Identification of sample annotation errors in gene expression datasets

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Abstract  The comprehensive transcriptomic analysis of clinically annotated human tissue has found widespread use in oncology, cell biology, immunology, and toxicology. In cancer research, microarray-based gene expression profiling has successfully been applied to subclassify disease entities, predict therapy response, and identify cellular mechanisms. Public accessibility of raw data, together with corresponding information on clinicopathological parameters, offers the opportunity to reuse previously analyzed data and to gain statistical power by combining multiple datasets. However, results and conclusions obviously depend on the reliability of the available information. Here, we propose gene expression-based methods for identifying sample misannotations in public transcriptomic datasets. Sample mix-up can be detected by a classifier that differentiates between samples from male and female patients.

Correlation analysis identifies multiple measurements of material from the same sample. The analysis of 45 datasets (including 4913 patients) revealed that erroneous sample annotation, affecting 40% of the analyzed datasets, may be a more widespread phenomenon than previously thought. Removal of erroneously labelled samples may influence the results of the statistical evaluation in some datasets. Our methods may help to identify individual datasets that contain numerous discrepancies and could be routinely included into the statistical analysis of clinical gene expression data.

Keywords  Gene expression · Microarray · Misannotation · Quality control · Male–female classifier

Introduction

The generation of large gene expression datasets presents a logistic challenge that extends from the initial procurement and storage of tissue samples, through laboratory procedures, to bioinformatic data processing and analysis. Although anticipated to be low, little is known about the actual frequency of sample mix-up during this multi-step process. The reasons for sample identity being swapped between individuals are diverse, and these events are difficult to pinpoint retrospectively with absolute certainty. In datasets with roughly balanced frequencies of male and female individuals, it can be assumed that approximately half of the mix-ups will result in sex mislabeling. These cases can be identified by assessment of genes with male- or female-specific expression. Other commonly annotated clinicopathological parameters, such as tumor stage, would also be affected by mislabeling, but the lack of genes that exhibit for instance a reliable stage-specific expression...
pattern makes the standardized assessment of these parameters unsuitable.

Few attempts have been made to systematically identify sample mix-ups in public gene expression datasets. The MixupMapper software (Westra et al. 2011) requires DNA sequence data (SNP) in addition to gene expression data. However, the majority of previous studies are based exclusively on gene expression data. Recent approaches use the expression of the X-chromosomal gene \(XIST\) and genes located on the Y chromosome for the discrimination between male and female samples in the analysis of single datasets. However, these methods are not generalizable because of the lack of normalization across datasets (‘t Hoen et al. 2013; Broman et al. 2015).

To gain insight into frequencies of sample annotation discrepancies in publicly available gene expression datasets, we established a male–female classifier based on gene expression array data. In addition, correlations between expression values for pairs of samples were assessed to identify multiple measurements of tissue from the same individual, as this represents an additional hypothetical source of inconsistencies with regard to sample annotation.

### Methods

In this investigation, 45 publicly available MIAME-compliant sample collections were included (see Tables 1, 2 for details), all with accessible gene expression array data and available information on male or female sex for each study subject. In total, the studies comprised 4913 patients (3034 females, 1879 males). Gene expression array data and information on male or female sex for each study subject were accessed from the Gene Expression Omnibus (GEO) or directly from the authors’ Web site (Edgar et al. 2002; Shedden et al. 2008; Bild et al. 2006). Only datasets using the AffymetrixGeneChip® HG-U133A or HG-U133 Plus 2.0 were included in this analysis.

To construct the classifier, we proceeded in three steps: 1. selection of probe sets with male- or female-specific expression, 2. dataset normalization to enable analysis of unlabelled cohorts and cohorts comprising only female or only male patients, and 3. combination of evidence from male- and female-specific probe sets into a final classifier that categorizes each sample as “correctly classified,” “misclassified,” or “unconfident.” In each step (1–3) a likelihood-based strategy was applied that ensures robustness against outliers (Algorithms 1–3 in Suppl. material).

