

List of Figures

1	First principal component of methylation modules	2
2	Heatmap of methylation modules	3

1 Preprocessing

1.1 Gene expression array data

Gene expression array data for 344 AML patients was downloaded from GEO (GSE 14468). Normalization was performed in Gentles et al 2015 [1]. Annotation of the probes to gene with nearest transcription start site were available from GEO. We query for the Gene Ontology terms of the corresponding genes and filtered the probes based on their GO biological process annotation to obtain genes related to methylation and transcriptional regulation. After filter for genes that have keyword “methylation” or “transcrip”, 3417 out of 17788 are left for the downstream analysis.

1.2 Methylation array data

Methylation array data for 344 AML patients was downloaded from GEO (GSE 18700). Annotation of the probes to the nearest transcription start site allowing for a maximum distance of 5 kb from the TSS was provided along with the dataset. Any probes that were mapped to a gene are filtered for downstream analysis and in total 22759 out of 25626 probes are left after this step.

Standardization was performed on the probes across patients. PhenoGraph [3] were used to cluster probes. PhenoGraph first constructs a nearest-neighbor graph ($k = 30$) that has individual probe as a node. The edge weight between probe pairs are computed as the Jaccard similarity coefficients between their nearest neighbor set. Louvain community detection was performed in this algorithm to find the clustering assignment that maximizes the modularity on the graph. After applying PhenoGraph on the 22759 methylation probes, 20 communities (modules) were obtained. The counts of probes in each module along with the first principal component of each module are shown in the Figure 1.

