

Original article

curatedOvarianData: clinically annotated data for the ovarian cancer transcriptome

Benjamin Frederick Ganzfried^{1,2,†}, Markus Riester^{1,2,†}, Benjamin Haibe-Kains³, Thomas Risch¹, Svitlana Tyekucheva^{1,2}, Ina Jazic¹, Xin Victoria Wang^{1,2}, Mahnaz Ahmadifar¹, Michael J. Birrer⁴, Giovanni Parmigiani^{1,2}, Curtis Huttenhower² and Levi Waldron^{1,2,*}

¹Department of Biostatistics and Computational Biology, Dana-Farber Cancer Institute, Boston, MA, 02115, ²Department of Biostatistics, Harvard School of Public Health, Boston, MA, 02115, USA, ³Bioinformatics and Computational Genomics Laboratory, Institut de recherches cliniques de Montréal, Montréal, QC, H2W 1R7, Canada and ⁴Center for Cancer Research, Massachusetts General Hospital, Boston, MA, 02114, USA

[†]These authors contributed equally to this work.

*Corresponding author: Email: levi@jimmy.harvard.edu

Submitted 10 January 2013; Revised 16 February 2013; Accepted 19 February 2013

Citation details: Ganzfried,B.F., Riester,M., Haibe-Kains,B., et al. curatedOvarianData: clinically annotated data for the ovarian cancer transcriptome. *Database* (2013) Vol. 2013: article ID bat013; doi:10.1093/database/bat013

This article introduces a manually curated data collection for gene expression meta-analysis of patients with ovarian cancer and software for reproducible preparation of similar databases. This resource provides uniformly prepared microarray data for 2970 patients from 23 studies with curated and documented clinical metadata. It allows users to efficiently identify studies and patient subgroups of interest for analysis and to perform meta-analysis immediately without the challenges posed by harmonizing heterogeneous microarray technologies, study designs, expression data processing methods and clinical data formats. We confirm that the recently proposed biomarker *CXCL12* is associated with patient survival, independently of stage and optimal surgical debulking, which was possible only through meta-analysis owing to insufficient sample sizes of the individual studies. The database is implemented as the *curatedOvarianData* Bioconductor package for the R statistical computing language, providing a comprehensive and flexible resource for clinically oriented investigation of the ovarian cancer transcriptome. The package and pipeline for producing it are available from <http://bcb.dfci.harvard.edu/ovariancancer>.

Database URL: <http://bcb.dfci.harvard.edu/ovariancancer>

Introduction

A wealth of genomic data, in particular microarray data, is publicly available through diverse online resources. Major databases of gene expression data, e.g. the Gene Expression Omnibus (GEO) (1) and ArrayExpress (2), offer the potential to identify sets of genes predictive of cancer survival and of patient resistance to chemotherapy using thousands of samples from multiple laboratories. Such high numbers of samples are needed to robustly identify and validate gene signatures for incorporation into routine clinical practice (3). However, inconsistent formatting

among database interfaces, expression data storage and clinical metadata annotations present formidable obstacles to making efficient use of these resources.

Existing resources aiming to make large-scale high-dimensional analysis across multiple studies tend to serve only a few specifically targeted needs. To develop reproducible biomarker discovery methods appropriate for clinical translation, a data resource must be accurate and retain clinical variables of known importance as much as possible. The insilicoDB (4) project provides many curated gene expression data sets; however, it is not a focused resource in terms of retention or quality assurance of clinical

annotations, or retention of all relevant data sets and clinical variables for any one cancer type. The other major database of curated gene expression studies, the Gene Expression Atlas (2), provides machine- rather than manually annotated data, resulting in reduced consistency of annotation across studies. These are among the only databases that offer basics such as uniform gene identifiers to enable cross-study analysis, and then for only the most common microarray technologies. Carey *et al.* (5) describe a framework for the curation, annotation and storage of microarray and high-throughput data in general. This framework allows, for example, institutions to provide researchers access to in-house and public data in a standardized and convenient fashion. However, there is no existing database that provides these resources for ovarian cancer.

