

Variance Stabilizing Transformations for image-based compound profiling features

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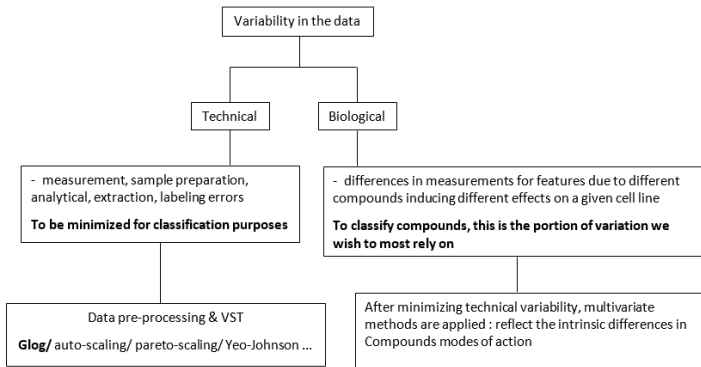
Image-based multi-parameteric compound profiling features

- Biological method used as a proxy for distinguishing compounds in the drug discovery chain using a range of features extracted from image-based assays by applying High Throughput Microscopy (HTM)
- The features provide information on
 - i. Intracellular biomarkers: texture, intensity, spatial distribution etc
 - ii. Cells: shape, geometry, quantity .. .
- Why
 - i. understand how compounds induce their desired properties and describe their mechanisms of action
 - ii. preferentially identify highly specific compounds having a desired effect on a given biological target
 - iii. early detection of undesired compound effects on cells + cellular activity: toxicity

.. introduction

It's all good, but Most of these features often

- are highly correlated: need to limit features used for analysis
- have non-normal distributions: mean-variance relationship present
 - multivariate classification methods hugely depend on variance



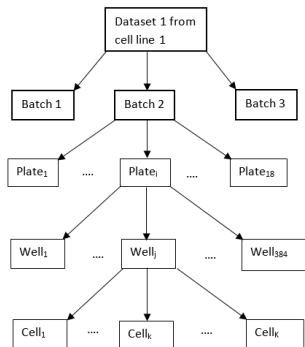
Aim of the analysis

To assess the:

- I. effect of glog transformation on separation of treatment replicates from non-replicates
- II. effect of glog transformation on proportion of actively-called treatments
- III. performance of a glog transformation on treatments separation when applied at cell- or well-level

♣ Treatment: a compound at a given concentration (a compound can have 4 or 5 concentration levels - 1 μ M (microMolar), 3 μ M, 3.34 μ M, 9 μ M and 11.1 μ M)

Data, the log transformation and data pre-processing



Data

- i. 2 cancer cell lines: Liver & Colon
- ii. a plate had btwn 134909 & 281177 (152679 & 330117) in 1st(2nd) data
- iii. a well had btwn 73 & 1812 (95 & 2120) cells in the 1st(2nd) data
- iv. 311 compounds including DMSO control
- v. we tested a total of 1253 treatments
- vi. 462 features extracted from each cell

..Data, the glog transformation and data pre-processing

Glog transformation

→ formula

$$z = \text{Log}(y - \alpha + \sqrt{(y - \alpha)^2 + \lambda})$$

→ where

- z: glog-transformed data
- y: untransformed data
- α : feature mean across DMSO controls
- λ : transformation parameter

Data pre-processing

→ Aggregation - calculating mean for each feature per well

→ Normalization -

$$\frac{\text{feature}_{value} - \text{mean.feature}_{DMSO}}{\text{pooled.SD.feature}_{across.plates}}$$

→ Feature selection

- MRMR: identify set of features with low pairwise correlation & high reproducibility among replicates.
- AUC value for btwn 2-75 features
- optimal feature: maximizes separation of treatment replicates within 1 Std error of AUC

→ Active calling: treatments with $\geq 50\%$ active replicates

Methodology

★ Hotelling's T^2 method

~> measures difference in 2 multivariate means

~> formula

$$T^2 = \frac{(\bar{\mathbf{X}}_1 - \bar{\mathbf{X}}_2)'(\bar{\mathbf{X}}_1 - \bar{\mathbf{X}}_2)}{\mathbf{S}_p\left(\frac{1}{n_1} + \frac{1}{n_2}\right)}$$

~> normality assumptions for optimal results

~> Only actively-called treatments in pre- & post-transformation used

~> + shift in T^2 distribution indicate improved treatment separation

★ AUC method

2-steps involved in AUC-calculation

~> Pearson correlation btwn pairs of replicates (& non-replicates) were calculated & distributions plotted