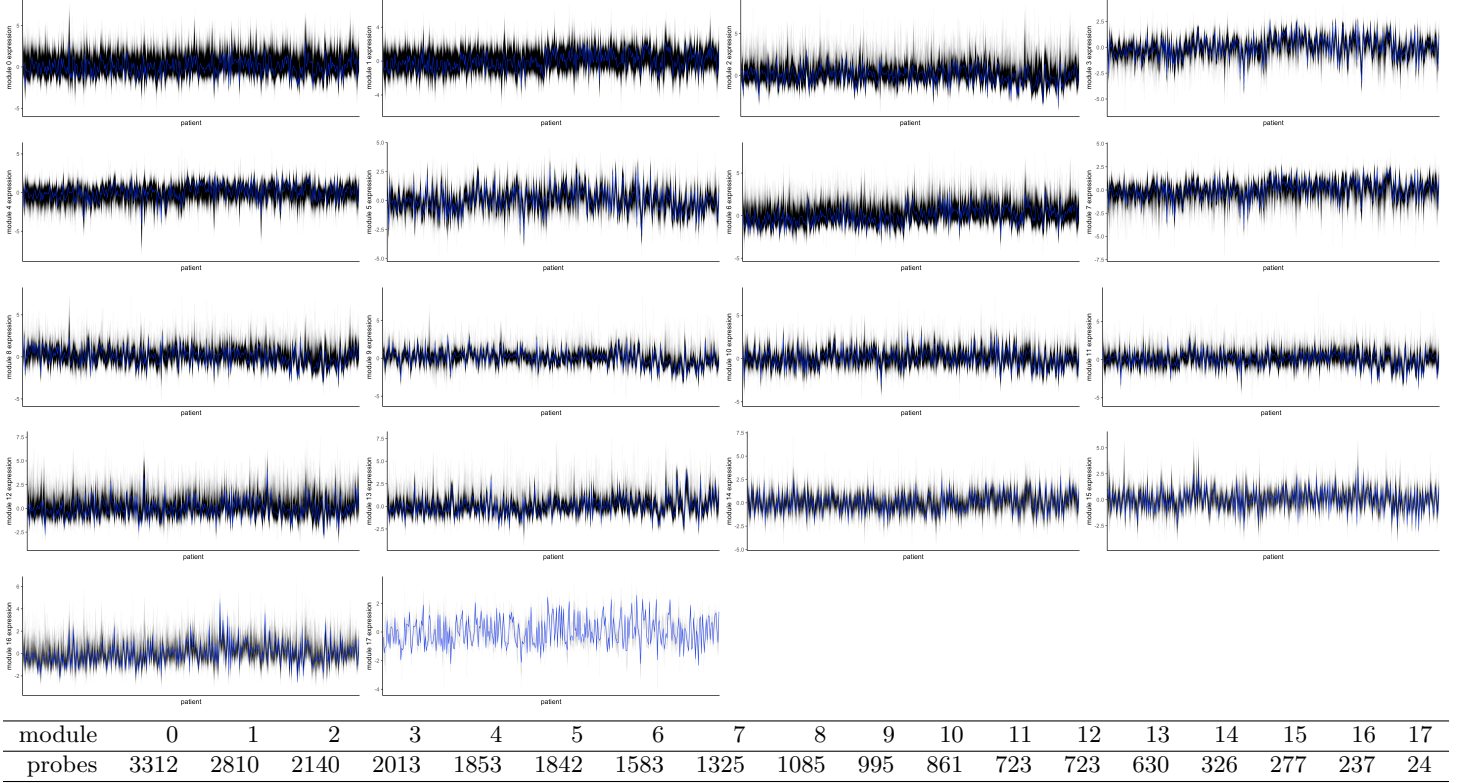


Figure 1: **First principal component of methylation modules** Each panel represent a module and the black lines represent the normalized methylation level of a probe across 344 patients. The blue lines show the first principal component of each module. The size of each module is shown in the table

In addition, the heatmap of expression level of the clusters are shown in Appendix Figure 2.

A Methylation array heatmap

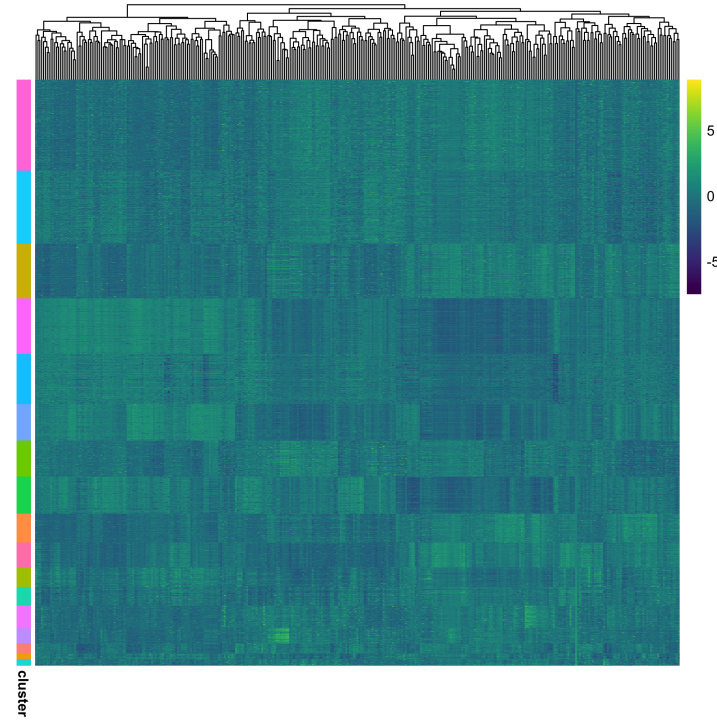


Figure 2: **Heatmap of methylation modules** Heatmap of methylation array expression. The rows are annotated with the PhenoGraph clusters and columns are arranged by hierarchical clustering on patients using Euclidean distance.

References

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- [2] M. Ashburner, C. A. Ball, J. A. Blake, D. Botstein, H. Butler, J. M. Cherry, A. P. Davis, K. Dolinski, S. S. Dwight, J. T. Eppig, *et al.*, “Gene ontology: tool for the unification of biology,” *Nature genetics*, vol. 25, no. 1, p. 25, 2000.
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