Discovering gene regulatory control using ChIP-chip and ChIP-seq

Part 1

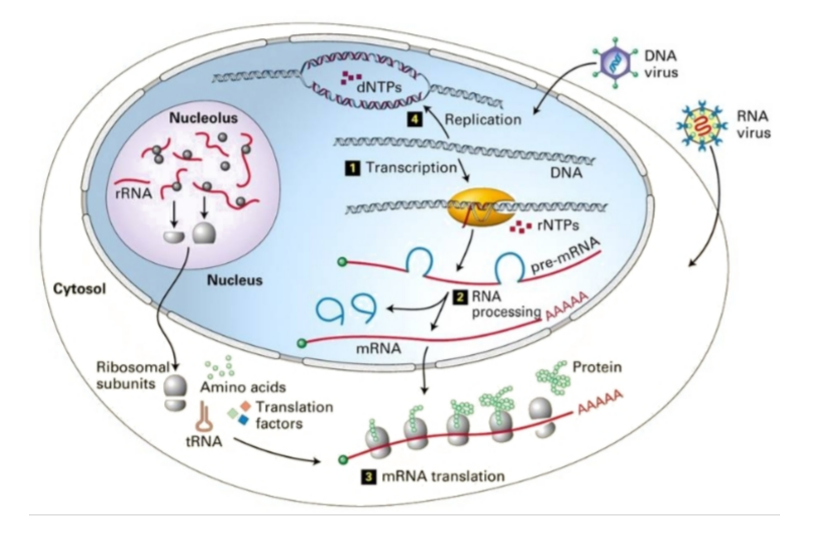
"An introduction to gene regulatory control, concepts and methodologies"