Ovarian cancer is the fifth-leading cause of cancer deaths among women (6) and has been the focus of numerous clinical transcriptome investigations. The *curatedOvarianData* database is the result of a focused effort to enable meta-analysis of these studies and to provide the highest quality and most comprehensive gene expression data resource for any cancer. It provides standardized gene expression and clinical data for 2970 ovarian cancer patients from 23 studies spanning 11 gene expression measurement platforms, in the form of documented ExpressionSet objects for R/Bioconductor (7). Gene expression data were collected from public databases and author websites, processed in a consistent manner and mapped uniformly to official Human Gene Nomenclature Committee (HGNC) (8) gene symbols. Curation of clinical annotations was machine-checked for correctness of syntax and human-checked by two individuals to ensure accuracy. This data package is geared primarily towards bioinformatic and statistical researchers, providing an ideal resource for development and assessment of algorithms for high-dimensional classification, clustering and survival analysis. It will also be valuable to ovarian cancer researchers for biomarker identification and validation. In addition to providing all publicly available gene expression studies with patient survival in common forms of ovarian cancer, it includes tumours of rare histologies, normal tissues and uncommon early-stage tumours. Special effort is made to retain the most important clinical variables from author-provided metadata and from the original publications: overall survival, optimal debulking surgery and tumour stage, grade and histology.

We also developed a software pipeline for automated and reproducible production of this and comparable data libraries. The pipeline includes a controlled language for curation of clinical annotations, defined by a template, which is intuitive for non-programmers to create and edit, but which is also used directly for machine syntax checking of curated annotations. The pipeline handles all steps of the process including data download, microarray preprocessing,

merging of duplicate probe sets and sample technical replicates, up-to-date probe-set to gene mapping and building of the R/Bioconductor objects and package.

One important application of the database is testing of hypothesized prognostic markers of ovarian cancer using multiple independent studies. We validated a recently proposed independent prognostic indicator of ovarian cancer, *CXCL12* (9), using 13 published studies, demonstrating for this biomarker that numerous studies are needed to overcome the lack of power in individual studies of smaller sample size. We provide code in the documentation of the *curatedOvarianData* package demonstrating how this comprehensive analysis, which was previously impractical to achieve, is a straightforward application of the database.

Methods and implementation

The pipeline for creating the data package from public databases (Table 1) is fully automated, with the exceptions of manual curation of clinical annotations (Figure 1). This manual curation was integrated in the pipeline with short R scripts that reformat user-provided annotations into a standardized template, which largely follows the format of The Cancer Genome Atlas (29). This template is provided in Table 2 and used as a unit test in the *curatedOvarianData* package, i.e. the curation is automatically checked for valid values in the package building process. Downloading phenotype data and expression data from GEO (1), syntax validation of curated clinical metadata, microarray data preprocessing, normalization, gene mapping and the creation of Bioconductor 'ExpressionSet' objects, which link gene expression data and phenotype annotations, were fully automated. The generation of the package is reproducible using the pipeline provided at <https://bitbucket.org/lwaldron/curatedovariandata>.

Data acquisition and curation

Our search for clinically annotated ovarian cancer microarray studies identified 21 published studies, which provided 23 publicly available data sets from various sources (Table 1). The search not only targeted studies of primary tumours annotated with patient survival but also included studies providing other potentially valuable clinical annotation. Other main factors of interest included drug resistance, outcome of the primary tumour debulking surgery, histology, stage and grade. We excluded studies not measuring gene expression (i.e. studies of genomic copy number), studies of cell lines, animal models, or non-primary tumours, and data sets not providing clinical information. Expression and clinical data were obtained from the two major public repositories GEO (i) and ArrayExpress (ii), otherwise from supplementary data of the original publications. Data from GEO were obtained using the GEOquery package (31). Clinical annotations

