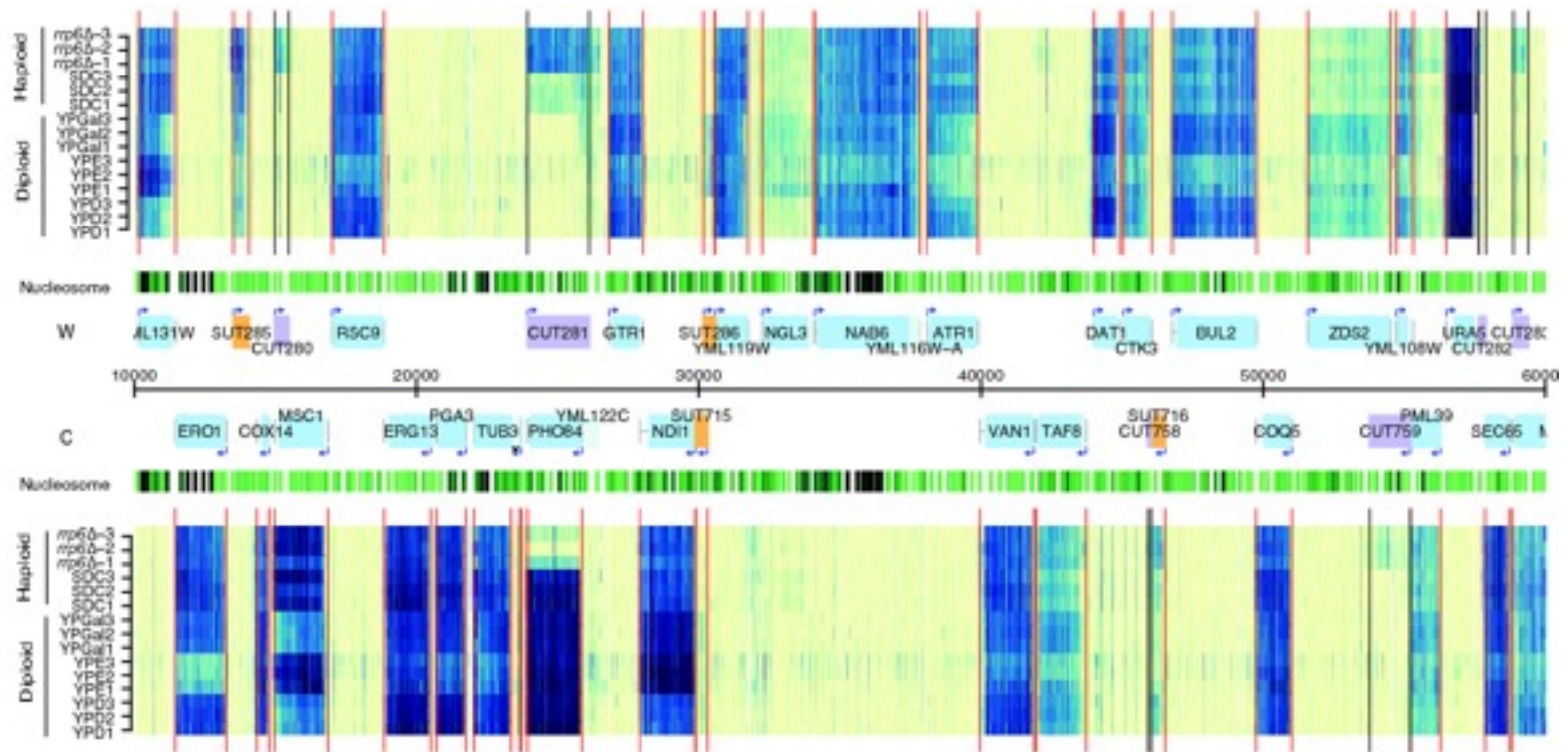


What you still might want to know about microarrays



Brixen, 2 July 2012
Wolfgang Huber
EMBL

Brief history

Late 1980s: Lennon, Lehrach: cDNAs spotted on nylon membranes

1990s: Affymetrix adapts microchip production technology for in situ oligonucleotide synthesis (commercial, patent-fenced)

1990s: Brown lab in Stanford develops two-colour spotted array technology (open and free)

1998: Yeast cell cycle expression profiling on spotted arrays (Spellmann) and Affymetrix (Cho)

1999: Tumor type discrimination based on mRNA profiles (Golub)

2000-ca. 2004: Affymetrix dominates the microarray market

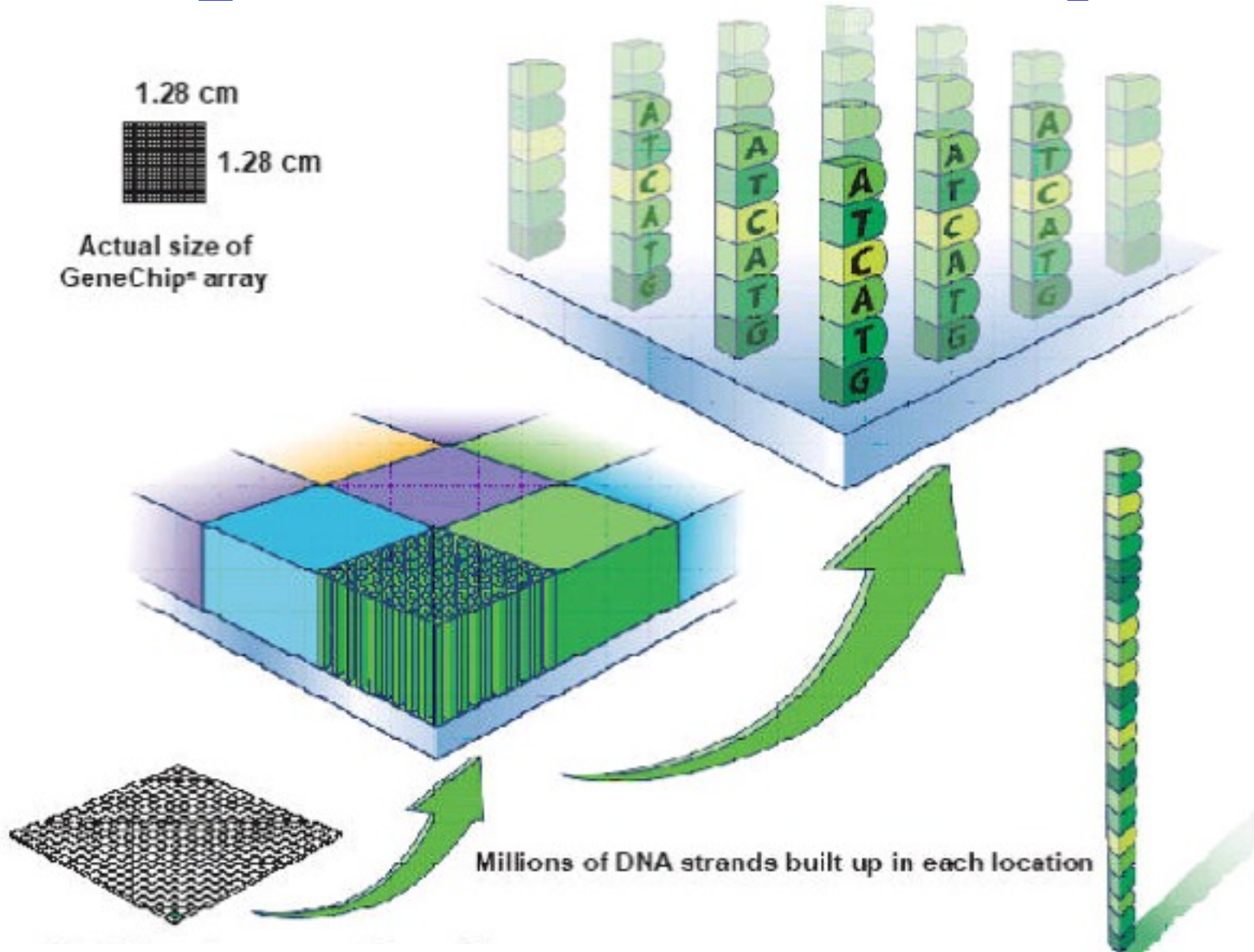
Since ~2003: Nimblegen, Illumina, Agilent (and others)

Throughout 2000's: CGH, CNVs, SNPs, ChIP, tiling arrays

Since ~2007: 2nd-generation sequencing (454, Solexa)

Oligonucleotide microarrays

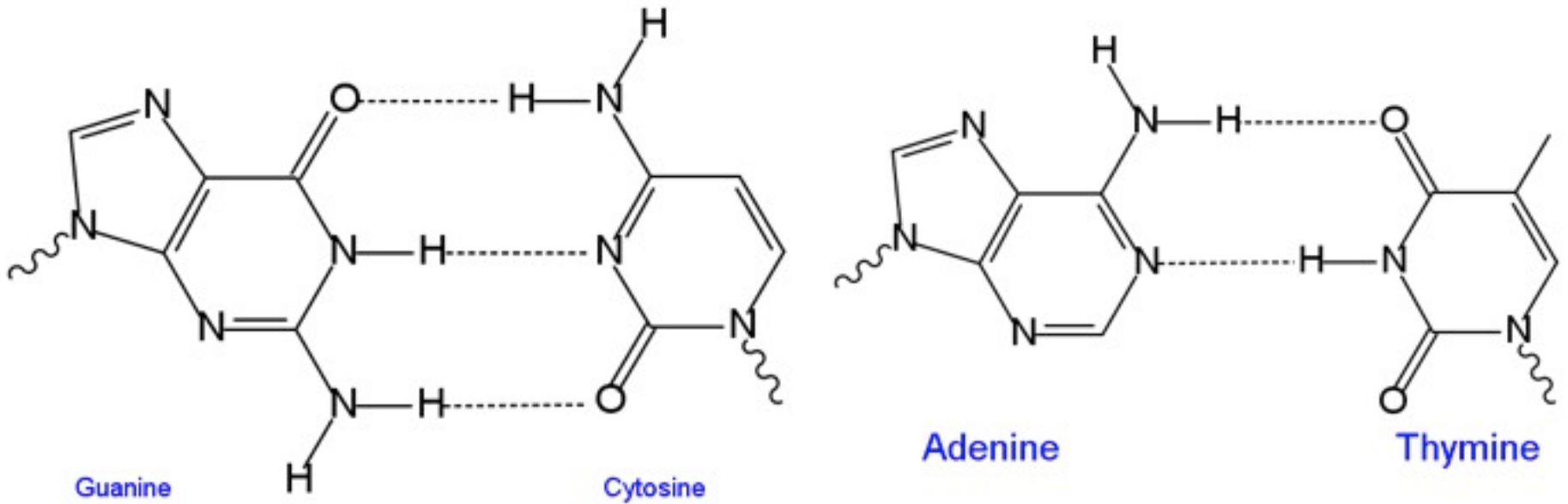
1.28 cm
1.28 cm
Actual size of
GeneChip® array



5 μm , $2560^2 = 6.5$ Mio features

Actual strand = 25 base pairs

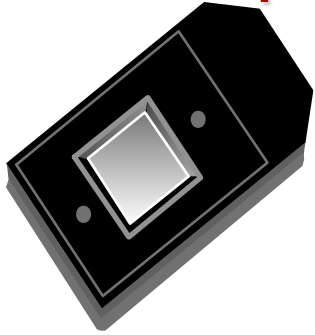
Base Pairing



Ability to use hybridisation for constructing specific + sensitive probes at will is unique to DNA (cf. proteins, RNA, metabolites)

Oligonucleotide microarrays

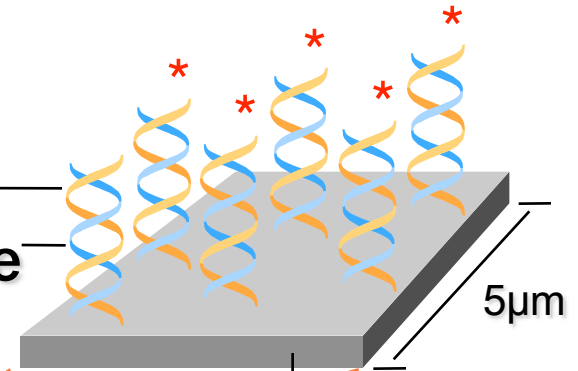
GeneChip



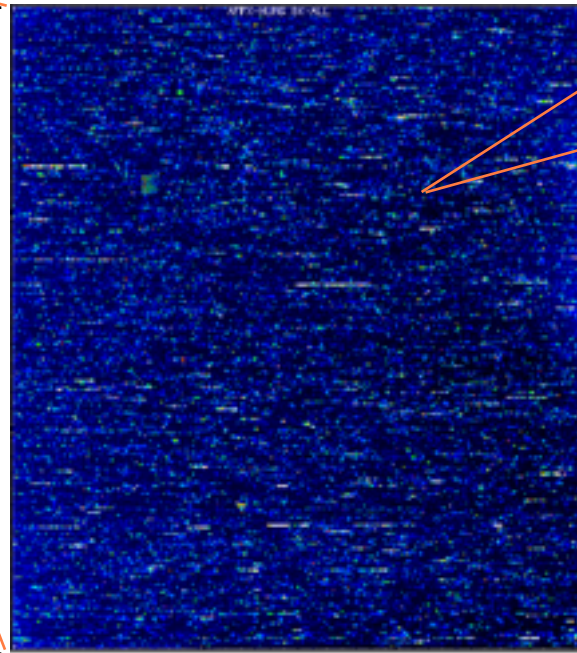
Target - single stranded cDNA

oligonucleotide probe

Hybridized Probe Cell



1.28cm



millions of copies of a specific oligonucleotide probe molecule per cell

up to 6.5 Mio different probe cells

Image of array after hybridisation and staining

Probe sets

GeneChip® Expression Array Design

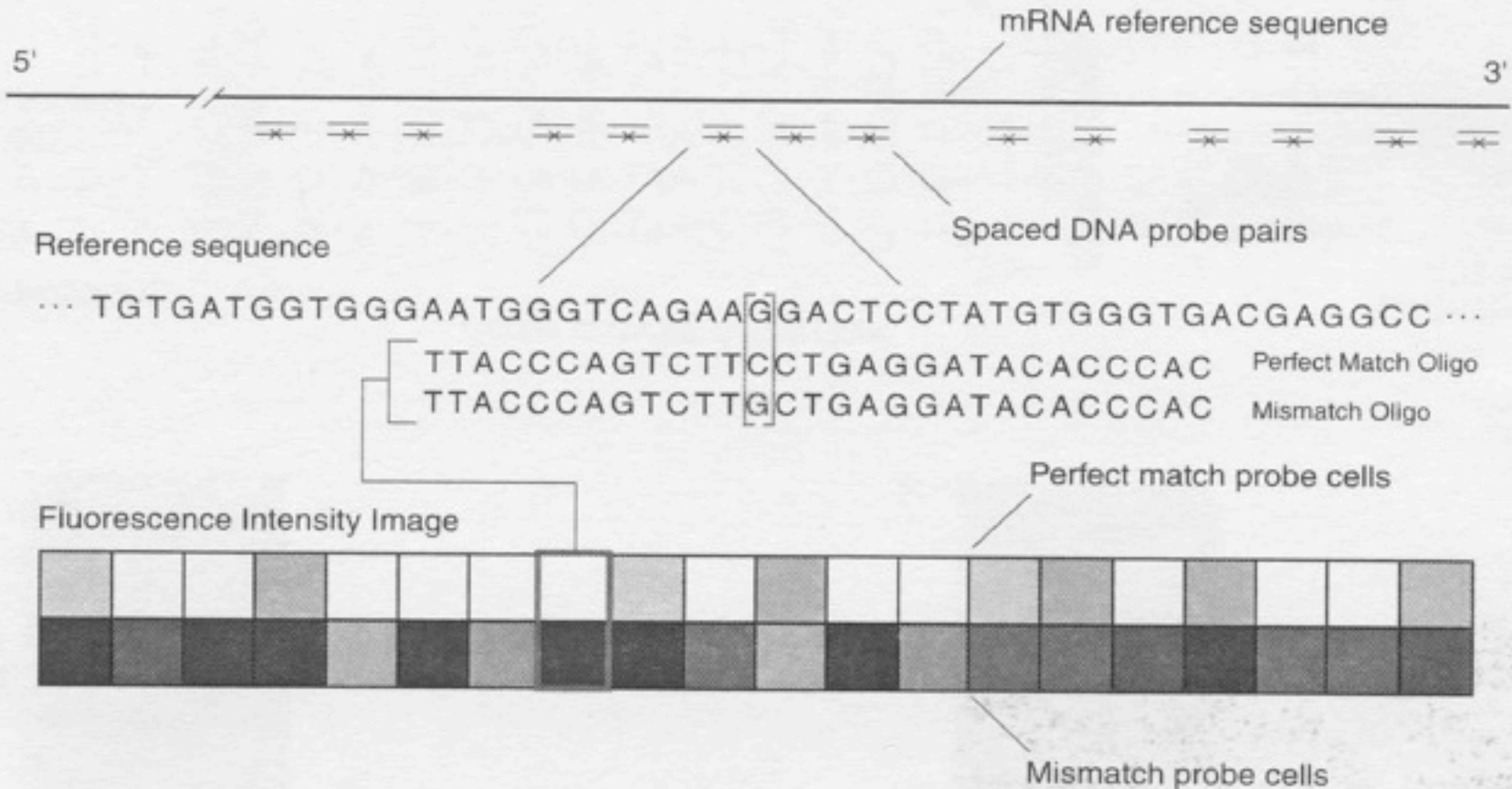


Figure 1-3 Expression tiling strategy

Terminology for transcription arrays

Each target molecule (transcript) is represented by several oligonucleotides of (intended) length 25 bases

