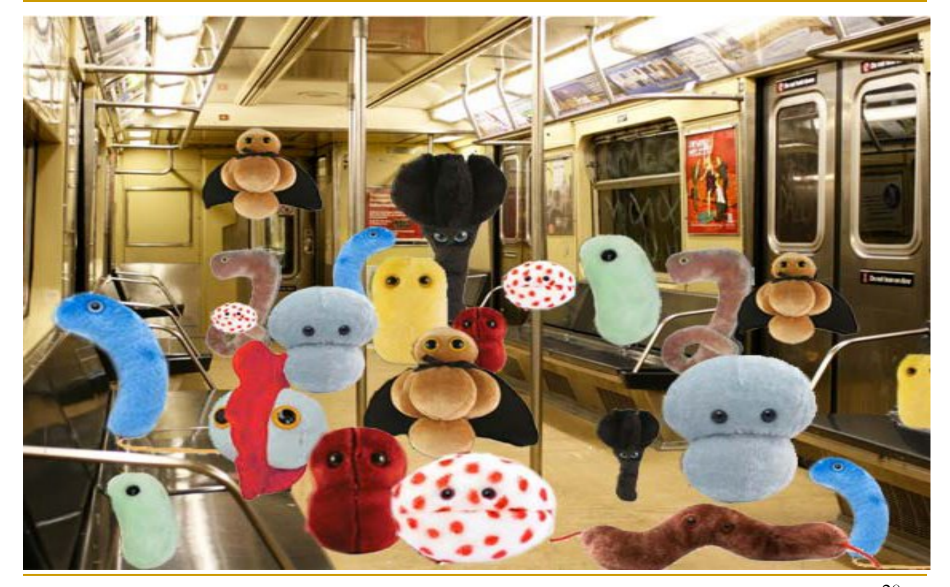
Deletion in the short arm of chromosome 16 (16p11.2)



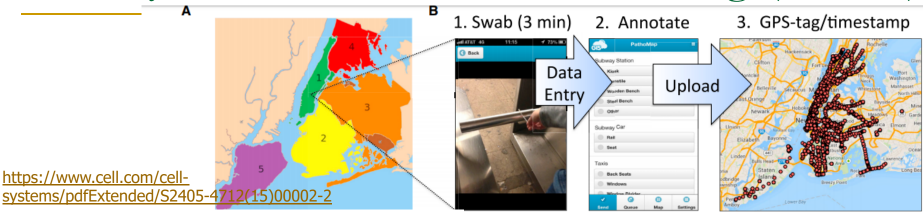
Duplication in the short arm of chromosome 16 (16p11.2)



## 3- City-Scale Microbiome Profiling



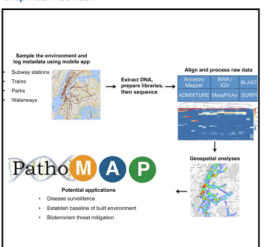
## 3- City-Scale Microbiome Profiling (cont'd)



#### Cell Systems

#### Geospatial Resolution of Human and Bacterial Diversity with City-Scale Metagenomics

Graphical Abstract



#### Authors

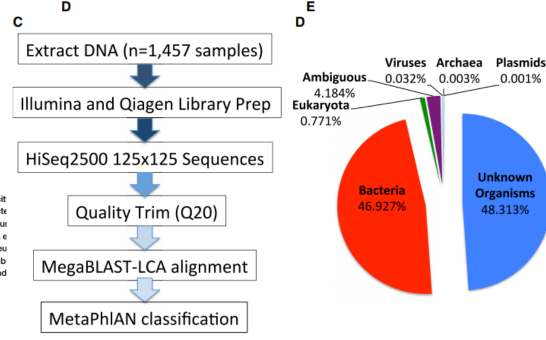
Ebrahim Afshinnekoo, Cem Meydan, ..., Shawn Levy, Christopher E. Mason

#### Correspondence

chm2042@med.cornell.edu

#### In Brief

Afshinnekoo et al. describe a cit molecular profile of DNA collecte city's subway system, public sur and one waterway. These data e baseline analysis of bacterial, eu and aracheal organisms in the b environment of mass transit and life.





(A) The tive boroughs of NYC include (1) Manhattan (green), (2) Brooklyn (yellow), (3) Queens (orange), (4) Bronx (red), (5) Staten Island (lavender).

(B) The collection from the 466 subway stations of NYC across the 24 subway lines involved three main steps: (1) collection with Copan Elution swabs, (2) data entry into the database, and (3) uploading of the data. An image is shown of the current collection database, taken from <a href="http://pathomap.giscloud.com">http://pathomap.giscloud.com</a>. (C) Workflow for sample DNA extraction, library preparation, sequencing, quality trimming of the FASTQ files, and alignment with MegaBLAST and MetaPhlAn to

## Plague in New York Subway System?

### Plague (Yersinia Pestis)



#### What Is It?

Published: December, 2018

Plague is caused by Yersinia pestis bacteria. It can be a life-threatening infection if not treated promptly. Plague has caused several major epidemics in Europe and Asia over the last 2,000 years. Plague has most famously been called "the Black Death" because it can cause skin sores that form black scabs. A plague epidemic in the 14th century killed more than one-third of the population of Europe within a few years. In some cities, up to 75% of the population died within days, with fever and swollen skin sores.

## Plague in New York Subway System?

## Plague (Yersi₁<sup>®</sup>

#### What Is It?

Published: December, 2018

Plague is caused by Yersinia treated promptly. Plague has last 2,000 years. Plague has cause skin sores that form b than one-third of the popul the population died within

## The New Hork Times Bubonic Plague in the Subway System? Don't Worry About It



In October, riders were not deterred after reports that an Ebola-infected man had ridden the subway just before he fell ill. Robert Stolarik for The New York Times

https://www.nytimes.com/2015/02/07/nyregion/bubonic-plague-in-the-subway-system-dont-worry-about-it.html

The findings of Yersinia Pestis in the subway received wide coverage in the lay press, causing some alarm among New York residents

#### Failure of Bioinformatics



data. Rob Knight, a professor in the department of pediatrics at the University of California, San Diego, calls this type of error "a failure of bioinformatics," in that Mason had assumed the gene fragments were unique to the pathogens, when in fact they can also be detected in other

Living in a microbial world

**Charles Schmidt** 

*Nature Biotechnology*, **volume 35**, pages401–403 (2017)

https://www.nature.com/articles/nbt.3868

There is a critical need for **fast** and **accurate** genome analysis.

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## Genome Analysis



## No machine can read the *entire* content of a genome



.....

## How Long is DNA?



## The Genetic Similarity Between Species



Human ~ Human 99.9%



Human ~ Chimpanzee 96%



Human ~ Cat 90%



Human ∼ Cow 80%



Human ∼ Banana 50-60%

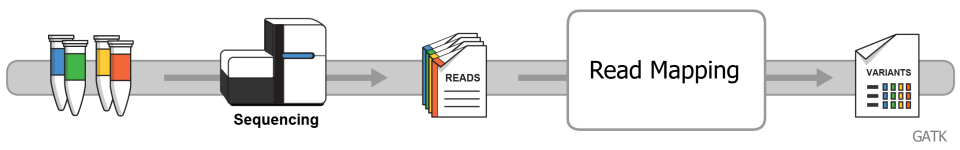
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## Cracking the 1st Human Genome Sequence

■ **1990-2003:** The Human Genome Project (HGP) provides a complete and accurate sequence of all **DNA base pairs** that make up the human genome and finds 20,000 to 25,000 human genes.



## Vast Improvement in Sequencing



**CCCCCTATATATACGTACTAGTACGT** 

ACGACTTTAGTACGTACGT TATATACGTACTAGTACGT

ACGTACG CCCCTACGTA
TATATACGTACTAGTACGT

ACGACTTTAGTACGTACGT TATATATACGTACTAAAGTACGT TATATATACGTACTAGTACGT

ACG TTTTTAAAACGTA
TATATATACGTACTACGT

**ACGAC GGGGAGTACGT** 



1x10<sup>12</sup> bases\*



44 hours\*



<1000 \$

\* NovaSeq 6000

## High-Throughput Sequencers



Illumina MiSeq



Illumina NovaSeq 6000



Pacific Biosciences Sequel II



Pacific Biosciences RS II





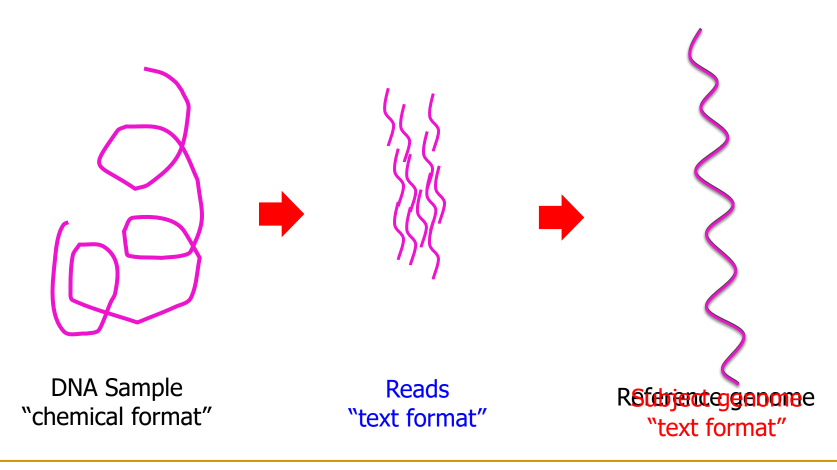
Oxford Nanopore MinION



... and more! All produce data with different properties.

### Genome Analysis

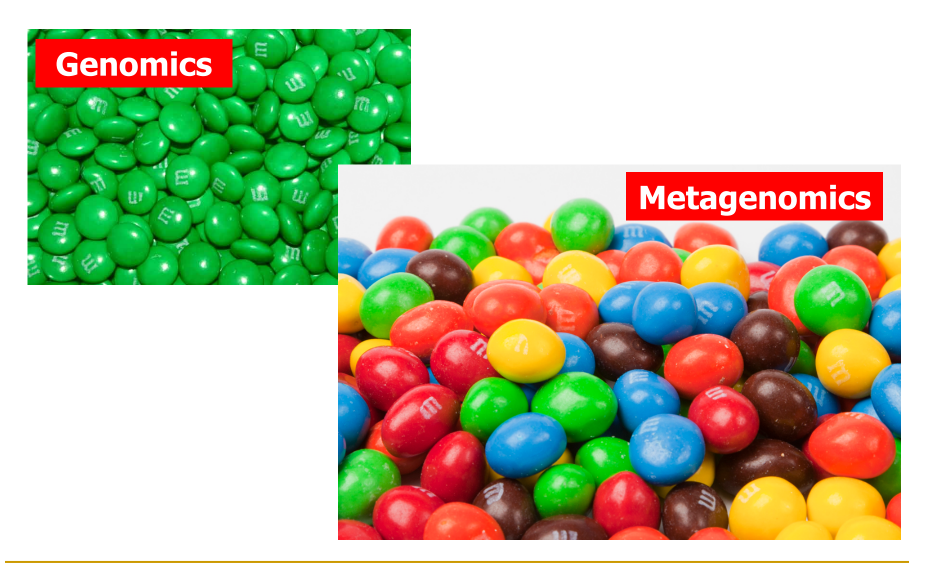
Map reads to a known reference genome with some minor differences allowed



## Metagenomics Analysis

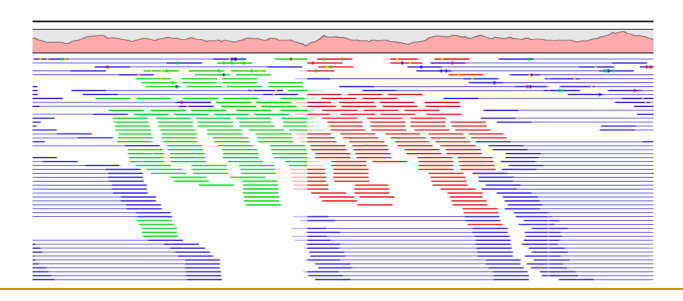
Reads from different unknown donors at sequencing time are mapped to many known reference genomes genetic material recovered directly from environmental Reads Reference samples "text format" Database

## Genomics vs. Metagenomics



## Challenges in Read Mapping

- Need to find many mappings of each read
- Need to tolerate small variances/errors in each read
- Need to map each read very fast (i.e., performance is important, life critical in some cases)



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## Read Mapping: A Brute Force Algorithm

#### Reference



Read

## Very Expensive! $O(m^2kn)$

m: read length

k: no. of reads

*n*: reference genome length

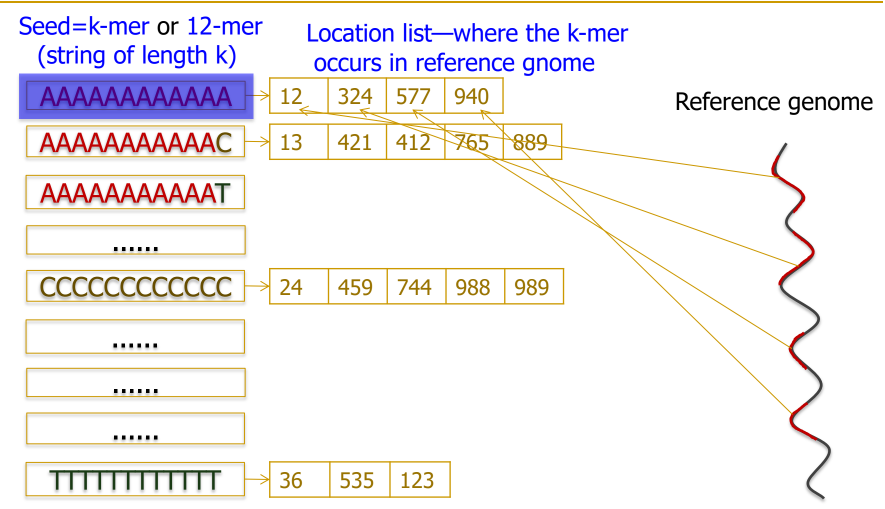
## Similar to Searching Yellow Pages!

