Pathway Fingerprinting: a working example

Gabriel Altschuler

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This document demonstrates how to use the pathprint package to analyze a dataset using Pathway Fingerprints. The pathprint package takes gene expression data and processes this into discrete expression scores (+1,0,-1) for a set of 633 pathways. For more information, see the pathprint website.

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1 Summary

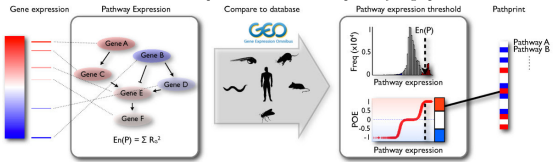
Systems-level descriptions of pathway activity across gene expression repositories are confounded by platform, species and batch effects. Pathprinting integrates pathway curation, profiling methods, and public repositories, to represent any expression profile as a ternary score (-1, 0, +1) in a standardized pathway panel. It provides annotation and a robust framework for global comparison of gene expression data.

2 Background

New strategies to combat complex human disease require systems approaches to biology that integrate experiments from cell lines, primary tissues and model organisms. We have developed Pathprint, a functional approach that compares gene expression profiles in a set of pathways, networks and transcriptionally regulated targets. It can be applied universally to gene expression profiles across species. Integration of large-scale profiling methods and curation of the public repository overcomes platform, species and batch effects to yield a standard measure of functional distance between experiments. A score of 0 in the final pathprint vector represents pathway expression at a similar level to the majority of arrays of the same platform in the GEO database, while scores of 1 and -1 reflect significantly high and low expression respectively.

3 Method

Below we describe the individual steps used to construct the pathway fingerprint.



Rank-normalized gene expression is mapped to pathway expression. A distribution of expression scores across the Gene Expression Omnibus (GEO is used to produce a probability of expression (POE) for each pathway. A pathprint vector is derived by transformation of the signed POE distribution into a ternary score, representing pathway activity as significantly underexpressed (-1), intermediately expressed (0), or overexpressed (+1).

4 Pathway sources

Canonical pathway gene sets were compiled from Reactome, Wikipathways, and KEGG (Kyoto Encyclopedia of Genes and Genomes), which were chosen because they include pathways relating to metabolism, signaling, cellular processes, and disease. For the major signaling pathways, experimentally derived transcriptionally upregulated and downregulated gene sets were obtained from Netpath. We have supplemented the curated pathways with non-curated sources of interactions by including highly connected modules from a functional-interaction network, termed 'static modules.' The modules cover 6,458 genes, 1,542 of which

are not represented in any of the pathway databases. These static modules offer the opportunity to examine the activity of less studied or annotated biological processes, and also to compare their activity with that of the canonical pathways.

Pathprinting: An integrative approach to understand the functional basis of disease Gabriel M Altschuler, Oliver Hofmann, Irina Kalatskaya, Rebecca Payne, Shannan J Ho Sui, Uma Saxena, Andrei V Krivtsov, Scott A Armstrong, Tianxi Cai, Lincoln Stein and Winston A Hide Genome Medicine (2013) 5:68 DOI: 10.1186/gm472

5 Initial data processing

An existing GEO sample on the Human Affy ST 1.0 chip will be used as en example. The dataset GSE26946 profiles expression data from iPS and human ES cells. The R package GEOquery can be used to retrieve the data. An 'exprs' object, i.e. a dataframe with row names corresponding to probe or feature IDs and column names corresponding to sample IDs is required by pathprint. In addition, we need to know the GEO reference for the platform, in this case GPL6244, and the species, which is 'human' or "Homo sapiens' (both styles of name work).

6 Pathway fingerprinting

6.1 Fingerprinting from new expression data

Now the data has been prepared, the pathprint function exprs2fingerprint can be used to produce a pathway fingerprint from this expression table.

```
> library(pathprint)
> library(SummarizedExperiment)
> library(pathprintGEOData)
> # load the data
```

```
> data(compressed_result)
> # load("chipframe.rda")
> ds = c("chipframe", "genesets", "pathprint.Hs.gs", "platform.thresholds", "pluripotents.frame")
> data(list = ds)
> # extract GEO.fingerprint.matrix and GEO.metadata.matrix
> GEO.fingerprint.matrix = assays(result)$fingerprint
> GEO.metadata.matrix = colData(result)
> GSE26946.fingerprint <- exprs2fingerprint(exprs = GSE26946.exprs,
                                           platform = GSE26946.platform,
                                           species = GSE26946.species,
                                           progressBar = FALSE
[1] "Running fingerprint"
> GSE26946.fingerprint[1:5, 1:3]
                                                 GSM663450
Glycolysis / Gluconeogenesis (KEGG)
Citrate cycle (TCA cycle) (KEGG)
                                                         1
Pentose phosphate pathway (KEGG)
                                                         1
Pentose and glucuronate interconversions (KEGG)
                                                         1
Fructose and mannose metabolism (KEGG)
                                                         1
                                                 GSM663451
Glycolysis / Gluconeogenesis (KEGG)
Citrate cycle (TCA cycle) (KEGG)
                                                         1
Pentose phosphate pathway (KEGG)
                                                         1
Pentose and glucuronate interconversions (KEGG)
Fructose and mannose metabolism (KEGG)
                                                 GSM663452
Glycolysis / Gluconeogenesis (KEGG)
Citrate cycle (TCA cycle) (KEGG)
Pentose phosphate pathway (KEGG)
                                                         1
Pentose and glucuronate interconversions (KEGG)
                                                         1
Fructose and mannose metabolism (KEGG)
```