Probe: one of these 25-mer oligonucleotides

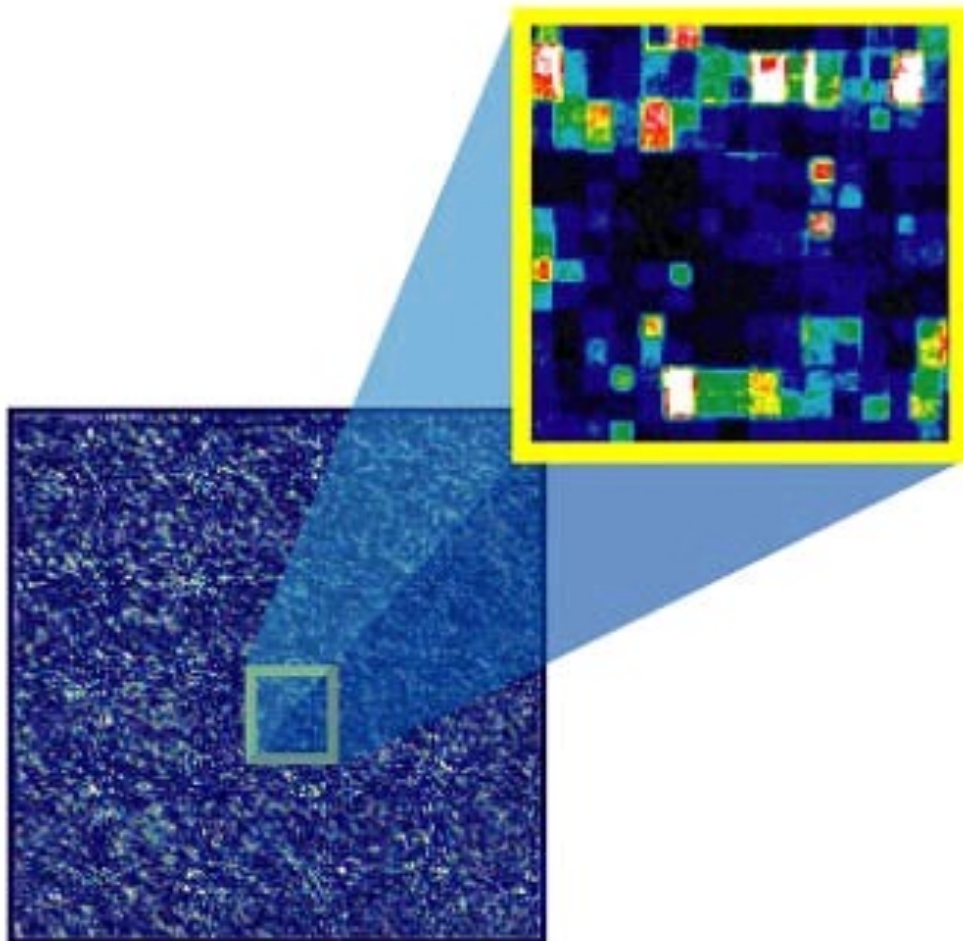
Probe set: a collection of probes (e.g. 11) targeting the same transcript

MGED/MIAME: „probe“ is ambiguous!

Reporter: the sequence

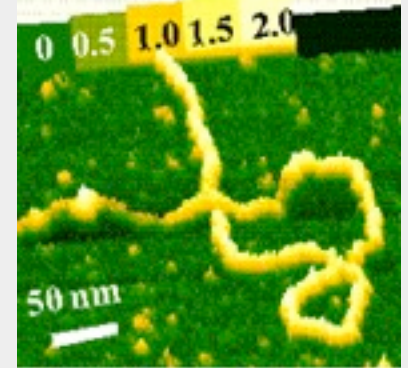
Feature: a physical patch on the array with molecules intended to have the same reporter sequence (one reporter can be represented by multiple features)

Image analysis



- **several dozen pixels per feature**
 - **segmentation**
 - **summarisation into one number representing the intensity level for this feature**
- **CEL file**

μ array data



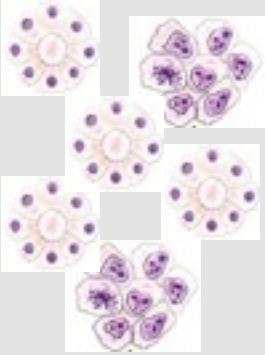
arrays:

probes =

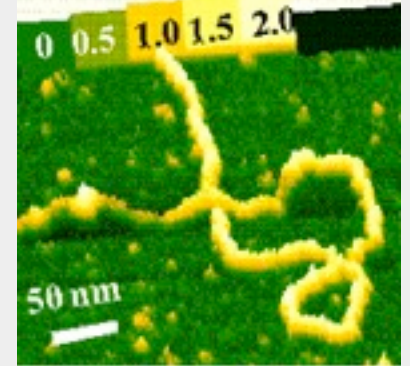
gene-specific

DNA strands

μ array data

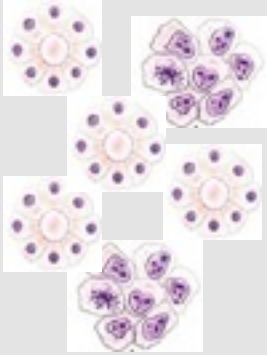


samples:
mRNA from
tissue
biopsies,
cell lines

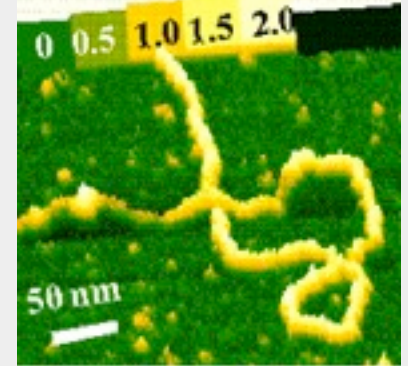
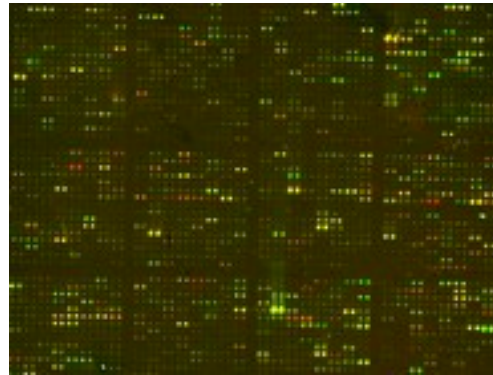


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μ array data

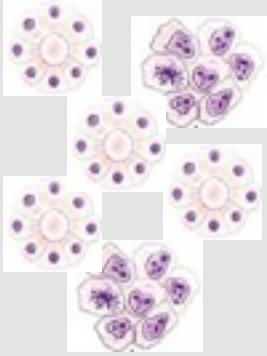


samples:
mRNA from
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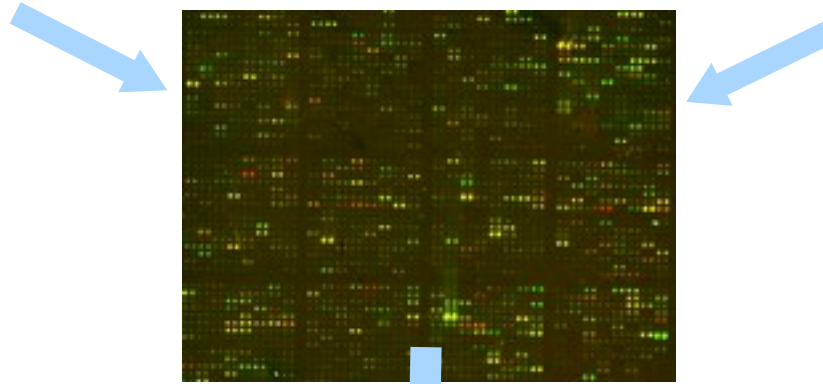


arrays:
probes =
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DNA strands

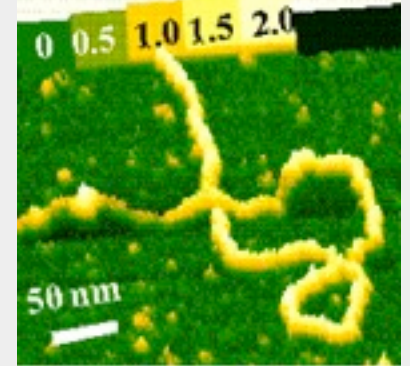
μ array data



samples:
mRNA from
tissue
biopsies,
cell lines



**fluorescent detection
of the amount of
sample-probe binding**



arrays:
probes =
gene-specific
DNA strands

	tissue A	tissue B	tissue C
ErbB2	0.02	1.12	2.12
VIM	1.1	5.8	1.8
ALDH4	2.2	0.6	1.0
CASP4	0.01	0.72	0.12
LAMA4	1.32	1.67	0.67
MCAM	4.2	2.93	3.31

Microarray Infrastructure in Bioconductor

Platform-specific data import and initial processing

Affymetrix 3' IVT (e.g. Human U133 Plus 2.0, Mouse 430 2.0):

`affy`

Affymetrix Exon (e.g. Human Exon 1.0 ST):

`oligo, exonmap, xps`

Affymetrix SNP arrays:

`oligo`

Nimblegen tiling arrays (e.g. for ChIP-chip):

`Ringo`

Affymetrix tiling arrays (e.g. for ChIP-chip):

`Starr`

Illumina bead arrays:

`beadarray, lumi`

<http://www.bioconductor.org/docs/workflows/oligoarrays>

Flexible data import

Using generic R I/O functions and constructors

Biobase

limma

Chapter *Two Color Arrays* in the useR-book.

limma user guide

Normalisation and quality assessment

preprocessCore

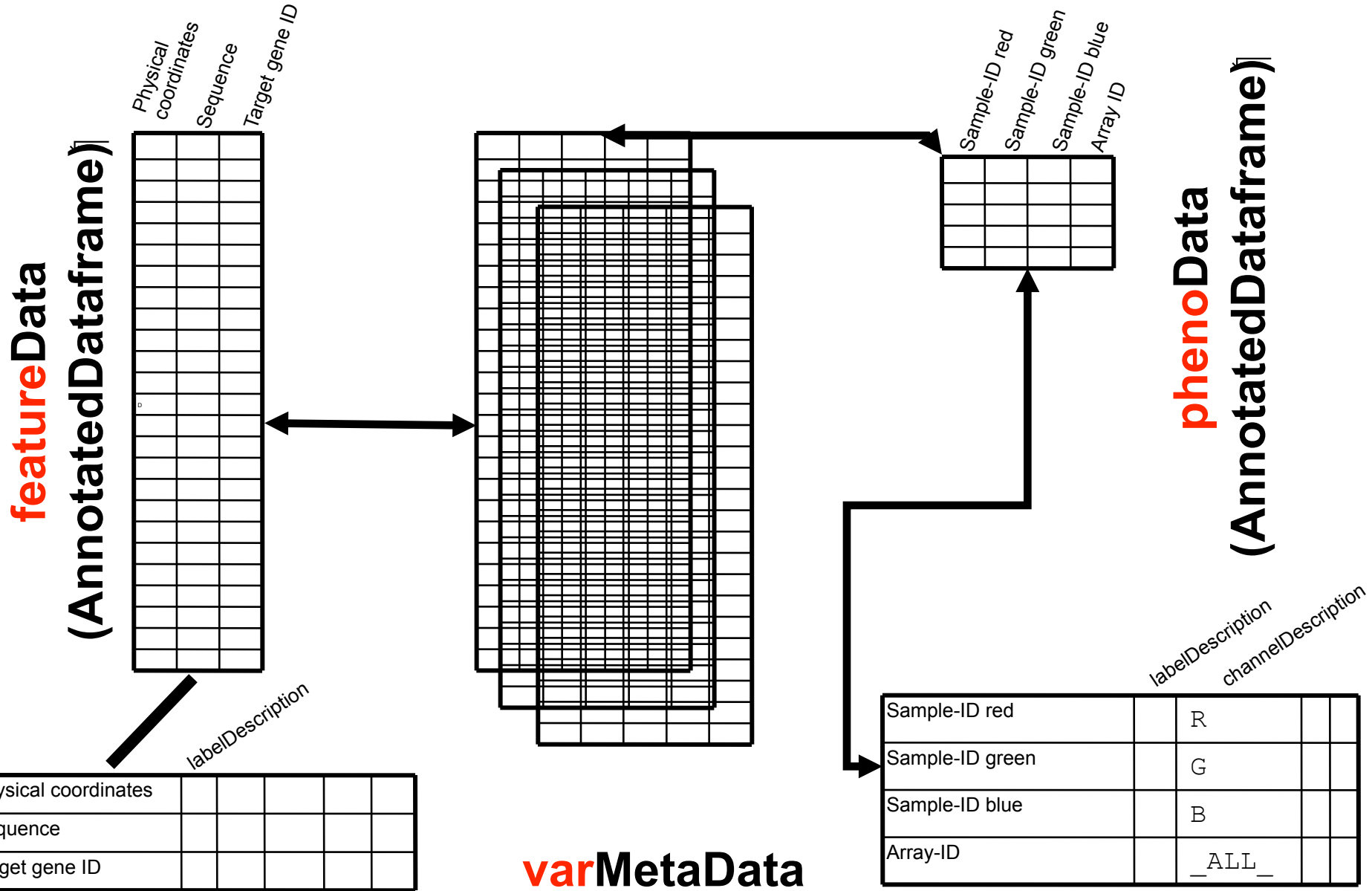
limma

vsn

arrayQualityMetrics

NChannelSet

assayData can contain $N=1, 2, \dots$, matrices of the same size



Annotation / Metadata

Keeping data together with the metadata (about reporters, target genes, samples, experimental conditions, ...) is one of the major principles of Bioconductor

- avoid alignment bugs
- facilitate discovery

Often, the same microarray design is used for multiple experiments. Duplicating that metadata every time would be inefficient, and gene annotation for reporters can get out of date ⇒

instead of `featureData`, just keep a pointer to an annotation package.

(In principle, one could also want to do this for samples.)

Annotation infrastructure for Affymetrix

For `affy`:

`hgu133plus2probe` nucleotide sequence of the features (for preprocessing e.g. `gcrma`; for own annotation)

`hgu133plus2cdf` maps the physical features on the array to probe sets

`hgu133plus2.db` maps probe sets to target genes and provides target gene annotation collected from public databases

Genotyping

cr1mm Genotype Calling (CRLMM) and Copy Number Analysis tool for Affymetrix SNP 5.0 and 6.0 and Illumina arrays.

snpMatrix

.... others

See also:

Genome-wide association study of CNVs in 16,000 cases of eight common diseases and 3,000 shared controls, The Wellcome Trust Case Control Consortium, Nature 464 (2010), p. 713-720(Box 1).

Gene expression analysis with microarrays

Microarray Analysis Tasks

Data import

reformatting and setup/curation of the metadata

Normalisation

Quality assessment & control

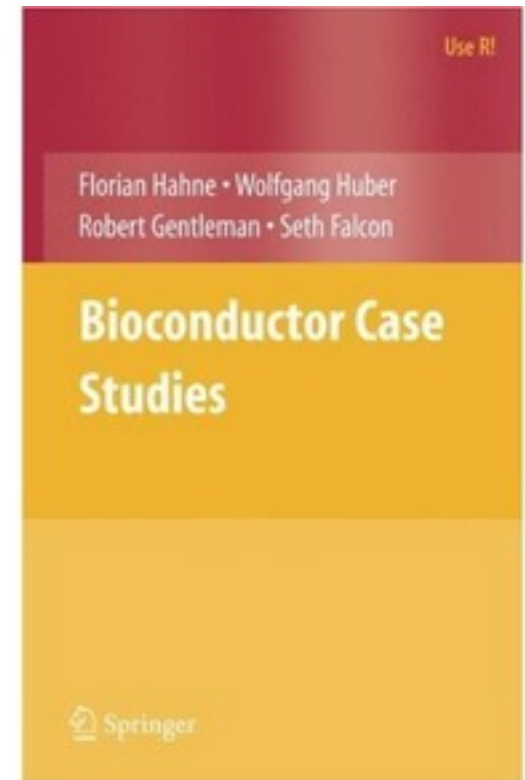
Differential expression

Using gene-level annotation

Gene set enrichment analysis

Clustering & Classification

Integration of other datasets



▶ What is wrong with microarray data?