Step 1: Get the page number from the book's index using a small portion of the name (e.g., 1st letter).

Step 2: Retrieve the page(s).

Step 3: Look for the full name & get the phone number.

## Step 1: Indexing the Reference Genome



We can query the table with substrings from reads to quickly find a list of possible mapping locations



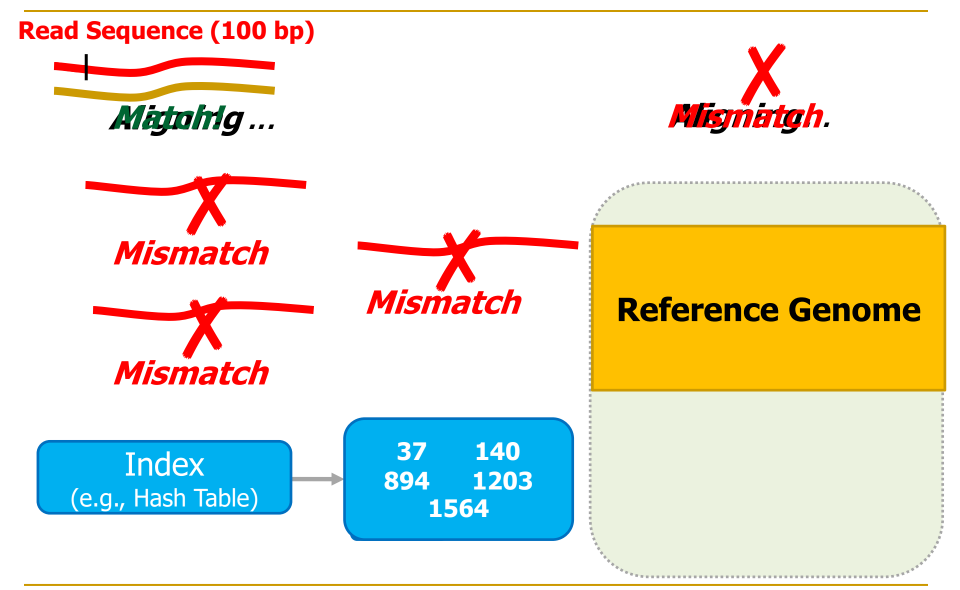
## Genome Index Properties

- The index is built only once for each reference.
- Seeds can be overlapping, non-overlapping, spaced, adjacent, non-adjacent, minimizers, compressed, ...

Tool	Version	Index Size*	Indexing Time		
mrFAST	2.2.5	16.5 GB	20.00 min		
minimap2	0.12.7	7.2 GB	3.33 min		
BWA-MEM	0.7.17	4.7 GB	49.96 min		

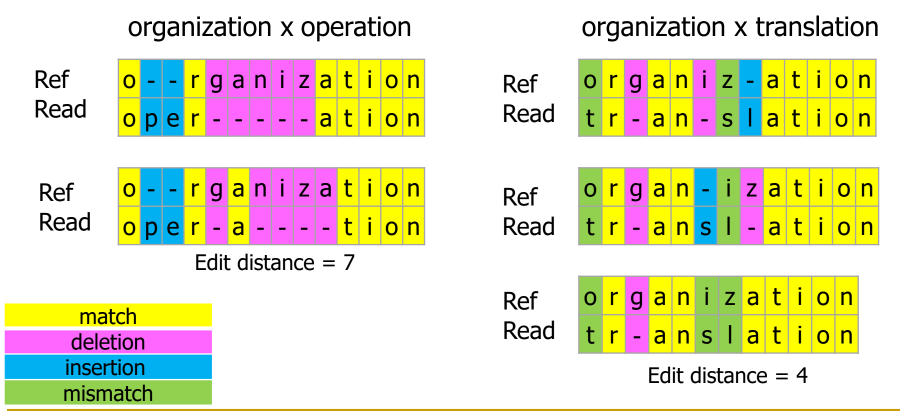
<sup>\*</sup>Human genome = 3.2 GB

## Step 2: Query the Index Using Read Seeds



## Step 3: Read Alignment (Verification)

 <u>Edit distance</u> is defined as the minimum number of edits (i.e. insertions, deletions, or substitutions) needed to make the read exactly match the reference segment.



### An Example of Hash Table Based Mappers

- + Guaranteed to find all mappings → very sensitive
- + Can tolerate up to e errors



https://github.com/BilkentCompGen/mrfast

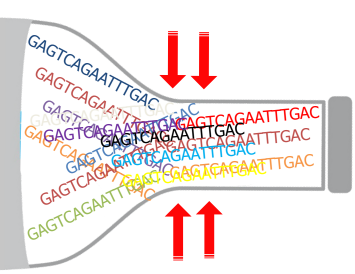
# Personalized copy number and segmental duplication maps using next-generation sequencing

Can Alkan<sup>1,2</sup>, Jeffrey M Kidd<sup>1</sup>, Tomas Marques-Bonet<sup>1,3</sup>, Gozde Aksay<sup>1</sup>, Francesca Antonacci<sup>1</sup>, Fereydoun Hormozdiari<sup>4</sup>, Jacob O Kitzman<sup>1</sup>, Carl Baker<sup>1</sup>, Maika Malig<sup>1</sup>, Onur Mutlu<sup>5</sup>, S Cenk Sahinalp<sup>4</sup>, Richard A Gibbs<sup>6</sup> & Evan E Eichler<sup>1,2</sup>

## Bottlenecked in Read Alignment!!

378 Million bases/minute

Read Sequencing \*\*



2 Million bases/minute

Read Mapping\*

### 150x slower

<sup>\*</sup> BWA-MEM

<sup>\*\*</sup> NovaSeq 6000, MinION

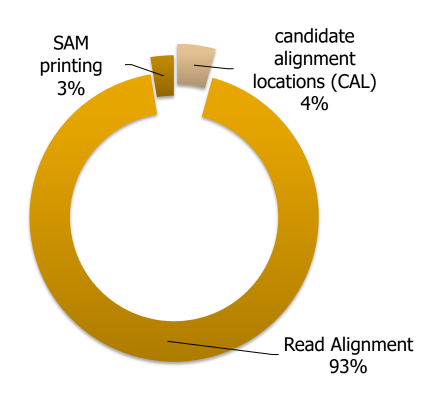
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## What Makes Read Mapper Slow?

Key Observation # 1

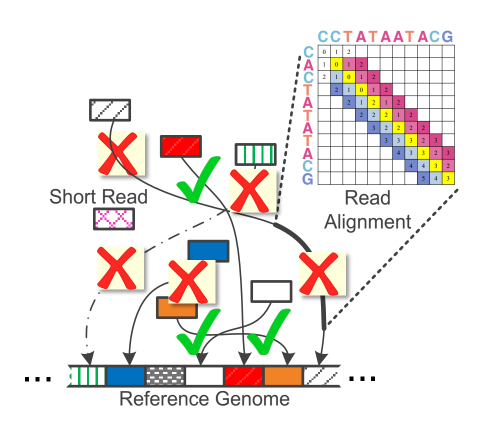
93%
of the read mapper's
execution time is spent
in read alignment.



Alser et al, Bioinformatics (2017)

## What Makes Read Mapper Slow? (cont'd)

### Key Observation # 2



98% of candidate locations have high dissimilarity with a given read.

Cheng et al, BMC bioinformatics (2015) Xin et al, BMC genomics (2013)

## What Makes Read Mapper Slow? (cont'd)

### Key Observation # 3

Quadratic-time dynamicprogramming algorithm WHY?!

Enumerating all possible prefixes

NETHERLANDS x SWITZERLAND

NETHERLANDS x S

NETHERLANDS x SW

**NETHERLANDS x SWI** 

**NETHERLANDS x SWIT** 

**NETHERLANDS x SWITZ** 

**NETHERLANDS x SWITZE** 

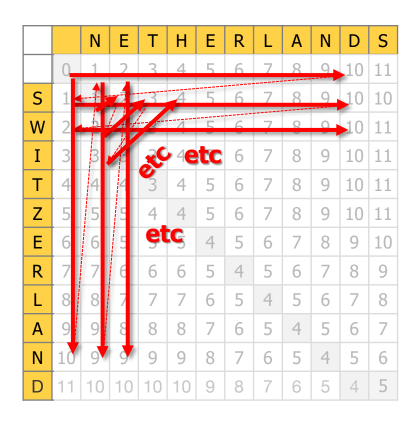
NETHERLANDS x SWITZER

**NETHERLANDS x SWITZERL** 

NETHERLANDS x SWITZERLA

**NETHERLANDS x SWITZERLAN** 

NETHERLANDS x SWITZERLAND



## What Makes Read Mapper Slow? (cont'd)

### Key Observation # 3

Quadratic-time dynamicprogramming algorithm

Enumerating all possible prefixes

 Data dependencies limit the computation parallelism

Processing row (or column) after another

Entire matrix is computed even though strings can be dissimilar.

			_	_		_	_		_		_	
		N	Е	Т	Н	Е	R	L	Α	N	D	S
	0	1	2	3	4	5	6	7	8	9	10	11
S	1	1	2	3	4	5	6	7	8	9	10	10
W	2	2	2	3	4	5	6	7	8	9	10	11
Ι	3	3	3	3	4	5	6	7	8	9	10	11
Т	4	4	4	3	4	5	6	7	8	9	10	11
Z	5	5	5	4	4	5	6	7	8	9	10	11
Е	6	6	5	5	5	4	5	6	7	8	9	10
R	7	7	6	6	6	5	4	5	6	7	8	9
L	8	8	7	7	7	6	5	4	5	6	7	8
Α	9	9	8	8	8	7	6	5	4	5	6	7
N	10	9	9	9	9	8	7	6	5	4	5_	6
D	11	10	10	10	10	9	8	7	6	5	4	5

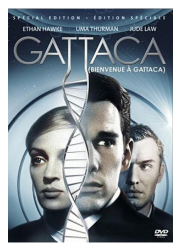
Number of differences is computed only at the backtraking step.

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### Our Goal

 Our goal is to significantly reduce the time spent on calculating the optimal alignment in genome analysis from hours to mere seconds using both new algorithms & hardware accelerators, given limited computational resources (i.e., personal computer or small hardware).