Ian Simpson

ian.simpson@.ed.ac.uk http://bit.ly/bio2links

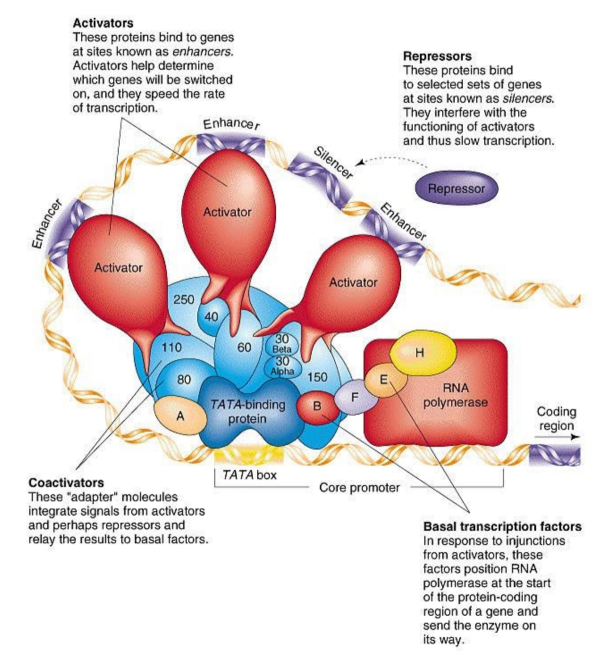


#### The Central Dogma of Molecular Biology



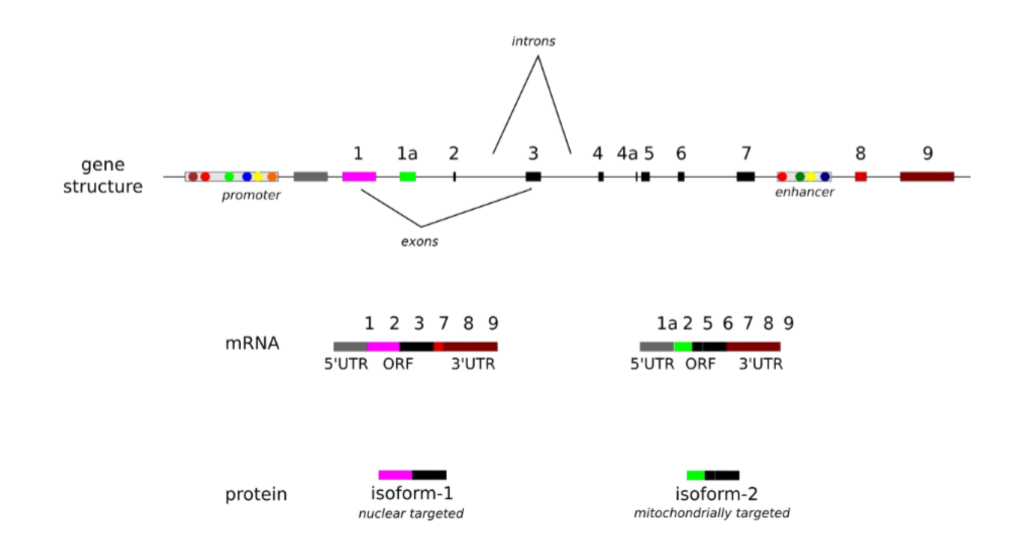


#### Transcription





#### Gene structure



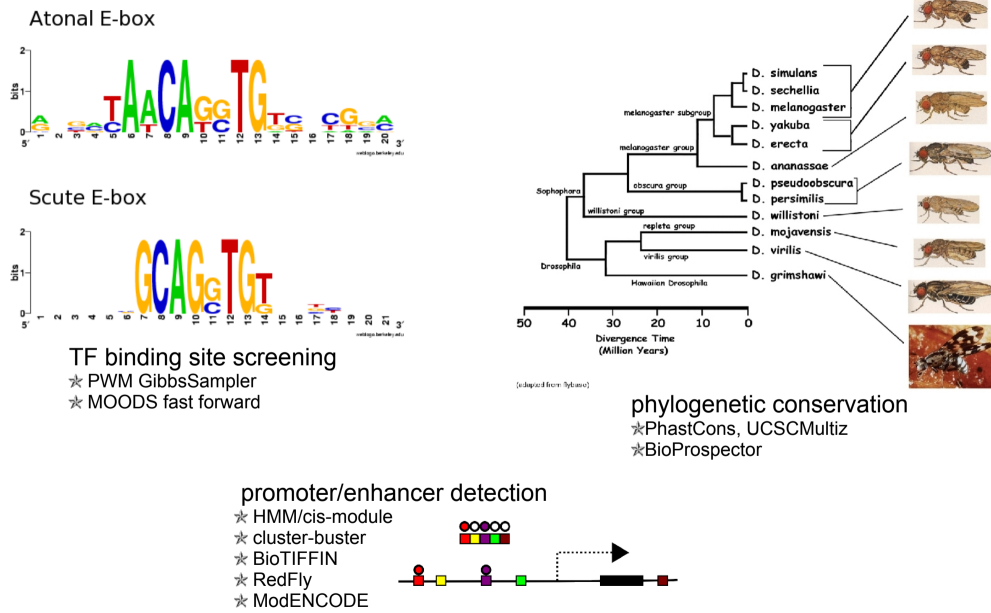


Example PWM for the human P53 protein

CONSENSUS	R	R	R	С	w	w	G	Y	Y	Y	R	R	R	С	w	w	G	Y	Y	Y	
p53 target																					score
GADD45A	G	Α	А	С	Α	т	G	т	С	т	Α	Α	G	С	Α	т	G	С	т	G	241
MDM2_1	G	Α	А	С	G	т	G	т	С	т	Α	Α	Α	С	т	т	G	Α	С	С	221
MDM2_2	Α	G	А	С	Α	А	G	т	С	Α	G	G	Α	С	т	т	Α	Α	С	т	226
BAX	G	С	С	С	А	С	G	С	С	С	Α	G	G	С	т	т	G	т	С	т	233
MMP2	A	G	А	С	А	А	G	С	С	т	G	A	Α	С	т	т	G	т	С	т	245
GDF15_1	A	G	А	С	Α	Α	G	т	С	т	G	G	G	С	Α	Α	G	Α	т	G	246
GDF15_2	A	G	С	С	А	т	G	С	С	С	G	G	G	С	Α	Α	G	Α	Α	С	241
GTSE1	A	G	G	С	А	А	G	С	С	С	С	Α	Α	С	т	т	G	С	т	С	230
CDKN1A	G	Α	А	С	Α	т	G	т	С	С	С	Α	Α	С	Α	т	G	т	т	G	244
GML	G	G	А	С	Α	т	G	С	С	т	G	G	G	С	Α	Α	G	С	Α	т	251
SCARA3	G	G	G	С	Α	А	G	С	С	С	Α	G	Α	С	Α	Α	G	т	т	G	249
RRM2B	т	G	А	С	А	т	G	С	С	С	Α	G	G	С	Α	т	G	т	С	т	259
PMAIP1	A	G	G	С	т	т	G	С	С	С	С	G	G	С	Α	Α	G	т	т	G	242
TP53INP1	G	Α	А	С	т	т	G	G	G	G	G	Α	Α	С	Α	т	G	т	т	т	211
TNFRSF10B	G	G	G	С	А	т	G	т	С	С	G	G	G	С	Α	Α	G	А	С	G	258
P53AIP1	т	С	т	С	т	т	G	С	С	С	G	G	G	С	т	т	G	т	С	G	237
TP53/3	G	Α	G	С	А	т	G	G	G	т	G	G	G	С	Α	Α	G	С	т	G	223
BBC3	G	G	А	С	Α	А	G	т	С	Α	G	G	Α	С	т	т	G	С	Α	G	246
TNFRSF6	т	G	G	С	т	т	G	т	С	Α	G	G	G	С	т	т	G	т	С	С	242
IGFBP3	Α	G	G	С	т	т	G	G	С	Α	G	G	т	С	т	т	G	С	С	С	227
SFN	G	С	Α	т	т	Α	G	С	С	С	Α	G	Α	С	Α	т	G	т	С	С	222
р53_PWM																					
A	7	5	11	-177	14	7	-177	-177	-177	4	6	6	9	-177	12	7	1	5	3	-177	]
С	-177	3	2	20	-177	1	-177	10	19	10	3	-177	-177	21	-177	-177	-177	6	10	6	
G	11	13	7	-177	1	-177	21	3	2	1	12	15	11	-177	-177	-177	20	-177	-89	9	
т	3	-177	1	1	6	13	-177	8	-177	6	-89	-177	1	-177	9	14	-177	10	8	6	

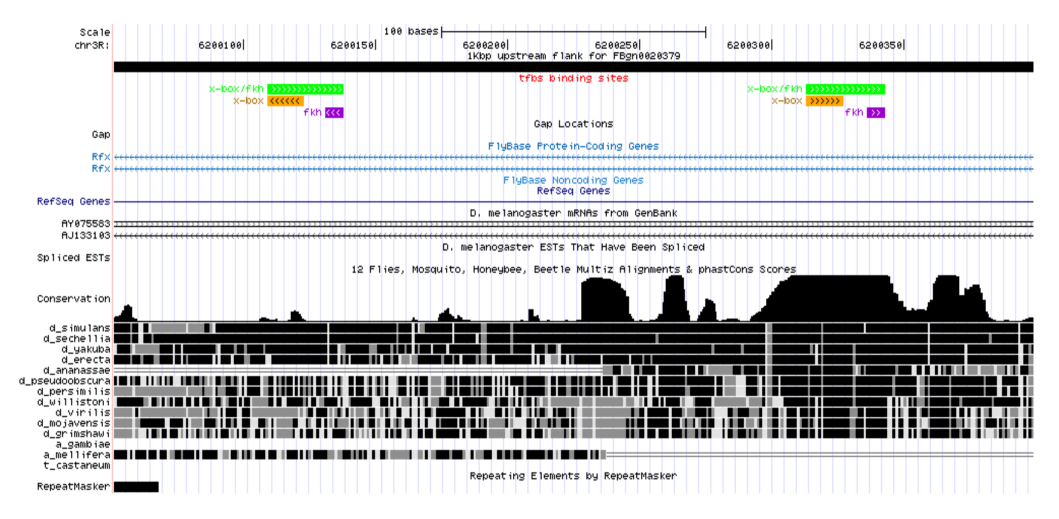


### The classic footprinting method





# Classic phylogenetic footprinting approach





# Limitations of the classical approach to finding TFBSs

•The number and quality of binding site sequences is low

•There is no explicit relation between conservation and function i.e. sites are often conserved, but conserved sites do not necessarily function

