

BRIEF COMMUNICATION

The Doppelgänger Effect: Hidden Duplicates in Databases of Transcriptome Profiles

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Abstract

Whole-genome analysis of cancer specimens is commonplace, and investigators frequently share or re-use specimens in later studies. Duplicate expression profiles in public databases will impact re-analysis if left undetected, a so-called “doppelgänger” effect. We propose a method that should be routine practice to accurately match duplicate cancer transcriptomes when nucleotide-level sequence data are unavailable, even for samples profiled by different microarray technologies or by both microarray and RNA sequencing. We demonstrate the effectiveness of the method in databases containing dozens of datasets and thousands of ovarian, breast, bladder, and colorectal cancer microarray profiles and of matching microarray and RNA sequencing expression profiles from The Cancer Genome Atlas (TCGA). We identified probable duplicates among more than 50% of studies, originating in different continents, using different technologies, published years apart, and even within the TCGA itself. Finally, we provide the *doppelgangR* Bioconductor package for screening transcriptome databases for duplicates. Given the potential for unrecognized duplication to falsely inflate prediction accuracy and confidence in differential expression, doppelgänger-checking should be a part of standard procedure for combining multiple genomic datasets.

Sufficient germ-line sequence markers provide a “fingerprint” that can be matched uniquely in a database of genotypes (1). Publicly available human genomic data is therefore normally summarized at a level that cannot be identified uniquely to protect patient privacy. Cancer transcriptomes undergo alterations that are highly distinctive but much more difficult to identify uniquely in summarized form. Re-use of tissue specimens is widespread in clinical genomic studies, creating a “doppelgänger effect” in publicly available datasets: hidden duplicates that, if left undetected, can inflate statistical significance or apparent accuracy of genomic models when combining data from different studies (Figure 1A). The proposed method relies on exhaustive comparisons of dataset pairs and sample pairs to empirically estimate the distribution of pairwise transcriptome correlations between biological replicates within a dataset or between two datasets where potentially different profiling technologies were

used. The key aspects to identifying duplicates in a pair of datasets are 1) using transcript identifiers available in both datasets, 2) batch correction (2), 3) calculating Pearson’s Correlation Coefficient (PCC) between every sample in one dataset against every sample in the other dataset, and 4) duplicate-oriented outlier detection. The background distribution of pairwise PCC values varies depending on the tissue assayed and the technologies used, and must be estimated for every dataset pair. Doppelgängers can be identified as outliers at the high end of the distribution of batch-corrected correlations. The detailed methodology of package development and validation can be found in the [Supplementary Material](#) (available online).

We studied databases of ovarian, breast, bladder, and colorectal cancers and of cell lines and assessed their accuracy against a “gold standard” of duplicated samples generated through further manual inspection of expression data, clinical annotations, and