1997 2015

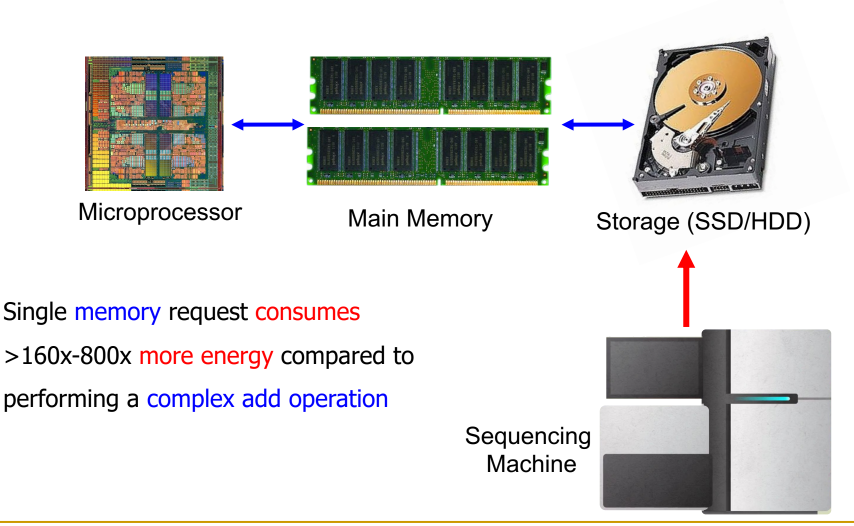
## Open Questions

How and where to enable

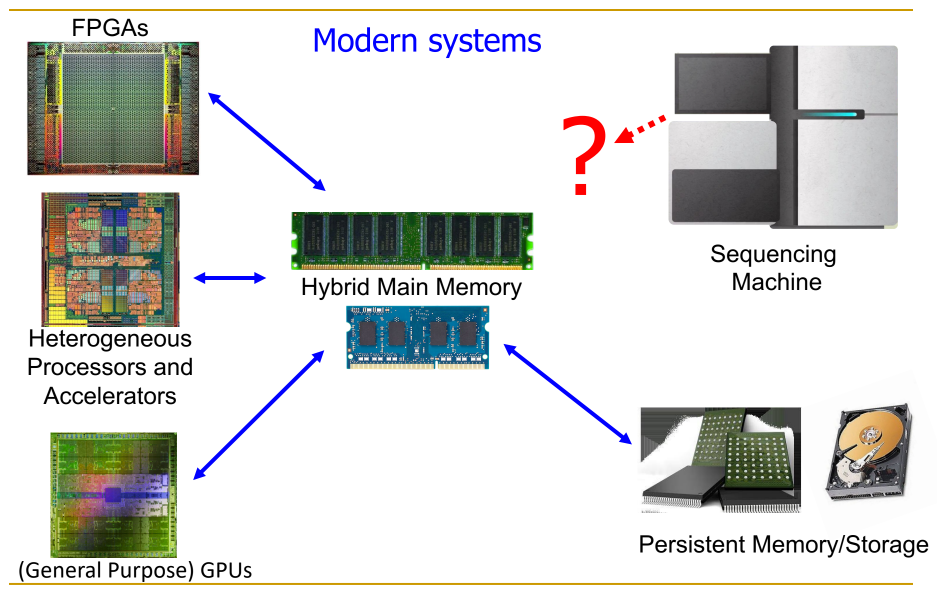
fast, accurate, cheap,

privacy-preserving, and exabyte scale analysis of genomic data?

## Pushing Towards New Architectures

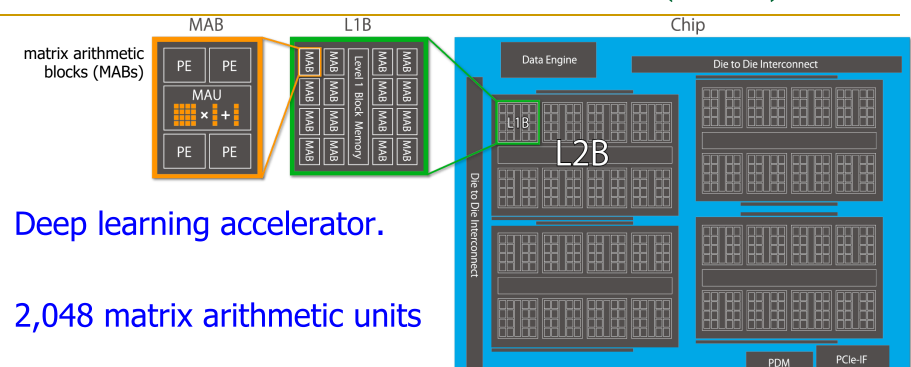


### Processing Genomic Data Where it Makes Sense



Most speedup comes from parallelism enabled by novel architectures and algorithms

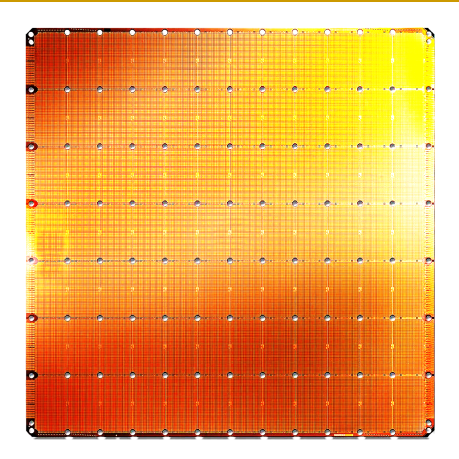
## Preferred Networks' MN-Core (2018)



Fabrication process	TSMC 12nm
Estimated power consumption (W)	500
Peak performance (TFLOPS)	32.8 (DP) / 131 (SP) / 524 (HP)
Estimated performance per watt (TFLOPS / W)	0.066 (DP) / 0.26 (SP) / 1.0 (HP)

(Notes) DP: double precision, SP: single precision, HP: half precision.

## Cerebras's Wafer Scale Engine (2019)



The largest ML accelerator chip

**400,000** cores



### **Cerebras WSE**

1.2 Trillion transistors 46,225 mm<sup>2</sup>

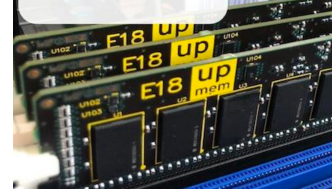
**Largest GPU** 

21.1 Billion transistors 815 mm<sup>2</sup>

https://www.cerebras.net/cerebras-wafer-scale-engine-why-we-need-big-chips-for-deep-learning/

## UPMEM Processing-in-DRAM Engine (2019)

- Processing in DRAM Engine
- Includes standard DIMM modules, with a large number of DPU processors combined with DRAM chips.
- Replaces standard DIMMs
  - DDR4 R-DIMM modules
    - 8GB+128 DPUs (16 PIM chips)
    - Standard 2x-nm DRAM process



Large amounts of compute & memory bandwidth





https://www.anandtech.com/show/14750/hot-chips-31-analysis-inmemory-processing-by-upmem

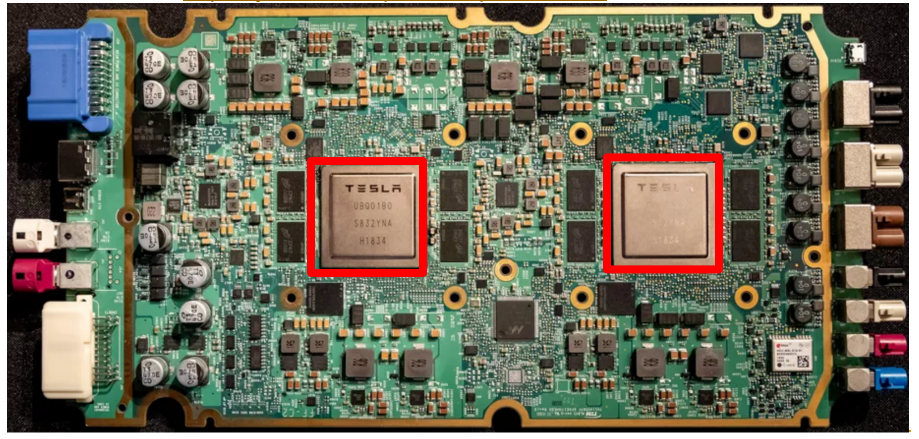
## TESLA Full Self-Driving Computer (2019)

ML accelerator: 260 mm<sup>2</sup>, 6 billion transistors,
 600 GFLOPS GPU, 12 ARM 2.2 GHz CPUs.



Two redundant chips for better safety.

https://youtu.be/Ucp0TTmvqOE?t=4236



### Illumina + Edico Genome

**PRESS RELEASE** 

# Illumina Acquires Edico Genome to Accelerate Genomic Data Analysis

Edico's DRAGEN® Bio-IT Platform Delivers Faster, Streamlined Output for Next-Generation Sequencing

SAN DIEGO--(BUSINESS WIRE)--May 15, 2018-- Illumina, Inc. (NASDAQ: ILMN) today announced that it acquired Edico Genome, the leading provider of data analysis acceleration solutions for next-generation sequencing (NGS). Edico Genome's DRAGEN® Bio-IT Platform (DRAGEN) uses field programmable gate array (FPGA) technology in conjunction with proprietary software algorithms to reduce both data footprint and time to results.

https://www.illumina.com/company/news-center/press-releases/2018/2349147.html

### Illumina + PacBio

PRESS RELEASE

Illumina to Acquire Pacific
Biosciences for Approximately \$1.2
Billion, Broadening Access to Long-Read Sequencing and Accelerating
Scientific Discovery

- Brings Together Highly Accurate Short- and Long-Read Sequencing Technologies, Paving the Path to a More Perfect View of a Genome
- Pacific Biosciences' Recent Advances with its Sequel SMRT<sup>®</sup> Technology, Combined with Illumina's Infrastructure, will Expand Biological Discovery and Clinical Insight
- Long-Read Sequencing Market Opportunity Expected to Grow to \$2.5B by 2022

SAN DIEGO & MENLO PARK, Calif .-- (BUSINESS WIRE) -- Nov. 1, 2018 -- Illumina, Inc. (NASDAQ: ILMN)

https://www.illumina.com/company/news-center/press-releases/press-release-details.html?newsid=2374913

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## Ongoing Directions

### Seed Filtering Technique:

- Goal: Reducing the number of seed (k-mer) locations.
  - Heuristic (limits the number of mapping locations for each seed).
  - Supports exact matches only.

### Pre-alignment Filtering Technique:

- Goal: Reducing the number of invalid mappings (>E).
  - Supports both exact and inexact matches.
  - Provides some falsely-accepted mappings.

### Read Alignment Acceleration:

- Goal: Performing read alignment at scale.
  - Limits the numeric range of each cell in the DP table and hence supports limited scoring function.
  - May not support backtracking step due to random memory accesses.

## An Example of Ongoing Directions

Read Sequence (100 bp) 2) Pre-Alignment Filtering... Match! 3) Rapid Alignment Allgbing...



**Reference Genome** 

1) Seed Filtering ...





Hash Table

37 140 894 1203 1564

## Ongoing Directions

### Seed Filtering Technique:

- Goal: Reducing the number of seed (k-mer) locations.
  - Heuristic (limits the number of mapping locations for each seed).
  - Supports exact matches only.

### Pre-alignment Filtering Technique:

- Goal: Reducing the number of invalid mappings (>E).
  - Supports both exact and inexact matches.
  - Provides some falsely-accepted mappings.

### Read Alignment Acceleration:

- Goal: Performing read alignment at scale.
  - Limits the numeric range of each cell in the DP table and hence supports limited scoring function.
  - May not support backtracking step due to random memory accesses.

### FastHASH

- Goal: Reducing the number of seed (k-mer) locations.
  - Heuristic (limits the number of mapping locations for each seed).
  - Supports exact matches only.

