A perspective on the current state of genomics

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Sequencing Platforms

 1986 - Dye terminator Sanger sequencing, peaking at about 900kb/day in early 2000s





'Next' Generation

 2005 – 'Next Generation Sequencing' as Massively parallel sequencing, both throughput and speed advances. The first was the Genome Sequencer (GS) instrument developed by 454 life Sciences (later acquired by Roche), Pyrosequencing 1.5Gb/day



Illumina (Solexa)

NovaSeq

 2006 – The second 'Next Generation Sequencing' platform. Now the dominant platform with 75% market share of sequencer and and estimated >90% of all bases sequenced are from an Illumina machine, Sequencing by Synthesis > 1600Gb/day.



HiSeq

Complete Genomics

 2006 – Using DNA nanoball sequencing, has been a leader in Human genome resequencing, having sequenced over 20,000 genomes to date. In 2013 purchased by BGI and is now set to release their first commercial sequencer, the Revolocity. Throughput on par with HiSeq

DEFUNCT

Human genome/exomes only.

10,000 Human Genomes per year



Bench top Sequencers

- **❖** Life Technologies
- Ion Torrent
- Ion Proton
- Gene Studio S5







- **❖**Illumina
- MiSeq
- MiniSeq
- iSeq 100







The 'Next' Generation Sequencers (3rd Generation)

 2009 – Single Molecule Read Time sequencing by Pacific Biosystems, most successful third generation sequencing platforms, RSII ~2Gb/day, newer Pac Bio Sequel ~14Gb/day, near 100Kb reads.

SMRT Sequencing





Iso-seq on Pac Bio possible, transcriptome without 'assembly'

Oxford Nanopore



• 2015 – Another 3rd generation sequencer, founded in 2005 and currently in beta testing. The sequencer uses nanopore technology developed in the 90's to sequence single molecules. Throughput is about 500Mb per flowcell, capable of near 200kb reads.

Fun to play with but results are highly variable

Nanopore Sequencing





FYI: 4th generation sequencing is being described as In-situ sequencing

Bioinformatics

Old Way of thinking about Bioinformatics



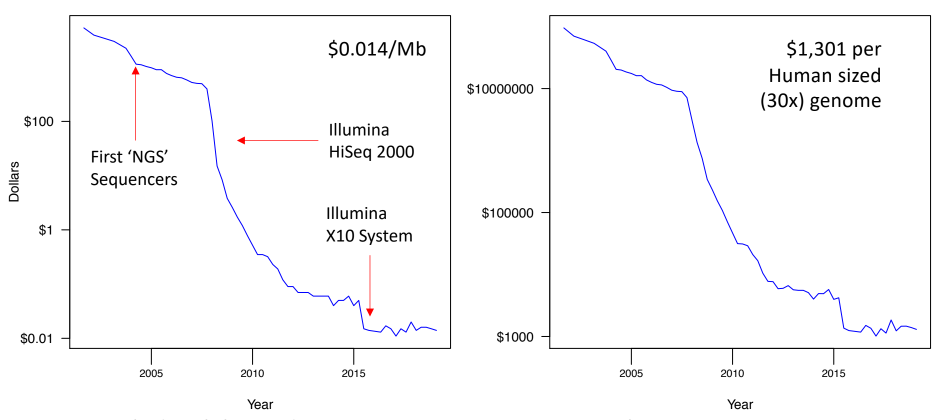
Appro Cluster

Sequencing Costs

February 2019

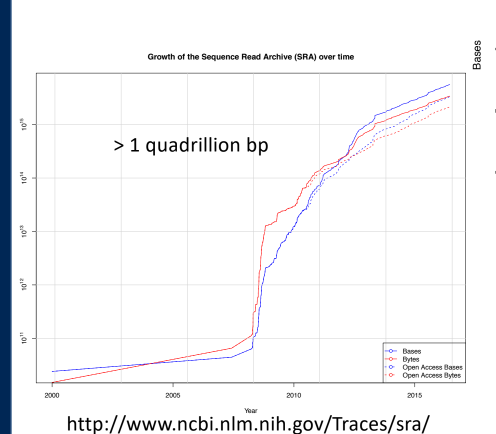
Cost per Megabase of Sequence

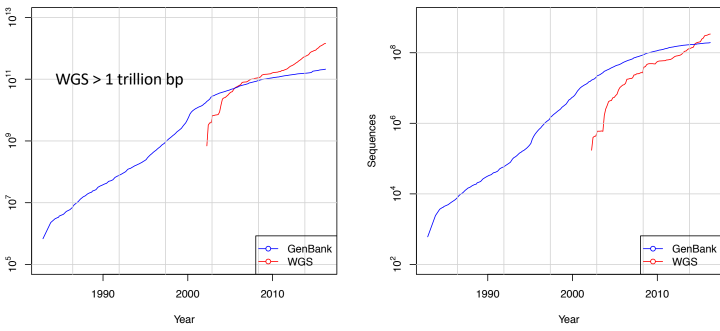
Cost per Human Sized Genome @ 30x (Illumina)