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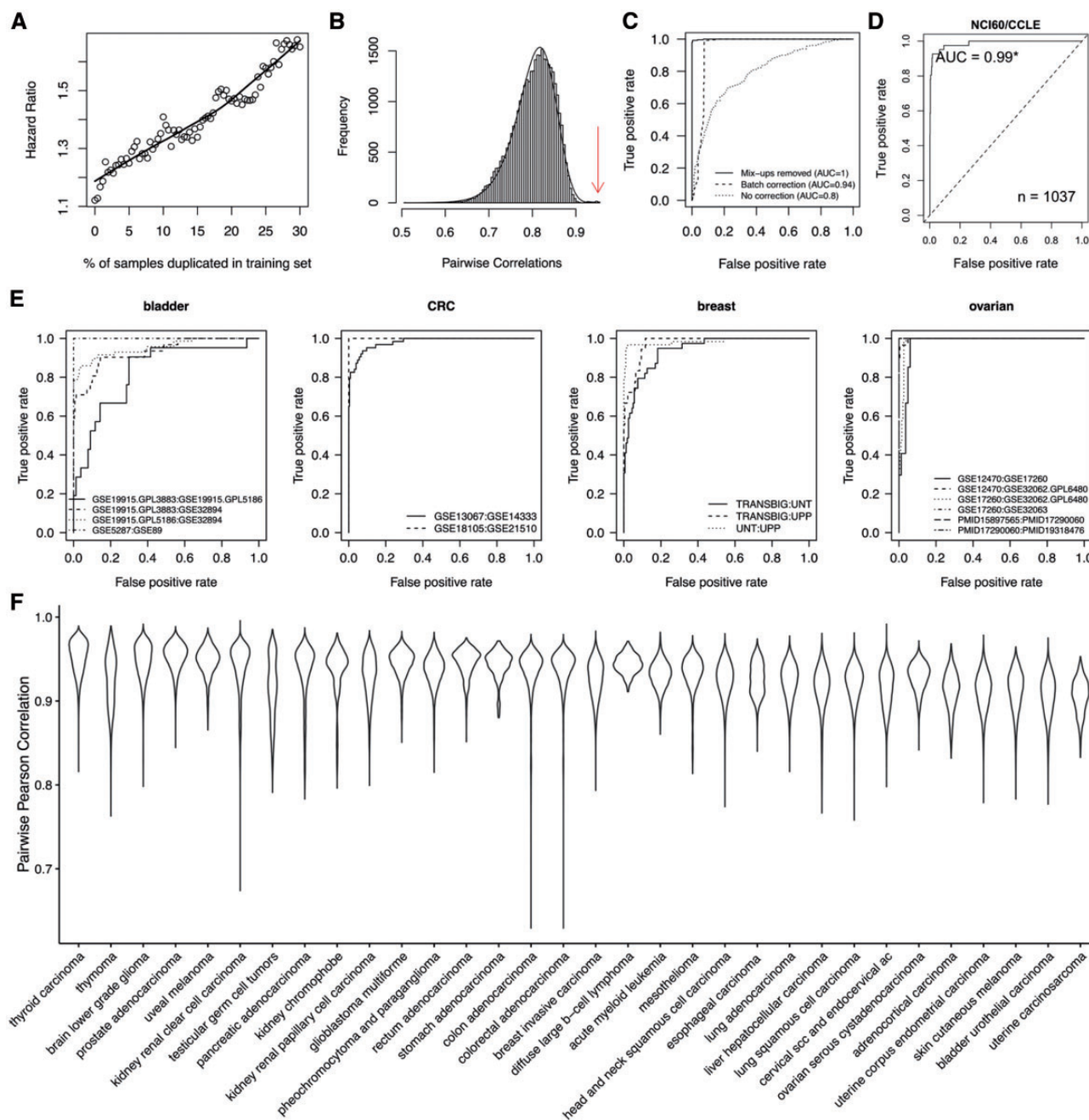


Figure 1. Demonstration and benchmarking of the *doppelgangR* method for identifying expression profiles of the same biological specimen. **A)** The “*doppelgänger*” effect: hidden duplicates can inflate the apparent accuracy of predictive and prognostic models. Models of overall survival for high-grade, serous ovarian cancer were trained and then validated in two studies containing duplicates identified by *doppelgangR* (see [Supplementary Methods](#), available online). Validation set hazard ratio (HR) was calculated with duplicates incrementally removed so that between 0% and 30% cross-study duplication of samples remained. Thirty percent duplication inflates the apparent hazard ratio from 1.1 to 1.7. **B)** *doppelgangR* identifies duplicate expression profiles as outliers with unusually high pairwise correlation compared with other pairs of unrelated expression profiles. This histogram is the diagnostic plot produced by *doppelgangR* software, showing the best fit to the distribution of pairwise correlations, with **vertical darklines** showing outliers that are probable duplicates in the UNT (3) and Miller et al. breast cancer datasets (4). **C)** Batch correction allows RNA-seq profiles to be matched accurately to Affymetrix microarray profiles in the The Cancer Genome Atlas (TCGA) ovarian cancer dataset. True positives are tumors whose RNA-seq and microarray profiles are more highly correlated to each other than to any other profile. Batch correction increases area under the receiver operating characteristic plot (AUC) from 0.79 to AUC = 0.94, and removing 50 microarray profiles incorrectly labeled by TCGA further increases AUC to > 0.995. **D and E)** Benchmarking. We estimated the accuracy of the *doppelgangR* approach by applying it to pairs of datasets with confirmed duplicates (see [Supplementary Methods](#), available online). **D)** Shows AUC for identifying the 43 cell lines present in two different panels (CCLE [n = 1037] and NCI60 [n = 59]). **E)** The performance on primary tumor data in four cancer types. The AUC averaged across the four cancer types is 0.97. **F)** Suitability to TCGA cancer types. The *doppelgangR* approach only works for cancer types in which expression profiles of individual tumors are sufficiently distinct. Violin plots depict distributions of Pearson Correlation Coefficients (PCCs) for all pairs of expression profiles within each TCGA dataset, in order (left to right) of increasing distinctiveness. In cancer types with high pairwise PCCs, such as thyroid carcinoma, patients have very similar expression profiles and are hard to distinguish based on expression data only. In contrast, in cancer types with low PCCs, such as bladder cancer, extensive genomic alterations generate unique expression fingerprints that make *doppelgänger* identification possible.