~> separation btwn the 2 distribution quantified by constructing an ROC curve using a series of correlation thresholds & calculating an AUC value

~> transformations leading to higher AUC values --> improved treatments separation compared to corresponding untransformed data

Results: EDA

No. of replicates(% of sample treatments)

9 (%)	36 (%)	45 (%)
1192(95.208)	28(2.236)	32(2.556)

- ★ DMSO control replicated across 1512 wells

- ★ For both data sets

- ★ Implications

- ★ For calculation of Hotelling's T^2 , a limited number of selected features was used to maximize its power
- ★ 10 highest ranked features from MRMR used to calculate T^2

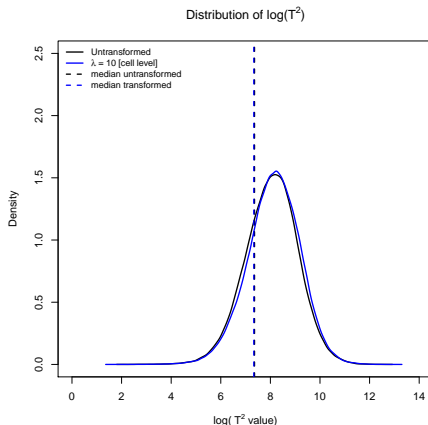
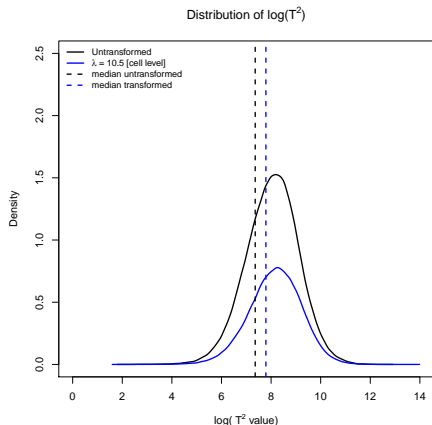
Transformations effects on treatments separation

Prologue

- ◇ Only glog transformations of λ equal to 0.1 and, 0.5 to 25 at 0.5 interval investigated for both T^2 and AUC methods
- ◇ Each transformed data compared to its corresponding untransformed data defined by actively-called treatments present both pre- and post-transformation
- ◇ Improved treatments separation shown by +ve shifts in distribution (and/or associated statistics) of T^2 for transformed compared to untransformed, and/or higher AUC values
- ◇ Results presented for the first cell line only since results largely led to similar conclusions for both cell lines

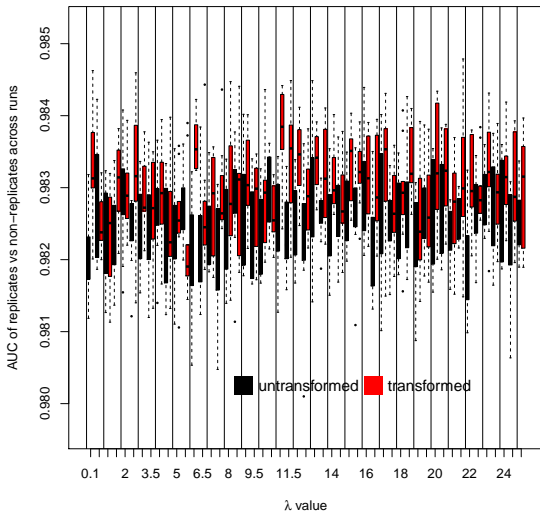
Transformations effects on treatments separation - T^2

- ✱ Presence of very high [very different] and very low [highly similar] values
- ✱ Some led to slight but negligible improvements (e.g $\lambda = 10.5$)
- ✱ Others led to no improvements (e.g $\lambda = 10$)