- Includes: labor, administration, management, utilities, reagents, consumables, instruments (amortized over 3 years), informatics related to sequence productions, submission, indirect costs.
- http://www.genome.gov/sequencingcosts/

Growth in Public Sequence Database



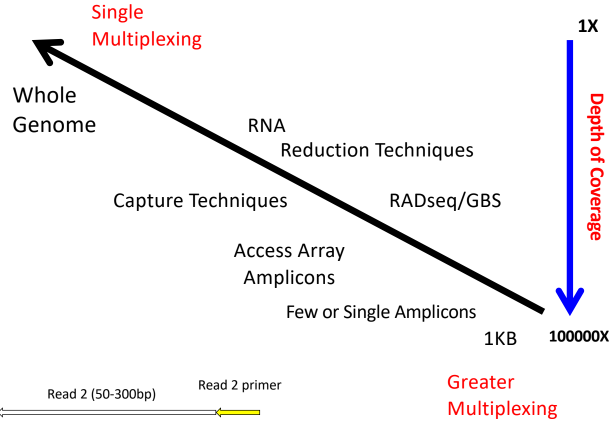


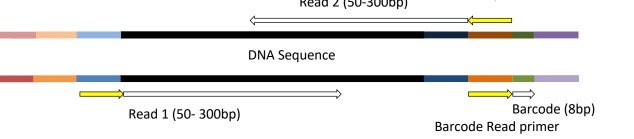
http://www.ncbi.nlm.nih.gov/genbank/statistics

Illumina's Flexibility

Genomic reduction allows for greater coverage and multiplexing of samples.

You can fine tune your depth of coverage needs and sample size with the reduction technique





Sequencing Libraries: MLA-seq

DNA-seq DNase-seq tagRNA-seq EnD-seq

RNA-seq ATAC-seq PAT-seq Pool-seq

Amplicons MNase-seq Structure-seq G&T-seq

CHiP-seq FAIRE-seq MPE-seq Tn-Seq

MeDiP-seq Ribose-seq STARR-seq BrAD-seq

RAD-seq STARR-seq Brad-seq

smRNA-seg Mod-seg SLAF-seg

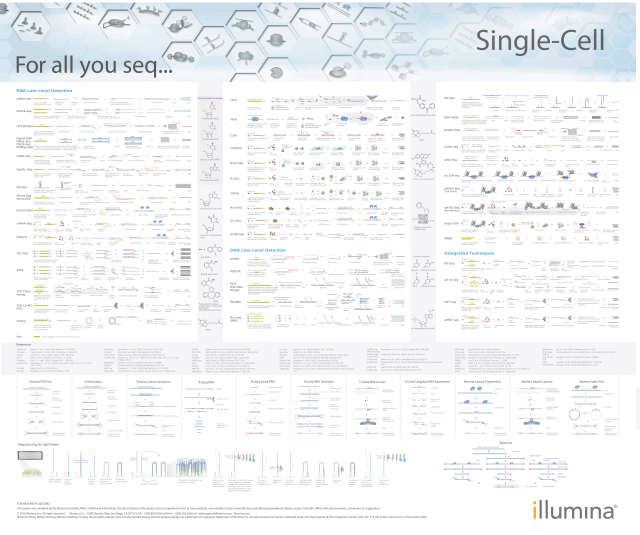
ddRAD-seq Simurated Mod seq Sexus

Methods. 2018 Jun 11. pii: S1046-2023(18)30064-1. doi: 10.1016/j.ymeth.2018.06.004. [Epub ahead of print]

fCLIP-seq for transcriptomic footprinting of dsRNA-binding proteins: lessons from DROSHA.

Kim B¹, Kim VN².

For all you seq



For all you seq...