•Assumptions have to be made about where to look and how to score

•Extremely biased information, low number of experiments to determine sites

•Non-physiological conditions used during assessment

•Measurements made only in specific tissue or cells at specific times local solutions to the PWM problem, may be wrong for other conditions



#### Problems with the available data sources

#### \*Main source of site specific data remains pattern or PWM (or HMM)

Common name	Binary nomenclature	Number of PWMs
human	Homo sapiens	476
mouse	Mus musculus	423
rat	Rattus norvegicus	253
chick	Gallus gallus	133
clawed frog	Xenopus laevis	84
<u>fruit fly</u>	Drosophila melanogaster	<u>68</u>
thale cress	Arabidopsis thaliana	45
yeast	Saccharomyces cerevisiae	39
monkey	Cercopithecus aethiops	29
gibbon ape	Hylobates lar	24
cattle	Bos taurus	23
domestic pig	Sus scrofa	20
zebra fish	Brachydanio rerio	19
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# Replacing classical prediction with direct localisation

What do we need

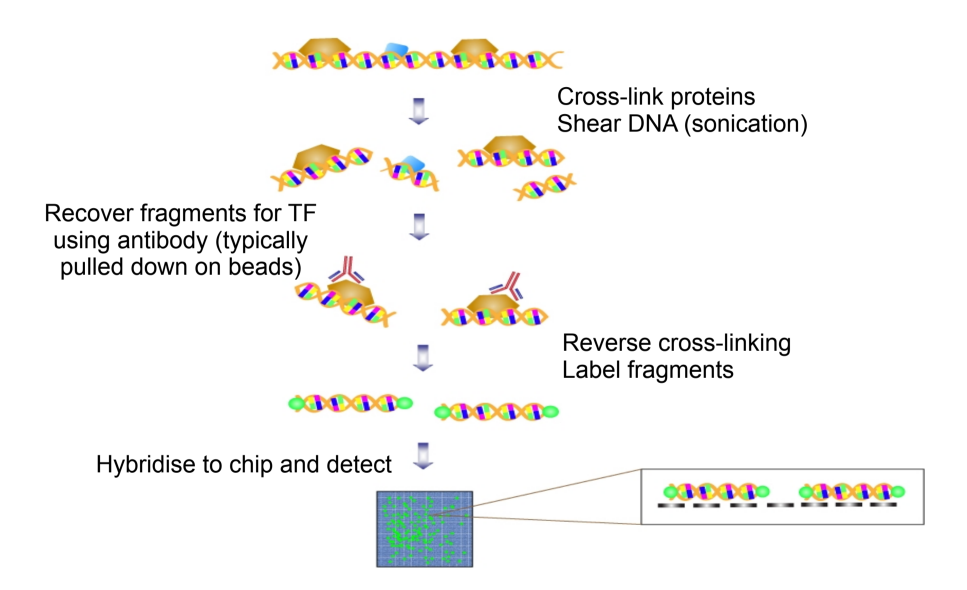
•Assays that cover the whole genome (aren't biased)

•Applicable to all transcription factors (good coverage)

- •Can be measured in lots of different conditions (condition specific, biologically relevant)
- •Can be mapped onto precise (and small) genome locations (high resolution)
- •Cost effective, accurate and reliable



Chromatin immuno-precipitation (ChIP)





How do we get from populations of DNA fragments to positions on chromosomes ?

Currently there are two main choices

ChIP-chip	Hybridisation onto a genomic tiling array
Chip-seq	Direct sequencing of the bound (now released) fragments

#### ChIP-chip

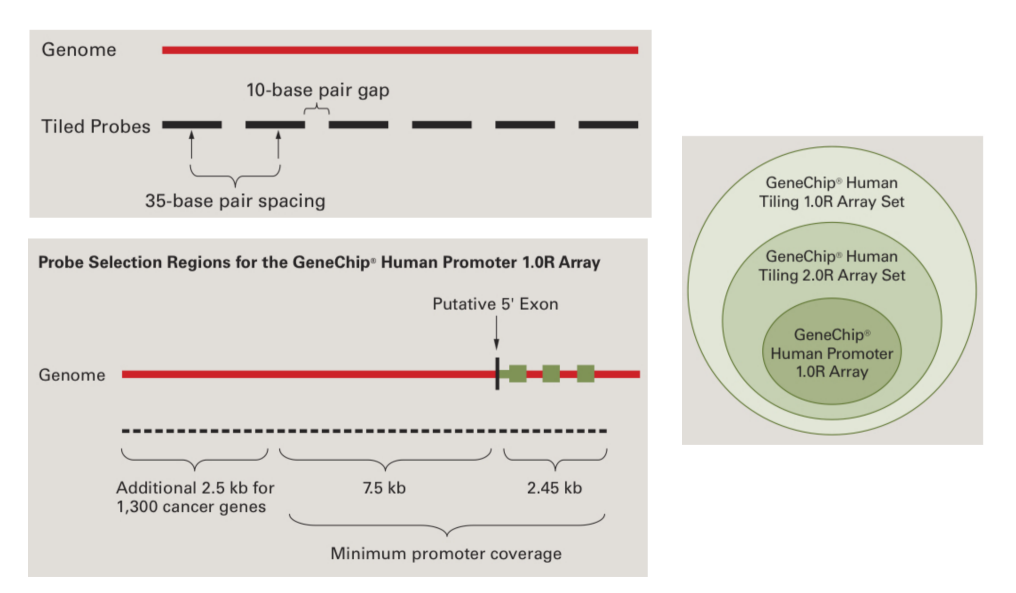
Here a manufactured slide is used in which fragments spanning the genome have been synthesised and attached to the slide surface in a geometric Arrangement. We label our TF retrieved fragments, hybridise them to the slide and then read fluorescence from the features.