Many data are measured in definite units:

- time in seconds
- lengths in meters
- energy in Joule, etc.

Climb Mount Plose (2465 m) from Brixen (559 m) with weight of 76 kg, working against a gravitation field of strength 9.81 m/s^2 :

$$\begin{aligned} & (2465 - 559) \cdot 76 \cdot 9.81 \text{ m kg m/s}^2 \\ & = 1\,421\,037 \text{ kg m}^2 \text{ s}^{-2} \\ & = 1\,421.037 \text{ kJ} \end{aligned}$$

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A complex measurement process lies between mRNA concentrations and intensities

- RNA degradation
- amplification efficiency
- reverse transcription efficiency
- hybridization efficiency and specificity
- labeling efficiency
- quality of actual probe sequences (vs intended)
- scratches and spatial gradients on the array
- cross-talk across features
- cross-hybridisation
- optical noise
- image segmentation
- signal quantification
- signal "preprocessing"

A complex measurement process lies between mRNA concentrations and intensities

- RNA degradation

- quality of actual probe sequences

- image segmentation

- a
eff

- r
tra
eff

- h
eff
specimen

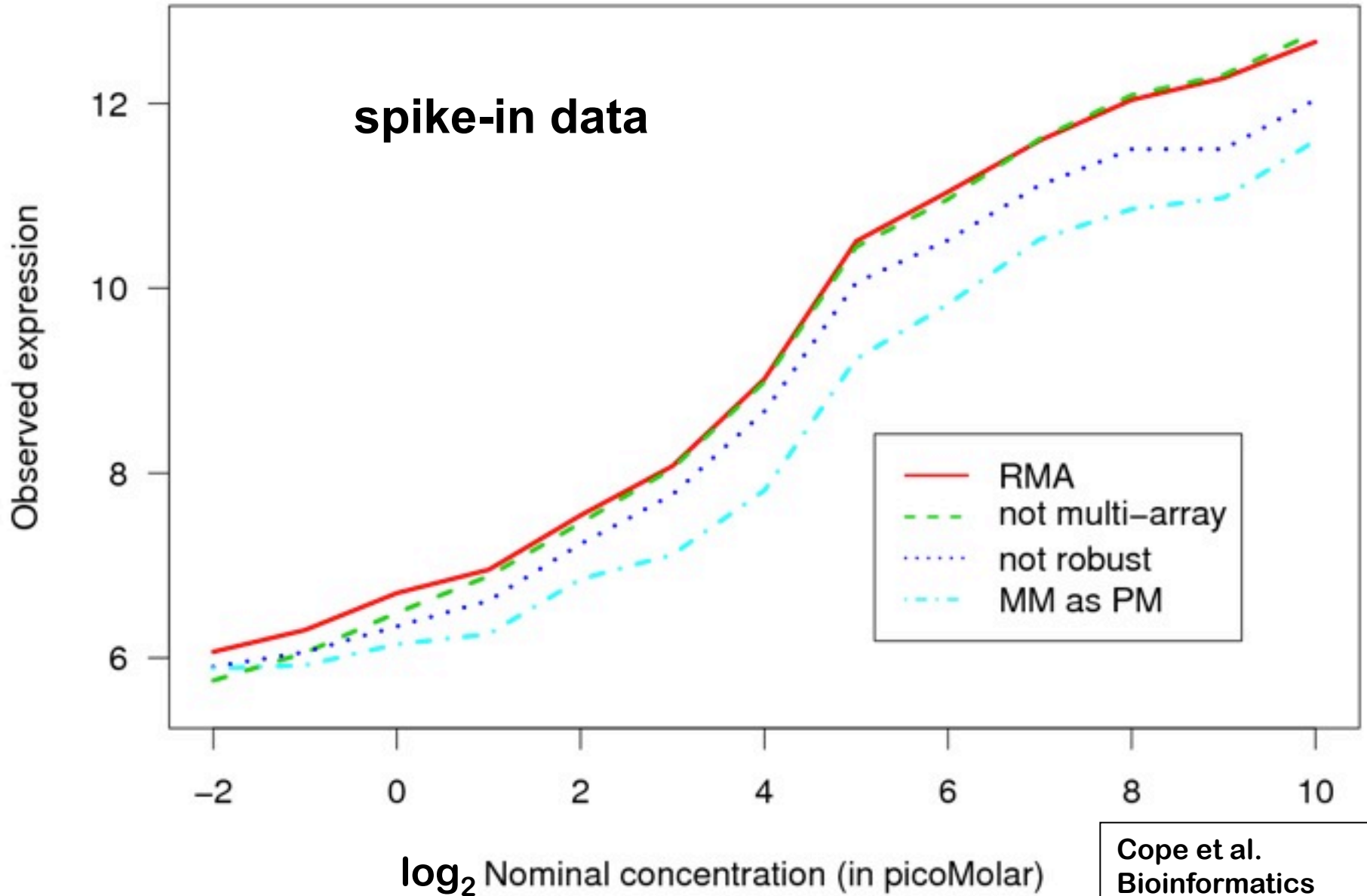
- labeling efficiency

- optical noise

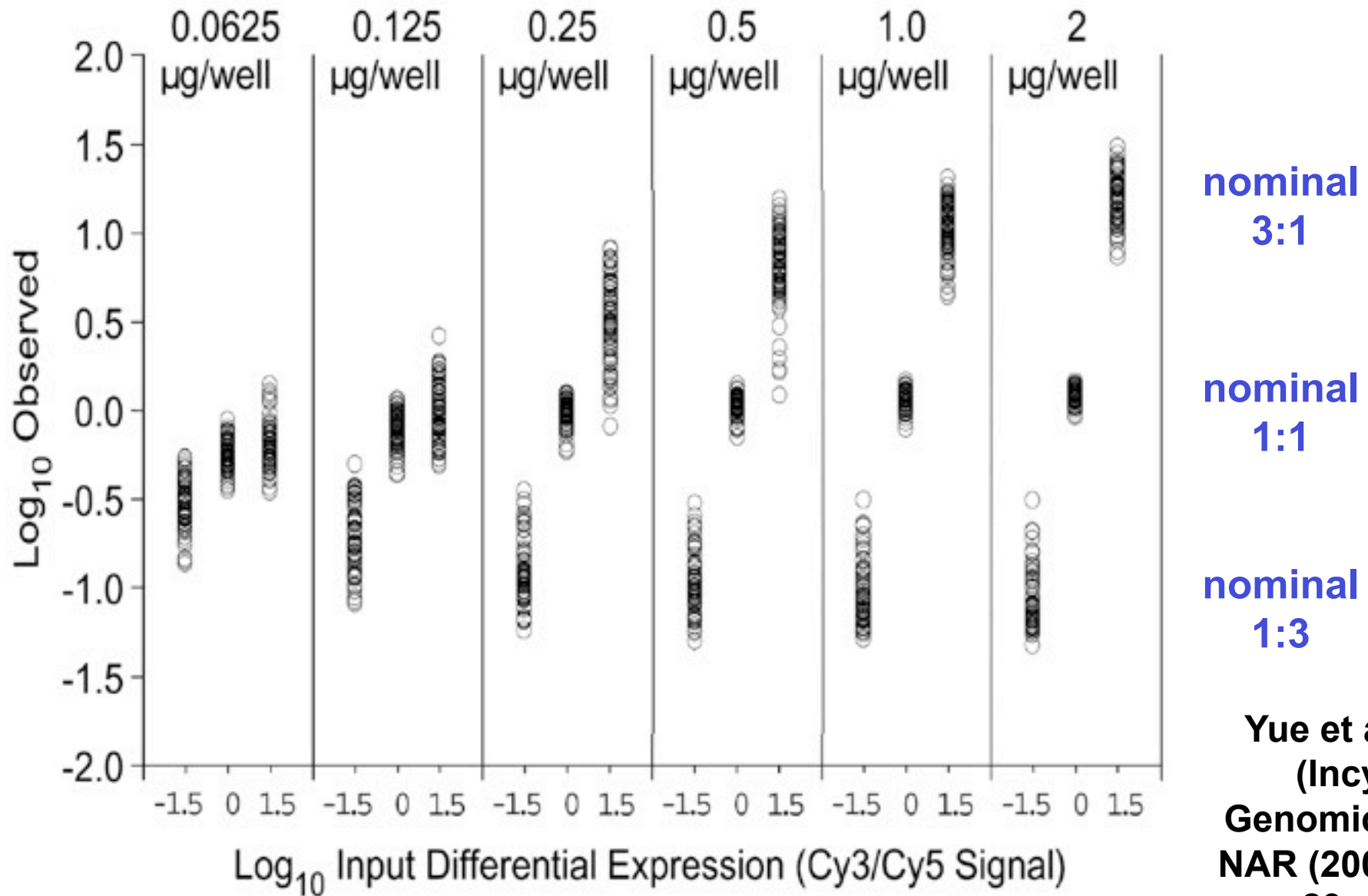
The problem is less that these steps are 'not perfect'; it is that they vary from array to array, experiment to experiment.

Background signal and non-linearities

“mild” non-linearity



► ratio compression



Preprocessing Terminology

Calibration, normalisation: adjust for systematic drifts associated with dye, array (and sometimes position within array)

Background correction: adjust for the non-linearity at the lower end of the dynamic range

Transformation: bring data to a scale appropriate for the analysis (e.g. logarithm; variance stabilisation)

Log-ratio: adjust for unknown scale (units) of the data

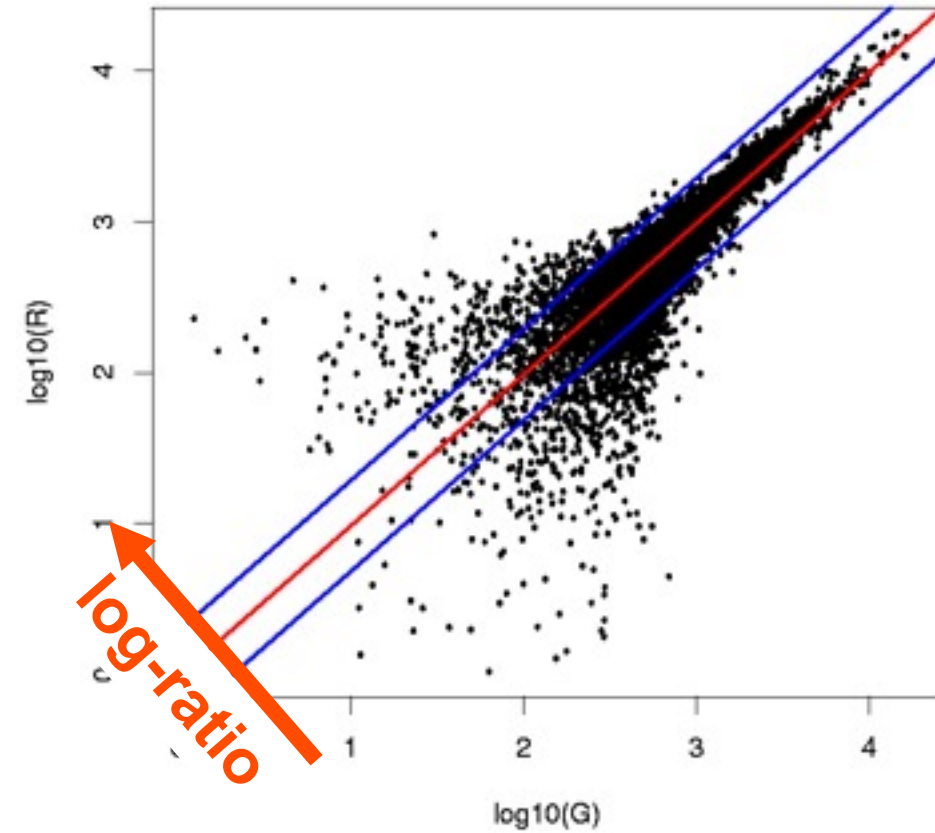
Existing approaches differ in the order in which these steps are done, some are exactly stepwise („greedy“), others aim to gain strength by doing things simultaneously.

Statistical issues



► Which genes are differentially transcribed?

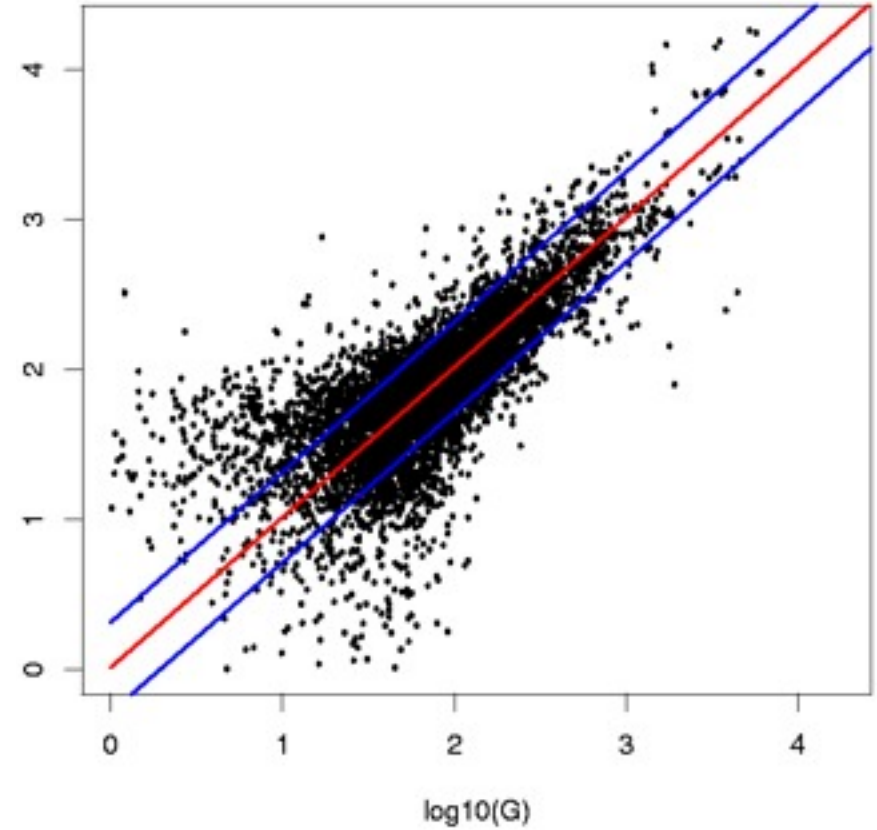
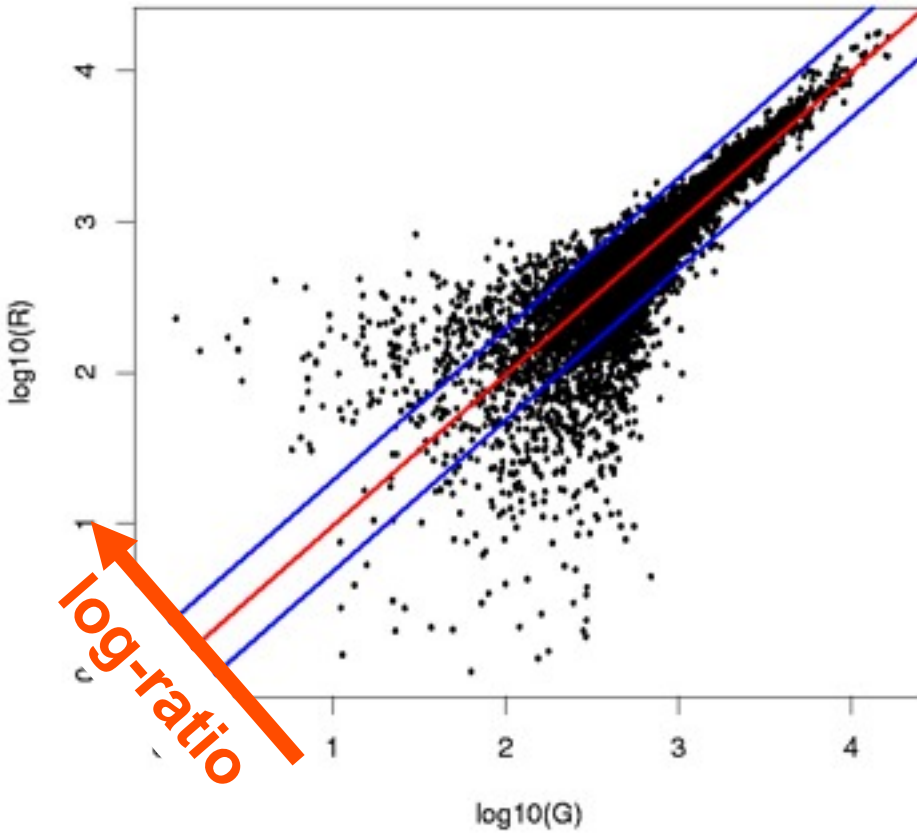
same-same