sample identifiers (Supplementary Table 1, available online). Confirmed doppelgängers were identified in more than half of all studies (Table 1). For example, among the 1467 breast cancer gene expression profiles, *doppelgangR* identifies 59 samples present in both the Sotiriou et al. (3) and Miller et al. (4) studies (Figure 1B; additional samples are duplicated by the TRANSBIG dataset, see Table 1). Although these studies were published by Belgian and Singaporean groups, respectively, careful reading of the papers reveals that their datasets shared a cohort of samples originating from Uppsala County, Sweden. Such international collaborations are beneficial to the cancer research community, but pose challenges to investigators developing independent validations and meta-analyses. In the ovarian cancer database, which we have inspected in great detail (5), we identified 17% of records as nonunique, including duplicates in different datasets originating from the same institution (6,7), between the TCGA dataset and datasets of institutions that contributed samples to the TCGA project (8,9) and within the TCGA dataset itself. In approximately 75% of duplicate pairs, samples matched by expression data had identical or compatible clinical and tumor data, but in the other 25% of cases the clinical data were discordant (10). Previous work on identifying duplicate microarray profiles has been limited to matching identical raw data files (11), and this would not identify any of these duplicates.

In addition to identifying samples reprofiled by microarray, our approach accurately aligns microarray and log-transformed RNA-seq profiles for the same patients with area under the receiver operating characteristic curve (AUC) greater than 0.9 for

10 of 12 TCGA cancer types where both microarray and RNA-seq are available (Supplementary Table 2, available online). AUC is less than 0.9 for two cancer types among the 10 “quietest” genomes (Figure 1F), kidney renal clear cell and papillary cell carcinoma. Batch correction across datasets is critical: For example, in ovarian cancer, batch correction increases AUC from 0.80 to 0.94. Further inspection of the remaining errors reveals that almost all were because of an experimental RNA mix-up in the original TCGA Affymetrix microarray dataset, resulting in erroneously duplicated profiles attributed to different patients. Correction of the mixed-up samples increases AUC for matching RNA-seq to microarray profiles to greater than 0.995 (Figure 1C). We reported this sample mix-up to the TCGA Data Coordinating Center, which in turn removed these 50 profiles on August 25, 2015 (12).

Our approach of duplicate identification reliably works when individual tumors have distinctive expression profiles, as is the case for cell line panels (Figure 1D) and for primary tumors from breast, ovarian, bladder, and colorectal cancers (Figure 1E). We expected it to be more prone to false positives for less differentiated expression profiles such as low-grade and early-stage tumors, and, generally observed, this where sufficient numbers of annotated samples were available: Samples falsely identified as duplicates were enriched for low-grade (CRC: 95% confidence interval [CI] = 1.2 to 2.2; ovarian: 95% CI = 1.0 to 1.5) and early-stage (bladder: 95% CI = 1.3 to 4.2; CRC: 95% CI = 1.6 to 2.6). The exception was early-stage ovarian cancer samples, for which *doppelgangR* was extremely effective. These samples have distinctive profiles, and their rate of sharing was high, possibly