#### ChIP-seq

Taking advantage of high throughput sequencing technology (so called next-gen) we attempt to sequence all the fragments. This is quantitative.

In both cases we have issues with mapping, signal processing (noise) and significance testing







Features of genome tiling arrays

•Generally resolution can be as low as ~3kb, Tfs bind to on average 6-8bp

•How do we know which gene to map to ? (meta-data) microarray, gene proximity, functional annotation, in-vivo expression comparison to true positive

•Redundancy probes map to more than one location

 Coverage, cannot cover the genome. This introduces bias. even in Drosophila commonly only 50% of genome possible 2 human chromosomes at 35bp resolution → 1 million features

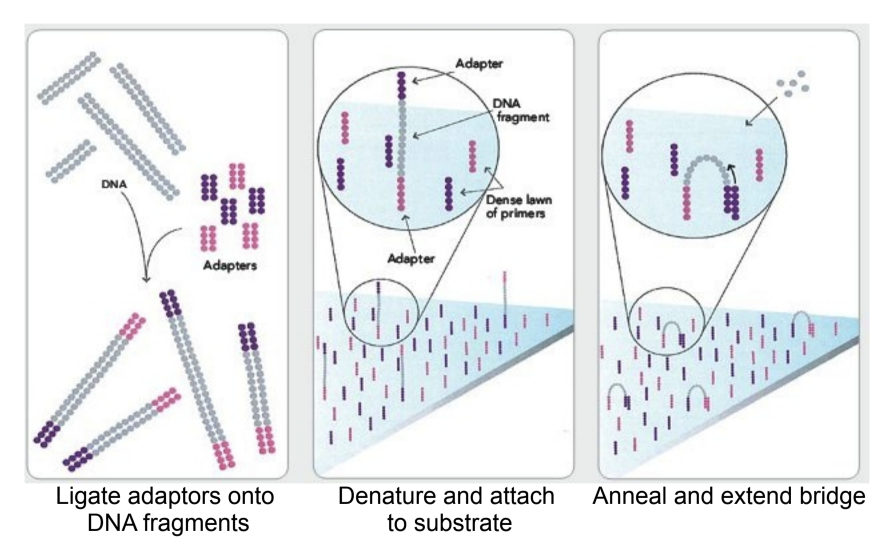
•Can estimate site occupancy frequency

•Cross-hybridisation can be big problem with repetitive DNA (~5% human genome)

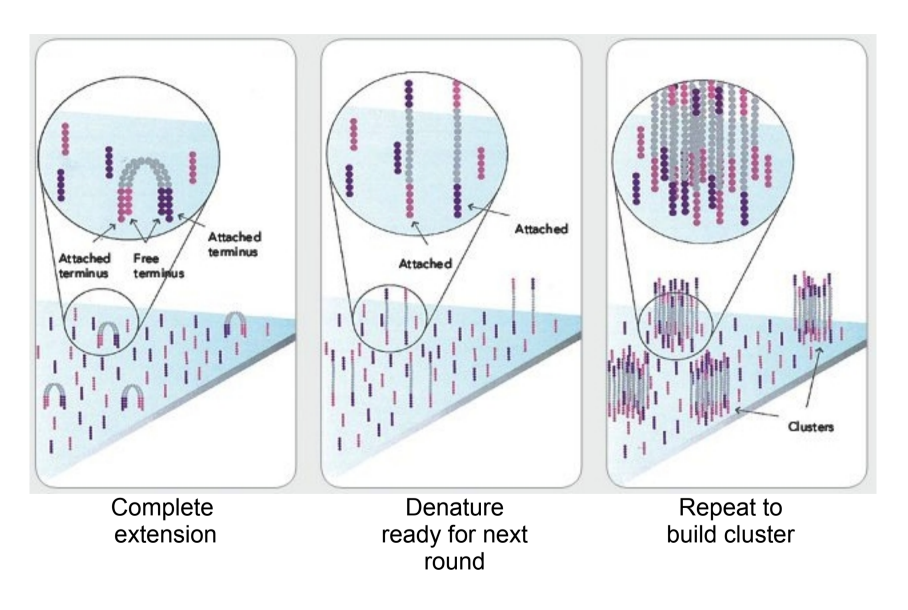
•Processed just like a gene expresison microarray SAM, limma (modelled error, tight control of FDR)