Xin et al. BMC Genomics 2013, **14**(Suppl 1):S13 http://www.biomedcentral.com/1471-2164/14/S1/S13



### **PROCEEDINGS**

**Open Access** 

### Accelerating read mapping with FastHASH

Hongyi Xin<sup>1</sup>, Donghyuk Lee<sup>1</sup>, Farhad Hormozdiari<sup>2</sup>, Samihan Yedkar<sup>1</sup>, Onur Mutlu<sup>1\*</sup>, Can Alkan<sup>3\*</sup>

From The Eleventh Asia Pacific Bioinformatics Conference (APBC 2013) Vancouver, Canada. 21-24 January 2013



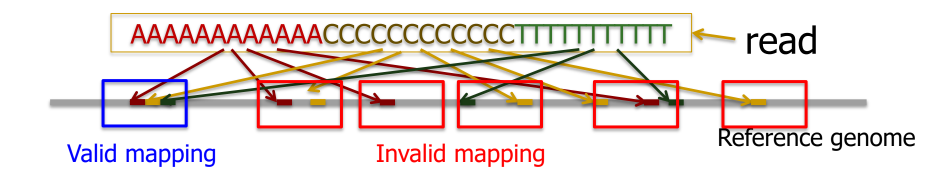
### FastHASH Conclusion

- Problem: Existing read mappers perform poorly in mapping billions of short reads to the reference genome, in the presence of errors
- Observation: Most of the verification calculations are unnecessary → filter them out
- Key Idea: To reduce the cost of unnecessary verification
  - Select Cheap and Adjacent k-mers.
- Key Result: FastHASH obtains up to 19x speedup over the state-of-the-art mapper without losing valid mappings

## Key Observations

### Observation 1 (Adjacent k-mers)

- Key insight: Adjacent k-mers in the read should also be adjacent in the reference genome
- Key idea: 1) sort the location list based on their number of locations and 2) search for adjacent locations in the k-mers' location lists



## Key Observations

### Observation 1 (Adjacent k-mers)

- Key insight: Adjacent k-mers in the read should also be adjacent in the reference genome
- Key idea: 1) sort the location list based on their number of locations and 2) search for adjacent locations in the k-mers' location lists

### Observation 2 (Cheap k-mers)

- Key insight: Some k-mers are cheaper to verify than others because they have shorter location lists (they occur less frequently in the reference genome)
- Key Idea: Read mapper can choose the cheapest k-mers and verify their locations

### Cheap K-mer Selection

occurrence threshold = 500read GGGTATGGCTAG AAGGTTGAGAGC CTTAGGCTTACC AAGCTCAATTIC CCTCCTTAATTIT TCCTCTTAAGAA 326 338 350 376 388 1231 Location 151 1470 4414 2 loc. 2 loc. 9219 Number of Locations 4 loc. Cheapest 3 k-mers 1K loc. 2K loc. 1K loc. Expensive 3 k-mers Previous work needs FastHASH verifies only: to verify: 8 locations 3004 locations

### More on FastHASH

- Download source code and try for yourself
  - Download link to FastHASH

Xin et al. BMC Genomics 2013, **14**(Suppl 1):S13 http://www.biomedcentral.com/1471-2164/14/S1/S13



### **PROCEEDINGS**

**Open Access** 

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From The Eleventh Asia Pacific Bioinformatics Conference (APBC 2013) Vancouver, Canada. 21-24 January 2013



## Ongoing Directions

### Seed Filtering Technique:

- Goal: Reducing the number of seed (k-mer) locations.
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### Read Alignment Acceleration:

- Goal: Performing read alignment at scale.
  - Limits the numeric range of each cell in the DP table and hence supports limited scoring function.
  - May not support backtracking step due to random memory accesses.

# Pre-alignment Filtering Technique

### Read Alignment is expensive

Our goal is to reduce the need for dynamic programming algorithms

# Ideal Filtering Algorithm

Step 2 Step 3 Query Read the Alignment Index

- Filter out most of incorrect mappings.
- 2. Preserve all correct mappings.
- 3. Do it quickly.

# Shifted Hamming Distance (SHD)

Bioinformatics, 31(10), 2015, 1553-1560

doi: 10.1093/bioinformatics/btu856

Advance Access Publication Date: 10 January 2015

**Original Paper** 



Sequence analysis

# Shifted Hamming distance: a fast and accurate SIMD-friendly filter to accelerate alignment verification in read mapping

Hongyi Xin<sup>1,\*</sup>, John Greth<sup>2</sup>, John Emmons<sup>2</sup>, Gennady Pekhimenko<sup>1</sup>, Carl Kingsford<sup>3</sup>, Can Alkan<sup>4,\*</sup> and Onur Mutlu<sup>2,\*</sup>

Xin+, "Shifted Hamming Distance: A Fast and Accurate SIMD-friendly Filter to Accelerate Alignment Verification in Read Mapping", Bioinformatics 2015.

# Shifted Hamming Distance

### Key observation:

If two strings differ by E edits, then every bp match can be aligned in at most 2E shifts.

### Key idea:

- Compute "Shifted Hamming Distance": AND of 2E Hamming Distances of two strings, to identify invalid mappings
  - Uses bit-parallel operations that nicely map to SIMD instructions

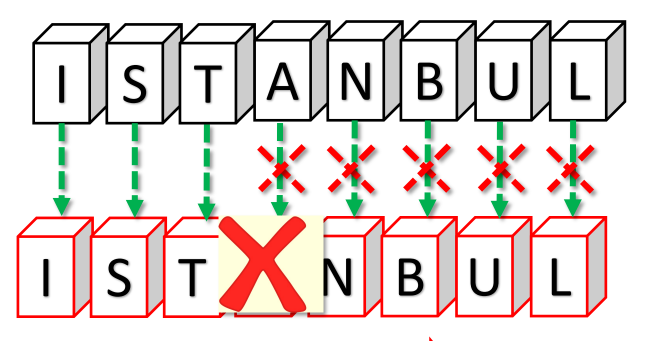
### Key result:

- SHD is 3x faster than SeqAn (the best implementation of Gene Myers' bit-vector algorithm), with only a 7% false positive rate
- The fastest CPU-based filtering (pre-alignment) mechanism

# Hamming Distance ( $\Sigma \oplus$ )

3 matches 5 mismatches

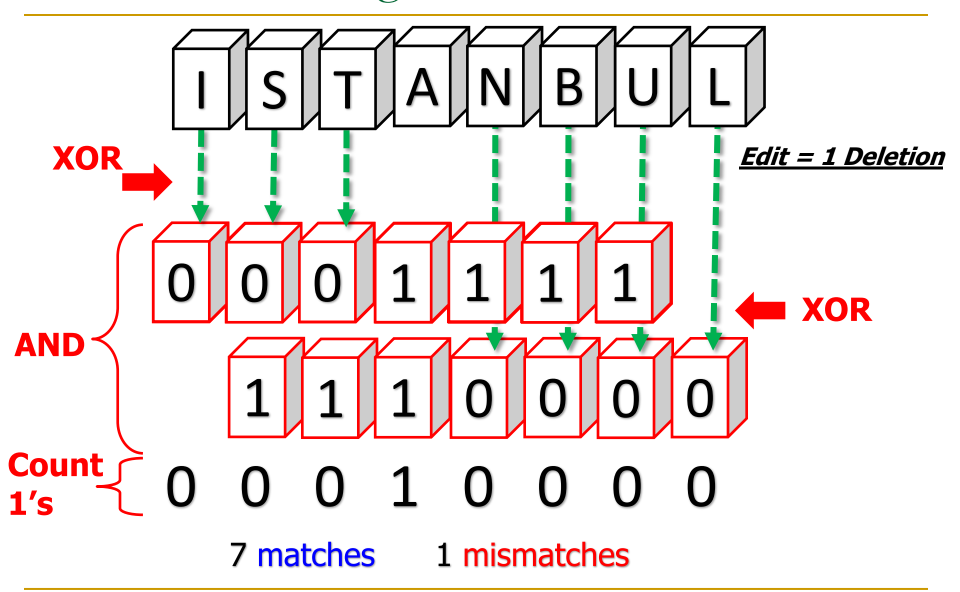
<u>Edit = 1 Deletion</u>





To cancel the effect of a deletion, we need to shift in the *right* direction

## Shifted Hamming Distance



### More on SHD

- Download and test for yourself
- https://github.com/CMU-SAFARI/Shifted-Hamming-Distance

Bioinformatics, 31(10), 2015, 1553–1560 doi: 10.1093/bioinformatics/btu856 Advance Access Publication Date: 10 January 2015

Original Paper



Sequence analysis

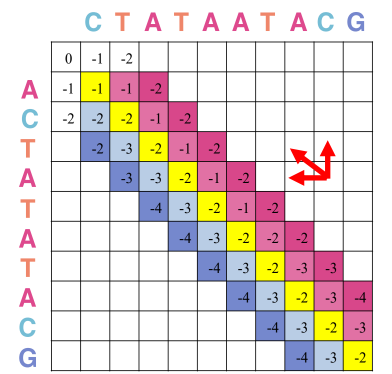
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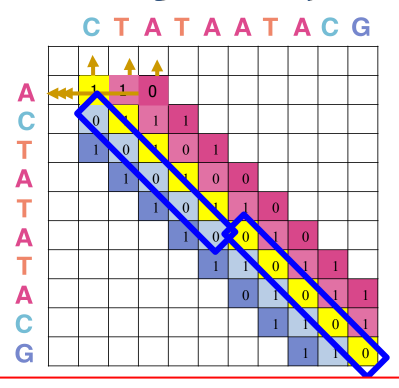
# Can we Speed up SHD Computations?

# Alignment Matrix vs. Neighborhood Map





#### **Neighborhood Map**



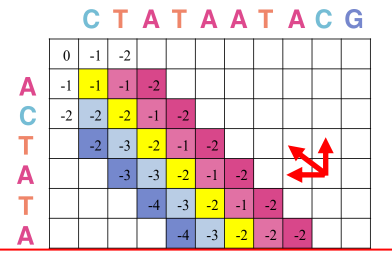
Our goal to track the diagonally consecutive matches in the neighborhood map.

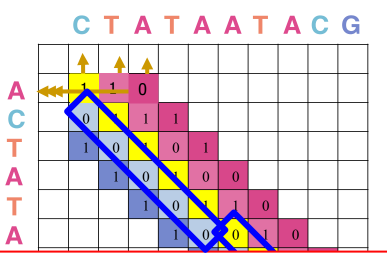
86

# Alignment Matrix vs. Neighborhood Map

#### Needleman-Wunsch

#### Neighborhood Map

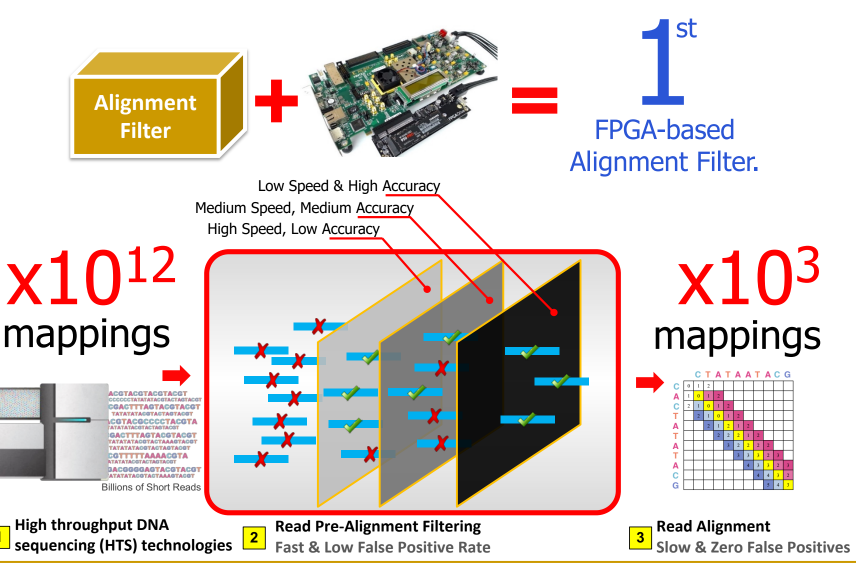




Independent vectors can be processed in parallel using hardware technologies



# Our Solution: GateKeeper



### GateKeeper Walkthrough

Generate 2E+1 masks

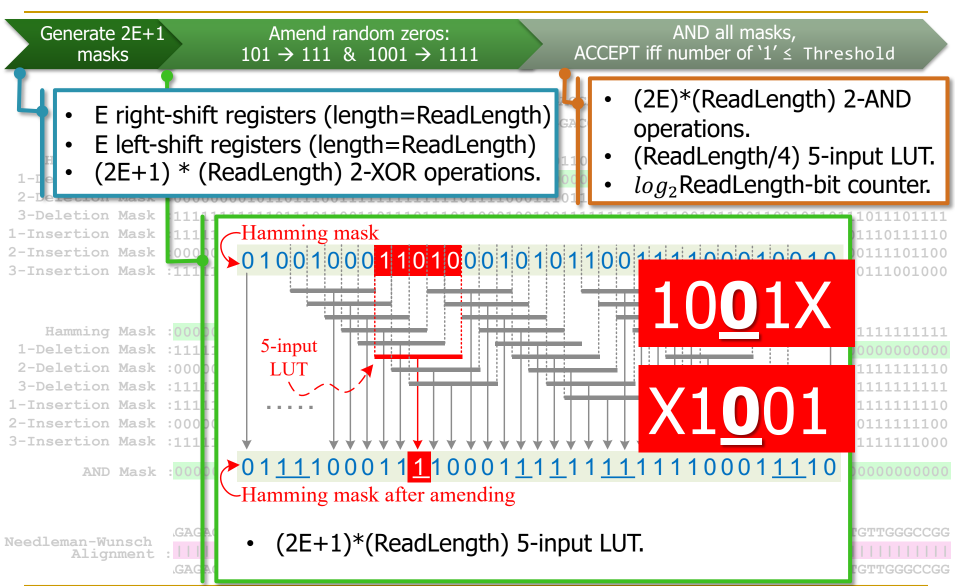
Amend random zeros:  $101 \rightarrow 111 \& 1001 \rightarrow 1111$ 