The initial probe set selection was based on 10 publicly available non-small cell lung cancer (NSCLC) gene expression datasets analyzed on the AffymetrixGeneChip® HG-U133A or HG-U133 Plus 2.0 array (Suppl. material: Algorithm 1). For each sample, sex information and gene expression measurements for 22,277 probe sets were available. Only seven probe sets achieved median male–female classification accuracy above 75 % and only five above 90 %. The top four probe sets were included in the classifier (Table 3). Two of them map to the \(XIST\) (221728_x_at and 214218_s_at), located on the X chromosome, and the other two to \(RPS4Y1\) (201909_at) and \(DDX3Y\) (205000_at), respectively, both located on the Y chromosome. \(XIST\) is expressed from the inactive female X chromosome and silenced in men. This is illustrated in one NSCLC dataset (GSE31210), with high expression of \(XIST\) (221728_x_at) observed in all patients labelled as female (Fig. 1), but only in one sample labelled as male. Hence, this exception

### Table 1 Overview of analyzed datasets

| Type                  | Cohorts                                                                 | Sample size (female/male) |
|-----------------------|--------------------------------------------------------------------------|---------------------------|
| Non-small cell lung cancer | GSE37745, Shedden, GSE31547, GSE29013, GSE14814, GSE4573, GSE31210, GSE19188, GSE31546, GSE10445 | 1338 (594/744)            |
| Colon cancer          | GSE33113, GSE12945, GSE31595, GSE4271, GSE1433, GSE17536, GSE17537      | 769 (358/411)             |
| Other cancer          | GSE5720, GSE4107, GSE42952, GSE34111, GSE31684                          | 200 (64/136)              |
| Non-cancer            | GSE19027, GSE17913, GSE23343, GSE25462, GSE7821, GSE20950, GSE24427   | 408 (219/189)             |
| Breast cancer         | GSE11121, GSE2034, TRANSBIG (GSE7390/GSE6532), GSE16446, GSE20194, GSE20271, GSE22093, GSE23988 | 1373 (1373/0)             |
| Ovarian cancer        | Bild, GSE14764, GSE19829, GSE26712                                      | 426 (426/0)               |
| Prostate cancer       | GSE17951, GSE25136, GSE3325, GSE8218                                    | 399 (0/399)               |

Tissue collections and gene array datasets analyzed by the male–female classifier, if available identified by their Gene Expression Omnibus (GEO) Series (GSE) number.
### Table 2 Detailed description of analyzed datasets