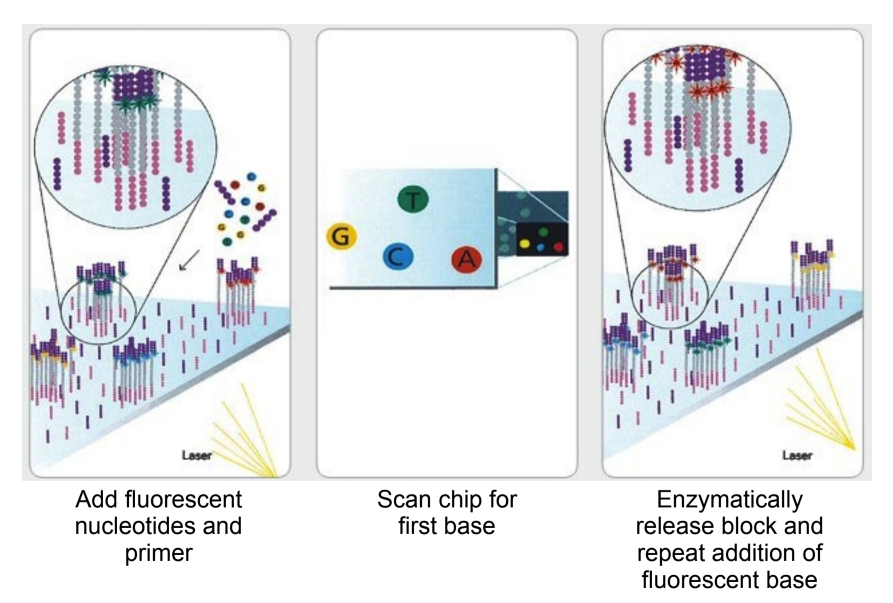
Illumina/Solexa SBS sequencing system



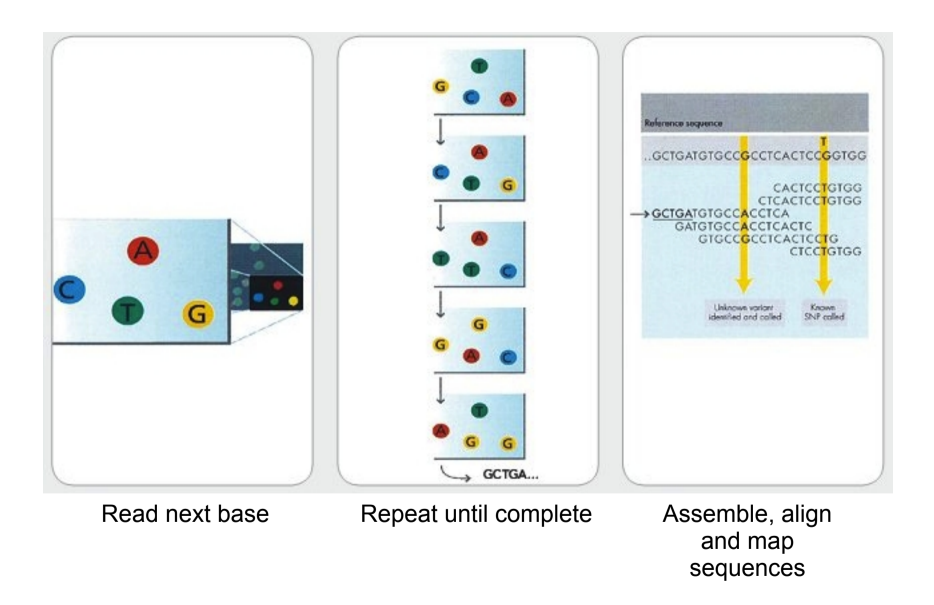














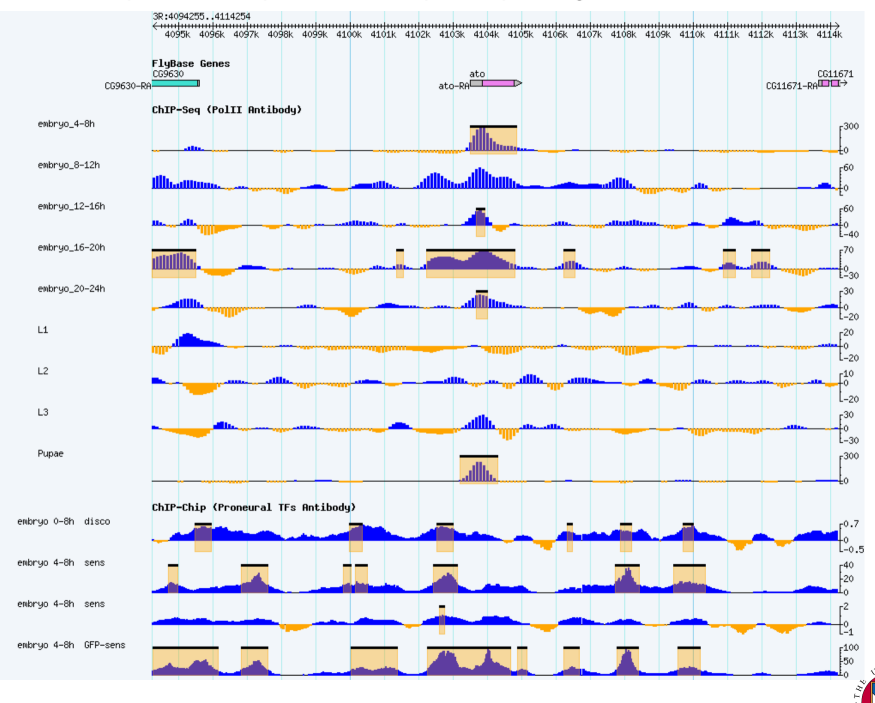
## Features of high-throughput sequence data

•Very high resolution, typically 25-mers with mid-spacing ~35bp

- •Huge datasets, many Gb of sequence, assembly non-trivial
- •Complete genome coverage, no assumption, no bias
- •Generally superior at identifying bound sites beyond expectation (this is related to a more accurate ability to discriminate signal from noise)
- •Sequences are counted to determine the frequency of site occupancy (better than chip, here seq num is proportional to bound sites)
- •Sequences are mapped and converted into signal peaks (typical sizes of bound peaks can range from 50bp-1kb)
- •Strong correlation between statistical significance of peak and presence of binding motif (might seem obvious!)



# Example ChIP-Chip and ChIP-seq data spanning the atonal locus



Ian Simpson, Institute for Adaptive and Neural Computation, School of Informatics, University of Edinburgh

NIV

#### Real world examples of ChIP-chip and ChIP-seq in use



#### Genome-Wide Mapping of in Vivo Protein-DNA Interactions David S. Johnson, et al. Science 316, 1497 (2007); DOI: 10.1126/science.1141319

Developmental Cell 10, 797-807, June, 2006 ©2006 Elsevier Inc. DOI 10.1016/j.devcel.2006.04.009

#### A Temporal Map of Transcription Factor Activity: Mef2 Directly Regulates Target Genes at All Stages of Muscle Development

Thomas Sandmann,<sup>1</sup> Lars J. Jensen,<sup>1</sup> Janus S. Jakobsen,<sup>1</sup> Michal M. Karzynski,<sup>1</sup> Michael P. Eichenlaub,<sup>1</sup> Peer Bork,<sup>1</sup> and Eileen E.M. Furlong<sup>1,\*</sup> <sup>1</sup> European Molecular Biology Laboratory D-69117 Heidelberg Germany