AND all masks, ACCEPT iff number of `1' ≤ Threshold

```
:GAGAGAGATATTTAGTGTTGCAGCACTACAACACAAAAGAGGACCAACTTACGTGTCTAAAAAGGGGGAACATTGTTGGGCCGGA
 Reference
   GAGAGAGATAGTTAGTGTTGCAGCCACTACAACACAAAAGAGGACCAACTTACGTGTCTAAAAGGGGAGACATTGTTGGGCCGG
--- Masks after amendment ---
'TTAGTGTTGCAG-CACTACAACACAAAAGAGGACCAACTTACGTGTCTAAAAGGGGGGAACATTGTTGGGCCGG
```

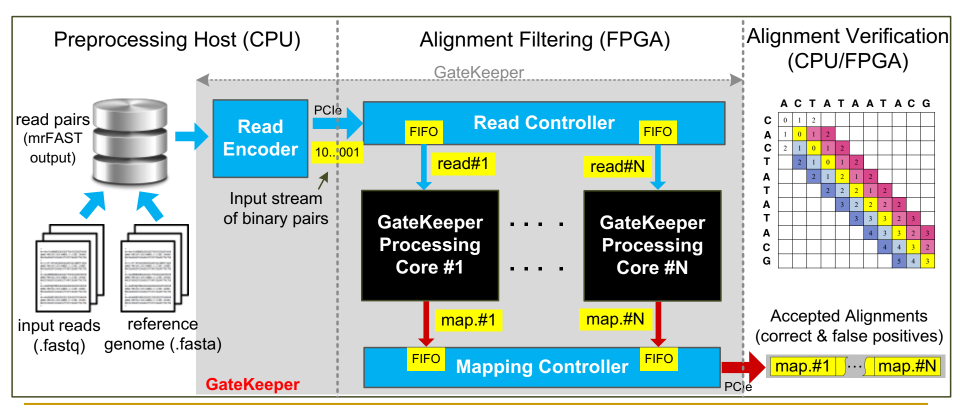
Needleman-Wunsch Alignment

# GateKeeper Walkthrough (cont'd)

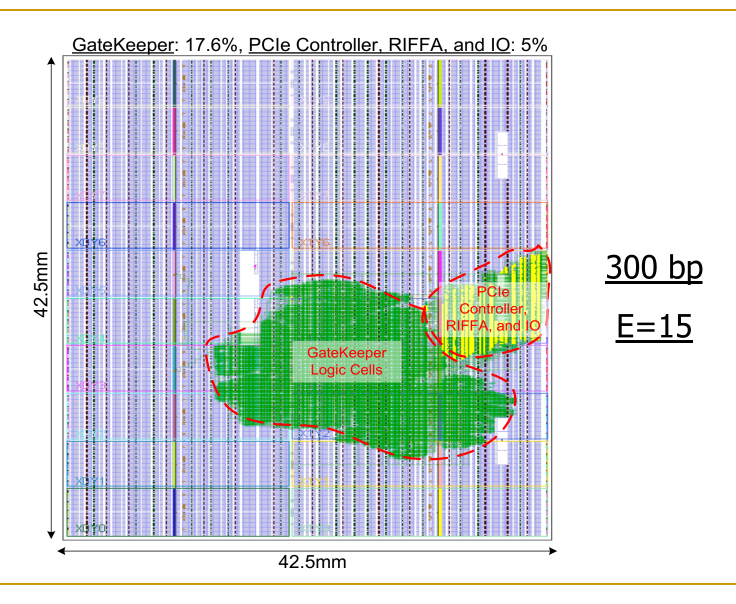


### GateKeeper Accelerator Architecture

- Maximum data throughput =~13.3 billion bases/sec
- Can examine 8 (300 bp) or 16 (100 bp) mappings concurrently at 250 MHz
- Occupies 50% (100 bp) to 91% (300 bp) of the FPGA slice LUTs and registers



# FPGA Chip Layout



# GateKeeper: Speed & Accuracy Results

# 90x-130x faster filter

than SHD (Xin et al., 2015) and the Adjacency Filter (Xin et al., 2013)

# 4x lower false accept rate

than the Adjacency Filter (Xin et al., 2013)

# 10x speedup in read mapping

with the addition of GateKeeper to the mrFAST mapper (Alkan et al., 2009)

# Freely available online

github.com/BilkentCompGen/GateKeeper

# GateKeeper Conclusions

- FPGA-based pre-alignment greatly speeds up read mapping
  - 10x speedup of a state-of-the-art mapper (mrFAST)

- FPGA-based pre-alignment can be integrated with the sequencer
  - It can help to hide the complexity and details of the FPGA
  - Enables real-time filtering while sequencing

### More on GateKeeper

Download and test for yourself
 <a href="https://github.com/BilkentCompGen/GateKeeper">https://github.com/BilkentCompGen/GateKeeper</a>

Alser+, "GateKeeper: A New Hardware Architecture for Accelerating Pre-Alignment in DNA Short Read Mapping", Bioinformatics, 2017.

### Sequence analysis

# GateKeeper: A New Hardware Architecture for Accelerating Pre-Alignment in DNA Short Read Mapping

Mohammed Alser<sup>1,\*</sup>, Hasan Hassan<sup>2</sup>, Hongyi Xin<sup>3</sup>, Oğuz Ergin<sup>2</sup>, Onur Mutlu<sup>4,\*</sup>, and Can Alkan<sup>1,\*</sup>

Can we do better? Scalability?

# Shouji (障子)

Bioinformatics, 2019, 1–9

doi: 10.1093/bioinformatics/btz234

Advance Access Publication Date: 28 March 2019

Original Paper



#### Sequence alignment

# Shouji: a fast and efficient pre-alignment filter for sequence alignment

Mohammed Alser<sup>1,2,3,\*</sup>, Hasan Hassan<sup>1</sup>, Akash Kumar<sup>2</sup>, Onur Mutlu<sup>1,3,\*</sup> and Can Alkan<sup>3,\*</sup>

<sup>1</sup>Computer Science Department, ETH Zürich, Zürich 8092, Switzerland, <sup>2</sup>Chair for Processor Design, Center For Advancing Electronics Dresden, Institute of Computer Engineering, Technische Universität Dresden, 01062 Dresden, Germany and <sup>3</sup>Computer Engineering Department, Bilkent University, 06800 Ankara, Turkey

\*To whom correspondence should be addressed.

Associate Editor: Inanc Birol

Received on September 13, 2018; revised on February 27, 2019; editorial decision on March 7, 2019; accepted on March 27, 2019

Alser+, "Shouji: a fast and efficient pre-alignment filter for sequence alignment", Bioinformatics 2019,

https://doi.org/10.1093/bioinformatics/btz234



# Shouji

### Key observation:

- Correct alignment always includes long identical subsequences.
- Processing the entire mapping at once is ineffective for hardware design.

### Key idea:

 Use overlapping sliding window approach to quickly and accurately find all long segments of consecutive zeros.

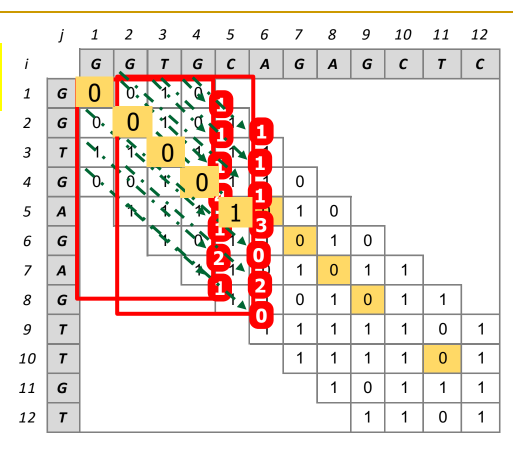
### Key result:

- Shouji on FPGA is at least 160x faster than its CPU implementation.
- Shouji accelerates best-performing CPU read aligner Edlib
  (Bioinformatics 2017) by up to 18.8x using 16 filtering units that
  work in parallel.
- Shouji is 2.4x to 467x more accurate than GateKeeper (Bioinformatics 2017) and SHD (Bioinformatics 2015).

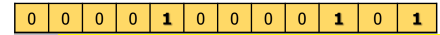
# Shouji Walkthrough

Building the Neighborhood Map

Finding all common subsequences (diagonal segments of consecutive zeros) shared between two given sequences.



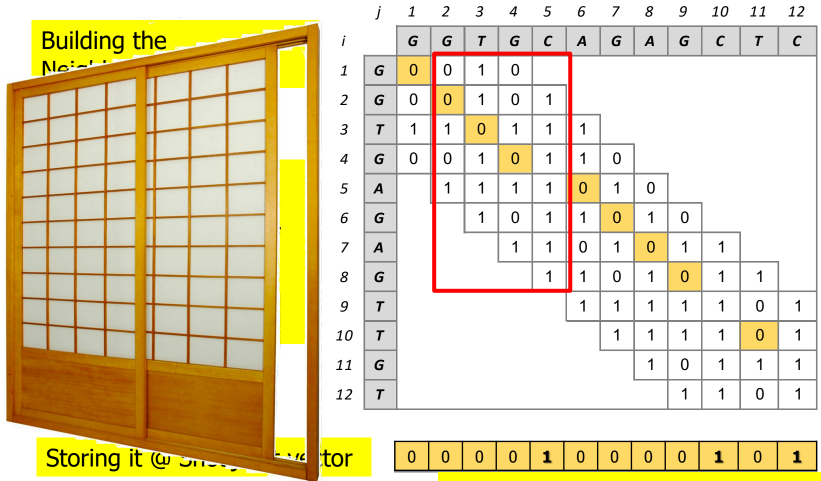
Storing it @ Shouji Bit-vector



ACCEPT iff number of '1' ≤ Threshold

Shouji: a fast and efficient pre-alignment filter for sequence alignment, *Bioinformatics* 2019, https://doi.org/10.1093/bioinformatics/btz234

# Shouji Walkthrough

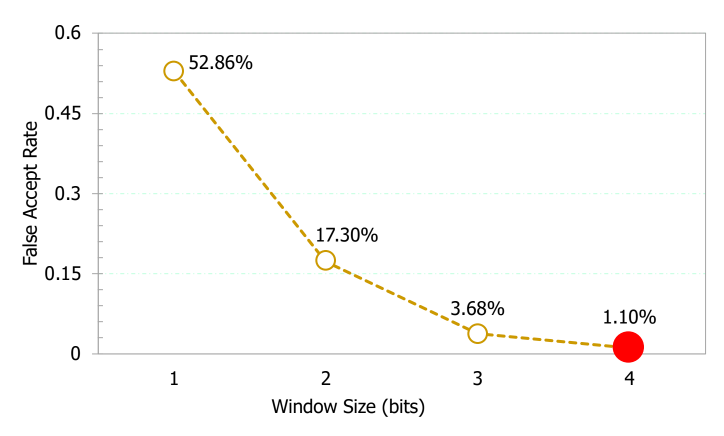


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Shouji: a fast and efficient pre-alignment filter for sequence alignment, *Bioinformatics* 2019, https://doi.org/10.1093/bioinformatics/btz234

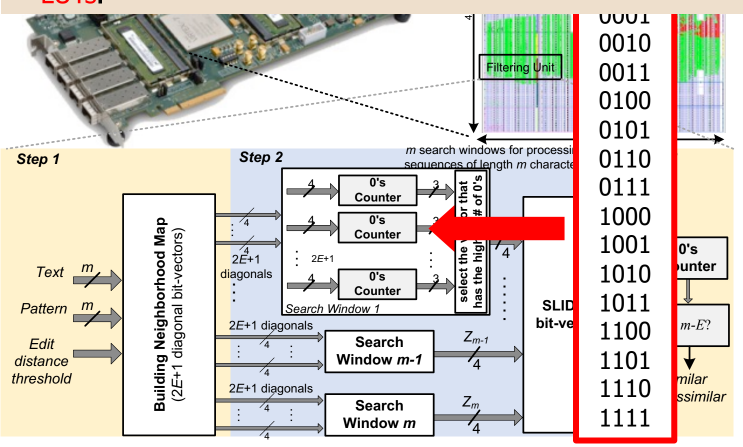
# Sliding Window Size

 The reason behind the selection of the window size is due to the minimal possible length of the identical subsequence that is a single match (e.g., such as `101').