| Cohort | # Female | # Male | # Total | Type (disease or subject of study) |
|--------|----------|--------|---------|-----------------------------------|
| GSE37745 | 89       | 107    | 196     | NSCLC                             |
| Shedden | 220      | 223    | 443     | NSCLC                             |
| GSE31547 | 36       | 14     | 50      | NSCLC + controls                  |
| GSE29013 | 17       | 38     | 55      | NSCLC                             |
| GSE14814 | 23       | 67     | 90      | NSCLC                             |
| GSE4573  | 47       | 82     | 129     | NSCLC                             |
| GSE31210 | 109      | 95     | 204     | NSCLC                             |
| GSE19188 | 23       | 59     | 82      | NSCLC                             |
| GSE31546 | 14       | 3      | 17      | NSCLC                             |
| GSE10445 | 16       | 56     | 72      | NSCLC                             |
| GSE4107  | 12       | 10     | 22      | Colorectal cancer                 |
| GSE33113 | 48       | 42     | 90      | Colorectal cancer                 |
| GSE31595 | 22       | 15     | 37      | Colorectal cancer                 |
| GSE12945 | 28       | 34     | 62      | Colorectal cancer                 |
| GSE14333 | 106      | 120    | 226     | Colorectal cancer                 |
| GSE17536 | 81       | 96     | 177     | Colorectal cancer                 |
| GSE17537 | 29       | 26     | 55      | Colorectal cancer                 |
| GSE4271  | 32       | 68     | 100     | Other cancer: glioma              |
| GSE31684 | 25       | 68     | 93      | Other cancer: bladder             |
| GSE34111 | 6        | 24     | 30      | Other cancer: gastrointestinal    |
| GSE5720  | 24       | 30     | 54      | Other cancer: 9 different tissues |
| GSE42952 | 9        | 14     | 23      | Other cancer: pancreatic          |
| GSE19027 | 11       | 48     | 59      | Bronchial epithelium of (non-) smokers with and without lung cancer |
| GSE17913 | 38       | 40     | 78      | Smoking                           |
| GSE23343 | 7        | 10     | 17      | Insulin resistance/type 2 diabetes |
| GSE25462 | 28       | 22     | 50      | Insulin resistance/type 2 diabetes |
| GSE7821  | 28       | 12     | 40      | Healthy twins                     |
| GSE20950 | 27       | 12     | 39      | Insulin resistance/obesity        |
| GSE24427 | 80       | 45     | 125     | Multiple sclerosis                |
| GSE11121 | 200      | 0      | 200     | Breast cancer                     |
| GSE2034  | 286      | 0      | 286     | Breast cancer                     |
| TRANSBIG (GSE7390/GSE6532) | 280 | 0 | 280 | Breast cancer |
| GSE16446 | 114      | 0      | 114     | Breast cancer; chemo response     |
| GSE20194 | 247      | 0      | 247     | Breast cancer; chemo response     |
| GSE20271 | 139      | 0      | 139     | Breast cancer; chemo response     |
| GSE22093 | 47       | 0      | 47      | Breast cancer; chemo response     |
| GSE23988 | 60       | 0      | 60      | Breast cancer; chemo response     |
| Bild    | 133      | 0      | 133     | Ovarian cancer                    |
| GSE14764 | 80       | 0      | 80      | Ovarian cancer                    |
| GSE19829 | 28       | 0      | 28      | Ovarian cancer                    |
| GSE26712 | 185      | 0      | 185     | Ovarian cancer                    |
| GSE17951 | 0        | 153    | 153     | Prostate cancer                   |
| GSE25136 | 0        | 79     | 79      | Prostate cancer                   |
| GSE3325  | 0        | 19     | 19      | Prostate cancer                   |
| GSE8218  | 0        | 148    | 148     | Prostate cancer                   |

Overview over the studied tissue collections and gene array data
was clearly located in the female XIST expression range. RPS4Y1 and DDX3Y showed the opposite behavior, with high expression values observed in male patients. RPS4Y1 encodes a structurally conserved ribosomal protein with putative function during spermatogenesis (Lopes et al. 2010), whereas DDX3Y is primarily expressed in testis and is involved in germ-line translation control (Rauschendorf et al. 2011). Probe sets with low discriminating power were not included in the classifier.

The expression levels of the four selected probe sets were evaluated in 35 additional datasets, including seven colon cancer, five other cancer, and seven non-cancer datasets containing samples from both male and female subjects, as well as eight breast cancer, four ovarian cancer, and four prostate cancer datasets. A plot of raw expression values for the probe set 201909_at (RPS4Y1) across all datasets showed high male–female classification accuracy per dataset, but large overall expression shifts between datasets (Fig. 2a). After normalizing expression values with a linear transformation to median values of 0 and 1 for the low and high expression groups, respectively (Suppl. material: Algorithm 2), expression levels were reliably comparable across cohorts (Fig. 2b).