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PLOS GENETICS

# Combinatorial Binding Leads to Diverse Regulatory Responses: Lmd Is a Tissue-Specific Modulator of Mef2 Activity

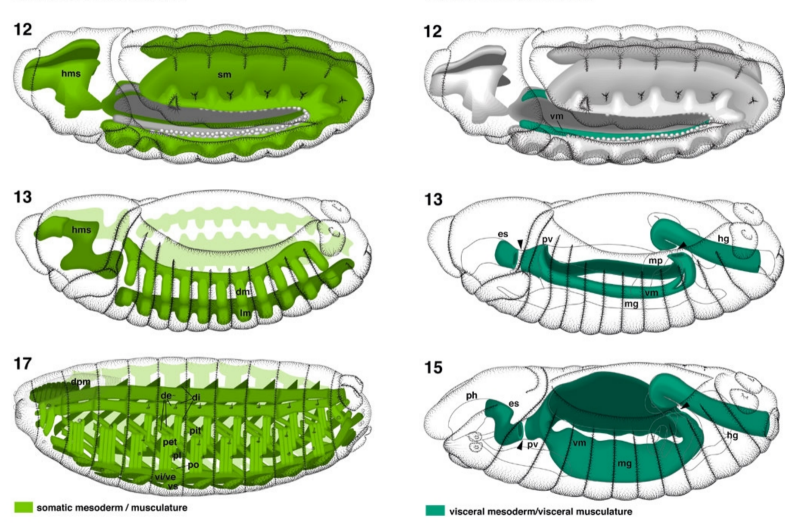
Paulo M. F. Cunha<sup>®</sup>, Thomas Sandmann<sup>®¤a</sup>, E. Hilary Gustafson, Lucia Ciglar, Michael P. Eichenlaub<sup>¤b</sup>, Eileen E. M. Furlong<sup>\*</sup>

European Molecular Biology Laboratory, Heidelberg, Germany



# Studying Drosophila musculature development using ChIP-chip

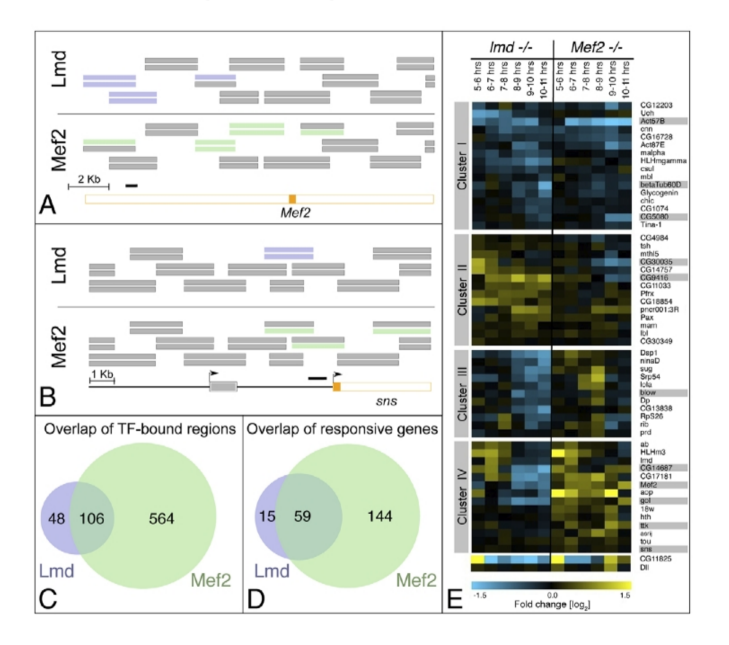
**Visceral Musculature** 



#### Somatic Musculature

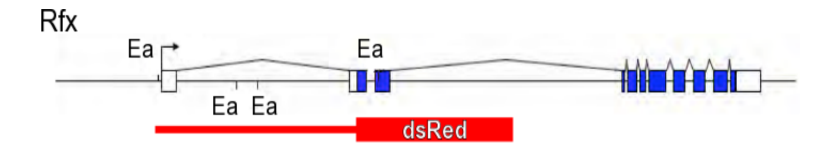


ChIP-chip blocks integrated with gene expression data for Mef2 and Lmd

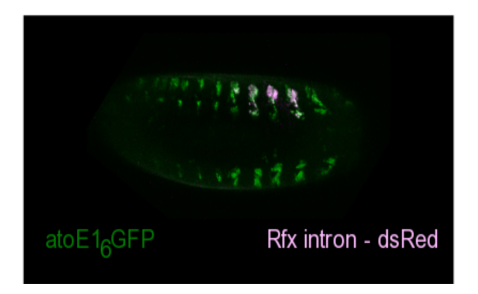




Validation of enhancers and TF binding sites

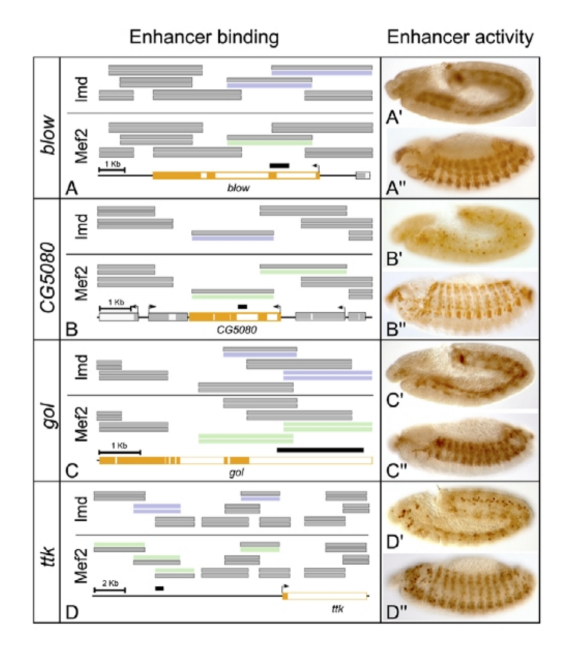


Dmel ATGCTTACCACCACTATTCGAA**CAGCTG**TGAGCGTTGCCACTTGTCTTGAGGATTAACCAA Dsec ATGCTTACCAGCACTGTTCGAA**CAGCTG**TGAGCGTTGCCACTTGTCTTGGGGGGTTAACCAG Dere -TGCTAGCCAACACTGTTCGAA**CAGCTG**GGAGCGTTGCCACTTGTGCTCGCGGGCTAACCAA Dyak ATGTTAACCAACACTGTTCGCA**CAGCTG**TAAGCGTTGCCACTTGTGCTTGCGAATTAGCCAG