# Hardware Implementation

 Counting is performed concurrently for all bit-vectors and all sliding windows in a single clock cycle using multiple 4-input LUTs.



## More on Shouji

#### Download and test for yourself

https://github.com/CMU-SAFARI/Shouji

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Original Paper



Sequence alignment

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Alser+, "Shouji: a fast and efficient pre-alignment filter for sequence alignment", Bioinformatics 2019,

https://doi.org/10.1093/bioinformatics/btz234

### SneakySnake: A Fast and Accurate Universal Genome Pre-Alignment Filter for CPUs, GPUs, and FPGAs

Mohammed Alser<sup>1,3</sup>, Taha Shahroodi<sup>1</sup>, Juan Gómez-Luna<sup>1</sup>, Can Alkan<sup>3</sup>, and Onur Mutlu<sup>1,2,3</sup>

<sup>1</sup>Department of Computer Science, ETH Zurich, Zurich 8006, Switzerland <sup>2</sup>Department of Electrical and Computer Engineering, Carnegie Mellon University, Pittsburgh 15213, PA, USA <sup>3</sup>Department of Computer Engineering, Bilkent University, Ankara 06800, Turkey

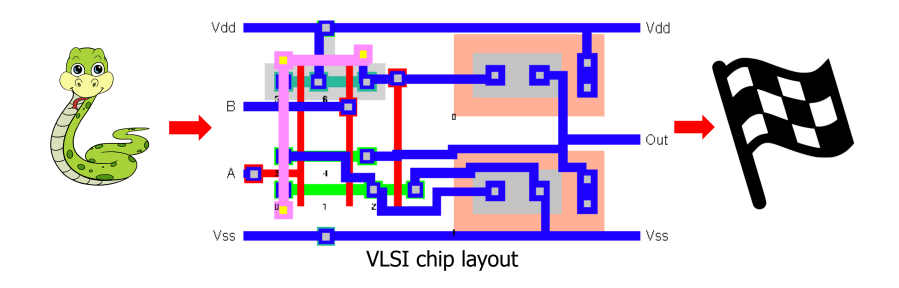
Alser + "SneakySnake: A Fast and Accurate Universal Genome Pre-Alignment Filter for CPUs, GPUs, and FPGAs." arXiv preprint (2019).

### Key observation:

Correct alignment is a sequence of non-overlapping long matches.

### Key idea:

 Approximate edit distance calculation is similar to Single Net Routing problem in VLSI chip.



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Correct alignment is a sequence of non-overlapping long matches.

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### Key result:

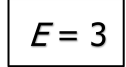
- SneakySnake is up to four orders of magnitude more accurate than Shouji (Bioinformatics'19) and GateKeeper (Bioinformatics'17).
- SneakySnake accelerates the state-of-the-art CPU-based sequence aligners, Edlib (Bioinformatics'17) and Parasail (BMC Bioinformatics'16), by up to 37.6× and 43.9× (>12× on average), respectively, without requiring hardware acceleration, and by up to 413× and 689× (>400× on average), respectively, using hardware acceleration.

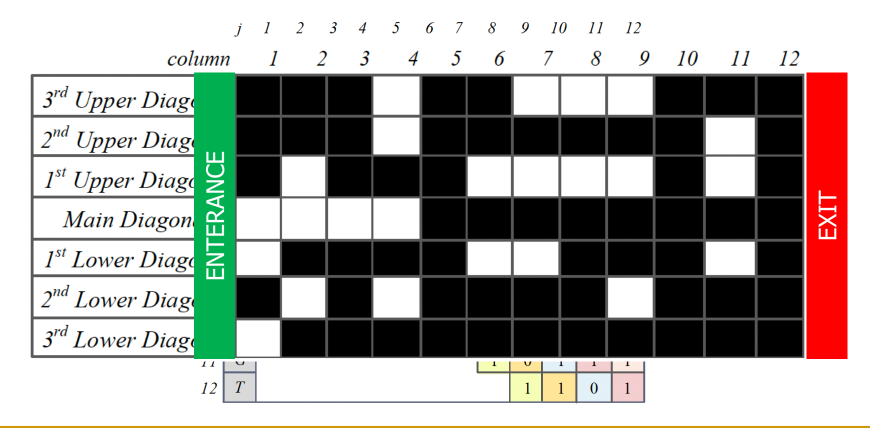
# SneakySnake Walkthrough

**Building Neighborhood Map** 

Finding the Optimal Routing Path

Examining the Snake Survival





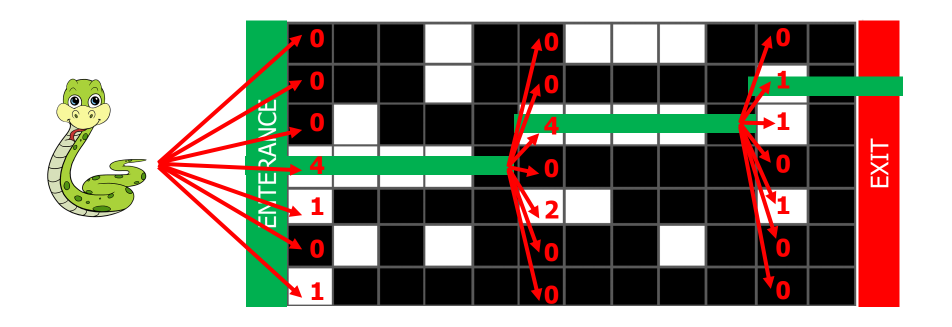
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# SneakySnake Walkthrough

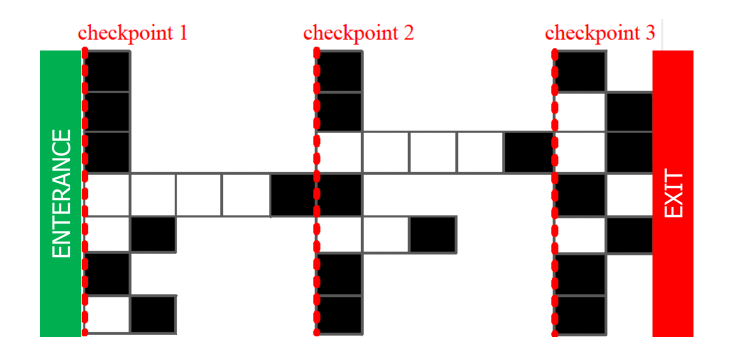
**Building Neighborhood Map** 

Finding the Routing Travel Path

Examining the Snake Survival

This is what you actually need to build and it can be done on-the-fly!





## FPGA Resource Analysis

 FPGA resource usage for a single filtering unit of GateKeeper, Shouji, and Snake-on-Chip for a sequence length of 100 and under different edit distance thresholds (E).

	<i>E</i> (bp)	Slice LUT	Slice Register	No. of Filtering Units
GateKeeper	2	0.39%	0.01%	16
	5	0.71%	0.01%	16
Shouji	2	0.69%	0.08%	16
	5	1.72%	0.16%	16
Snake-on-Chip	2	0.68%	0.16%	16
	5	1.42%	0.34%	16

### SneakySnake: A Fast and Accurate Universal Genome Pre-Alignment Filter for CPUs, GPUs, and FPGAs

Mohammed Alser<sup>1,3</sup>, Taha Shahroodi<sup>1</sup>, Juan Gómez-Luna<sup>1</sup>, Can Alkan<sup>3</sup>, and Onur Mutlu<sup>1,2,3</sup>

<sup>1</sup>Department of Computer Science, ETH Zurich, Zurich 8006, Switzerland <sup>2</sup>Department of Electrical and Computer Engineering, Carnegie Mellon University, Pittsburgh 15213, PA, USA <sup>3</sup>Department of Computer Engineering, Bilkent University, Ankara 06800, Turkey

### Download and test for CPU, GPU, and FPGA:

https://github.com/CMU-SAFARI/SneakySnake

Alser + "SneakySnake: A Fast and Accurate Universal Genome Pre-Alignment Filter for CPUs, GPUs, and FPGAs." arXiv preprint (2019).

# Read Mapping & Filtering

- Problem: Heavily bottlenecked by Data Movement
- Shouji performance limited by DRAM bandwidth [Alser+, Bioinformatics 2019]
- GateKeeper performance limited by DRAM bandwidth [Alser+, Bioinformatics 2017]
- Ditto for SHD [Xin+, Bioinformatics 2015]
- Solution: Processing-in-memory can alleviate the bottleneck

# Read Mapping & Filtering in Memory

We need to design mapping & filtering algorithms that fit processing-in-memory

#### GRIM-Filter

Jeremie S. Kim, Damla Senol Cali, Hongyi Xin, Donghyuk Lee, Saugata Ghose, Mohammed Alser, Hasan Hassan, Oguz Ergin, Can Alkan, and Onur Mutlu, "GRIM-Filter: Fast Seed Location Filtering in DNA Read Mapping Using Processing-in-Memory Technologies" to appear in <u>BMC Genomics</u>, 2018. Proceedings of the <u>16th Asia Pacific Bioinformatics Conference</u> (APBC), Yokohama, Japan, January 2018. arxiv.org Version (pdf)

## GRIM-Filter: Fast Seed Location Filtering in DNA Read Mapping Using Processing-in-Memory Technologies

Jeremie S. Kim<sup>1,6\*</sup>, Damla Senol Cali<sup>1</sup>, Hongyi Xin<sup>2</sup>, Donghyuk Lee<sup>3</sup>, Saugata Ghose<sup>1</sup>, Mohammed Alser<sup>4</sup>, Hasan Hassan<sup>6</sup>, Oguz Ergin<sup>5</sup>, Can Alkan<sup>\*4</sup>, and Onur Mutlu<sup>\*6,1</sup>

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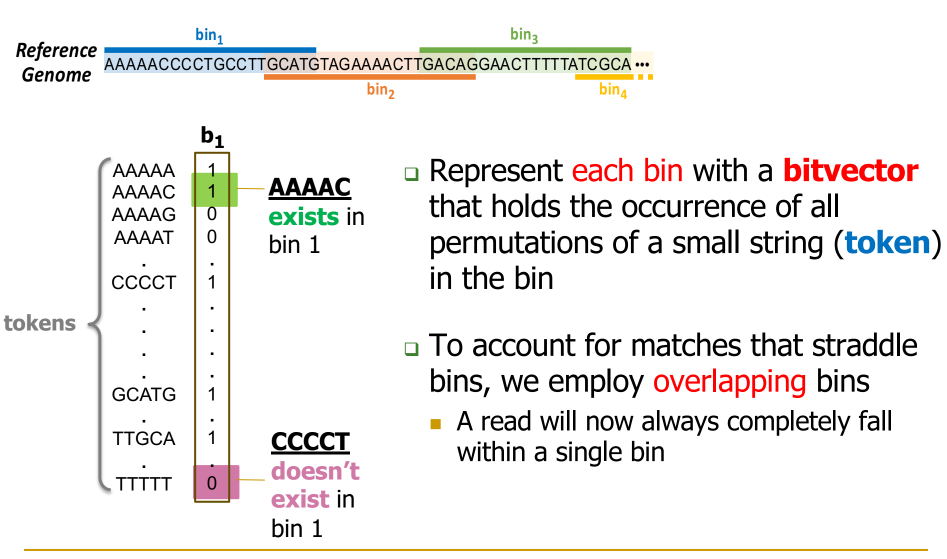
#### GRIM-Filter

- Key observation: FPGA and GPU accelerators are Heavily bottlenecked by Data Movement.
- Key idea: exploiting the high memory bandwidth and the logic layer of 3D-stacked memory to perform highly-parallel filtering in the DRAM chip itself.

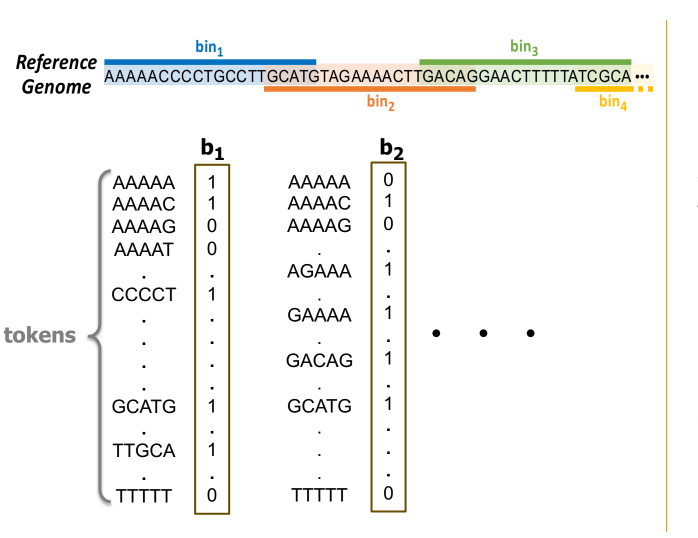
#### Key results:

- We propose an algorithm called GRIM-Filter
- GRIM-Filter with processing-in-memory is 1.8x-3.7x (2.1x on average) faster than FastHASH filter (BMC Genomics'13) across real data sets.
- GRIM-Filter has 5.6x-6.4x (6.0x on average) lower falsely accepted pairs than FastHASH filter (BMC Genomics'13) across real data sets.

