Subset Seeds on a Reconfigurable Architecture LAW 2007

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Motivations	B /I			
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Specialised architecture

Conclusion

Overview

Motivations

Subset seeds

Where are we now?

Specialised architecture

Conclusion

Subset seeds

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Specialised architecture

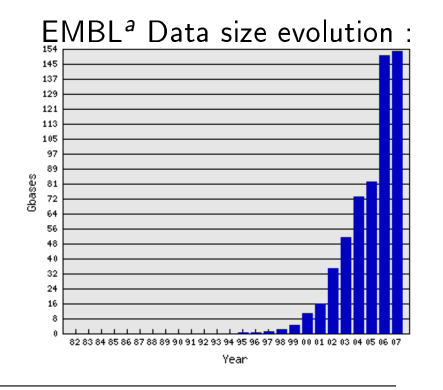
Conclusion

Where are we now?

Specialised architecture

Conclusion

Biological Data evolution



Data type evolution :

• Take the "*Junk DNA*" into account

^ahttp ://www.ebi.ac.uk/embl/ -Europe's primary nucleotide sequence resource

Large augmentation of amount of data

Main goal of the study

Biological similarities detection

Improvements of BLAST¹-like programs :

- Faster execution
- Larger amount of data
- More sensitive results

¹Altschul, S.; Gish, W.; Miller, W.; Myers, E. & Lipman, D. Basic Local Alignment Search Tool *Journal of Molecular Biology*, **1990**, 215, 403-410

Subset seeds

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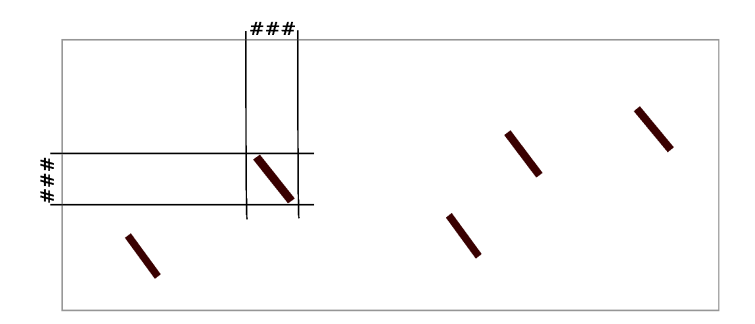
Conclusion

Seeds - Basic Ideas



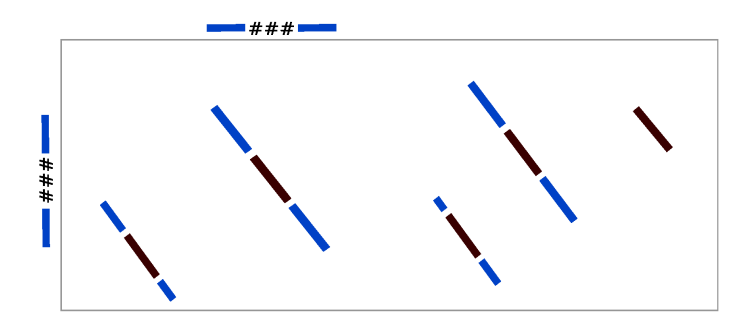
Computing local alignments, using dynamic programming

Seeds - Basic Ideas



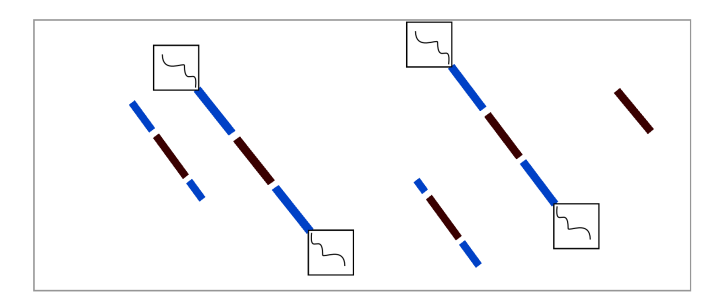
1. Detect matching seeds

Seeds - Basic Ideas



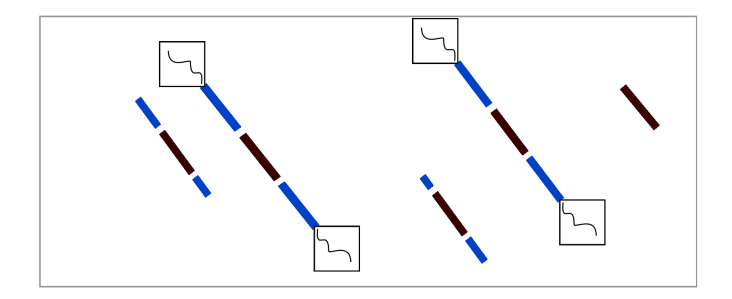
- 1. Detect matching seeds
- 2. Extend to neighbors