In a final step, the four sex-specific probe sets were combined to categorize each sample as “correctly classified,” “misclassified,” or “unconfident” (Suppl. material: Algorithm 3). First, for each cohort and for each probe set, the expression values were clustered into two groups of low and high values and a normal distribution was fitted to the low expression group, estimating location and scale with robust measures (median and Rousseeuw–Croux estimator Qn (Rousseeuw and Croux 1993)). Next, the expression value of the probe set for each sample was compared to the 99.9 % quantile of the fitted normal distribution. A value above this cut point is inconsistent with the typical range for the low expression group and thus provides strong evidence that the corresponding sample belongs to the high expression group. For each individual sample, a female-evidence score was then defined for each of the two XIST probe sets. As high XIST expression is inconsistent with male sex, the female-evidence score was set to 1 if the corresponding XIST expression value was above the cut point. Analogously, for DDX3Y and RPS4Y1, respectively, a male-evidence score was set to 1 if the expression value of the probe set was above the corresponding cut point. Taking the evidence scores of all four probe sets into account, a sample was classified as male if at least one male-evidence score was 1 and both female-evidence scores were 0. Vice versa, a sample was classified as female if at least one female-evidence score was 1 and both male-evidence scores were 0. Finally, the new classifications were compared to the original sex annotations, categorizing each sample as “correctly classified,” “misclassified,” or “unconfident.” Samples with both at least one positive female-evidence score and at least one positive male-evidence score, or with no positive evidence score, were classified as “unconfident.”

### Table 3: Probe sets included in the male–female classifier

| Affymetrix ID | Gene | Chromosome | Cut point (99% quantile) | Evidence (male/female) |
|---------------|------|------------|--------------------------|------------------------|
| 221728_x_at   | XIST | X          | >0.389                   | Female                 |
| 214218_s_at  | XIST | X          | >0.385                   | Female                 |
| 201909_at     | RPS4Y1 | Y          | >0.431                   | Male                   |
| 205000_at     | DDX3Y | Y          | >0.276                   | Male                   |

Probe sets included into the male–female classifier, with corresponding cut points for evidence whether a sample originates from a male or a female.
Results and discussion

The male–female classifier was applied to all 45 cohorts, categorizing 4913 patients (3034 females, 1879 males) (Fig. 3). In total 54 patients (1.1 %) were categorized as “misclassified” and 149 (3.0 %) were labelled “unconfident.” The direction of sex mislabeling was nearly balanced, with 29 female samples mislabeled as male and 25 male samples mislabeled as female. Overall, in 18 of the 45 cohorts (40 %) at least one “misclassified” sample was detected. The proportion of “correctly classified” samples was 100 % in 15 cohorts, below 90 % in five cohorts, and in between for the remaining 25 cohorts. Note that these numbers are probably overoptimistic, as 16 cohorts included in the study consisted of breast, ovarian, or prostate cancer patients, with lower risk of sex mislabeling. Still, one breast cancer patient in the cohort TRANSBIG (comprising node-negative untreated patients of GSE7390 and GSE6532) was classified as male (Fig. 3).

The prevalence of sample identity inconsistencies in public data repositories can be anticipated to be at least twice as high as indicated by the male–female classifier, as mix-up may occur also between samples from individuals of the same sex. To visualize the expression-based sex assignment per cohort, we plotted mean normalized expression values of the two X-chromosomal probe sets and the
Fig. 3 Application of the male–female classifier. Application of the male–female classifier to all cohorts, cohorts grouped by cancer type. Green “correctly classified,” red “misclassified,” and orange “unconfident” samples.

Fig. 4 Visualization of the male–female classifier with mean expression values of the two probe sets for XIST on the x-axis and DDX3Y and RPS4Y1 on the y-axis. The points represent individual patients. The point clouds on the left are characteristic for males and females, respectively. Colors indicate classification accuracy samples. Green “correctly classified,” red “misclassified,” and orange “unconfident.”

a Results for the Uppsala cohort (GSE37745): One female patient clearly mislabeled as male, and two samples are labeled “unconfident.”

b Results for GSE33113 with clear discrimination between males and females and no sex misannotations.

c Results for GSE5720 with two misclassified samples and large number of samples classified as “unconfident.”

d Results for a breast cancer dataset (TRANSBIG) with one male patient assigned to the category “misclassified.”
two Y-chromosomal probe sets against each other (Fig. 4). For most cohorts, two clearly distinguishable groups representing males and females can be recognized, and category assignment by visual inspection is well in agreement with our likelihood-based classifier.