#### **GRIM-Filter:** Bitvectors



#### **GRIM-Filter:** Bitvectors

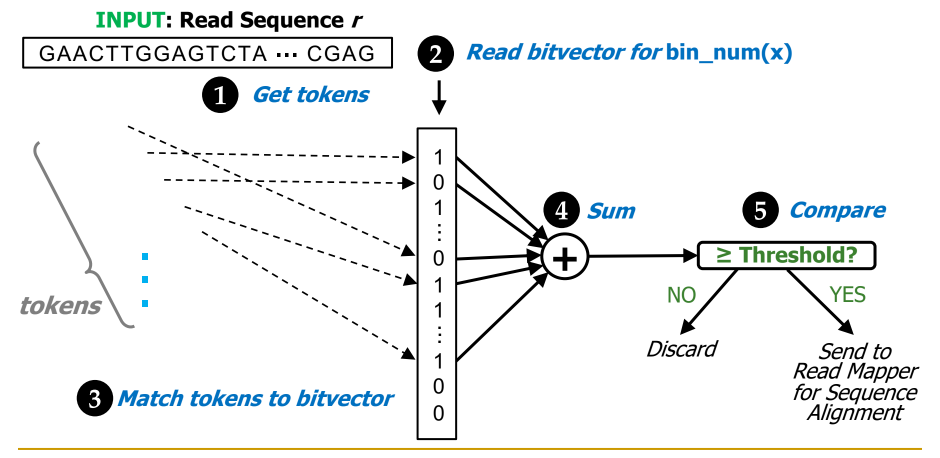


Storing all bitvectors requires  $\underline{4^n * t}$  bits in memory, where t = number of bins & n = token length.

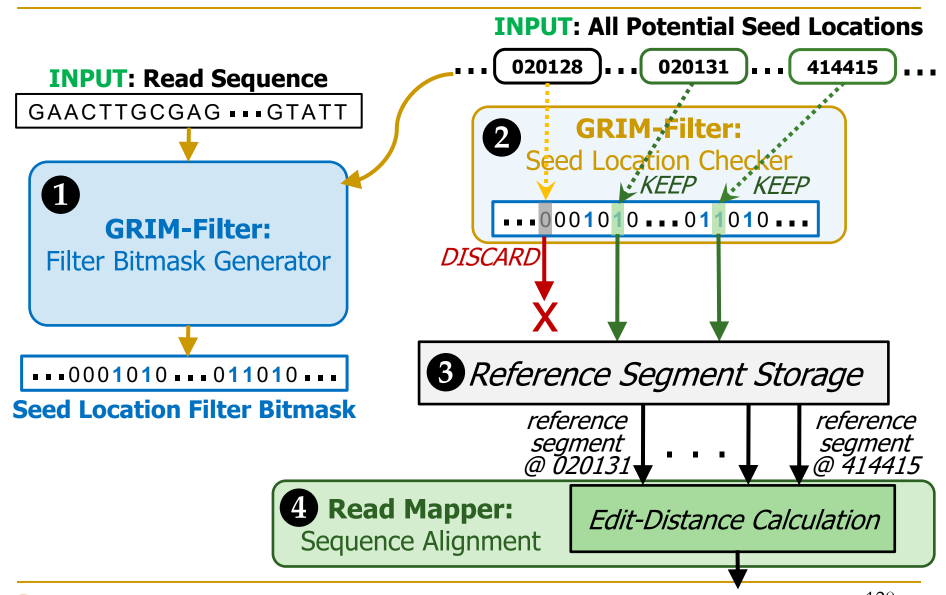
For **bin size** ~200, and **n** = 5, **memory footprint** ~3.8 GB

## GRIM-Filter: Checking a Bin

How GRIM-Filter determines whether to **discard** potential match locations in a given bin **prior** to alignment



### Integrating GRIM-Filter into a Read Mapper



SAFARI

**OUTPUT:** Correct Mappings

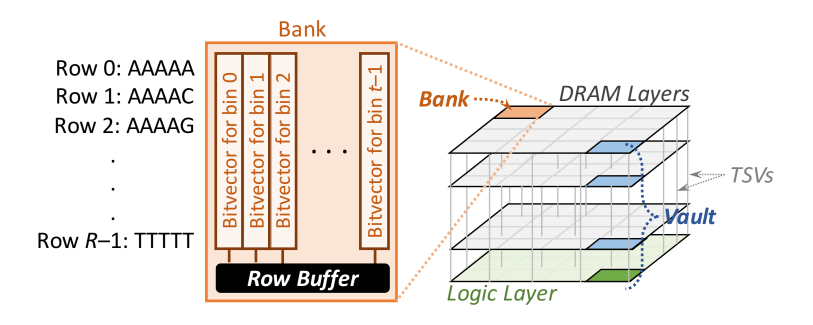
## Key Properties of GRIM-Filter

#### 1. Simple Operations:

- To check a given bin, find the sum of all bits corresponding to each token in the read
- Compare against threshold to determine whether to align
- 2. Highly Parallel: Each bin is operated on independently and there are many many bins
- 3. Memory Bound: Given the frequent accesses to the large bitvectors, we find that GRIM-Filter is memory bound

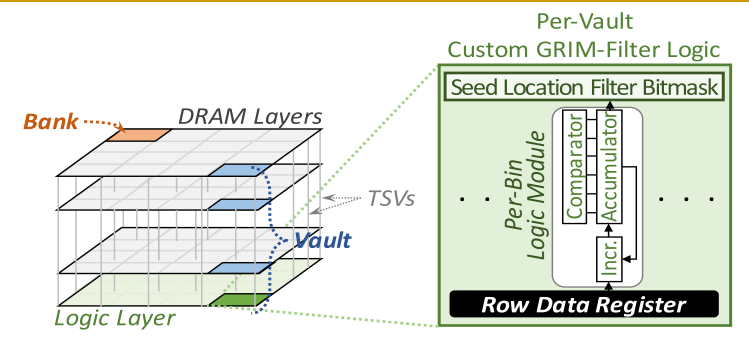
These properties together make GRIM-Filter a good algorithm to be run in 3D-Stacked DRAM

#### GRIM-Filter in 3D-Stacked DRAM



- Each DRAM layer is organized as an array of banks
  - A bank is an array of cells with a row buffer to transfer data
- The layout of bitvectors in a bank enables filtering many bins in parallel

#### GRIM-Filter in 3D-Stacked DRAM



- Customized logic for accumulation and comparison per genome segment
  - Low area overhead, simple implementation
  - For HBM2, we use 4096 incrementer LUTs, 7-bit counters, and comparators in logic layer

#### More on GRIM-Filter

Jeremie S. Kim, Damla Senol Cali, Hongyi Xin, Donghyuk Lee, Saugata Ghose, Mohammed Alser, Hasan Hassan, Oguz Ergin, Can Alkan, and Onur Mutlu, "GRIM-Filter: Fast Seed Location Filtering in DNA Read Mapping Using Processing-in-Memory Technologies" to appear in <u>BMC Genomics</u>, 2018.
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<u>arxiv.org Version (pdf)</u>

## GRIM-Filter: Fast Seed Location Filtering in DNA Read Mapping Using Processing-in-Memory Technologies

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#### GenCache

## GenCache: Leveraging In-Cache Operators for Efficient Sequence Alignment

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#### GenCache

 Key observation: State-of-the-art alignment accelerators are still bottlenecked by memory.

#### Key ideas:

- Performing in-cache alignment + pre-alignment filtering by enabling processing-in-cache using previous proposal, ComputeCache (HPCA'17).
- Using different Pre-alignment filters depending on the selected edit distance threshold.

#### Results:

- GenCache on CPU is 1.36x faster than GenAx (ISCA 2018).
   GenCache in cache is 5.26x faster than GenAx.
- GenCache chip has 16.4% higher area, 34.7% higher peak power, and 15% higher average power than GenAx.

#### GenCache's Four Phases

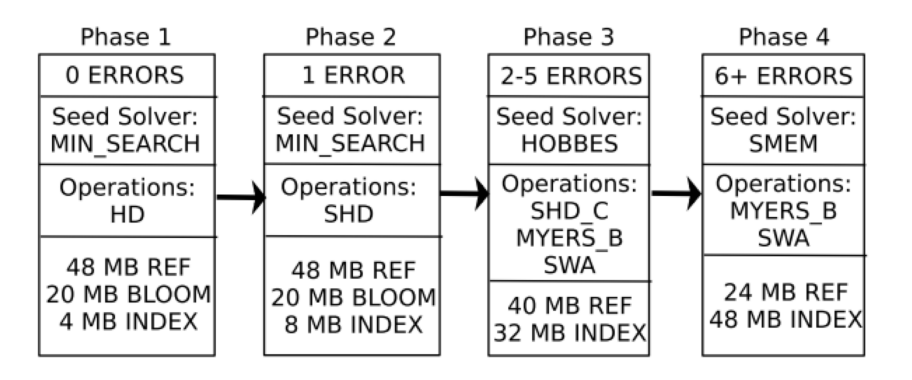


Figure 7: Four phases in the new alignment algorithm that exploits in-cache operators.

## Throughput Results

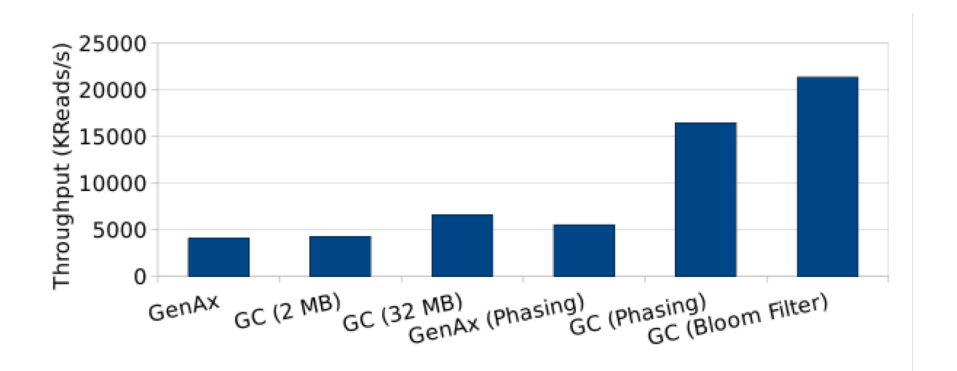


Figure 9: Throughput improvement of GenCache (Hardware & Software).

## Ongoing Directions

#### Seed Filtering Technique:

- Goal: Reducing the number of seed (k-mer) locations.
  - Heuristic (limits the number of mapping locations for each seed).
  - Supports exact matches only.

#### Pre-alignment Filtering Technique:

- Goal: Reducing the number of invalid mappings (>E).
  - Supports both exact and inexact matches.
  - Provides some falsely-accepted mappings.

#### Read Alignment Acceleration:

- Goal: Performing read alignment at scale.
  - Limits the numeric range of each cell in the DP table and hence supports limited scoring function.
  - May not support backtracking step due to random memory accesses.

#### Darwin

Session 3A: Programmable Devices and Co-processors

ASPLOS'18, March 24-28, 2018, Williamsburg, VA, USA

## Darwin: A Genomics Co-processor Provides up to 15,000× acceleration on long read assembly

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Seed filter: D-Soft

Read alignment accelerator: GACT ← We will cover this

Yatish+ "Darwin: A genomics co-processor provides up to 15,000 x acceleration on long read assembly." *ASPLOS* 2018. http://bejerano.stanford.edu/papers/p199-turakhia.pdf

#### Darwin: GACT Hardware Acceleration

#### Key observation:

 Data Dependencies limit accelerating the dynamic programming table calculation.

#### Key idea:

- Divide the dynamic programming table into overlapping tiles.
- Calculate each tile independently and in a systolic array fashion.
- Calculate many alignments concurrently.