Seeds - Basic Ideas



- 1. Detect matching seeds
- 2. Extend to neighbors
- 3. Perform alignments

Seeds - Basic Ideas



Sequences indexation

Representation, specificity and sensitivity

• ### : "classical" seed

Representation, specificity and sensitivity

- ### : "classical" seed
- # : Low specificity & High sensitivity (Slow and precise)

Representation, specificity and sensitivity

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- ########## : High specificity & Low sensitivity (Fast and imprecise)

Representation, specificity and sensitivity

- ### : "classical" seed
- # : Low specificity & High sensitivity (Slow and precise)
- ########## : High specificity & Low sensitivity (Fast and imprecise)

What is the good seed?

Main difficulty : design seeds to have best ratio specificity *v.s.* sensitivity.

Conclusion

Spaced² seeds

ATCAGTGCAATGCTCAAGA | | | | | . | | . | | | . | | | | ATCAGCGCGATGCGCAAGA

ATCAGTGCAATGCTCAAGA | | | | | . | | . | | | . | | | | ATCAGCGCGATGCGCAAGA ###--#-##

²Burkhardt, S., & Kärkkäinen, J. Better filtering with gapped q-grams. *Fundamenta Informaticae* 56, 1-2 (2003), 51-70. Preliminary version in Combinatorial Pattern Matching **2001**

Conclusion

Spaced² seeds

AT C A G T G C A A T G C T C A A G A | | | | | . | . | . | | | . | | | | | A T C A G C G C G A T G C G C A A G A

#######

ATCAGTGCAATGCTCAAGA | | | | | . | | . | | | | | | | ATCAGCGCGATGCGCAAGA ### - - # - ## ### - - # -

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Spaced seeds

ATCAGTGCAATGCTCAAGA ||||||||||| ATCAGCGCGATGCGCAAGA ###--#-## ###--#-## ###--#-##

Spaced seeds have better sensitivity

Multiple³ spaced seeds

Instead of a unique Spaced seed
{###- -#-##} (weight 6)

³M. Li, B. Ma, D. Kisman & J. Tromp PatternHunter II : Highly sensitive and fast homology search. *J. of Bioinformatics and Comp. Biol.*, 2(3), 417-439, 2004.

Conclusion

Multiple³ spaced seeds

Instead of a unique Spaced seed
{###- -#-##} (weight 6)

Use a set of spaced seeds { ####- -#-###, (weight 7) #- -###-#-##, (weight 7) #- - ###- -##-#, (weight 7)}

³M. Li, B. Ma, D. Kisman & J. Tromp PatternHunter II : Highly sensitive and fast homology search. *J. of Bioinformatics and Comp. Biol.*, 2(3), 417-439, 2004.

Multiple³ spaced seeds

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{###- -#-##} (weight 6)

Use a set of spaced seeds { ###+--#-###, (weight 7) #--###-#-##, (weight 7) #--###--##-#, (weight 7) ...}

Advantages and drawback

- :-) Better sensitivity
- :-(Higher memory usage, (and slower)

³M. Li, B. Ma, D. Kisman & J. Tromp PatternHunter II : Highly sensitive and fast homology search. *J. of Bioinformatics and Comp. Biol.*, 2(3), 417-439, 2004.