► Which genes are differentially transcribed?

same-same

tumor-normal



Sources of variation

amount of RNA in the biopsy

efficiencies of

-RNA extraction

-reverse transcription

-labeling

-fluorescent detection

probe purity and length

distribution

spotting efficiency, spot size

cross-/unspecific hybridization

stray signal

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Systematic

- similar effect on many measurements
- corrections can be estimated from data

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efficiencies of

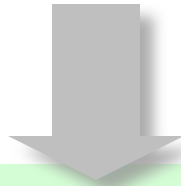
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Calibration

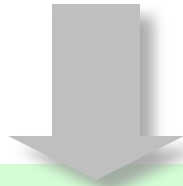
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Stochastic

- too random to be explicitly accounted for
- remain as “noise”

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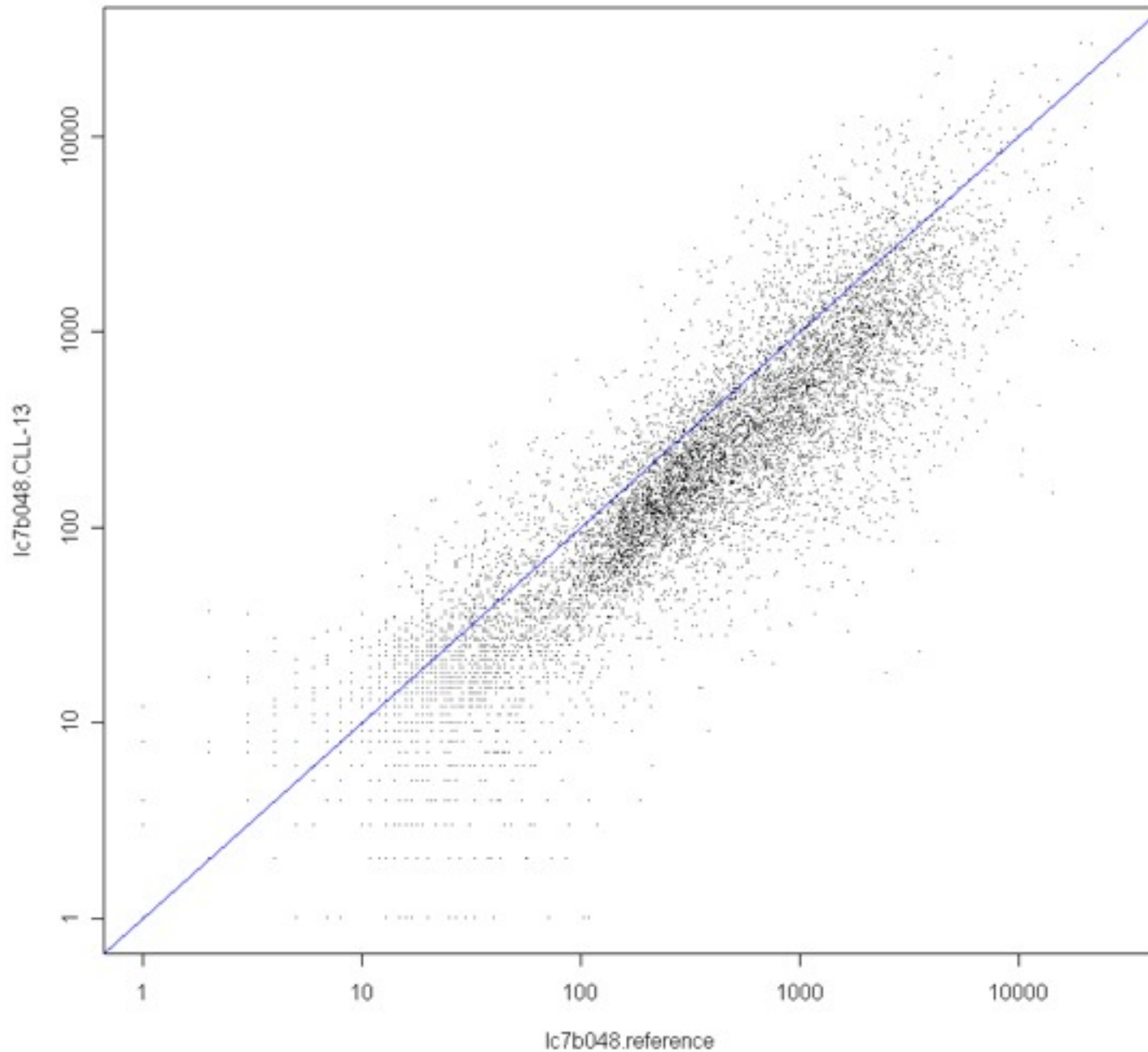
Stochastic

- too random to be explicitly accounted for
- remain as “noise”

Error model

**Why do you need
'normalisation'
(a.k.a. calibration)?**

Systematic effects



From: lymphoma
dataset

vsn package

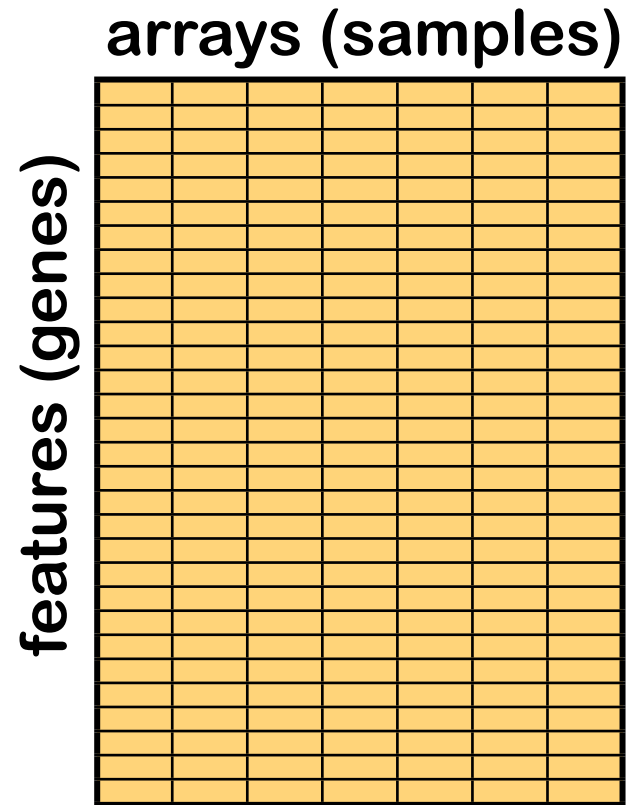
Alizadeh et al.,
Nature 2000

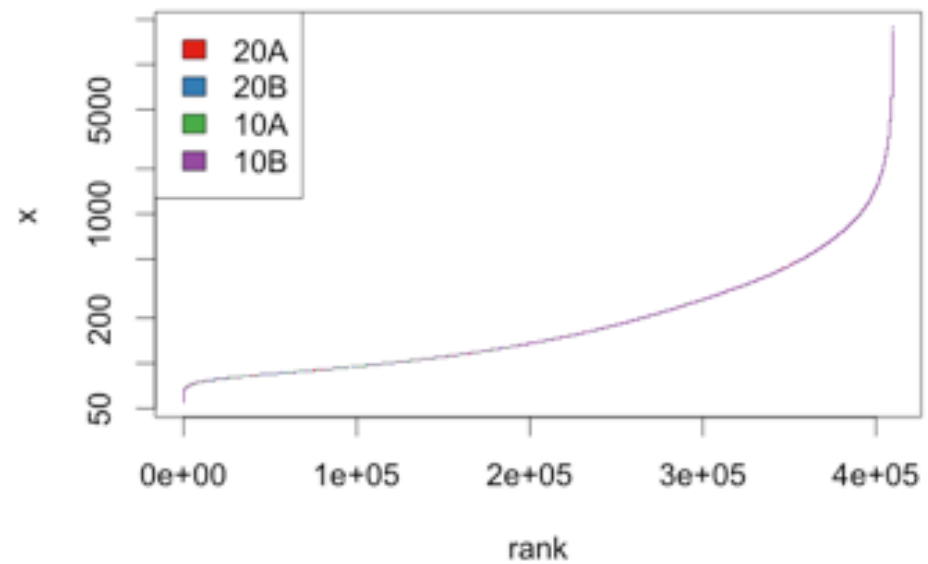
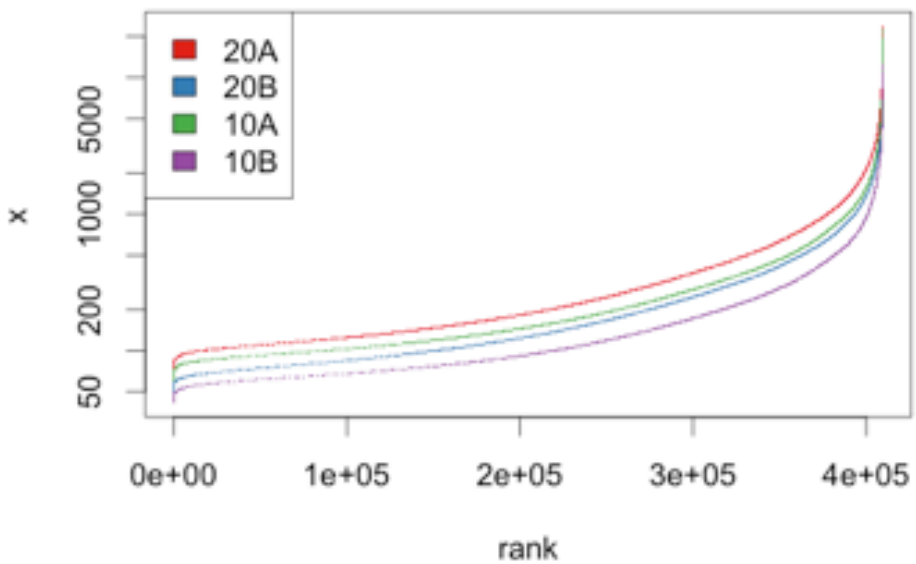
Quantile normalisation

Within each column (array),
replace the intensity values by
their rank

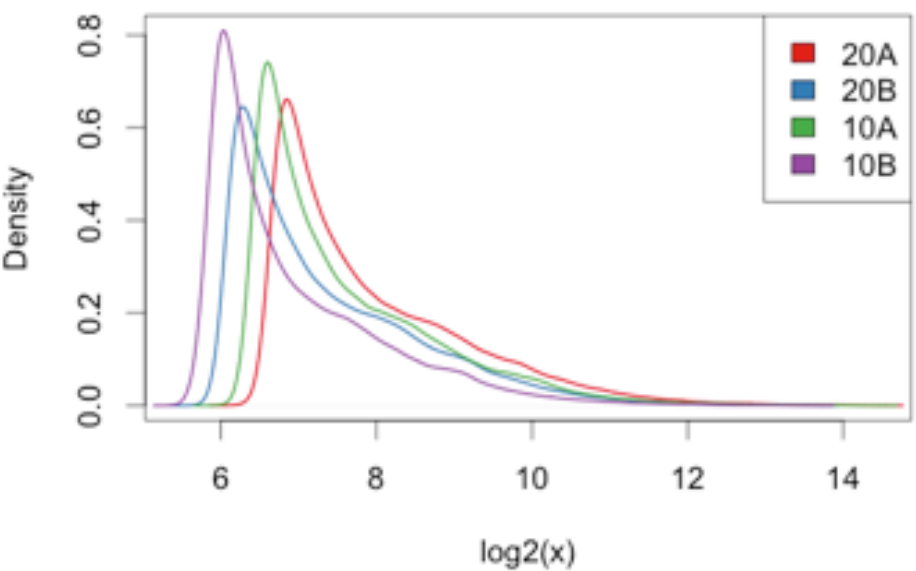
For each rank, compute the
average of the intensities with
that rank, across columns
(arrays)

Replace the ranks by those
averages

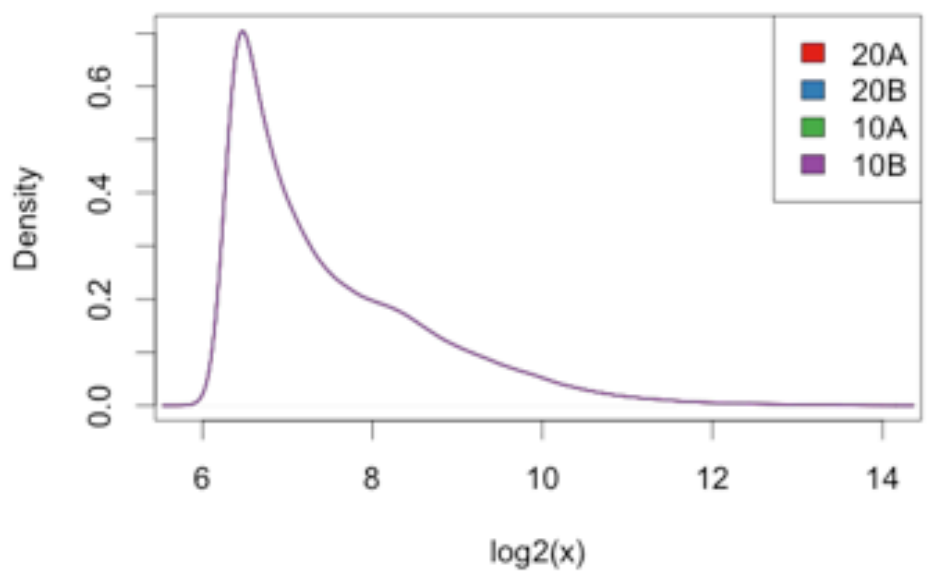




densities



densities



Quantile normalisation

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- + Simple, fast, easy to implement

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Quantile normalisation

- + Simple, fast, easy to implement**
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- May be conservative: rank transformation loses information - may yield less power to detect differentially expressed genes**

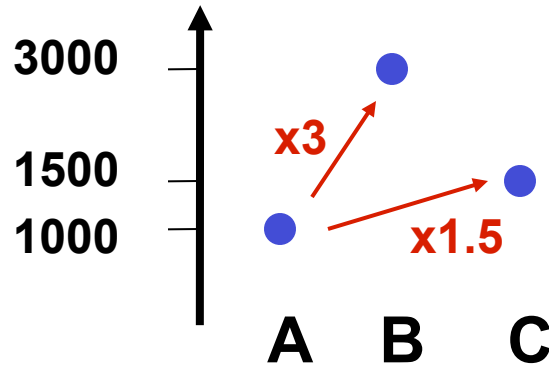
Quantile normalisation

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- Always "works", even if data are bad / inappropriate**
- May be conservative: rank transformation loses information - may yield less power to detect differentially expressed genes**
- Aggressive: if there is an excess of up- (or down) regulated genes, it removes not just technical, but also biological variation**

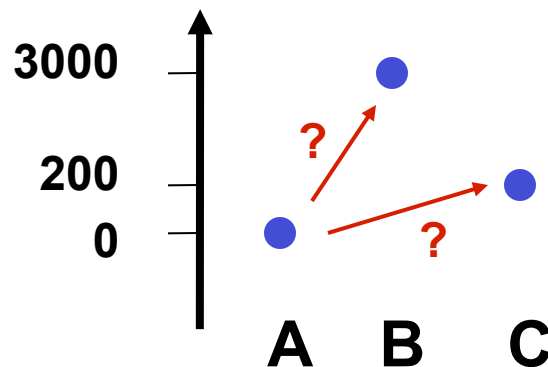
Estimating relative expression (fold-changes)

▶ ratios and fold changes

Fold changes are useful to describe continuous changes in expression



But what if the gene is “off” (below detection limit) in one condition?



▶ ratios and fold changes

The idea of the log-ratio (base 2)

0: no change

+1: up by factor of $2^1 = 2$

+2: up by factor of $2^2 = 4$

-1: down by factor of $2^{-1} = 1/2$

-2: down by factor of $2^{-2} = 1/4$

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A unit for measuring changes in expression: assumes that a change from 1000 to 2000 units has a similar biological meaning to one from 5000 to 10000.