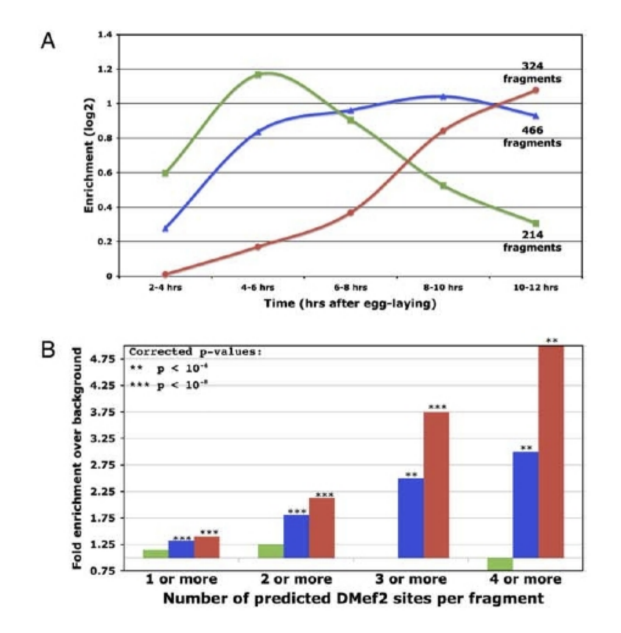


## Validation of enhancer activity for Mef2/Lmd candidate target genes



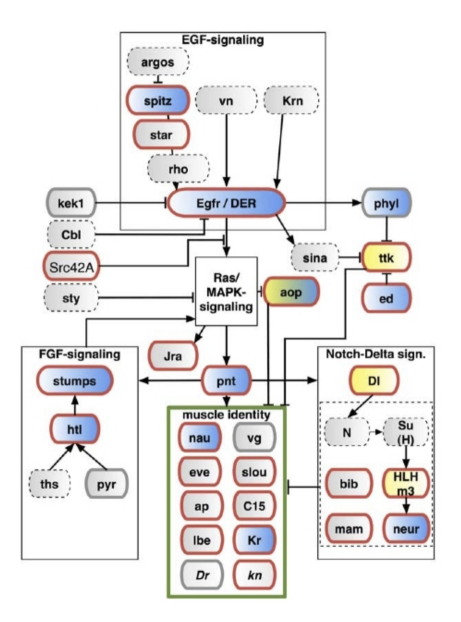


#### Temporal binding profiles of over-represented Mef2 bound blocks



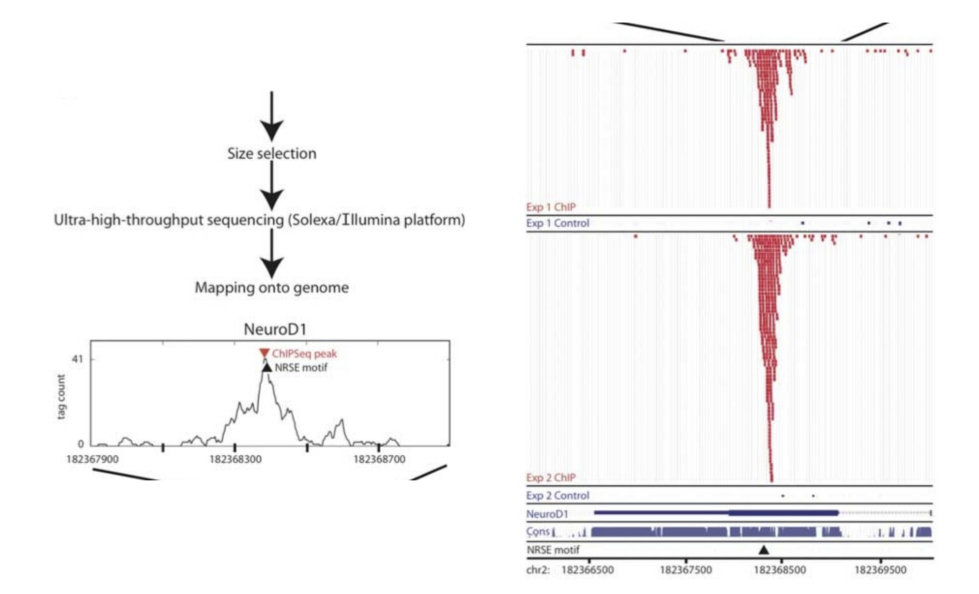


Synthesis of the target gene network and known myogensis pathway



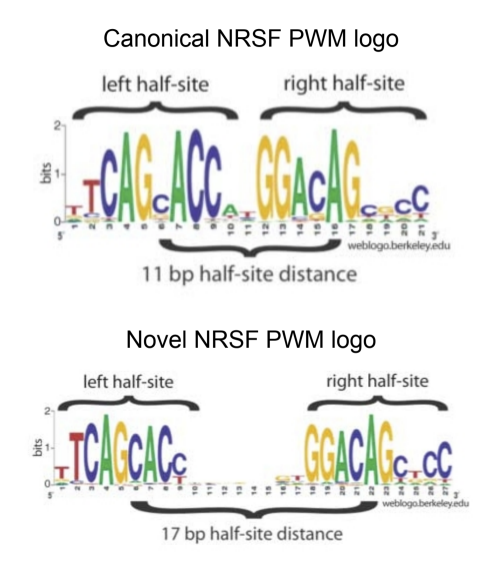


#### Chip-seq analysis of the neuron restrictive silencing factor (NRSF)



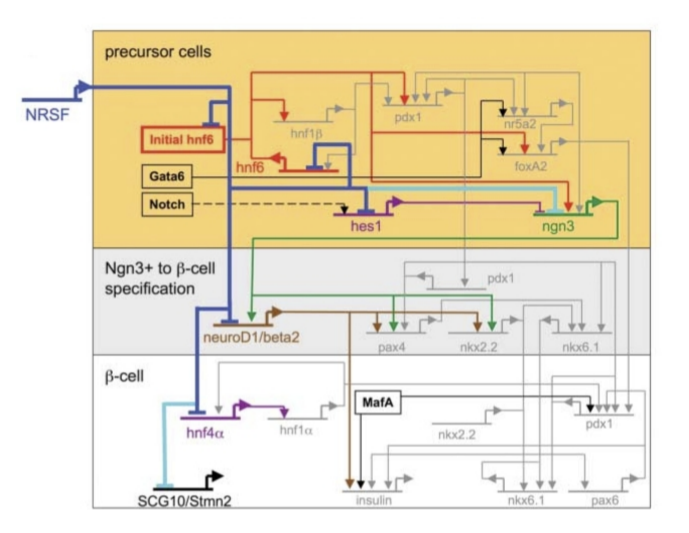


## ChIP-seq reveals new binding motif flexibility for NRSF





The gene regulatory network downstream of NRSF constructed from ChIP-seq data



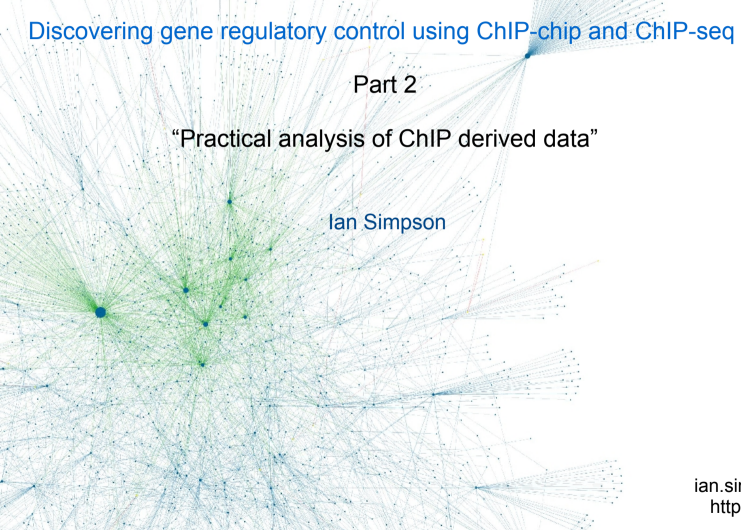


# Summary for ChIP based target prediction methods

•ChIP-chip and ChIP-seq allow for the first time physical identification of bound regions on the genomic scale

- •ChIP-seq presents higher resolution and is replacing ChIP-chip
- •Both methods require large data-processing and analysis
- •Novel methods have been developed to call bound regions from these data they are predominantly based on hidden markov models (HMM) and are naturally normally 2-state models (peak, non-peak)
- •The resulting regions can be used with classical methods to refine the nature of the regulatory element (PWM Gibbs/HMM profiling, motif detection, conservation)
- •Can also be refined by more precise experiments on the ChIP DNA such as targeted PCR
- •Revolutionises the analysis of gene regulatory networks by integration with gene expression data





ian.simpson@.ed.ac.uk http://bit.ly/bio2links



# **BIOINFORMATICS**



#### A hidden Markov model for analyzing ChIP-chip experiments on genome tiling arrays and its application to p53 binding sequences

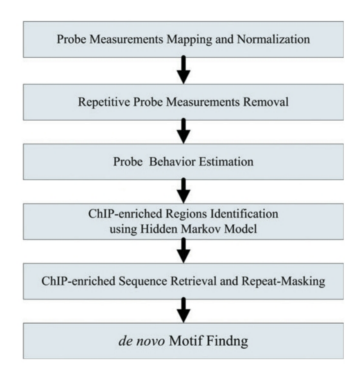