Subset seeds (protein example)

	C	S	Т	Р	Α	G	Ν	D	Ε	Q	Н	R	К	М	Ι	L	٧	F	Y	W	
С	9																				C
S	-1	4																			S
Т	-1	1	5																		Т
Ρ	-3	-1	-1	7																	Ρ
Α	0	1	0	-1	4																A
G	-3	0	-2	-2	0	6															G
Ν	-3	1	0	-2	-2	0	6														Ν
D	-3	0	-1	-1	-2	-1	1	6													D
Е	-4	0	-1	-1	-1	-2	0	2	5												Е
Q	-3	0	-1	-1	-1	-2	0	0	2	5											Q
Н	-3	-1	-2	-2	-2	-2	1	-1	0	0	8										Н
R	-3	-1	-1	-2	-1	-2	0	-2	0	1	0	5									R
К	-3	0	-1	-1	-1	-2	0	-1	1	1	-1	2	5								К
М	-1	-1	-1	-2	-1	-3	-2	-3	-2	0	-2	-1	-1	5							М
Ι	-1	-2	-1	-3	-1	-4	-3	-3	-3	-3	-3	-3	-3	1	4						I
L	-1	-2	-1	-3	-1	-4	-3	-4	-3	-2	-3	-2	-2	2	2	4					L
۷	-1	-2	0	-2	0	-3	-3	-3	-2	-2	-3	-3	-2	1	3	1	4				۷
F	-2	-2	-2	-4	-2	-3	-3	-3	-3	-3	-1	-3	-3	0	0	0	-1	6			F
Υ	-2	-2	-2	-3	-2	-3	-2	-3	-2	-1	2	-2	-2	-1	-1	-1	-1	3	7		Y
W	-2	-3	-2	-4	-3	-2	-4	-4	-3	-2	-2	-3	-3	-1	-3	-2	-3	1	2	11	W
	С	S	Т	Р	А	G	N	D	E	Q	Н	R	К	М	I	L	۷	F	Y	W	

Kucherov, G.; Noé, L. & Roytberg, M. A Unifying Framework for Seed Sensitivity and its Application to Subset Seeds *WABI*, **2005**, 3692 of LNCS, 251-263

Subset seeds (protein example)

	C	S	Т	Р	А	G	Ν	D	Е	Q	Н	R	К	М	Ι	L	٧	F	Y	W	
С	9																				С
S	-1	4																			S
Т	-1	1	5																	ĺ	Т
Ρ	-3	-1	-1	7																ĺ	Ρ
А	0	1	0	-1	4															ĺ	Α
G	-3	0	-2	-2	0	6														ĺ	G
Ν	-3	1	0	-2	-2	0	6														Ν
D	-3	0	-1	-1	-2	-1	1	6												ĺ	D
Е	-4	0	-1	-1	-1	-2	0	2	5											ĺ	Е
Q	-3	0	-1	-1	-1	-2	0	0	2	5										ĺ	Q
Н	-3	-1	-2	-2	-2	-2	1	-1	0	0	8										Н
R	-3	-1	-1	-2	-1	-2	0	-2	0	1	0	5								l	R
К	-3	0	-1	-1	-1	-2	0	-1	1	1	-1	2	5							ſ	К
М	-1	-1	-1	-2	-1	-3	-2	-3	-2	0	-2	-1	-1	5							М
Ι	-1	-2	-1	-3	-1	-4	-3	-3	-3	-3	-3	-3	-3	1	4					ĺ	Ι
L	-1	-2	-1	-3	-1	-4	-3	-4	-3	-2	-3	-2	-2	2	2	4				ĺ	L
۷	-1	-2	0	-2	0	-3	-3	-3	-2	-2	-3	-3	-2	1	3	1	4			[۷
F	-2	-2	-2	-4	-2	-3	-3	-3	-3	-3	-1	-3	-3	0	0	0	-1	6			F
Υ	-2	-2	-2	-3	-2	-3	-2	-3	-2	-1	2	-2	-2	-1	-1	-1	-1	3	7		Y
W	-2	-3	-2	-4	-3	-2	-4	-4	-3	-2	-2	-3	-3	-1	-3	-2	-3	1	2	11	W
	C	S	Т	Р	Α	G	Ν	D	E	Q	Н	R	К	М	Ι	L	٧	F	Y	W	

All amino acids are not equivalents

• Create groups of characters

Subset seeds (protein example)