Table 1. Data sets in the curatedOvarianData database

Data set	Reference	Platform	Samples	Late Stage ^a (%)	Serous Subtype (%)	Median Survival (Months)	Median Follow-up (Months)	Censoring (%)
E.MTAB.386	(10)	Ill. HumanRef-8 v2	129	99	100	42	55	43
GSE12418	(11)	SWEGENE v2.1.1_27k	54	100	100	N/A	N/A	N/A
GSE12470	(12)	Agilent G4110b	53	66	81	N/A	N/A	N/A
GSE13876	(13)	Operon Human v3	157	100	100	25	72	28
GSE14764	(14)	Affy U133a	80	89	85	54	37	74
GSE17260	(15)	Agilent G4112a	110	100	100	53	47	58
GSE18520	(16)	Affy U133 Plus 2.0	63	84	84	25	140	23
GSE19829.GPL570	(17)	Affy U133 Plus 2.0	28	N/A	N/A	47	62	39
GSE19829.GPL8300	(17)	Affy U95 v2	42	N/A	N/A	45	50	45
GSE20565	(18)	Affy U133 Plus 2.0	140	48	51	N/A	N/A	N/A
GSE2109	N/A	Affy U133 Plus 2.0	204	42	42	N/A	N/A	N/A
GSE26712	(19)	Affy U133a	195	96	95	46	90	30
GSE30009	(20)	TaqMan qRT-PCR 380	103	100	99	41	53	45
GSE30161	(21)	Affy U133 Plus 2.0	58	100	81	50	83	38
GSE32062.GPL6480	(22)	Agilent G4112a	260	100	100	59	56	53
GSE32063	(22)	Agilent G4112a	40	100	100	53	81	45
GSE6008	(23)	Affy U133a	99	54	41	N/A	N/A	N/A
GSE6822	(24)	Affy Hu6800	66	N/A	62	N/A	N/A	N/A
GSE9891	(25)	Affy U133 Plus 2.0	285	85	93	47	36	59
PMID15897565 ^b	(26)	Affy U133a	63	83	100	N/A	N/A	N/A
PMID17290060 ^c	(27)	Affy U133a	117	98	100	63	82	43
PMID19318476	(28)	Affy U133a	42	93	100	34	89	48
TCGA	(29)	Affy HT U133a	578	90	98	45	52	48

These data sets provide curated gene expression and clinical data for a total of 2970 samples, including all publicly ovarian cancer gene expression experiments with individual patient survival information at the time of press.

^aOnly FIGO Stages III and IV.

^bData set is a subset of the samples from the retracted paper PMID17290060, Dressman *et al.* (27).

^cPaper was retracted because of a misalignment of genomic and survival data (30); the corrected data are provided here.

N/A, not available.

were manually curated using one R script per data set, and original uncurated annotations were retained as a single field. Curated annotations were checked by syntax against a template, which standardized all the known clinically relevant indicators and allowable data values. Clinical data were twice independently curated (authors B.G. and T.R.), and all discrepancies were resolved for the final version. The availability of clinical data varied substantially across datasets (Figure 2).

Gene expression processing and gene mapping

Where raw data from Affymetrix U133a or U133 Plus 2.0 platforms were available, these were pre-processed by frozen Robust Multi-array Analysis (fRMA) (32), for other Affymetrix platforms by Robust Multi-array Average (RMA) (33), and otherwise we used pre-processed data as provided by the authors. Up-to-date maps from probe set

IDs to gene symbols were obtained from BioMart (34). Where BioMart maps were not available but target sequences were provided for the microarray platforms, we used the BLAST algorithm (35) to map these sequences against the human genome (build GRCh37) and to identify the gene transcript targeted by each probe. Otherwise, the annotations provided with the platform on GEO were used. In the *curatedOvarianData* version of the package, genes with multiple probe sets were represented by the probe set with the highest mean across all data sets of the sample platform (36), and this original probe set identifier was also stored in the ExpressionSet object (7). We selected the same representative probe set for all studies of a common microarray platform. Finally, we provide two alternative versions of the package: *NormalizerVcuratedOvarianData*, where redundant probe sets are averaged after filtering probe sets with low correlation to their redundant probe sets,