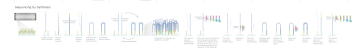








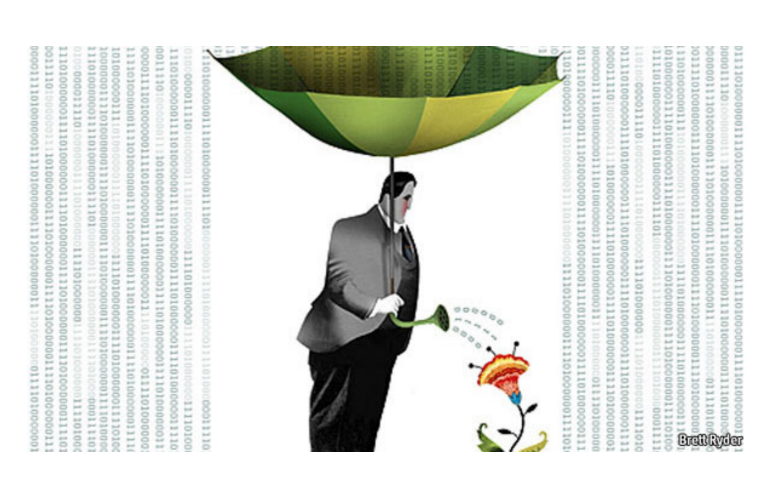








The data deluge



Plucking the biology from the Noise

Reality



• Its much more difficult than we may first think

Data Science

Data science is the process of formulating a quantitative question that can be answered with data, collecting and cleaning the data, analyzing the data, and communicating the answer to the question to a relevant audience.

7 Stages to Data Science

- 1. Define the question of interest
- 2. Get the data
- 3. Clean the data
- 4. Explore the data
- 5. Fit statistical models
- 6. Communicate the results
- 7. Make your analysis reproducible

1. Define the question of interest

Begin with the end in mind!

what is the question how are we to know we are successful what are our expectations

dictates

the data that should be collected the features being analyzed which algorithms should be use

- 2. Get the data
- 3. Clean the data
- 4. Explore the data

Know your data!

know what the source was technical processing in producing data (bias, artifacts, etc.)
"Data Profiling"



Data are never perfect but love your data anyway!

the collection of massive data sets often leads to unusual, surprising, unexpected and even outrageous.

5. Fit statistical models Machine Learning

Over fitting is a sin against data science!

Model's should not be over-complicated

 If the data scientist has done their job correctly the statistical models don't need to be incredibly complicated to identify important relationships

 In fact, if a complicated statistical model seems necessary, it often means that you don't have the right data to answer the question you really want to answer.

- 6. Communicate the results
- 7. Make your analysis reproducible

Remember that this is 'science'!

We are experimenting with data selections, processing, algorithms, ensembles of algorithms, measurements, models. At some point these *must all be tested for validity and applicability* to the problem you are trying to solve.