6.2 Using existing data

The pathprint package uses the object compressed_result, drawn from the data-package pathprintGEOData, which was constructed in 2012 and does not contain all the GEO data. When uncompressed yields GEO.fingerprint.matrix and GEO.metadata.matrix. GEO.fingerprint.matrix contains 188390 samples that have already been fingerprinted, along with their associated metadata, in the object GEO.metadata.matrix. As the above data record is publically available from GEO it is actually already in the matrix and we can compare this to the fingerprint processed above. It should be noted that occasionally there may be discrepancies in one or two pathways due to the way in which the threshold is applied.

```
> colnames(GSE26946.exprs) %in% colnames(GEO.fingerprint.matrix)
```

7 Fingerprint Analysis

7.1 Intra-sample comparisons

The fingerprint vectors can be used to compare the differntially expressed functions within the sample set. The most straight forward method to represent this is using a heatmap, removing rows for which there is no change in functional expression.

7.2 Using consensusFingerprint and fingerprinDistance, comparison to pluripotent arrays

We can also investigate how far in functional distance, these arrays are from other pluripotent fingerprints. This can be achieved using the set of pluripotent arrays included in the package, from which a consensus fingerprint can be created.

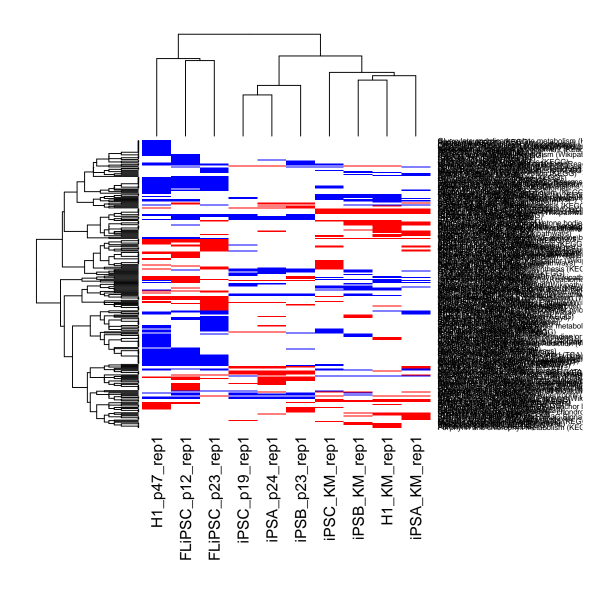


Figure 1: Heatmap of GSE26946 pathway fingerprints, blue = -1, white = 0, $\rm red = +1$

7.3 Identifying similar arrays

We can use the data contained within the GEO fingerprint matrix to order all of the GEO records according to distance from an experiment (or set of experiments, see below). This can be used, in conjunction with the metadata, to annotate a fingerprint with data from the GEO corpus. Here, we will identify experiments closely matched to the H1, embyonic stem cells within GSE26946

```
> GSE26946.H1<-consensusFingerprint(</pre>
   GSE26946.fingerprint[,grep("H1", GSE26946.names)], threshold=0.9)
> geo.H1Distance<-consensusDistance(</pre>
   GSE26946.H1, GEO.fingerprint.matrix)
[1] "Scaling against max length, 700"
> # look at top 20
> GEO.metadata.matrix[match(head(rownames(geo.H1Distance),20), rownames(GEO.metadata.matrix)),c("0
DataFrame with 20 rows and 3 columns
          <character> <character>
GSM663458
            GSE26946
                         GPL6244
             GSE26946
                          GPL6244
GSM663459
            GSE26946
                          GPL6244
GSM663455
GSM663453
            GSE26946
                          GPL6244
             GSE26946
                          GPL6244
GSM663451
GSM525414
            GSE21037
                          GPL6244
GSM697682
            GSE21655
                          GPL6244
GSM525413
            GSE21037
                          GPL6244
GSM525421
             GSE21037
                          GPL6244
GSM525420
             GSE21037
                          GPL6244
                                            Source
                                       <character>
GSM663458
                       Human Embryonic Stem Cells
GSM663459
                       Human Embryonic Stem Cells
GSM663455
                   induced pluripotent stem cells
GSM663453
                   induced pluripotent stem cells
GSM663451
                   induced pluripotent stem cells
. . .
GSM525414
                                iPSC-RTT clone 18
GSM697682 iPSC maintained under feeder conditions
GSM525413
                                iPSC-RTT clone 18
GSM525421
                                  iPSC-WT clone 2
GSM525420
                                  iPSC-WT clone 1
```

Distance from pluripotent consensus

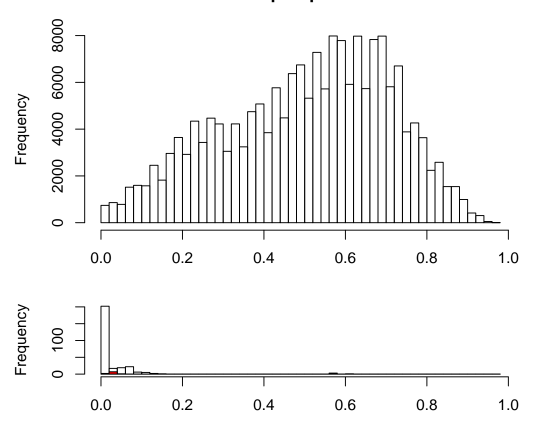


Figure 2: Histogram representing the distance from the pluripotent consensus fingerprint for all GEO (above), curated pluripotent samples (below), and GSE26946 samples (below, red)