Table 1. Overview of confirmed doppelgängers in all studies*

Dataset identifier by type of cancer	Total No. samples	No. of doppelgängers	Institutional source of doppelgängers
Bladder			
GSE1827, GSE13507, GSE31189, GSE31684, GSE37317, PMID: 17099711	570	0	Various, no doppelgängers identified
GSE19915, GSE32894	490	84	University Hospital of Lund, Sweden
GSE89, GSE5287	70	2	Aarhus University Hospital, Denmark
Breast			
MAINZ, NKI, VDX	881	0	Various, no doppelgängers identified
TRANSBIG, UNT, UPP	586	78	Uppsala County, Sweden
Colorectal			
GSE2109, GSE3964, GSE4045, GSE11237, GSE12225, GSE12945, GSE13294, GSE26682, GSE27544, GSE28702, GSE45270, TCGA (READ)	1275	0	Various, no doppelgängers identified
GSE13067, GSE14333	364	41	Royal Melbourne Hospital, Australia
GSE4526, GSE14095	225	37	Teikyo University School of Medicine, Japan
GSE14333, GSE17538	754	569	H. Lee Moffitt Cancer Center, USA
GSE18105, GSE21510, GSE21815	400	95	Tokyo Medical and Dental University Hospital, Japan
GSE26906, GSE39582	656	90	Various, France
GSE33113, TCGA (COAD)	226	2	Academic Medical Center, Netherlands
Ovarian			
GSE14764, GSE19829, GSE26712, GSE30161, GSE44104, GSE49997, GSE6008, GSE6822, GSE8842, GSE9891, GSE12418, GSE13876	1415	0	Various, no doppelgängers identified
E-MTAB-386, GSE18520	192	1	Brigham and Women's Hospital, USA
GSE12470, GSE17260, GSE32062, GSE32063	463	139	Niigata University, Japan
GSE20565, GSE26193	247	93	Resource Biological Center of the Institut Curie, France
TCGA, GSE2109, GSE51088, PMIDs: 15897565, 17290060, 19318476	1176	2	International Genomics Consortium (IGC)
		10	Cedars-Sinai Medical Center, USA
		88	Duke University Medical Center, USA

*Gene expression data were obtained from several R/Bioconductor packages (Supplementary Methods, available online), and the listed ids are the study ids given in these packages.

because of the rarity of early-stage ovarian cancer and the high importance of specimens.

We investigated the potential for applying this method to each of the 32 TCGA cancer types based on individual distinctiveness of transcriptomic aberrations. Using log-transformed level III RNA-seq data, which are summarized at the level of gene symbols, we calculated correlations between all sample pairs (Figure 1F). Our ranking of the cancer types for transcriptome distinctiveness, based on the 99.9th percentile of correlation between expression profiles, is very similar to Lawrence et al. (13), in which tumors were ranked by mutation rate. We do not expect our method to be effective for cancer types with “quiet” genomes and high correlation between nonduplicate expression profiles, such as kidney renal papillary cell carcinoma and the cancer types to its left in Figure 1F. We also note the existence of highly distinctive expression subtypes, such as in glioblastoma multiforme IDH1 mutant vs wild-type cases (14), which produce bimodal pairwise correlations that may complicate duplicate identification if these subtypes are not separated. Several other cancer types from TCGA exhibit distinctive subtypes that should be considered (Supplementary Figure 1, available online). Known subtypes in the datasets we examined in detail did not impact performance, however, such as prevalence of estrogen receptor–positive breast cancer tumors.

We note the potential utility of the doppelgänger approach also for identifying duplicates within a single study with as few as five samples (see Supplementary Results and Supplementary Figures 2–3, available online). In the databases we studied, within-study duplication was less common than between studies, but we found likely duplicates in six ovarian cancer studies and one CRC study. These profiles were such extreme outliers in terms of similarity of gene expression profiles that they are unlikely to have originated from different tumors, assuming the samples passed basic QC metrics (eg, coverage, tumor purity, RNA integrity scores). The approach reliably grouped together healthy tissues, which have much more homogeneous expression profiles than cancer tissues. Although our benchmarking is limited to microarray and RNA-seq data, we see no reason why this approach should not work for other quantitative mRNA assays such as nanoString and multiplexed quantitative real-time polymerase chain reaction, or even proteomic or other molecular profiles, provided that biological replicates are sufficiently distinct relative to technical replicates.

As genomic databases grow and collect tumor specimens from international collaborators, the chance of inter- and intrastudy duplication increases. Analysis of duplicate samples is a substantial concern that could alter the identification of subsets of patients with clinical differences or the development of specific gene signatures. While this approach can help identify duplicate profiles even when germ-line sequences are not available, we note some limitations. Automatic setting of the threshold defining “outliers” is a difficult problem to solve generally and should be reviewed using diagnostic histograms generated by the *doppelgangR* package. A thorough review

should take into consideration potential collaborations and/or multiple institution clinical management of individual patients, as well as suspiciously similar clinical patient data and identifiers.

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Notes

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