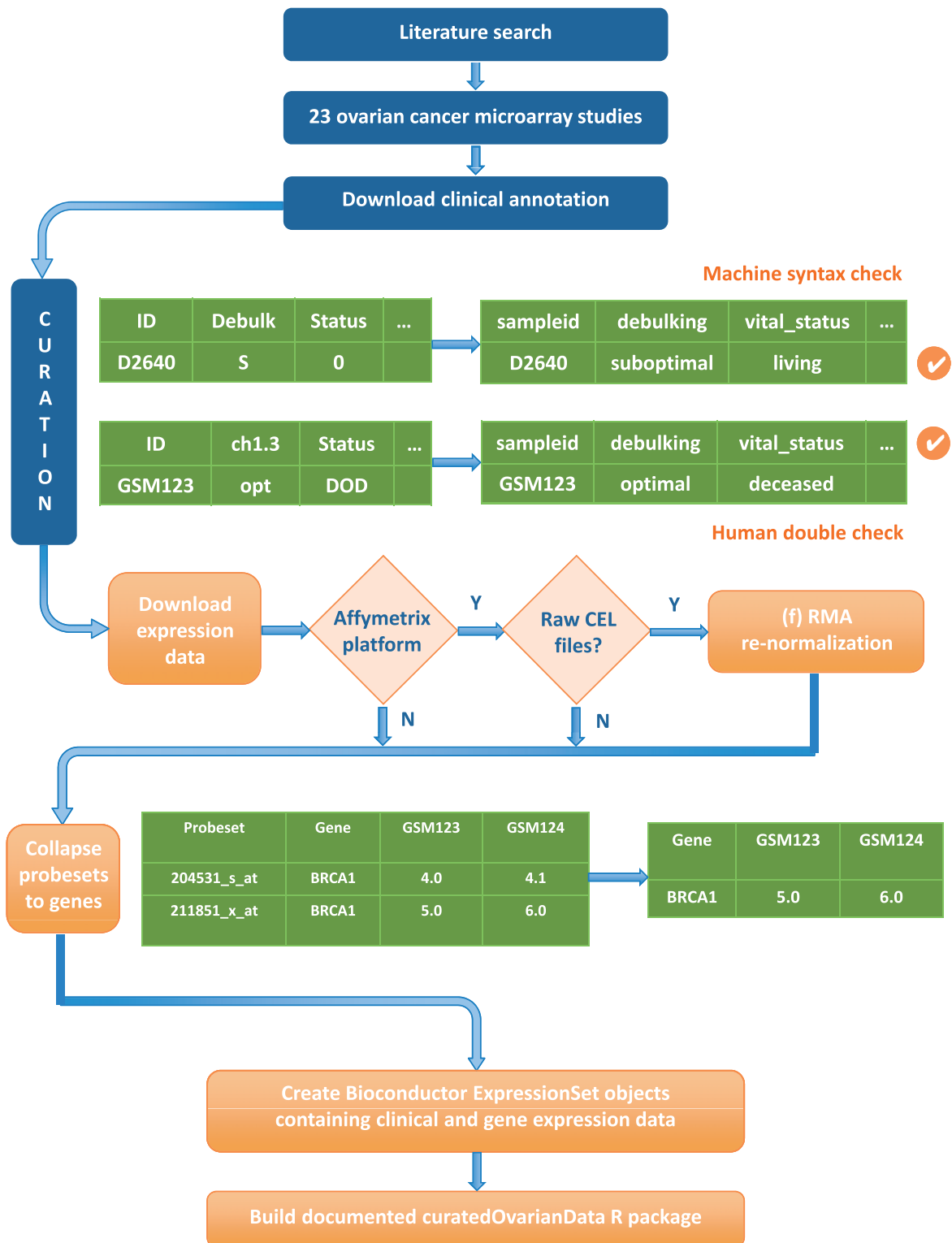


Figure 1. Flowchart of the data collection and curation pipeline. The software implementing this pipeline reproduces all steps from downloading of data to final packaging, requiring manual intervention only for identifying studies, curation of clinical metadata and documentation of the package.