	C	S	Т	Р	А	G	N	D	Ε	Q	Н	R	К	М	I	L	٧	F	Y	W	
С	9																				C
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D	-3	0	-1	-1	-2	-1	1	6													D
Е	-4	0	-1	-1	-1	-2	0	2	5												Е
Q	-3	0	-1	-1	-1	-2	0	0	2	5											Q
Н	-3	-1	-2	-2	-2	-2	1	-1	0	0	8										Н
R	-3	-1	-1	-2	-1	-2	0	-2	0	1	0	5									R
К	-3	0	-1	-1	-1	-2	0	-1	1	1	-1	2	5								К
М	-1	-1	-1	-2	-1	-3	-2	-3	-2	0	-2	-1	-1	5							М
Ι	-1	-2	-1	-3	-1	-4	-3	-3	-3	-3	-3	-3	-3	1	4						I
L	-1	-2	-1	-3	-1	-4	-3	-4	-3	-2	-3	-2	-2	2	2	4					L
۷	-1	-2	0	-2	0	-3	-3	-3	-2	-2	-3	-3	-2	1	3	1	4				۷
F	-2	-2	-2	-4	-2	-3	-3	-3	-3	-3	-1	-3	-3	0	0	0	-1	6			F
Υ	-2	-2	-2	-3	-2	-3	-2	-3	-2	-1	2	-2	-2	-1	-1	-1	-1	3	7		Y
W	-2	-3	-2	-4	-3	-2	-4	-4	-3	-2	-2	-3	-3	-1	-3	-2	-3	1	2	11	W
	С	S	Т	Р	А	G	N	D	E	Q	Н	R	К	М	I	L	۷	F	Y	W	

All amino acids are not equivalents

• Create groups of characters

- $\begin{array}{c|c} \# & A, C, D, E, F, G, H, I, K, L, M, N, P, Q, R, S, T, V, W, Y \\ & CFYWMLIVGPATSNHQEDRK \end{array}$
- **@**₁ **C**, **STPAG**, **NDEQ**, **HRK**, **MILV**, **FYW**
- @₂ *CFYWMLIV*, *GPATSNHQEDRK*
- # | A, C, D, E, F, G, H, I, K, L, M, N, P, Q, R, S, T, V, W, Y

Specialised architecture

Conclusion

So what?

Design good set(s) of subset (spaced) seeds

- Speed
- Memory
- Specificity
- Sensitivity

Subset seeds

Where are we now?

Specialised architecture

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Where are we now?

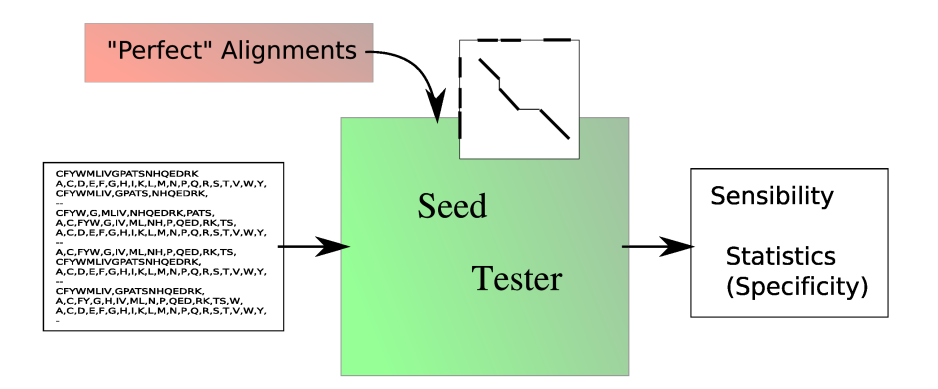
Specialised architecture

Conclusion

Motivations Subset seeds Where are we now? Specialised architecture Conclusion Seed tester CFYWMLIVGPATSNHQEDRK A,C,D,E,F,G,H,I,K,L,M,N,P,Q,R,S,T,V,W,Y, Sensibility CFYWMLIV, GPATS, NHQEDRK, Seed CFYW,G,MLIV,NHQEDRK,PATS, A,C,FYW,G,IV,ML,NH,P,QED,RK,TS, A,C,D,E,F,G,H,I,K,L,M,N,P,Q,R,S,T,V,W,Y, **Statistics** A,C,FYW,G,IV,ML,NH,P,QED,RK,TS, CFYWMLIVGPATSNHQEDRK, Tester A,C,D,E,F,G,H,I,K,L,M,N,P,Q,R,S,T,V,W,Y, (Specificity) CFYWMLIV, GPATSNHQEDRK, A,C,FY,G,H,IV,ML,N,P,QED,RK,TS,W, A,C,D,E,F,G,H,I,K,L,M,N,P,Q,R,S,T,V,W,Y,