Data science done well looks easy – and that's a big problem for data scientists

simplystatistics.org
March 3, 2015 by Jeff Leek

Bad data science (bioinformatics) also looks easy

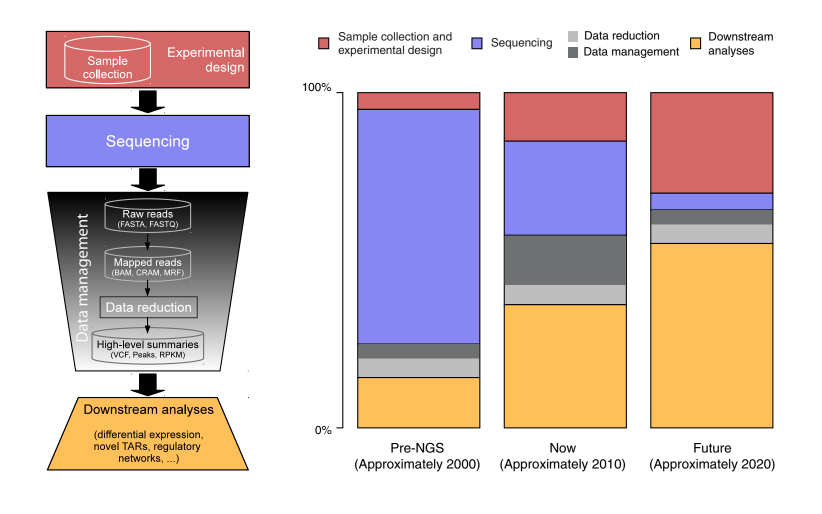
The Data Science in Bioinformatics

Bioinformatics is not something you are taught, it's a way of life

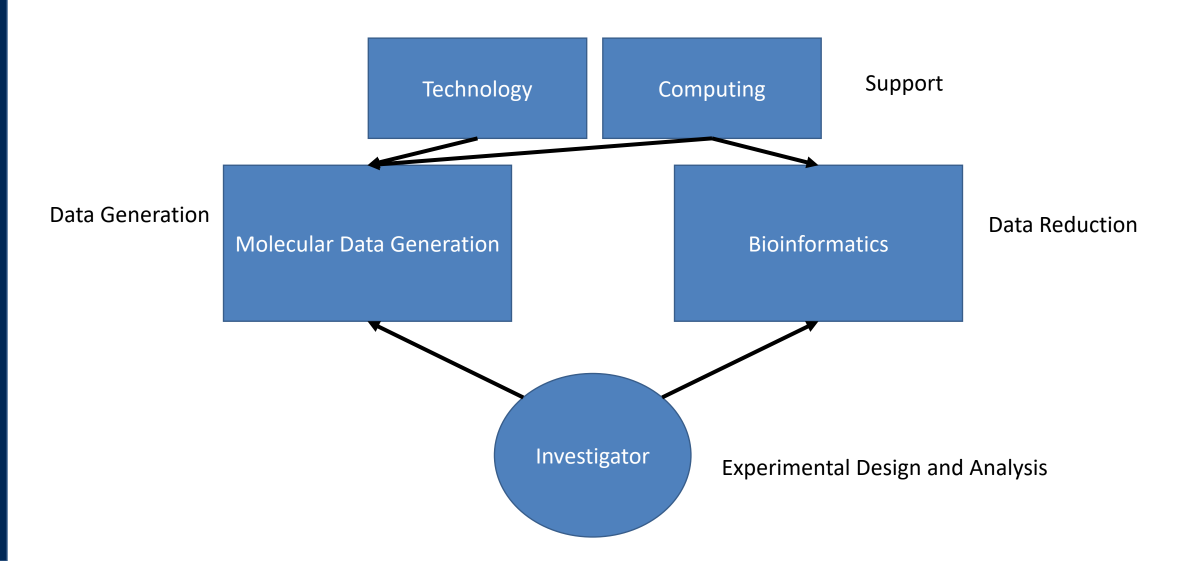
"The best bioinformaticians I know are **problem solvers** – they start the day not knowing something, and they enjoy finding out (themselves) how to do it. It's a great skill to have, but for most, it's not even a skill – it's a passion, it's a way of life, it's a thrill. It's what these people would do at the weekend (if their families let them)."

Mick Watson – Rosland Institute

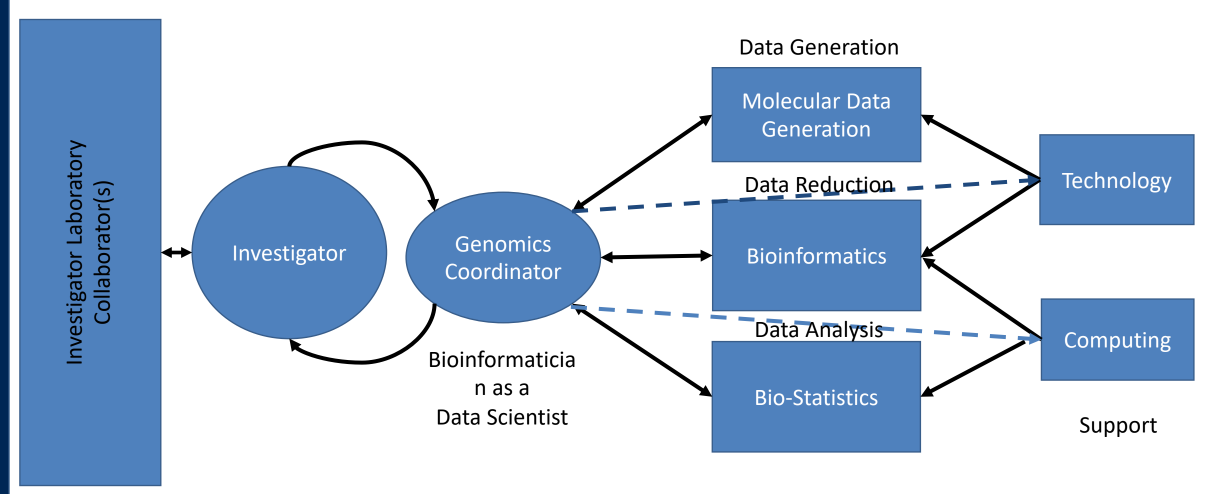
"The real cost of sequencing"