.... **data reduction**

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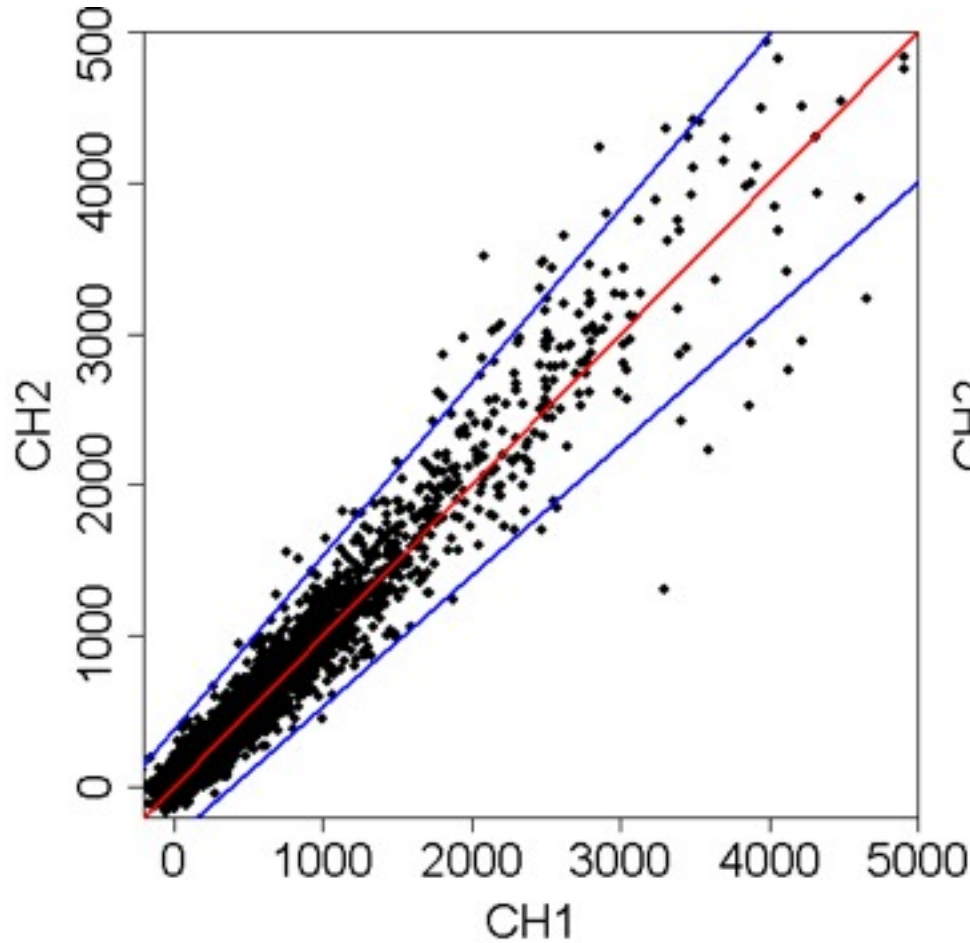
.... **data reduction**

What about a change from 0 to 500?

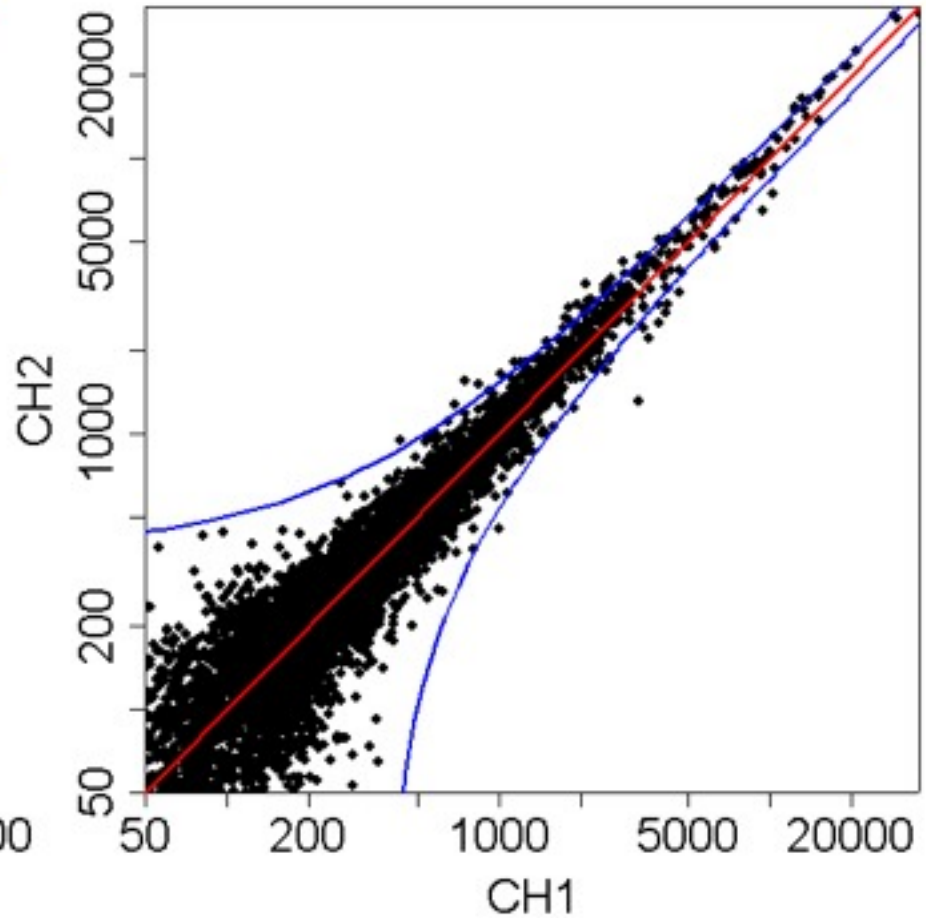
- conceptually

- noise, measurement precision

The two-component model for microarray data

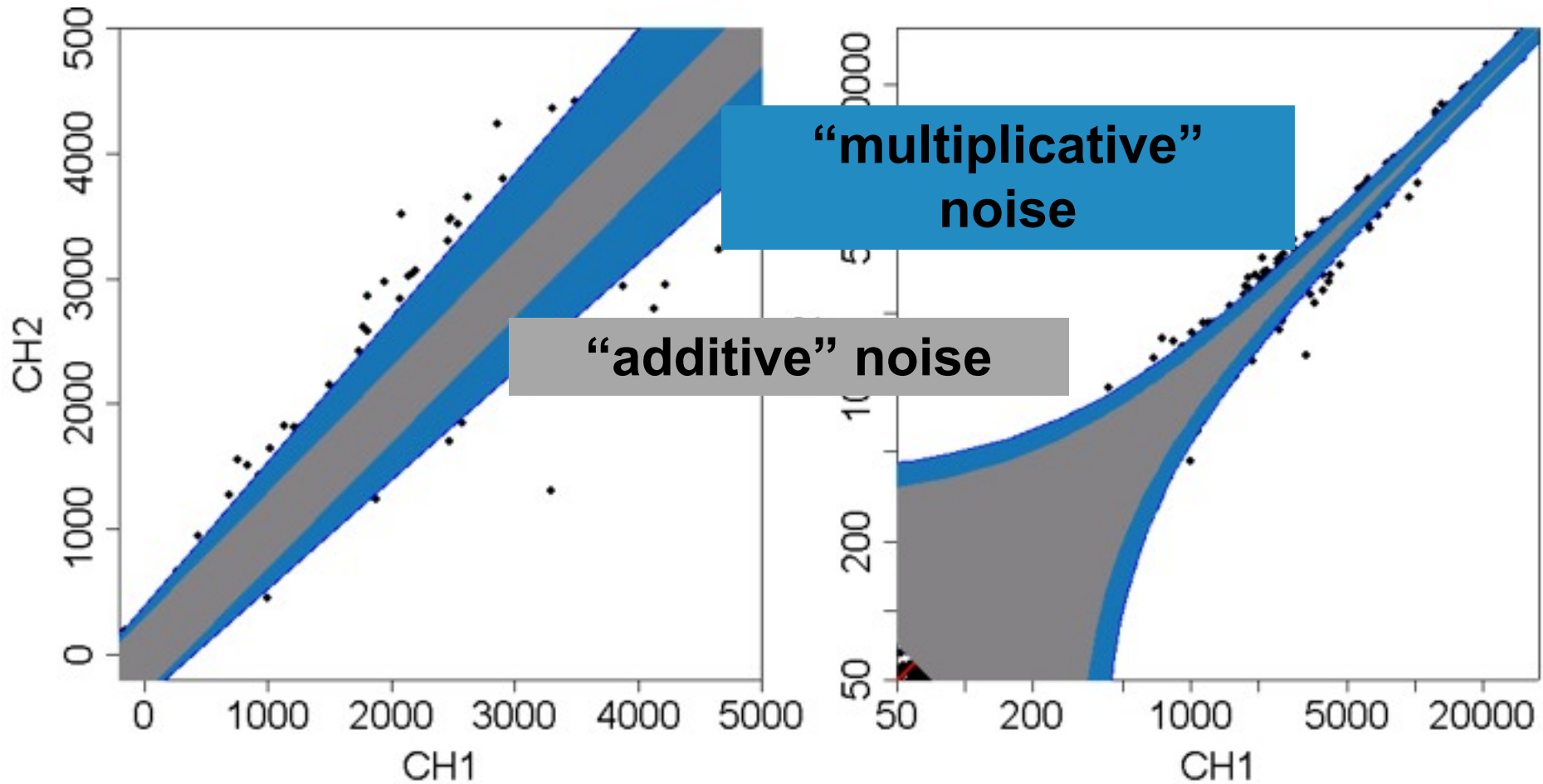


raw scale



log scale

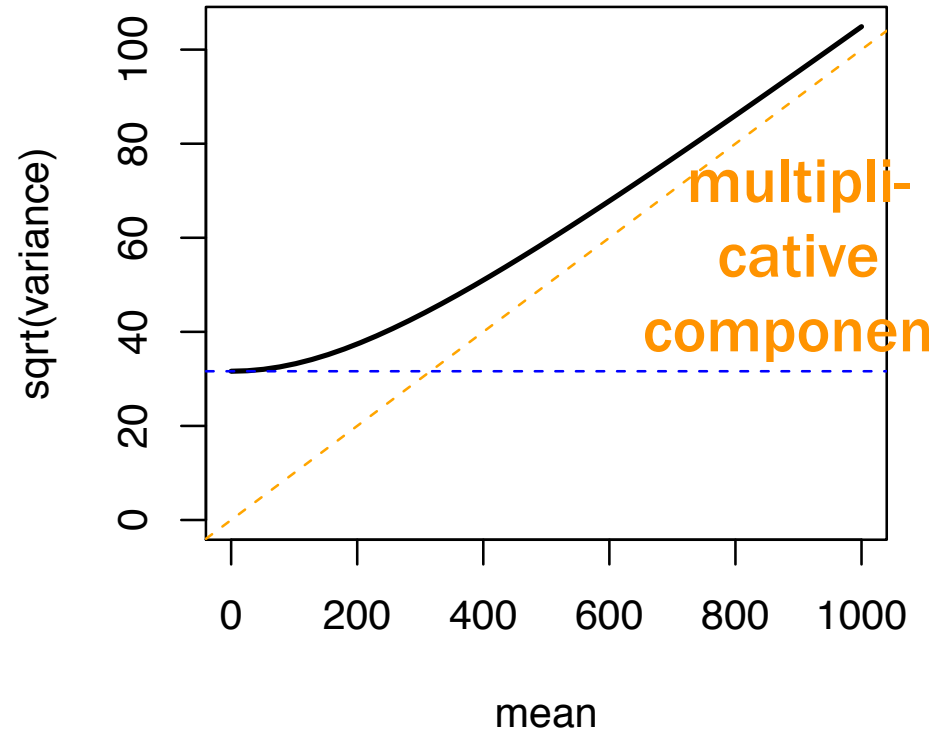
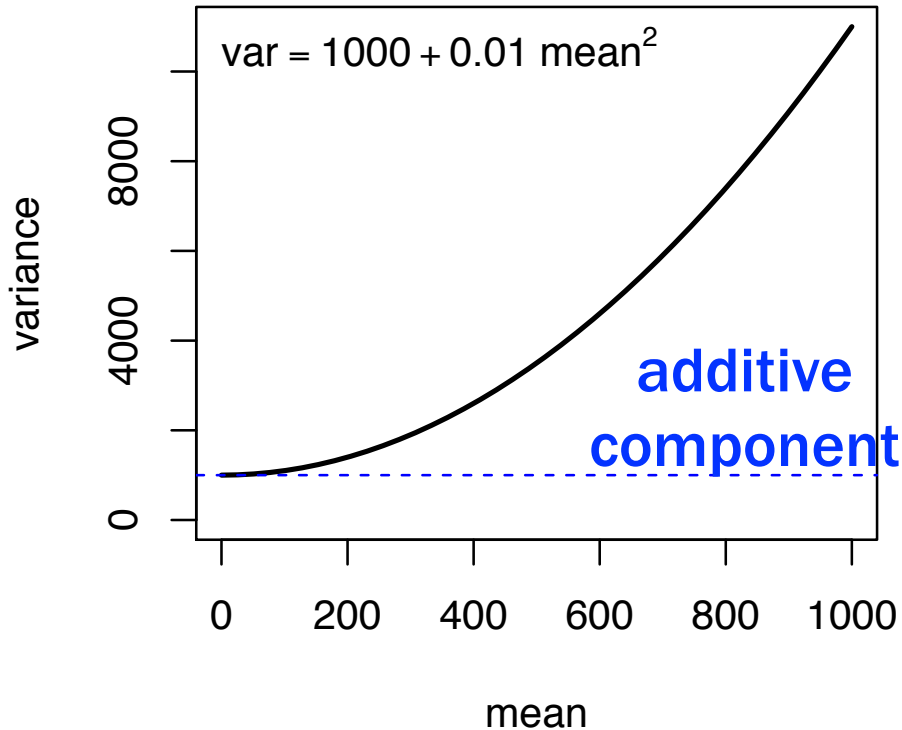
The two-component model for microarray data



raw scale

log scale

The additive-multiplicative error model



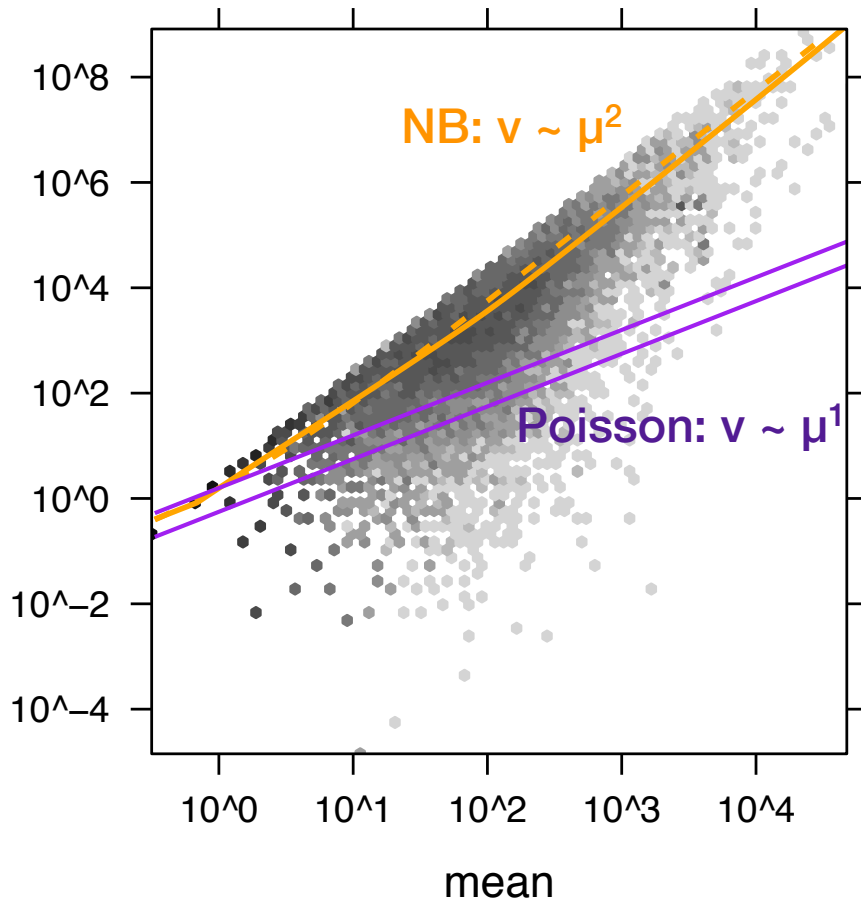
Trey Ideker et al.: JCB (2000)

David Rocke and Blythe Durbin: JCB (2001), Bioinformatics (2002)

For robust affine regression normalisation: W. Huber et al. Bioinformatics (2002)

For background correction in RMA: R. Irizarry et al., Biostatistics (2003)