Table 2. Curated clinical annotations

Characteristic	Allowed values	Description
sample_type	tumour, metastatic, cellline, healthy, adjacentnormal	Healthy, only from individuals without cancer; adjacentnormal, from individuals with cancer;
histological_type	ser, endo, clearcell, mucinous, other, mix, undifferentiated	ser, serous; endo, endometrioid; clearcell, mixture of ser + endo. Other includes sarcomatoid, endometrioid, papillary serous, adenocarcinoma, dysgerminoma
primarysite	ov, ft, other	Ov, ovary; ft, fallopian tube
arrayedsite	ov, ft, other	ov, ovary; ft, fallopian tube
summarygrade ^a	low, high	low, 1, 2, LMP (low malignant potential); high, 3, 2/3
summarystage	early, late	early, FIGO I, II, III; late, FIGO III, IV, IIII, IIII/IV
tumourstage	1, 2, 3, 4	FIGO Stage (I–IV, translated to 1–4 for R usage)
substage	a, b, c, d	Substage (abcd)
grade ^a	1, 2, 3	Grade (1–3)
age_at_initial_pathologic_diagnosis	1–99	Age at initial pathologic diagnosis in years
pltx	y/n	Patient treated with Platin
tax	y/n	Patient treated with Taxol
neo	y/n	Neoadjuvant treatment
days_to_tumour_recurrence	decimal	Time to recurrence or last follow-up in days
recurrence_status	recurrence, no recurrence	Recurrence censoring variable
days_to_death	decimal	Time to death or last follow-up in days
vital_status	living, deceased	Overall survival censoring variable
os_binary	short, long	Dichotomized overall survival time; as defined by study
relapse_binary	short, long	Dichotomized relapse variable; as defined by the study
site_of_tumour_first_recurrence	metastasis, locoregional, etc.	Site of the first recurrence
primary_therapy_outcome_success	completeresponse, etc.	Response to any kind of therapy
bulking	optimal, suboptimal	Amount of residual disease (optimal ≤ 1 cm)
percent_normal_cells	0–100+/-	Estimated percentage of normal cells; 20– ≤ 20%
percent_stromal_cells	0–100+/-	Estimated percentage of stromal cells
percent_tumour_cells	0–100+/-	Estimated percentage of tumour cells; 80+ ≥ 80%
batch	character	Hybridization date or other available batch variable
uncurated_author_metadata	character	All original, uncurated metadata

Additional study-specific details are provided in the package manual.

^aMost ovarian cancer pathologists follow the FIGO grading system, although some exceptions (15, 22, 25) are noted in the package manual.



Figure 2. Available clinical annotation. This heatmap visualizes for each curated clinical characteristic (rows) the availability in each data set (columns). Red indicates that the corresponding characteristic is available for at least one sample in the data set. See [Table 2](#) for descriptions of these characteristics.

using the Normalizer function of the Sleipnir library for computational functional genomics (37), and *FULLVcuratedOvarianData*, which does not collapse redundant probe sets targeting the same gene transcript but instead provides a probe set to gene symbol map in the featureData slot of each ExpressionSet.

Final packaging

Technical replicate samples were merged by averaging. Microarray expression data and clinical metadata were then represented as ExpressionSet objects (7) for each study. The ExpressionSet objects were also populated with citations, platform identifiers and details, data preprocessing methods and warnings of retracted papers (27) and specimens also used in other studies (26, 28, 29, 38). ExpressionSets were packaged as the *curatedOvarianData* R library, which provides a reference manual including

descriptions of the syntax template and summaries of the annotations, citation, microarray platform and other information for each study.