Wei Li, Clifford A. Meyer and X. Shirley Liu\*

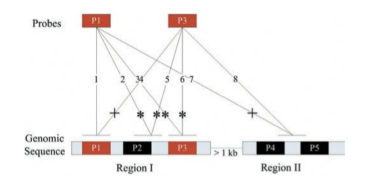
Department of Biostatistics and Computational Biology, Dana-Farber Cancer Institute, Harvard School of Public Health, Boston, MA 02115, USA

Received on January 15, 2005; accepted on March 27, 2005



#### HMMtiling





- (1) Initial probabilities: J/K for ChIP-enriched state, 1 J/K for non-enriched state.
- (2) Transition probabilities: J/K for transition to a different state, 1 J/K for staying in the same state.
- (3) Emission probability distribution of probe *i* in single dataset:  $N(\mu_i + 2\sigma_i, (1.5\sigma_i)^2)$  for ChIP-enriched state,  $N(\mu_i, \sigma_i^2)$  for non-enriched state. The parameters are based on the results on the Affymetrix SNP arrays (Lieberfarb *et al.*, 2003).
- (4) A probe *i*, with (PM-MM) value  $p_i$ , is defined as an outlier if its *Z*-value is >3 or <-2.5. We reassigned the *Z*-value of each outlier probe as 3 if *Z* > 3 and -2.5 if *Z* < -2.5.
- (5) If two adjacent probes are farther apart than 500 bp in the genome (usually due to a long repeat sequence between the two probes), in the forward and backward procedure, the enriched and non-enriched state probabilities of the latter probe are reset to the initial probabilities.



Qin et al. BMC Bioinformatics 2010, **11**:369 http://www.biomedcentral.com/1471-2105/11/369



METHODOLOGY ARTICLE

**Open Access** 

# HPeak: an HMM-based algorithm for defining read-enriched regions in ChIP-Seq data

Zhaohui S Qin<sup>\*1,2,3</sup>, Jianjun Yu<sup>3,4</sup>, Jincheng Shen<sup>1</sup>, Christopher A Maher<sup>2,3,4</sup>, Ming Hu<sup>1</sup>, Shanker Kalyana-Sundaram<sup>3,4</sup>, Jindan Yu<sup>5</sup> and Arul M Chinnaiyan<sup>2,3,4,6,7,8</sup>



# HPeak

#### Hypothetical DNA fragment