Two component error models



Microarrays

$$\text{var}(\mu) = b + c \cdot \mu^2$$

b: background

c: asymptotic coefficient of variation

Sequencing counts

early edgeR:

$$\text{var}(\mu) = \mu + \alpha \cdot \mu^2$$

μ : from Poisson

α : dispersion

DESeq

$$\text{var}(\mu) = \mu + \alpha(\mu) \cdot \mu^2$$

DESeq parametric option

$$\alpha(\mu) = a_1/\mu + a_0 \quad \Leftrightarrow$$

$$\text{var}(\mu) = \mu + a_1 \cdot \mu + a_0 \cdot \mu^2$$

► variance stabilizing transformations

X_u a family of random variables with

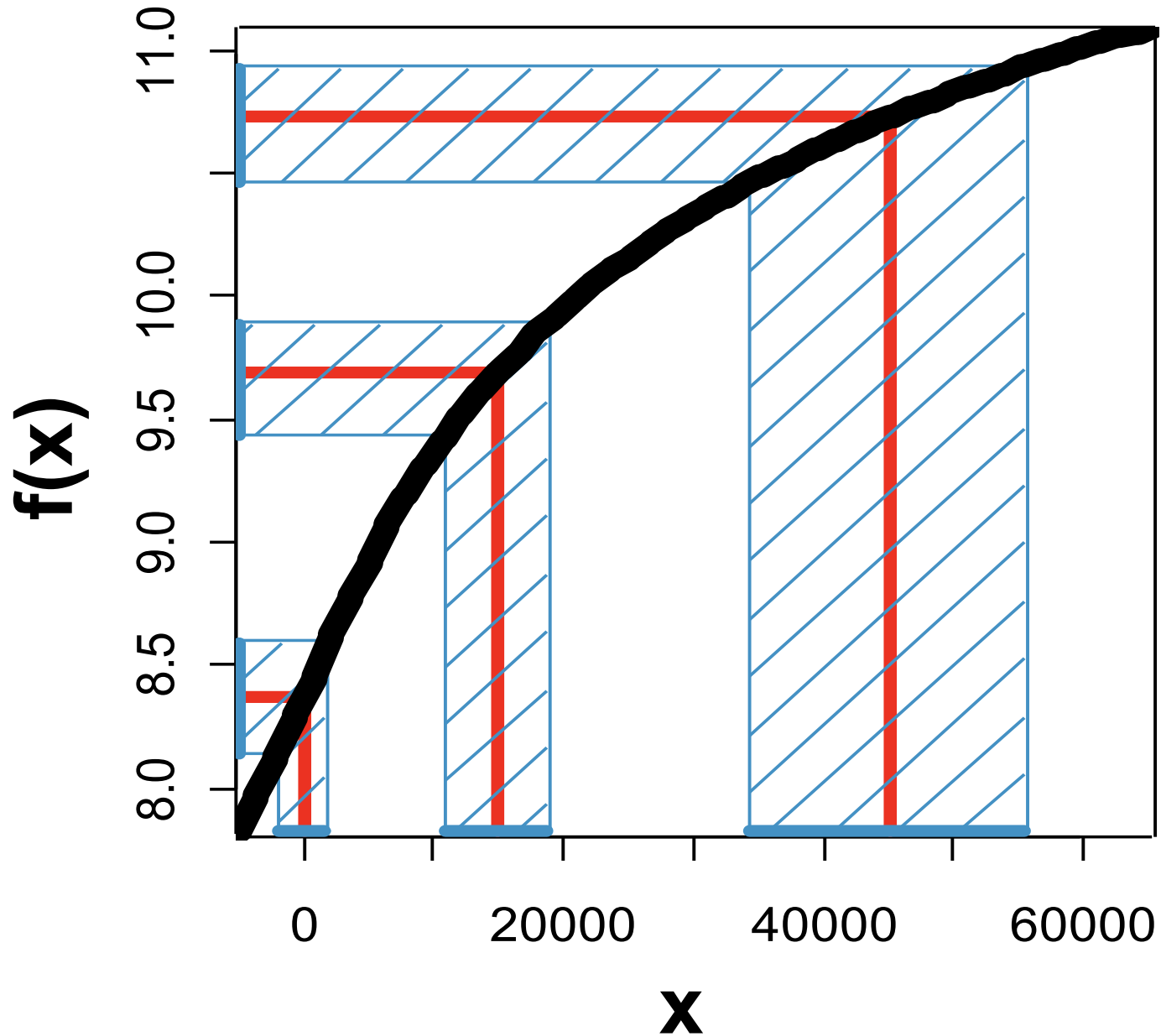
$E(X_u) = u$ and $\text{Var}(X_u) = v(u)$. Define

$$f(x) = \int^x \frac{du}{\sqrt{v(u)}}$$

Then, $\text{var } f(X_u) \approx$ does not depend on u

**Derivation: linear approximation,
relies on smoothness of $v(u)$.**

► variance stabilizing transformation



▶ variance stabilizing transformations

$$f(x) = \int^x \frac{1}{\sqrt{v(u)}} du$$

1.) constant variance ('additive') $v(u) = s^2 \Rightarrow f \propto u$

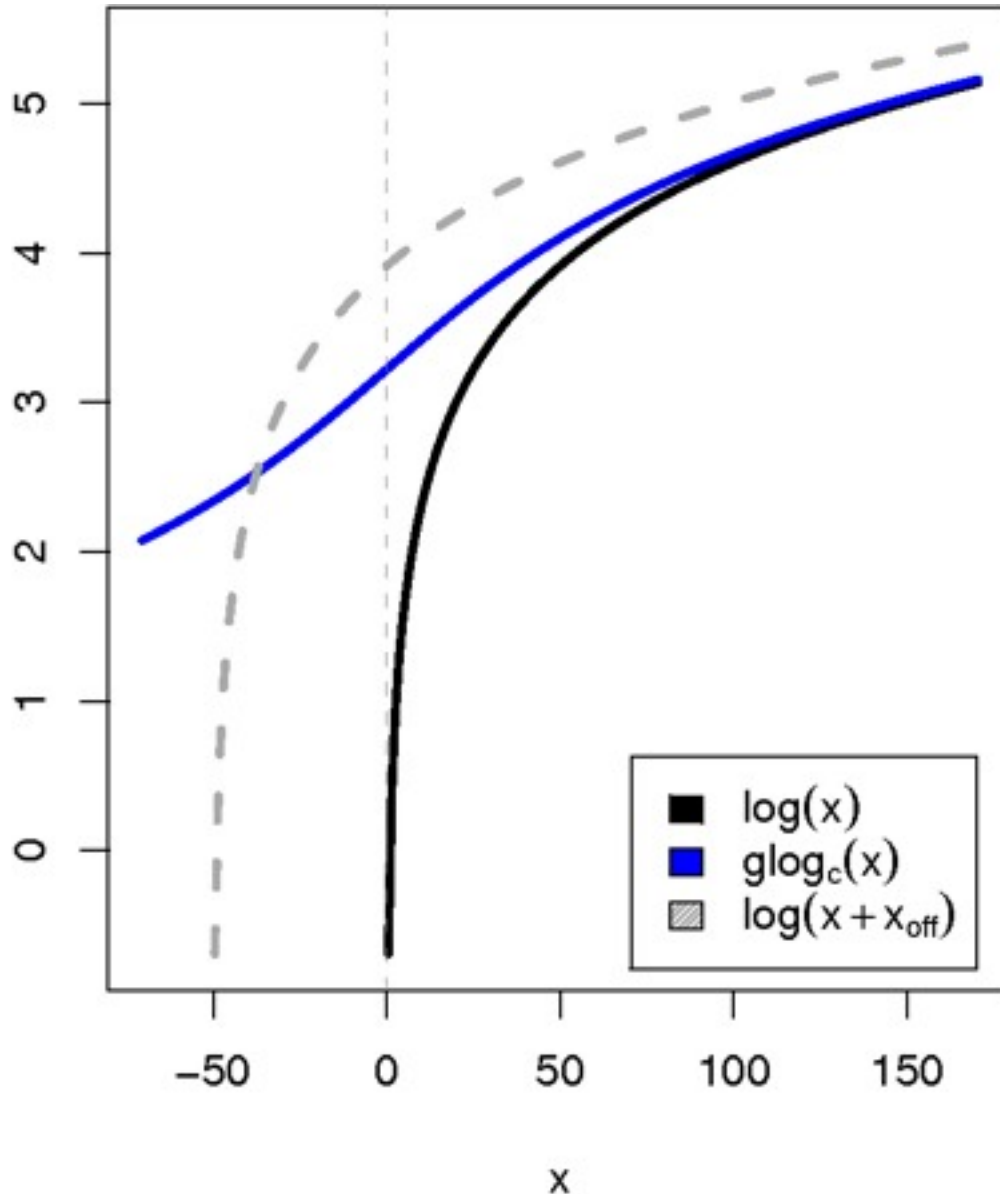
2.) constant CV ('multiplicative') $v(u) \propto u^2 \Rightarrow f \propto \log u$

3.) offset $v(u) \propto (u + u_0)^2 \Rightarrow f \propto \log(u + u_0)$

4.) additive and multiplicative

$$v(u) \propto (u + u_0)^2 + s^2 \Rightarrow f \propto \operatorname{arsinh} \frac{u + u_0}{s}$$

► the “glog” transformation



$$\text{glog}_2(x, c) = \log_2 \left(\frac{x + \sqrt{x^2 + c^2}}{2} \right)$$

$$\text{glog}_e(x, 1) + \log_e 2 = \text{arsinh}(x)$$

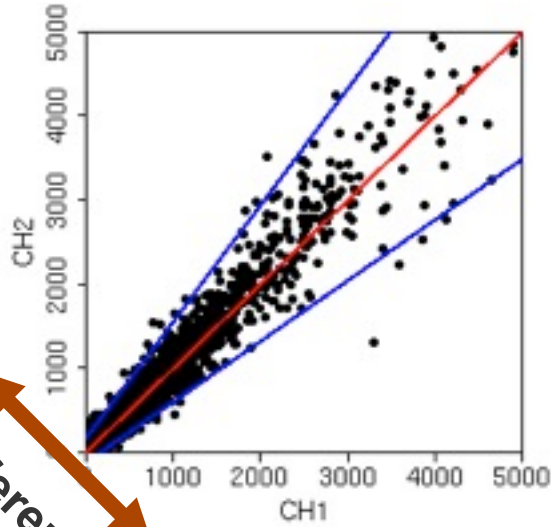
P. Munson, 2001

D. Rocke & B. Durbin,
ISMB 2002

W. Huber et al., ISMB
2002

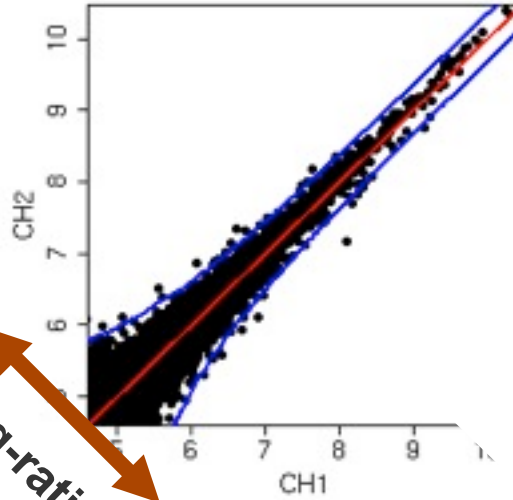
▶ glog

raw scale



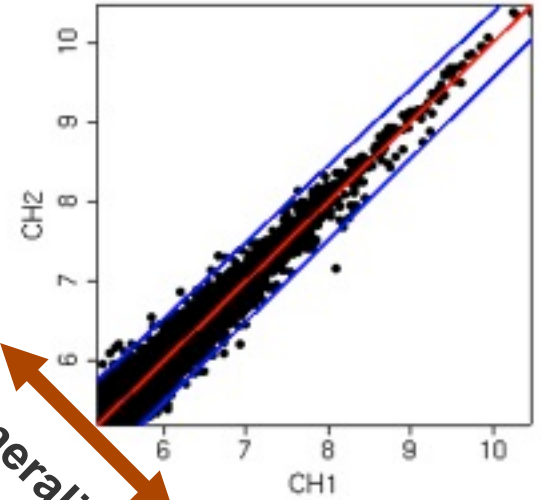
difference

log



log-ratio

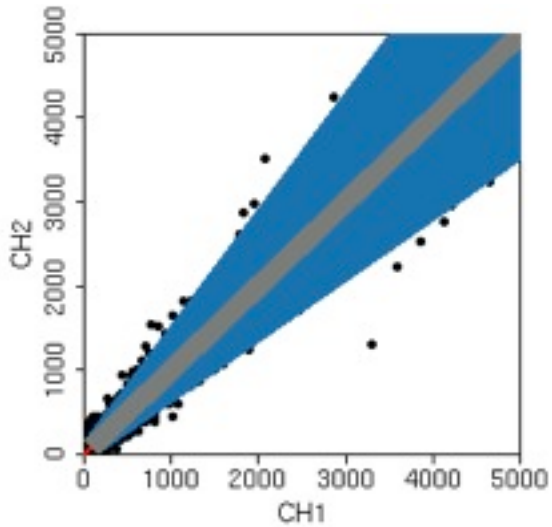
glog



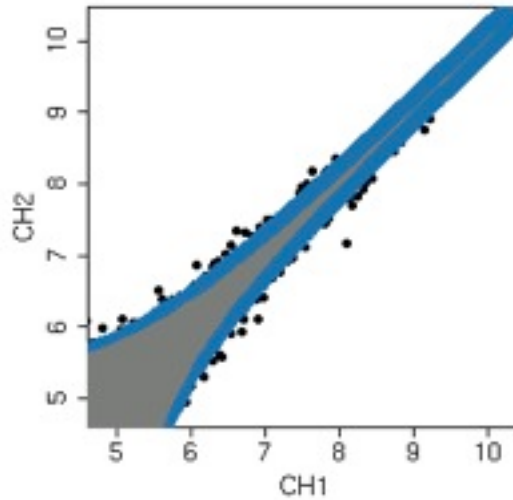
generalized
log-ratio

► glog

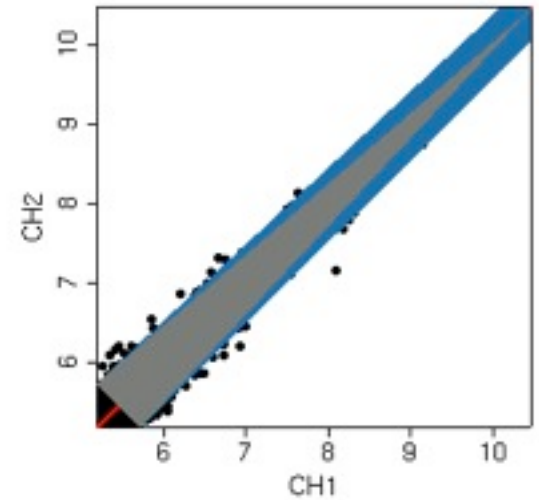
raw scale



log



glog



variance:



constant part



proportional part

vsn

$$\text{arsinh} \frac{y_{ki} - a_i}{b_i} = \mu_k + \varepsilon_{ki}, \quad \varepsilon_{ki} : N(0, c^2)$$

vsn

$$\text{arsinh} \frac{y_{ki} - a_i}{b_i} = \mu_k + \varepsilon_{ki}, \quad \varepsilon_{ki} : N(0, c^2)$$

measured intensity = offset + gain * true abundance

$$y_{ik} = a_{ik} + b_{ik} x_{ik}$$

$$a_{ik} = a_i + L_{ik} + \varepsilon_{ik}$$

a_i per-sample offset

L_{ik} local background
provided by image
analysis

$$\varepsilon_{ik} \sim N(0, b_i^2 s_1^2)$$

“additive noise”

$$b_{ik} = b_i b_k \exp(\eta_{ik})$$

b_i per-sample
normalization factor

b_k sequence-wise
labeling efficiency

$$\eta_{ik} \sim N(0, s_2^2)$$

“multiplicative noise”

vsn

$$\text{arsinh} \frac{y_{ki} - a_i}{b_i} = \mu_k + \varepsilon_{ki}, \quad \varepsilon_{ki} : N(0, c^2)$$

vsn

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- o **maximum likelihood estimator**: straightforward – but sensitive to deviations from normality

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- model holds for genes that are unchanged; differentially transcribed genes act as **outliers**.