Discussion

We introduce a data package for the R/Bioconductor statistical programming environment that includes all current major ovarian cancer gene expression data sets (Table 1). The process of downloading clinically annotated public genomic data and proceeding to a final computational analysis is, despite recent efforts (4, 5), still long and prone to errors. This is particularly true when the various data sets need to be comparable for meta-analyses, which requires a fully standardized annotation. Our data resource provides a comprehensive and highly curated resource for efficient meta-analysis of the ovarian cancer transcriptome, for

biological analysis and bioinformatic methods development. It additionally provides a complete computational pipeline to reproduce this process for other cancers or data sources.

Two common problems of publicly available genomic data are the scarcity of clinical annotation and inconsistent definitions of clinical characteristics across independent data sets (5). In our review of original papers and curation of clinical annotations, we were however able to retain, in most studies, the clinical variables of proven importance: overall survival, age, optimal debulking surgery, tumour histology, grade and stage (Figure 2). Other characteristics such as detailed treatment information or recurrence free survival times were rarely available; however, ovarian cancer has a relatively standard treatment regimen of platinum chemotherapy and no radiotherapy. The most important clinical variables were in general consistently defined between studies, with these definitions provided in Table 2. Notably, all studies used the Federation of Gynecology and Obstetrics (FIGO) staging system, and all but one study (11) defined suboptimal debulking surgery as residual tumour mass > 1 cm (Table 2). The relatively large number of well-annotated data sets in this database may allow interesting future work, addressing the problem of recovering missing annotations from genomic data only (40).

One important use of this database is the assessment of prognostic biomarkers. As a demonstration, we examined a recent study by Popple *et al.* (9), which analysed the expression of the chemokine protein CXCL12 using a tissue microarray of 289 primary ovarian cancers. CXCL12/CXCR4 is a chemokine/chemokine receptor axis that has previously been shown to be directly involved in cancer pathogenesis (41, 42). Ovarian cancer constitutively expresses CXCL12 and CXCR4, and both tumour CXCL12/CXCR4 expression and stroma-derived CXCL12 expression have been reported to be prognostic factors in human ovarian cancer (41). Popple *et al.* found that high levels of CXCL12 protein were associated with significantly poorer survival compared with patients whose tumours produce low amounts of this chemokine, independently of stage, residual disease (optimal debulking) and adjuvant chemotherapy. The patient cohort was heterogeneous, with various histologic types, grades and stages, leaving open the question of whether this biomarker would be generalizable to other patient populations. Furthermore, differences in protein abundance may not be associated with RNA abundance.

To investigate these questions, we analysed CXCL12 expression in all primary tumour samples included in *curatedOvarianData* for which overall survival information was available. To ensure that the expression values were on the same scale across studies, all data sets were centred by their means and scaled by their standard deviations. A population hazard ratio (HR) was then pooled with a fixed-effects model, in which the HR for each cohort was

weighted with the inverse of the standard error. This is visualized as a forest plot in Figure 3. Although the effect is only significant ($P < 0.05$) in three cohorts individually, the pooled HR is significantly larger than 1 (HR = 1.15, 95% CI 1.09–1.23). HR refers to the HR between patients differing by one standard deviation in CXCL12 expression. This confirms the hypothesis that upregulation of CXCL12 is associated with poor outcome in 2108 patients from 13 independent studies with mixed stage, grade and histologies. The effect is thus small but consistently detected, emphasizing the importance of biomarker validation in sufficiently large data collections. To assess the independence of CXCL12 with stage and residual disease, we also analysed the 1776 patients from 10 studies where both FIGO tumour stage and success of debulking surgery were known. Adjustment for these two established predictors in multivariate analysis had little effect on the observed association between CXCL12 and overall survival (HR = 1.13, 95% CI 1.05–1.21). These HRs are comparable in magnitude to that reported by Popple *et al.* for 'moderate' CXCL12 staining (HR = 1.215, 95% CI 0.892–1.655), but lower than reported for 'high' staining (HR = 1.684, 95% CI 1.180–2.404). This potentially reflects that the function of this gene is at the protein level. Consistent with previous reports (9, 38), we found no significant association of the receptor CXCR4 with overall survival (HR = 0.95, 95% CI 0.9–1.01, $P = 0.09$). These analyses are straightforward and fully reproduced as examples in the package documentation. Additional analyses limited to more homogeneous patient subsets, e.g. limited to tumours of the same histology, are needed, but they are another straightforward application of the package.