A further error that may occur during tissue processing is sample duplication. The same sample may be analyzed twice and the duplicate is erroneously labelled with the identification number of another patient. To identify such duplications, a correlation-based analysis strategy was applied. For each cohort, the 1000 probe sets with highest variance across all samples were selected and Pearson correlation coefficients between all pairs of samples in the cohort were calculated. The largest distance between all ordered values of correlations was identified to distinguish between duplicated measurements and pairs of measurements from different samples. In 15 of the 45 cohorts at least one duplicate was identified. In total 32 duplicates were detected. Comparing these duplicates with the results from the male–female classifier, nine of the 54 “misclassified” assignments (16.7 %) could be explained by duplicated measurements.

The general impact of misannotated samples on gene expression is difficult to assess. To illustrate the relevance of misannotations in gene expression studies, we re-analyzed six lung cancer cohorts with available survival times. Prognostic relevance of a gene was determined by fitting a univariate Cox model (Cox 1972) to its expression values. The number of significant genes (p value <0.01; not FDR-adjusted) was first calculated for the original datasets. Removing all unambiguously misannotated samples from the six datasets with misannotations, 12–53 % of the previously significant genes were not significant any more. In contrast, using only the reduced number of samples, the number of newly discovered genes was in the range of 9–39 % of the original number of significant genes (Table 4).

To elucidate the reason behind the sample mislabeling observed in our own non-small cell lung cancer cohort (GSE37745), one patient annotated as male in the original records and assigned as female by our classifier was re-analyzed. First, new DNA and RNA samples were prepared from the original biobanked tissue specimen. Male sex was then confirmed based on the analysis of STR marker distribution using the AmpFLSTR® Identifiler® PCR Amplification Kit according to the manufacturer’s instructions (Applied Biosystems, Foster City, USA), suggesting that sample mix-up in this case did not occur during sample collection and biobanking procedures. Subsequently, the gene expression array analysis was repeated for the misclassified sample and for five additional control samples from the previously analyzed cohort. The pairwise correlation between the new and old misclassified sample was only 0.464, strongly indicating that these two samples were derived from different individuals. In contrast, a striking correlation of 0.993 was detected between the misclassified sample and a sample from one other female patient in the previously analyzed cohort. The high correlation suggests that the mRNA sample from one female patient erroneously had been measured twice in the previous analysis. A second duplicated measurement was detected, with correlation 0.990 between the expression values of two patients with sex label male. In contrast, all correlations of the repeated control samples with the corresponding original measurements were high (correlation coefficients: 0.910–0.987).

| Dataset     | No. of patients | No. of misannotations and duplications | No. of significant genes (original scenario) | Percentage of genes no longer significant after removal of the misannotated samples | Percentage of genes newly significant after removal of the misannotated samples |
|-------------|-----------------|----------------------------------------|---------------------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| GSE37745    | 196             | 3                                      | 450                                         | 12.22                                                                           | 14.00                                                                           |
| Shedden     | 443             | 14                                     | 1354                                        | 15.66                                                                           | 8.79                                                                            |
| GSE29013    | 55              | 1                                      | 419                                         | 15.51                                                                           | 14.32                                                                           |
| GSE4573     | 129             | 5                                      | 189                                         | 26.63                                                                           | 38.62                                                                           |
| GSE31547    | 50              | 1                                      | 318                                         | 50.51                                                                           | 23.27                                                                           |
| GSE19188    | 82              | 8                                      | 190                                         | 53.16                                                                           | 34.374                                                                          |