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- **maximum likelihood estimator**: straightforward – but sensitive to deviations from normality
- model holds for genes that are unchanged; differentially transcribed genes act as **outliers**.
- **robust** variant of ML estimator, à la **Least Trimmed Sum of Squares** regression.
- works well as long as <50% of genes are differentially transcribed (and may still work otherwise)

“usual” log-ratio

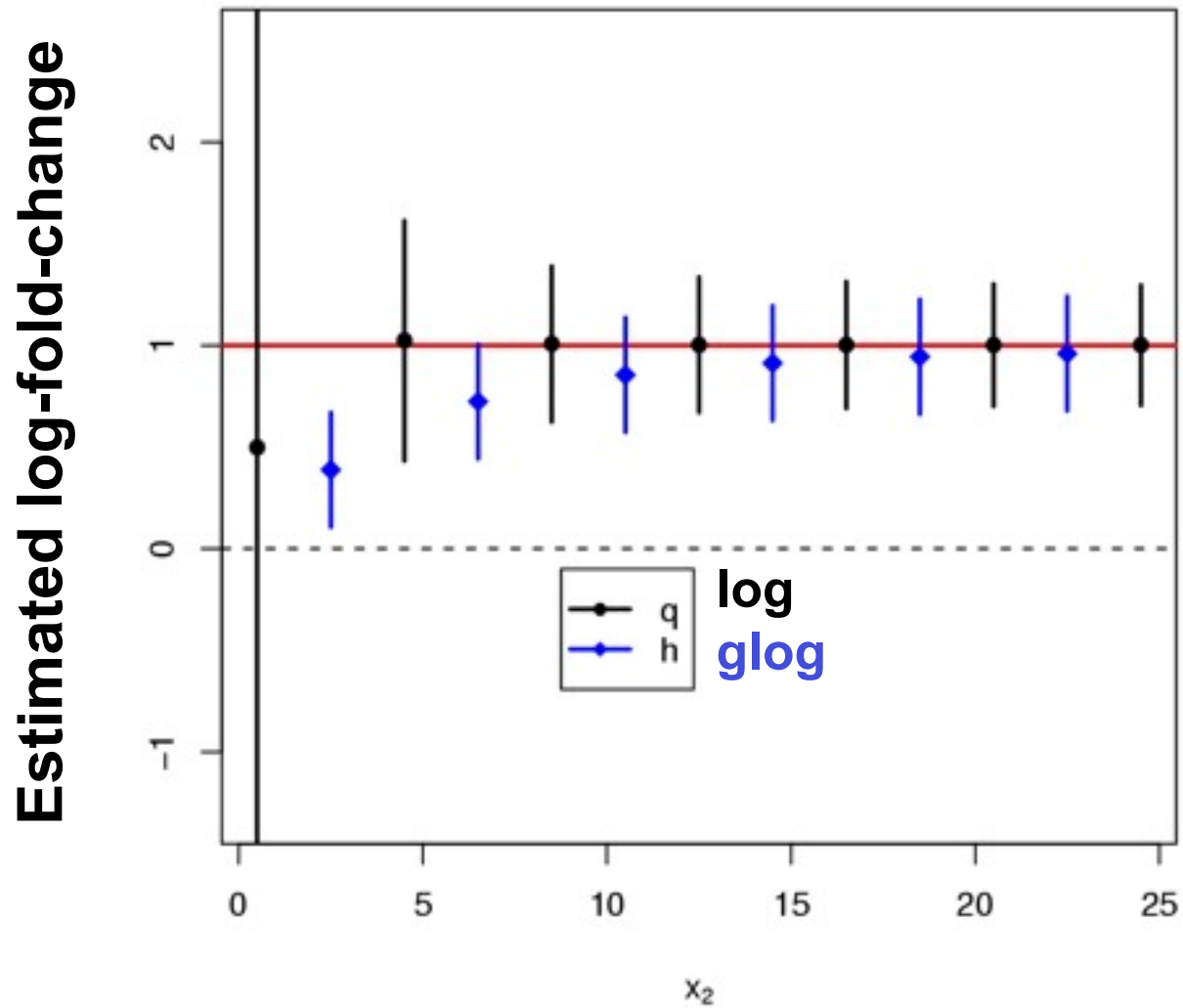
$$\log \frac{x_1}{x_2}$$

**'glog'
(generalized
log-ratio)**

$$\log \frac{x_1 + \sqrt{x_1^2 + c_1^2}}{x_2 + \sqrt{x_2^2 + c_2^2}}$$

c_1, c_2 are experiment specific parameters (~level of background noise)

► Variance Bias Trade-Off



Signal intensity

▶ Variance-bias trade-off and shrinkage estimators

Shrinkage estimators:

a general technology in statistics:

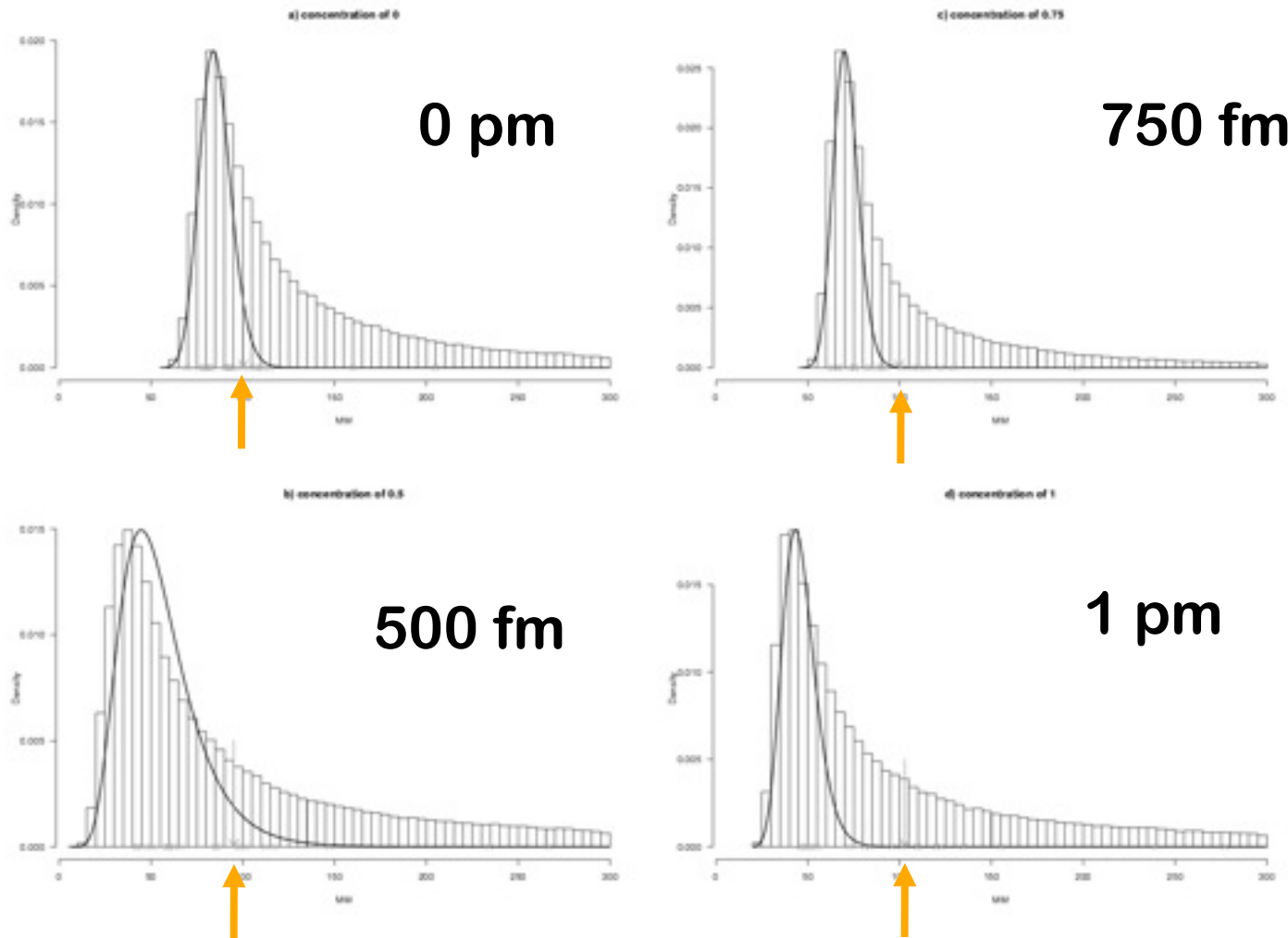
pay a small price in bias for a large decrease of variance, so overall the mean-squared-error (MSE) is reduced.

Particularly useful if you have few replicates.

Generalized log-ratio is a shrinkage estimator for log fold change

**further
background
correction
methods**

Background correction



Irizarry et al.
Biostatistics
2003

Fig. 5. Histograms of $\log_2(MM)$ for an array in which no probe-set was spiked along with the three arrays in which BioB-5 was spiked-in at concentrations of 0.5, 0.75, and 1 pM. The observed PM values for the 20 probes associated with BioB-5 are marked with crosses and the average with an arrow. The black curve represents the log normal distribution obtained from left-of-the-mode data.

RMA Background correction

$$PM = B + S$$

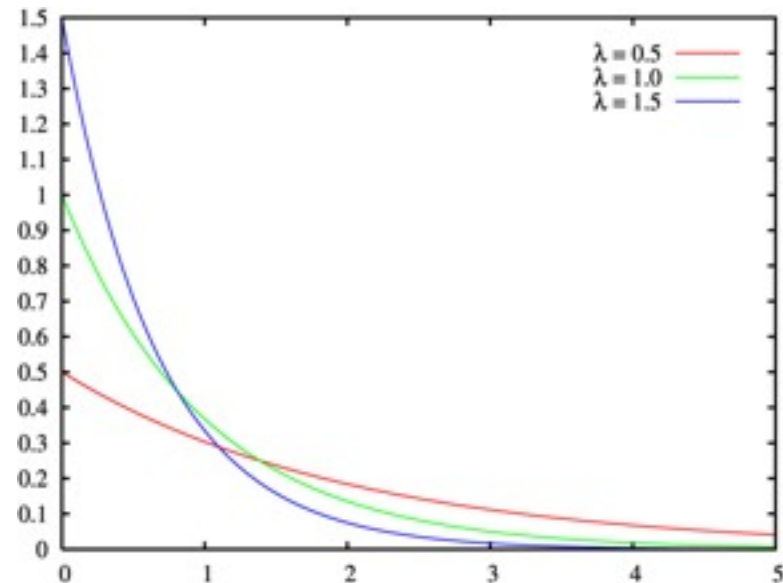
$B \sim$ log-normal with mean and sd read off MM values

$S \sim$ exponential

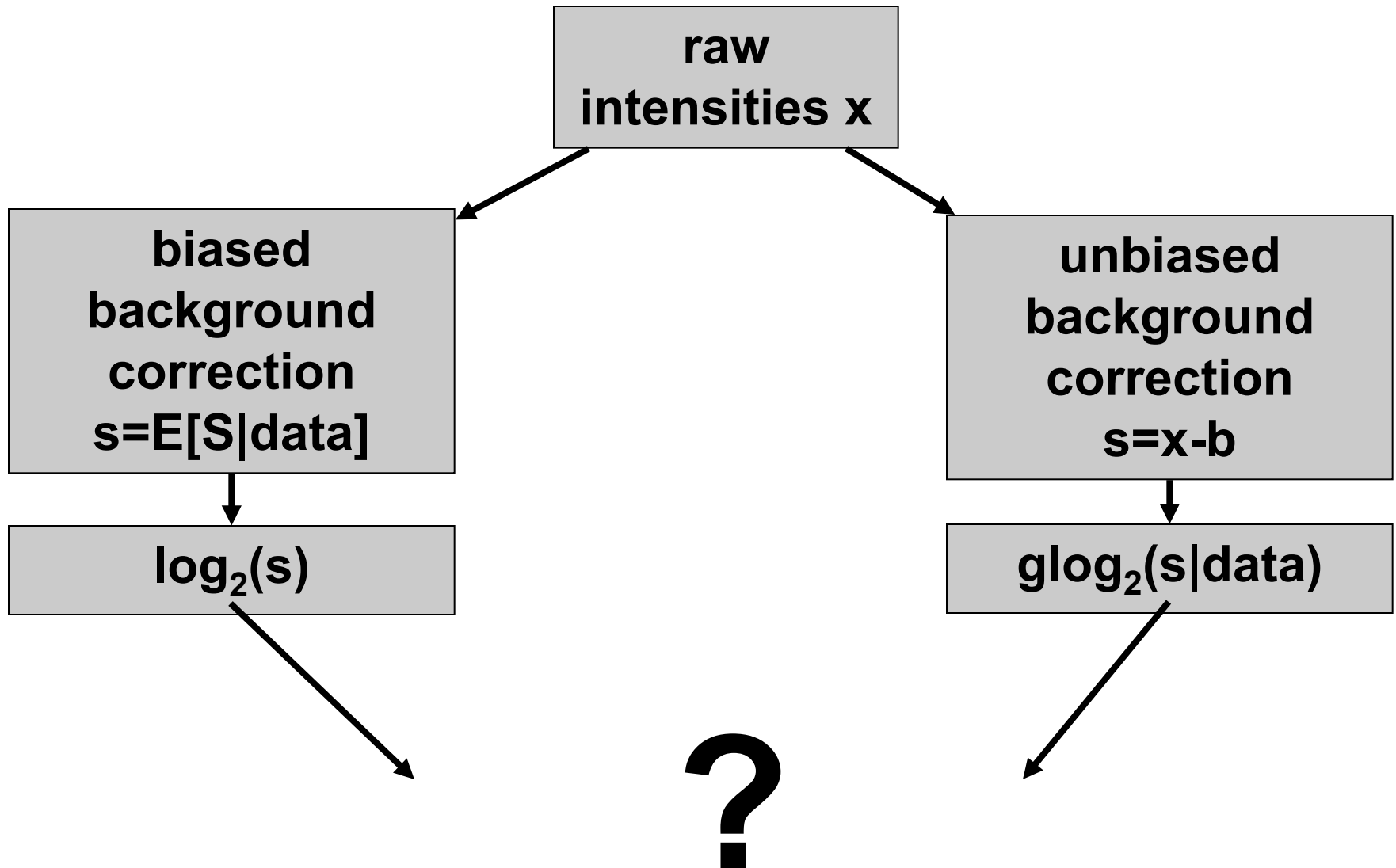
\Rightarrow closed form expression for $E[S | PM]$,
use this as \hat{s} (> 0).

(NB, $P[S > 0] = 1$ is not realistic)

Irizarry et al. (2002)

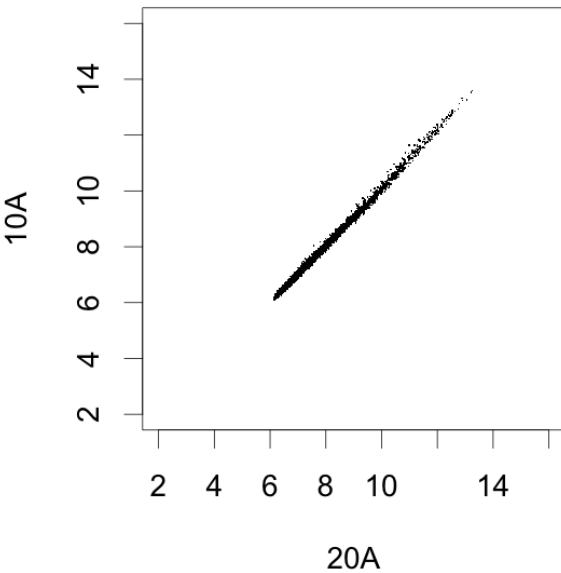


Background correction:

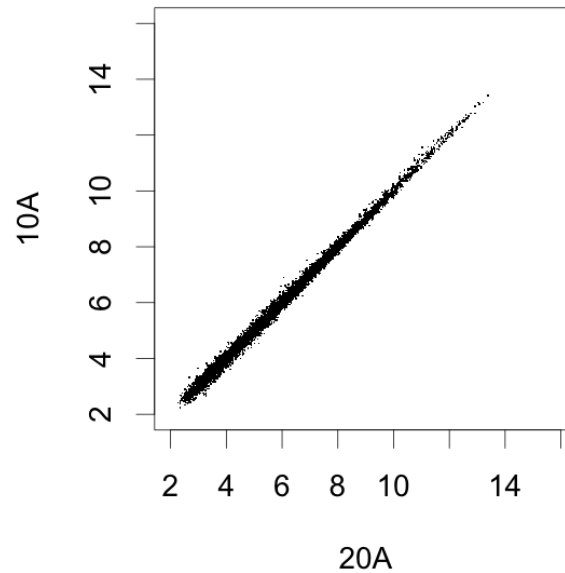


Comparison between RMA and VSN background correction

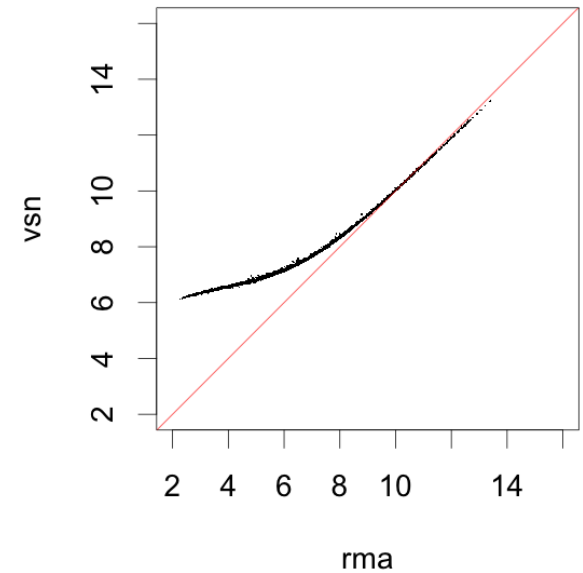
vsn: array 1 vs 3



rma: array 1 vs 3



array 1



Summaries for Affymetrix genechip probe sets

Data and notation

PM_{ikg} , MM_{ikg} = Intensities for perfect match and mismatch probe k for gene g on chip i

$i = 1, \dots, n$ one to hundreds of chips

$k = 1, \dots, J$ usually 11 probe pairs

$g = 1, \dots, G$ tens of thousands of probe sets.