Transformations effects on treatments separation - AUC

Evolution of AUC assessing replicability Vs transformation parameter

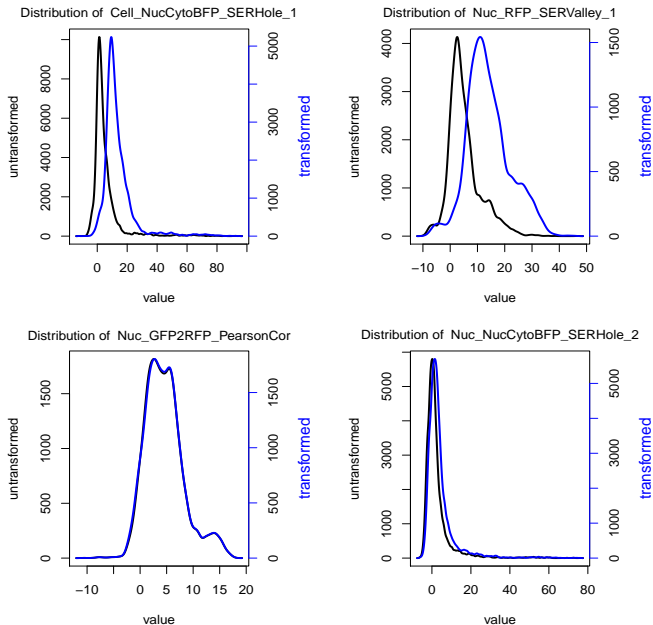


- high AUC b4-transformation
- some (e.g $\lambda = 0.1$) led to marginal increases
- others (e.g $\lambda = 5.5$) separated slightly poorer
- the differences were however very minimal & non-significant

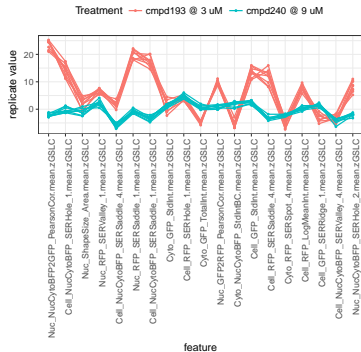
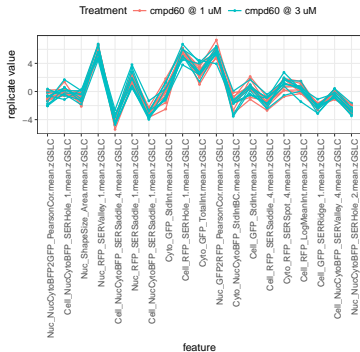
Transformations effects on treatments separation- Epilogue

- ◇ In both methods, minimal & insignificant differences were observed:
Transformations failed to improve treatments separation
- ◇ Why?

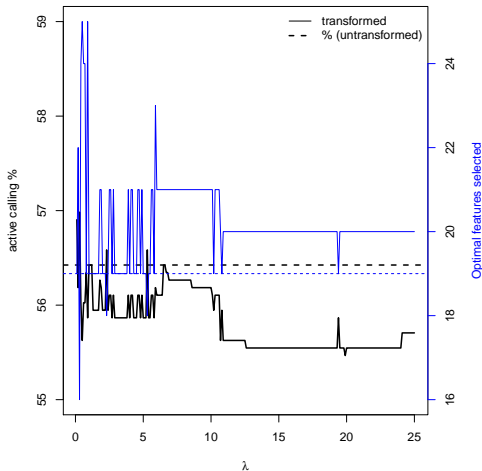
1. Transformation effect on features distributions



2. Differentiating ability of features selected (before transformation)

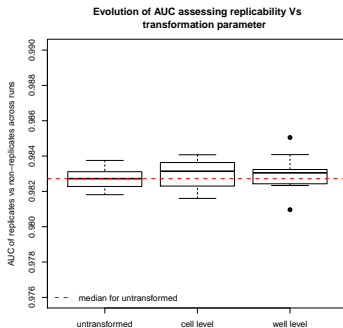
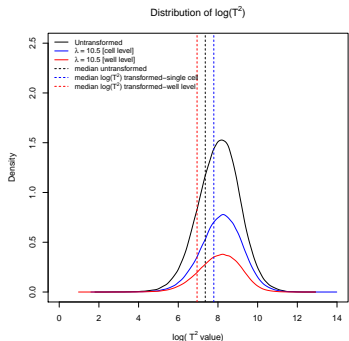


Effect of transformation on treatments active-calling



- lower % of active-calling
- intriguing relationship btwn number of features selected & prop. of active-calling
- similar relationship in 2nd cell line

Transform at cell- or well-level? [T^2 & AUC approaches]



- ~ For $\lambda = 10.5$
- ~ Minimal non-significant improvement in treatment separation when transformed at cell-level > well level
- ~ High AUC values pre-transformation
- ~ No clear preference for cell- or well-level transformation

From our study, we observed that:

- ~ Transformations did not improve treatments separation beyond what was seen pre-transformation
- ~ Transformations led to lower(higher) proportion of active-calling in 1st(2nd) data
- ~ Inverse relationship between proportion of active-calling and number of features selected was evident
- ~ There was no preference in transforming data at cell- or well-level