#### Key result:

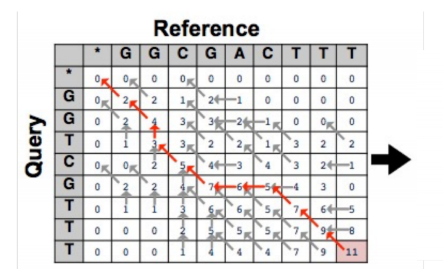
- It is simulated for TSMC 40nm CMOS process.
- It provides a speedup of up to 380x compared to GACT software.
- It is three orders of magnitude faster than Edlib (best-performing CPU read aligner).

#### Weaknesses:

It is not clear if tiling maintains the same accuracy as the original dynamic programming algorithm.

## Specialized Accelerator for Read Aligner

 Accelerating the read alignment algorithm as-is using specialized hardware (40 nm CMOS) provides a limited speedup (37x).



$$\begin{split} I(i,j) &= \max \left\{ H(i,j-1) - o, \, I(i,j-1) - e \right\} \\ D(i,j) &= \max \left\{ H(i-1,j) - o, \, D(i-1,j) - e \right\} \\ H(i,j) &= \max \left\{ \begin{matrix} 0 \\ I(i,j) \\ D(i,j) \\ H(i-1,j-1) + W(r_i,q_j) \end{matrix} \right. \end{split}$$

Dynamic programming for gene sequence alignment (Smith-Waterman)

VS.

**CPU-based read aligner** 

On 14nm CPU

35 ALU ops, 15 load/store

37 cycles

81nJ

Hardware accelerated read aligner

On 40nm Special Unit

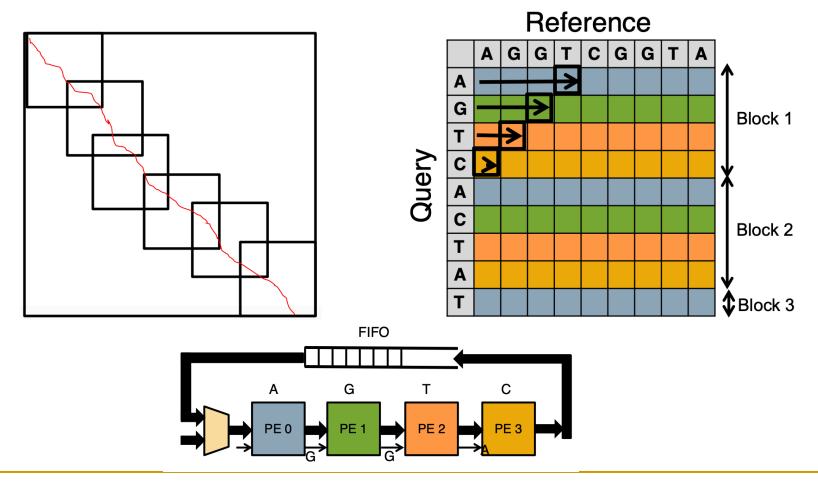
1 cycle (37x speedup)

3.1pJ (26,000x efficiency)

300fJ for logic (remainder is memory)

## GACT Alignment

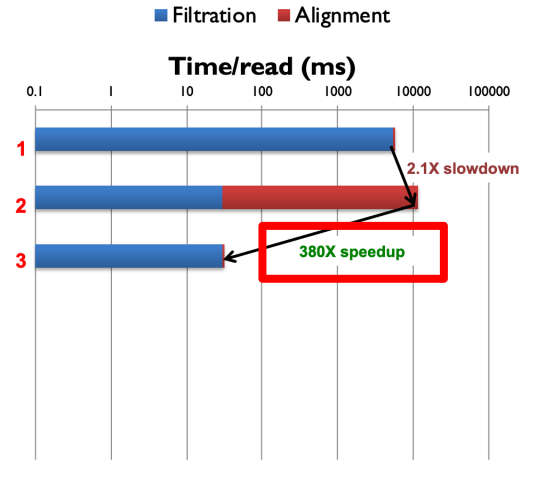
- Solution: Divide the table into overlapping tiles and compute them all independently using systolic arrays.
- Store the trace of each cell in an SRAM for traceback.



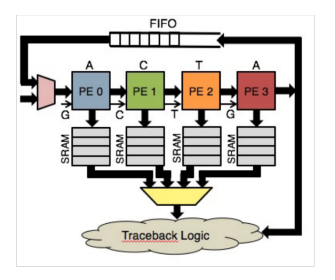
## Implementation Details

- It is simulated for TSMC 40 nm CMOS process.
- 64 systolic arrays are working concurrently.
- 64 PEs (processing elements) in each systolic array.
- Each entry of the dynamic programming table accommodates 16-bit value.
- Each systolic array requires 128 KB SRAM (each PE = 2 KB SRAM bank) for traceback purposes.

## GACT Hardware vs. Software Speedup



- Graphmap (software)
- Replace by D-SOFT and GACT (software)
- 3. GACT hardware-acceleration



#### GACT Hardware vs. Edlib

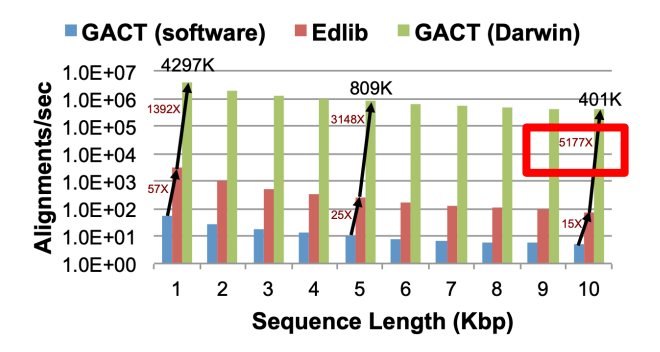


Figure 10: Throughput (alignments/second) comparison for different sequence lengths between a software implementation of GACT, Edlib library and the hardware-acceleration of GACT in Darwin.

#### More on Darwin

https://github.com/gsneha26/Darwin-WGA

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Yatish+ "Darwin: A genomics co-processor provides up to 15,000 x acceleration on long read assembly." *ASPLOS* 2018. <a href="http://bejerano.stanford.edu/papers/p199-turakhia.pdf">http://bejerano.stanford.edu/papers/p199-turakhia.pdf</a>

#### Disclaimer on Darwin

- Darwin is NOT developed in SAFARI group, but we developed BitMAC that is now <u>under review</u>.
- BitMAC = new read alignment algorithm + PIM specialized accelerator.
- BitMAC provides 2.1x better throughput per unit area and 59.2x better throughput per unit power when compared with GACT of Darwin.

## Conclusion on Ongoing Directions

- Read alignment can be substantially accelerated using computationally inexpensive and accurate pre-alignment filtering algorithms designed for specialized hardware.
- All the three directions are used by mappers today, but filtering has replaced alignment as the bottleneck.
- Pre-alignment filtering does not sacrifice any of the aligner capabilities, as it does not modify or replace the alignment step.

## Agenda for Today

- Why Genome Analysis?
- What is Genome Analysis?
- How we Map Reads?
- What Makes Read Mapper Slow?
- Algorithmic & Hardware Acceleration
  - Seed Filtering Technique
  - Pre-alignment Filtering Technique
  - Read Alignment Acceleration
- Where is Read Mapping Going Next?

## Where is Read Mapping Going Next?

Will 100% accurate genome-long reads alleviate/eliminate the need for read mapping?

Think about metagenomics, pan-genomics, ...

## Where is Read Mapping Going Next?

#### nature genetics

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Letter | Open Access | Published: 19 November 2018
```

## Assembly of a pan-genome from deep sequencing of 910 humans of African descent

```
Rachel M. Sherman ☑, Juliet Forman, [...] Steven L. Salzberg ☑

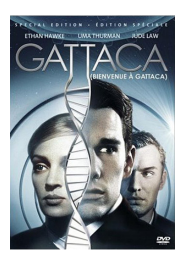
Nature Genetics 51, 30–35(2019) | Cite this article

39k Accesses | 29 Citations | 875 Altmetric | Metrics
```

African pan-genome contains ~10% more DNA than the current human reference genome.

#### Did we Achieve Our Goal?

 Our goal is to significantly reduce the time spent on calculating the optimal alignment in genome analysis from hours to mere seconds using both new algorithms & hardware accelerators, given limited computational resources (i.e., personal computer or small hardware).







1997

2015

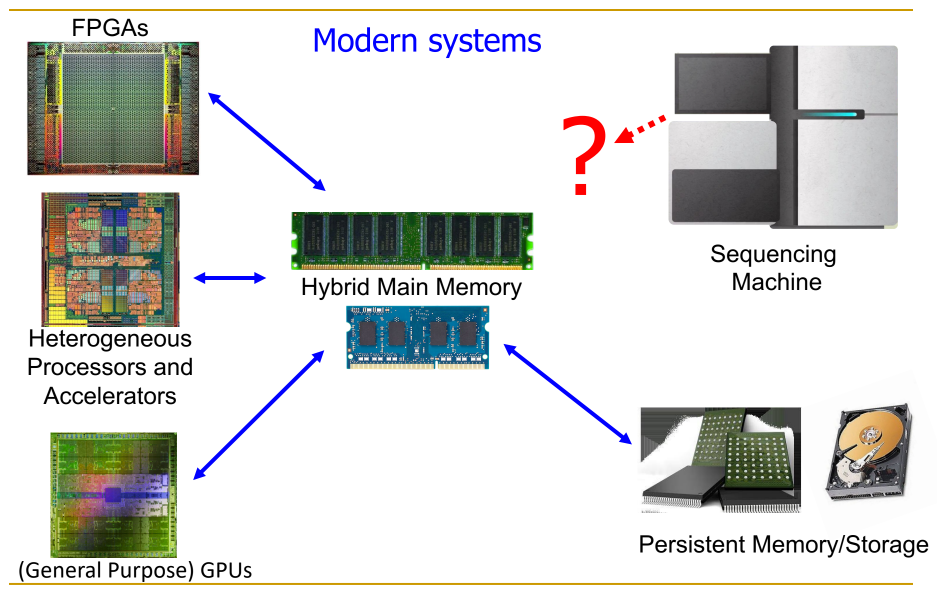
## Open Questions

How and where to enable

fast, accurate, cheap,

privacy-preserving, and exabyte scale analysis of genomic data?

### Processing Genomic Data Where it Makes Sense



#### Talk Conclusion

- System design for bioinformatics is a critical problem
  - It has large scientific, medical, societal, personal implications
- This talk is about accelerating a key step in bioinformatics: genome sequence analysis
  - In particular, read mapping
- Many bottlenecks exist in accessing and manipulating huge amounts of genomic data during analysis
- We cover various recent ideas to accelerate read mapping
  - A journey since September 2006

## Acknowledgments

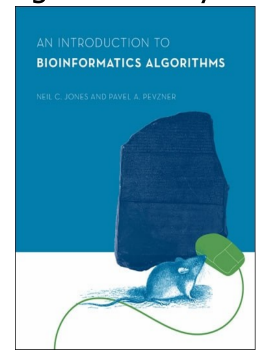
- Prof. Onur Mutlu, ETH Zurich
- Prof. Can Alkan, Bilkent University
- Many colleagues and collaborators
  - Damla Senol Cali, Jeremie Kim, Hasan Hassan, Donghyuk Lee, Hongyi Xin, ...

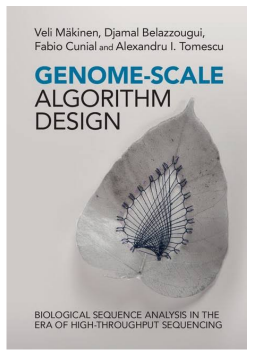
#### Funders:

- NIH and Industrial Partners (Alibaba, AMD, Google, Facebook, HP Labs, Huawei, IBM, Intel, Microsoft, Nvidia, Oracle, Qualcomm, Rambus, Samsung, Seagate, VMware)
- All papers, source code, and more are at:
  - https://people.inf.ethz.ch/omutlu/projects.htm

## Recommended Readings

- Jones, Neil C., Pavel A. Pevzner, and Pavel Pevzner. "An introduction to bioinformatics algorithms," MIT press, 2004.
- Mäkinen, Veli, Djamal Belazzougui, Fabio Cunial, and Alexandru I. Tomescu. "Genome-scale algorithm design," Cambridge University Press, 2015.





# Accelerating Genome Analysis Using New Algorithms and Hardware Designs

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#### **Preferred Networks**

16 December 2019