Table 4 Results of univariate Cox models

Results of univariate Cox models for six NSCLC datasets. Comparison between significance genes (p < 0.01) identified in the original cohort and significance genes identified in the reduced cohort after removal of misannotated and duplicated samples.
was detected. The easy-to-use classifier presented here, combined with correlation analysis to detect samples erroneously measured multiple times, helps to identify individual datasets that contain numerous discrepancies. Re-evaluation of gene expression array data demonstrated that sample mislabeling may have a considerable impact on the output of the statistical evaluation and allows inferences on the accuracy of biobanking. In conclusion, methods for identifying sample misannotations should be routinely included into the statistical analysis of clinical gene expression data.

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Compliance with ethical standards

Conflict of interest The authors declare no conflict of interest.

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References

Bild AH, Yao G, Chang JT, Wang Q, Potti A, Chasse D, Joshi MB, Harpole D, Lancaster JM, Berchuck A, Olson JA Jr, Marks JR, Dressman HK, West M, Nevins JR (2006) Oncogenic pathway signatures in human cancers as a guide to targeted therapies. Nature 439:353–357

Broman KW, Keller MP, Broman AT, Kendziorski C, Yandell BS, Sen S, Attie AD (2015) Identification and correction of sample mix-ups in expression genetic data: a case study. G3. Bethesda 5(10):2177–2186. doi:10.1534/g3.115.019778

Cox DR (1972) Regression models and life tables: a case study. J R Stat Soc Series B Stat Methodol 34(2): 187–220a. arXiv:1402.2633 [stat.AP]

Edgar R, Domrachev M, Lash AE (2002) Gene expression omnibus: NCBI gene expression and hybridization array data repository. Nucleic Acids Res 30:207–210

Lopes AM, Miguel RN, Sargent CA, Ellis PJ, Amorim A, Affara NA (2010) The human RPS4 paralogue on Yq11.223 encodes a structurally conserved ribosomal protein and is preferentially expressed during spermatogenesis. BMC Mol Biol 11:33

Petermann KB, Rozenberg GL, Zedek D, Groben P, McKinnon K, Buehler C, Kim WY, Shields JM, Penland S, Bear JE, Thomas NE, Serody JS, Sharpless NE (2007) CD200 is induced by ERK and is a potential therapeutic target in melanoma. J Clin Invest 117(12):3922–3929

Rauschendorf MA, Zimmer J, Hanstein R, Dickemann C, Vogt PH (2011) Complex transcriptional control of the AZFa gene DDX3Y in human testis. Int J Androl 34(1):84–96

Rousseeuw PJ, Croux C (1993) Alternatives to the median absolute deviation. J Am Stat Assoc 88(424):1273–1283

Rousseeuw PJ, Croux C (1993) Alternatives to the median absolute deviation. J Am Stat Assoc 88(424):1273–1283

Shedden K, Taylor JM, Enkemann SA, Tsao MS, Yeatman TJ, Gerald WL, Eschrich S, Jurisica I, Giordano TJ, Misek DE, Chang AC, Zhu CQ, Strumpf D et al (2008) Gene expression-based survival prediction in lung adenocarcinoma: a multi-site, blinded validation study. Nat Med 14(8):822–827

’t Hoen PA, Friedländer MR, Almlof J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Sammeth M, Pulyakhina I, Anvar SY, Kor不失 J, Samm

Verhaak RG, Tamayo P, Yang JY, Hubbard D, Zhang H, Creighton CJ, Fredrik O, Lawrence M, Carter SL, Mermel CH, Kostic AD, Etemadmoghadam D, Saksena G (2013) Prognostically relevant gene signatures of high-grade serous ovarian carcinoma. J Clin Invest 123(1):517–525

Westra HJ, Jansen RC, Fehmman RS, te Meerman GJ, van Heel D, Wijmenga C, Franke L (2011) MixupMapper: correcting sample mix-ups in genome-wide datasets increases power to detect small genetic effects. Bioinformatics 27(15):2104–2111