In constructing *curatedOvarianData*, we took several steps to minimize across-study batch effects. Where raw Affymetrix microarray data were available, we used a standardized pre-processing protocol. All data sets from the same platform were normalized with the same algorithms and parameters. For the Affymetrix U133A and U133 Plus 2.0, we chose the fRMA (32) normalizing algorithm, a variant of the standard RMA (33) algorithm that uses publicly available microarray databases to estimate probe-specific effects and variances, instead of using only the samples from the data set to be normalized. We provide example code in the database documentation for removing between-platform batch effects with the ComBat method (43). Such a batch effect removal is typically necessary when data sets are merged.

If different platforms are compared, then the mapping of probe sets to common identifiers such as gene symbols is a critical and error prone step. In particular when older platforms are considered, care must be taken to ensure that the probe sets target identical transcripts; gene identification is a persistent problem in genome-scale data integration. We used the BioMart database (34) to map

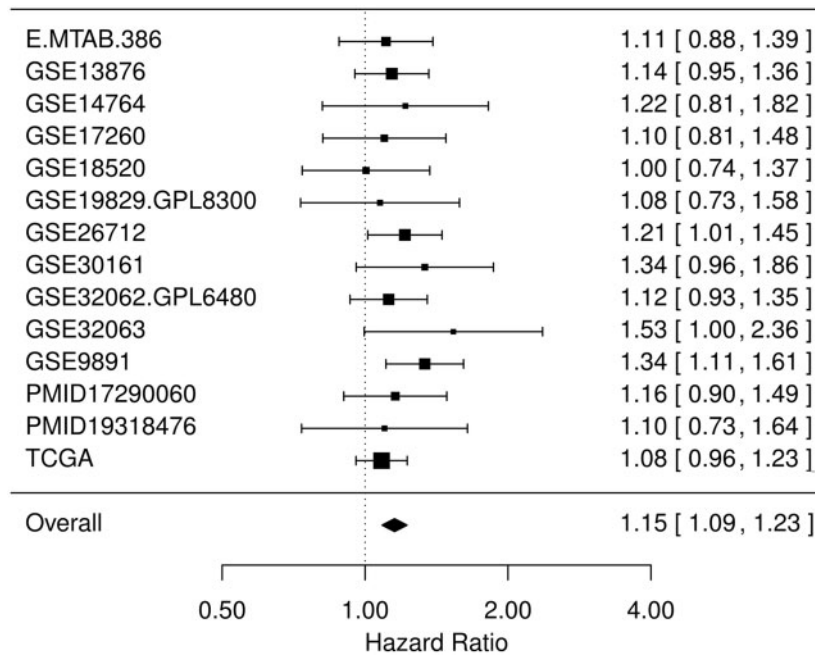


Figure 3. The database confirms *CXCL12* as prognostic of overall survival in patients with ovarian cancer. Forest plot of the expression of the chemokine *CXCL12* as a univariate predictor of overall survival, using all 14 data sets with applicable expression and survival information. HR indicates the factor by which overall risk of death increases with a one standard deviation increase in *CXCL12* expression. A summary HR significantly larger than 1 indicates that patients with high *CXCL12* levels had poor outcome and confirms in several lines of code the previously reported association between *CXCL12* abundance and patient survival (9). Consideration of important clinicopathological features such as stage, grade, histology and residual disease (optimal surgical debulking) is also straightforward; examples are provided in the package vignette.