Tasks:

calibrate (normalize) the measurements from different chips (samples)

summarize for each probe set the probe level data, i.e., 11 PM and MM pairs, into a single **expression measure**.

compare between chips (samples) for detecting differential expression.

Expression measures: MAS 4.0

Affymetrix GeneChip MAS 4.0 software used **AvDiff**, a trimmed mean:

$$AvDiff = \frac{1}{\#K} \sum_{k \in K} (PM_k - MM_k)$$

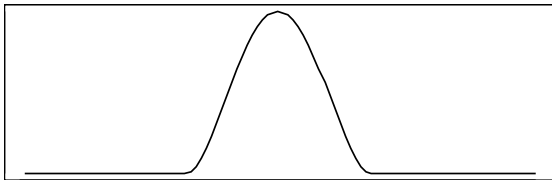
- sort $d_k = PM_k - MM_k$
- exclude highest and lowest value
- $K :=$ those pairs within 3 standard deviations of the average

Expression measures MAS 5.0

Instead of MM, use "repaired" version CT

$$\begin{aligned} \text{CT} &= \text{MM} && \text{if } \text{MM} < \text{PM} \\ &= \text{PM} / \text{"typical log-ratio"} && \text{if } \text{MM} \geq \text{PM} \end{aligned}$$

Signal = Weighted mean of the values $\log(\text{PM}-\text{CT})$
weights follow Tukey Biweight function
(location = data median,
scale a fixed multiple of MAD)



Expression measures: Li & Wong

dChip fits a model for each gene

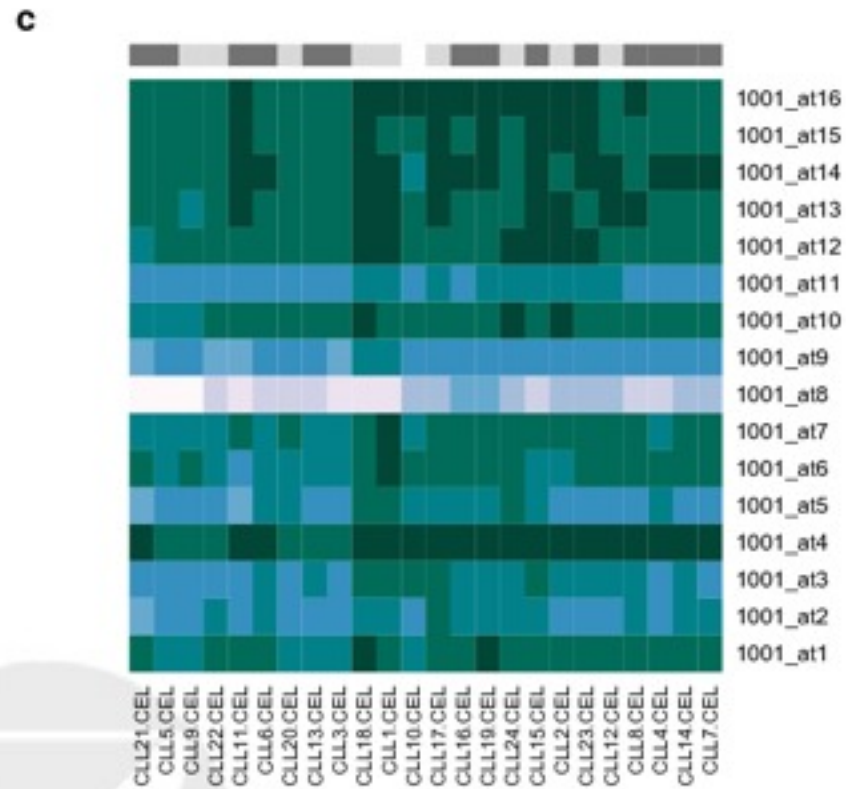
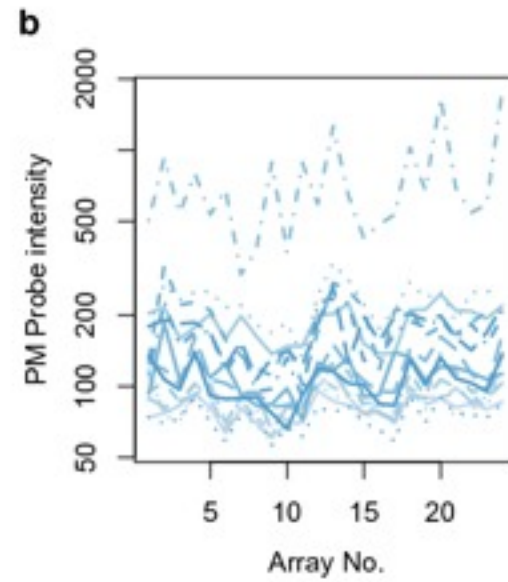
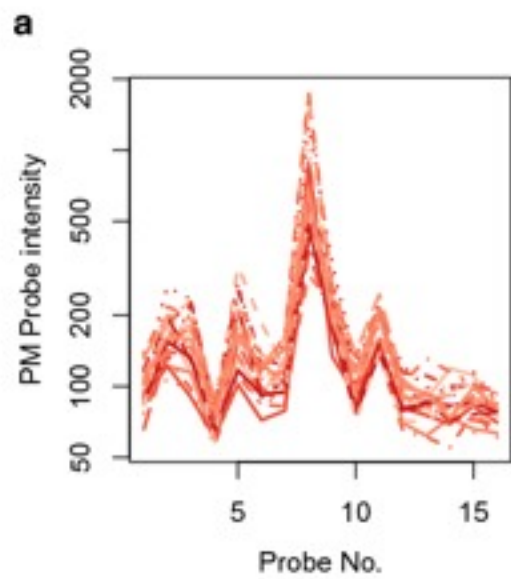
$$PM_{ki} - MM_{ki} = \theta_k \phi_i + \varepsilon_{ki}, \quad \varepsilon_{ki} \propto N(0, \sigma^2)$$

where

ϕ_i : **expression measure** for the gene in sample i

θ_k : **probe effect**

ϕ_i is estimated by maximum likelihood



Expression measures

RMA: Irizarry et al. (2002)

dChip

$$Y_{ki} = \theta_k \phi_i + \varepsilon_{ki}, \quad \varepsilon_{ki} \propto N(0, \sigma^2)$$

RMA

$$\log_2 Y_{ki} = a_k + b_i + \varepsilon_{ki}$$

b_i is estimated using the robust method **median polish** (successively remove row and column medians, accumulate terms, until convergence).

Quality assessment



Quality assessment



**arrayQualityMetrics
example quality report**

References

- Bioinformatics and computational biology solutions using R and Bioconductor, R. Gentleman, V. Carey, W. Huber, R. Irizarry, S. Dudoit, Springer (2005).**
- Variance stabilization applied to microarray data calibration and to the quantification of differential expression. W. Huber, A. von Heydebreck, H. Sültmann, A. Poustka, M. Vingron. Bioinformatics 18 suppl. 1 (2002), S96-S104.**
- Exploration, Normalization, and Summaries of High Density Oligonucleotide Array Probe Level Data. R. Irizarry, B. Hobbs, F. Collins, ..., T. Speed. Biostatistics 4 (2003) 249-264.**
- Error models for microarray intensities. W. Huber, A. von Heydebreck, and M. Vingron. Encyclopedia of Genomics, Proteomics and Bioinformatics. John Wiley & sons (2005).**
- Normalization and analysis of DNA microarray data by self-consistency and local regression. T.B. Kepler, L. Crosby, K. Morgan. Genome Biology. 3(7):research0037 (2002)**
- Statistical methods for identifying differentially expressed genes in replicated cDNA microarray experiments. S. Dudoit, Y.H. Yang, M. J. Callow, T. P. Speed. Technical report # 578, August 2000 (UC Berkeley Dep. Statistics)**
- A Benchmark for Affymetrix GeneChip Expression Measures. L.M. Cope, R.A. Irizarry, H. A. Jaffee, Z. Wu, T.P. Speed. Bioinformatics (2003).**

....many, many more...

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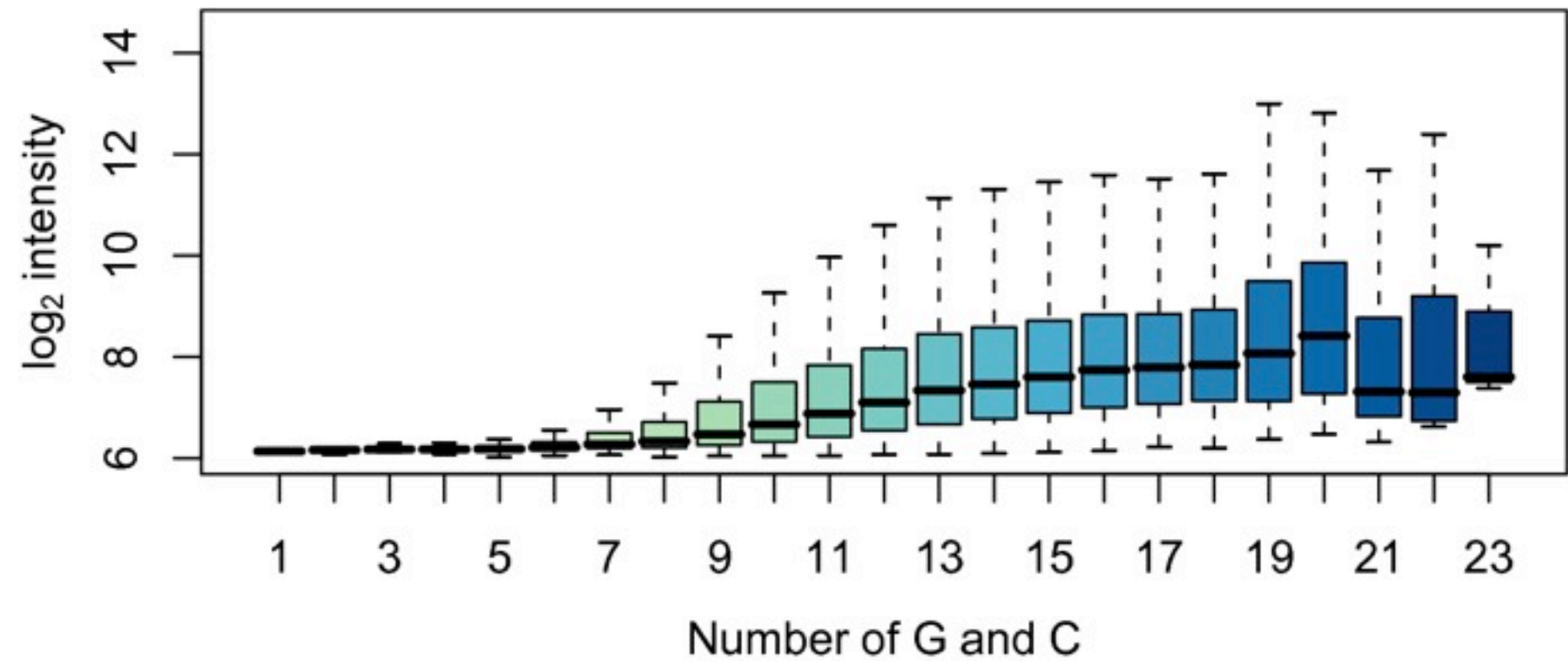
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► What about non-linear effects

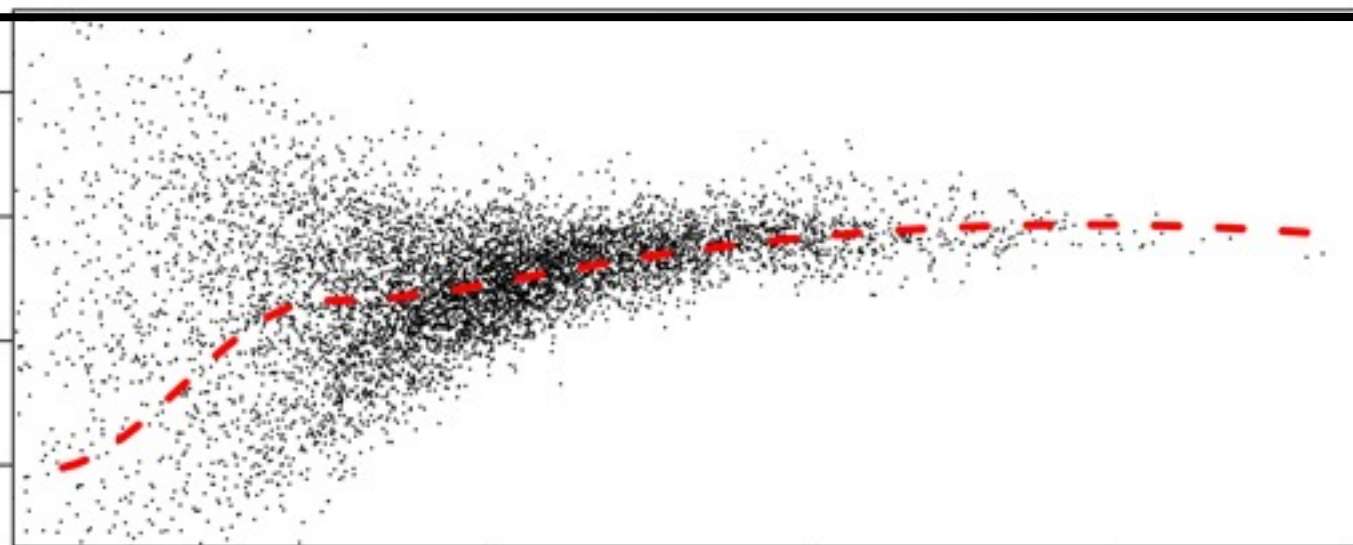
- Microarrays can be operated in a linear regime, where fluorescence intensity increases proportionally to target abundance (see e.g. Affymetrix dilution series)

Two reasons for non-linearity:

- At the high intensity end: saturation/quenching. This can (and should) be avoided experimentally - loss of data!
- At the low intensity end: background offsets, instead of $y=k \cdot x$ we have $y=k \cdot x+x_0$, and in the log-log plot this can look curvilinear. But this is an affine-linear effect and can be corrected by affine normalization. Local polynomial regression may be OK, but tends to be less efficient.

$$M = \log_2 R - \log_2 G$$

2
0
-2
-4



10

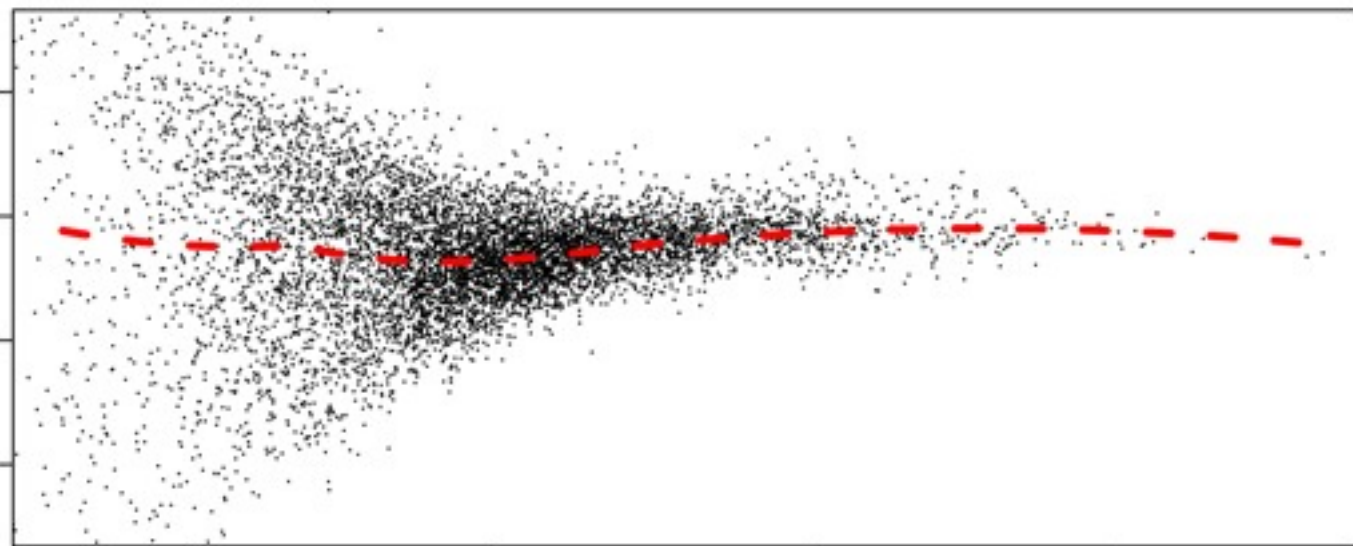
15

20

25

$$M = \log_2(R+c) - \log_2 G$$

2
0
-2
-4



10

15

20

25

$$A = \log_2(R+c) + \log_2 G$$