- Test quickly sensitivity (and possibly specificity)
- A few minutes
- Specificity is longer, \Rightarrow statistical computation is preferred

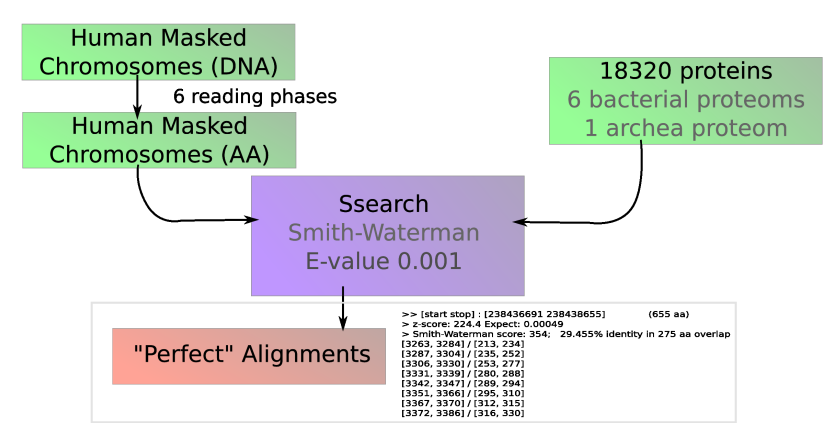
Seed tester



• Necessity to pre-compute *perfect* alignments

Conclusion

Seed tester



Today

- Chromosomes 1, 2 and 19 treated
- 3273 Alignments found

Lipman, D.J. & Pearson, W.R. Rapid and Sensitive Protein Similarity Searches *Science*, **1985**, 227, 1435-1441

First results

Selection of a Subset seeds :

- Find 98% of alignments while BLAST finds 96% of alignment.
- The 2% difference is biologically relevant
- Only 1 alignment found by BLAST is not found by subset seeds

Subset seeds

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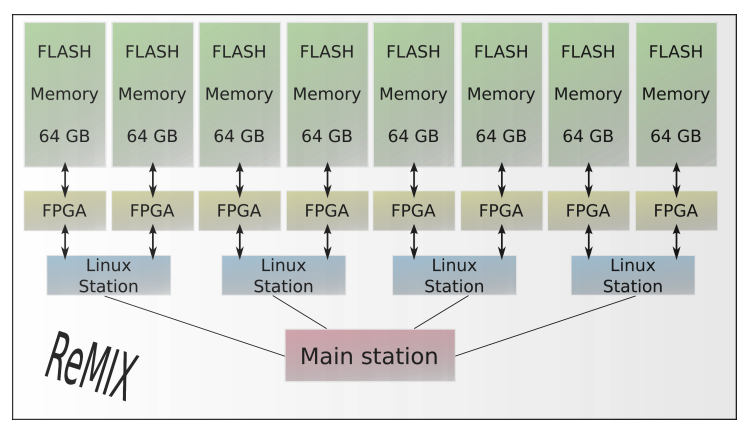
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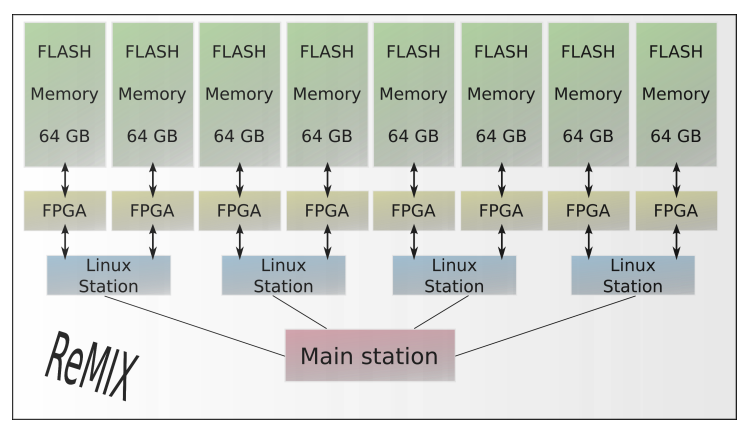
ReMIX, Overall presentation Lavenier, D.; Liu, X. & Georges, G. Seed-Based Genomic Sequence Comparison Using a FPGA/FLASH Accelerator IEEE FPT 2006.