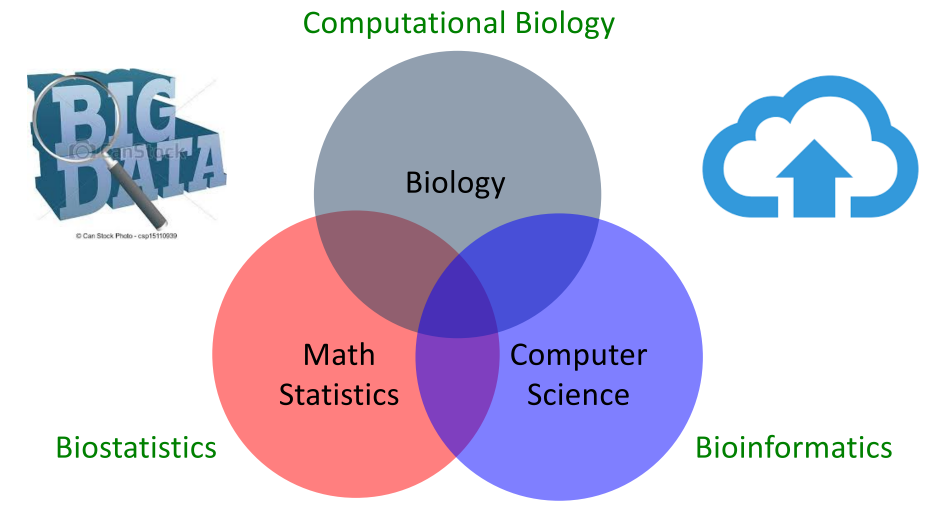
Old (Current) Model - Genomics



Needed Model - Genomics

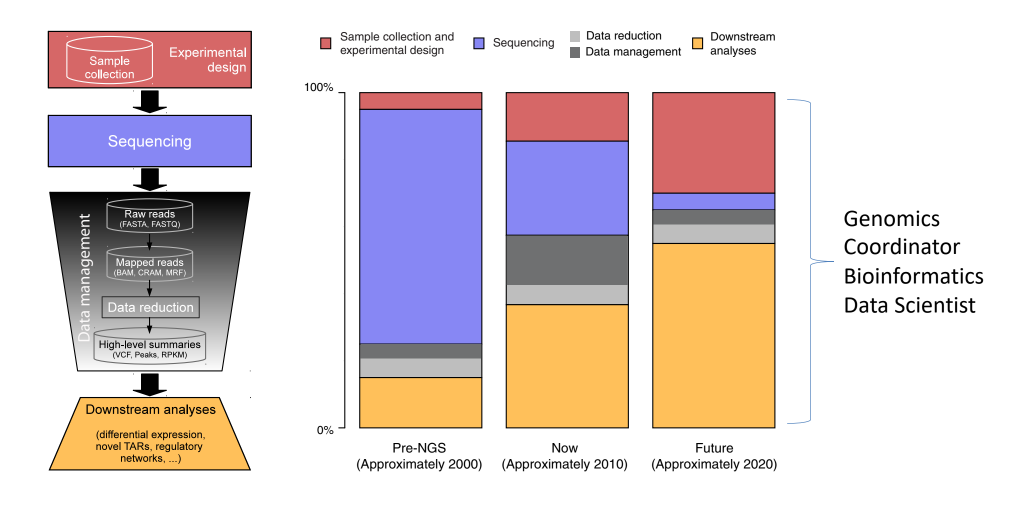


Genomics Coordinator – Data Scientist



'The data scientist role has been described as "part analyst, part artist."'
Anjul Bhambhri, vice president of big data products at IBM

"The real cost of sequencing"



Training - Models

- Workshops
 - Often enrolled too late
- Collaborations
 - More experience persons
- Apprenticeships
 - Previous lab personnel to young personnel
- Formal Education
 - Most programs are graduate level
 - Few Undergraduate

Substrate

Cloud Computing



Cluster Computing





BASTM



Laptop & Desktop





Environment

"Command Line" and "Programming Languages"











VS



Bioinformatics Software Suite





Prerequisites for doing Bioinformatics

- Access to a multi-core (24 cpu or greater), 'high' memory 64Gb or greater Linux server.
- Familiarity with the 'command line' and at least one programming language.
- Basic knowledge of how to install software
- Basic knowledge of R (or equivalent) and statistical programming
- Basic knowledge of Statistics and model building

The last mile



http://www.bikeblanket.com/blog/suisse

The Bottom Line: In Genomics

Spend the time (and money) planning and producing good quality, accurate and sufficient data.

Get to know to the data, develop and test expectations, explore and identify patterns.

Result, **spend much less time** (and less money) extracting biological significance and results with fewer failures and reproducible research.