The CTC saga: C3 comparison

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C3 comparison

This is the comparison between cancer tissue and normal blood. We are using the cancer tissue samples from the c1 comparison, i.e.:

- GSE22820: "Gene expression profiles were generated from 176 primary breast cancer patients and 12 normal breast samples", PMID: 21356353 (Agilent Whole Human Genome Microarray 4x44K G4112F): 176 cancer tissues (we don't take 2 metastatic cancers)
- GSE19783: "mRNA profiling from 115 breast cancer samples was performed", PMID: 21364938 (Agilent Whole Human Genome Microarray 4x44K G4112F): 113 cancer tissues
- GSE31364: "Seventy-two primary breast cancer tumor have been analyzed against a breast cancer reference pool.", PMID: 22384245 (Agendia_human_DiscoverPrint_v1 custom platform): 72 cancer tissues
- GSE18672: "Mammographic density and genetics A study of breast biopsies in relation to mammoraphic density", PMID: 20799965 (Agilent Whole Human Genome Oligo Microarray G4112A): 64 cancer tissues

Furthermore we are using the following datasets providing cancer samples:

- GSE15852: "Gene expression patterns distinguish breast carcinomas from normal breast tissues: the Malaysian context" (PMID: 20097481). Platform: Affymetrix genechip U133A. 43 cancer patients
- GSE12763: "In vivo antitumor activity of MEK and phosphatidylinositol 3-kinase inhibitors in basal-like breast cancer models" (PMID: 19567590). Platform: Affymetrix Human Genome U133 Plus 2.0 Array 30 human breast cancers

For the normal blood samples we are using the ones from (my) c2 comparison, that is:

- GSE27562: "In total, we collected blood from 57 women with a diagnosis of breast cancer and 37 with a benign diagnosis" (PMID: 21781289) Platform: Affymetrix: 31 normal blood samples
- GSE16443: "Diagnostic work-up revealed that 67 of these women had breast cancer while 54 had no malignant disease. Additionally, 9 samples from 6 healthy female controls (three pregnant women, one breast-feeding woman and two healthy controls at different timepoints in their menstrual cycle) were included." (PMID: 20078854). Platform: Applied Biosystems (ABI): 54 normal blood samples

The merge of these datasets result in 6421 (common) genes and 510 samples (85 normal blood samples and 425 cancer tissue).

> print.xtable(xtable(table(c3.data\$Origin, c3.data\$Study)), comment = FALSE)

	GSE12763	GSE15852	GSE16443	GSE19783	GSE22820	GSE27562	GSE31364
Cancer Tissue	30	43	0	113	176	0	72
Normal Blood	0	0	54	0	0	31	0

> plotMDS(c3.data, "Study", "Origin", main = "Distribution of samples in the merged datasets

Differential expression

We find 275 genes over-expressed in cancer samples.

Distribution of samples in the merged datasets

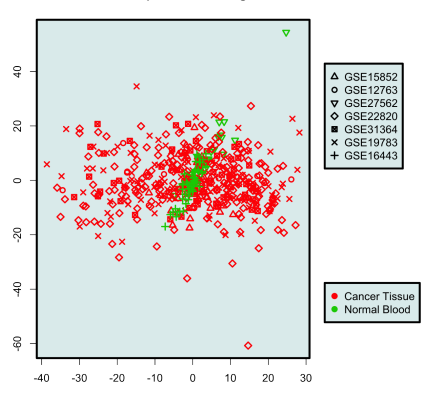


Figure 1: 2D (MDS) plot of the merged dataset