- 512 GB FLASH memory (indexation, step 1)
- FPGA : Compute approximatively 8×160 ungaped alignments simultaneously in 50 clock cycles (step 2)
- A clock cycle $\Rightarrow 25.10^{-9}$ seconds

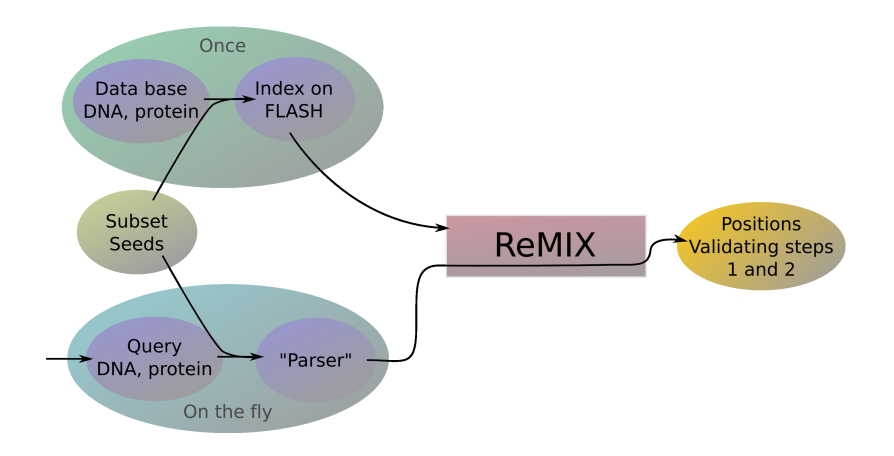
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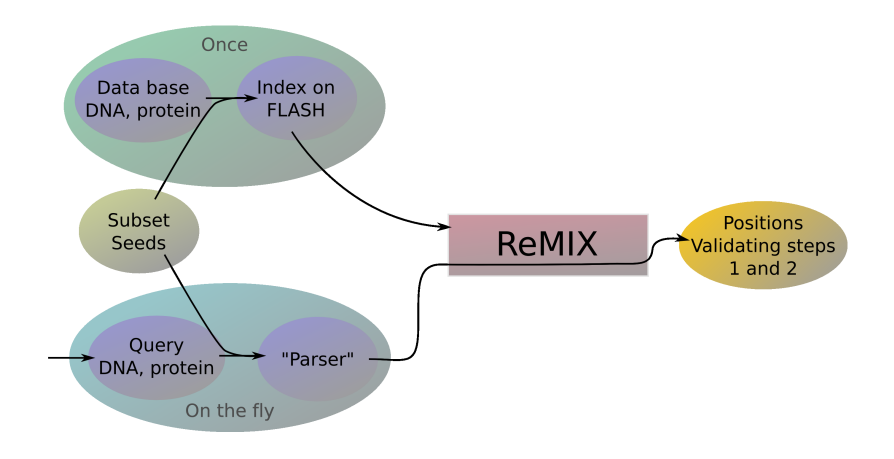
- 512 GB FLASH memory (indexation, step 1)
- FPGA : Compute approximatively 8×160 ungaped alignments simultaneously in 50 clock cycles (step 2)
- 1024 millions of ungaped alignments (neighbor) per second

ReMIX, Biological application



- Index computed once
- Query, parsed on the fly

ReMIX, Biological application



- Index computed once
- Query, parsed on the fly
- done seed ###
- todo all others...

Subset seeds

Where are we now?

Specialised architecture

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Where are we now?

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Conclusion

Goal

- BLAST-like programs :
 - Take larger amount of data
 - Increase speed
 - Increase sensitivity

Subset seeds

Where are we now?

Specialised architecture

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Conclusion

Goal

- BLAST-like programs :
 - Take larger amount of data
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 - Increase sensitivity

Done

- Subset seeds
- Framework for subset seeds testing
- (In progress) Implementation on ReMIX

Subset seeds

Where are we now?

Specialised architecture

Conclusion

Conclusion

Goal

- BLAST-like programs :
 - Take larger amount of data
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 - Increase sensitivity

Done

- Subset seeds
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To be done

- Investigation on subset seeds (98 %, 99 %, [99.9 %?])
- Feed the seed tester with new alignements
- Implementation, tests, distribution