stable manufacturer probe set identifiers or Genbank IDs to current standard gene symbols. For cases in which no stable identifiers were available, we used the BLAST algorithm (35) to identify gene symbols from the probe oligonucleotide sequences. When many genes are targeted by more than one probe set, several approaches of collapsing probe sets to single genes have been proposed (36, 44, 45). In the main version of the package, we selected the probe set with highest mean across all data sets from the same platform to represent each gene transcript, a method shown to perform well (36) and with the advantage of being traceable back to a single oligonucleotide probe sequence for each platform. We also provide two alternative packages with averaged and un-collapsed probe sets. The version with un-collapsed probe sets provides current HGNC symbols in the featureData slot of the ExpressionSet objects, which makes the application of alternative methods for collapsing probe sets to unique gene symbols straightforward, e.g. with the WGCNA R package (46).

We demonstrated meta-analytical use of the package by showing a survival association of the recently proposed prognostic biomarker *CXCL12* (9). Other possible uses include the validation of multi-gene signatures, and identification of novel gene signatures and biomarkers for patient survival and response to chemotherapy. Finally, this

package enables rigorous assessment of high-dimensional machine-learning algorithms in terms of their performance and computational requirements. We plan to continually include newly published ovarian cancer data sets in future versions of this package.

Conclusions

The *curatedOvarianData* package provides a comprehensive resource of curated gene expression and clinical data for the development and validation of ovarian cancer prognostic models, the investigation of ovarian cancer subtypes (10, 25, 29), and the comparative assessment of machine learning algorithms for gene expression data. This database greatly reduces the burden of time, expertise and error involved in assembling a compendium of curated gene expression data from tumours of known histopathology and from patients with known clinical progression. These advantages will be appealing to biostatisticians and bioinformaticians for development of analytical methods from high-dimensional genomic data, but the database will additionally provide a common, version-controlled and transparent platform for reproducible investigation of the ovarian cancer transcriptome. The pipeline for creating this database is published under an open license and will

facilitate creating similar resources for other cancers. As such, we hope this database and pipeline will provide one part of the solution to reproducibility in high-dimensional genomic research.

Acknowledgements

The authors thank Shaina Andelman for her contributions to graphic design, and also Steve Skates, Jie Ding and Dave Zhao.

Funding

National Cancer Institute at the National Institutes of Health [1RC4CA156551-01 to G.P. and M.B.]; the National Science Foundation [CAREER DBI-1053486 to C.H.]. M.R. acknowledges support from the National Cancer Institute initiative to found Physical Science Oncology Centers [U54CA143798]. Funding for open access charge: National Science Foundation [CAREER DBI-1053486 to C.H.].

Conflict of interest. None declared.

References

- Edgar,R., Domrachev,M. and Lash,A.E. (2002) Gene Expression Omnibus: NCBI gene expression and hybridization array data repository. *Nucleic Acids Res.*, **30**, 207–210.
- Parkinson,H., Sarkans,U., Kolesnikov,N. et al. (2011) ArrayExpress update—an archive of microarray and high-throughput sequencing-based functional genomics experiments. *Nucleic Acids Res.*, **39**, D1002–D1004.
- McDermott,U., Downing,J.R. and Stratton,M.R. (2011) Genomics and the continuum of cancer care. *N. Engl. J. Med.*, **364**, 340–350.
- Taminau,J., Steenhoff,D., Coletta,A. et al. (2011) inSilicoDb: an R/Bioconductor package for accessing human Affymetrix expert-curated datasets from GEO. *Bioinformatics*, **27**, 3204–3205.
- Carey,V.J., Gentry,J., Sarkar,R. et al. (2008) SGDI: system for genomic data integration. *Pac. Symp. Biocomput.*, 141–152.
- Siegel,R., Naishadham,D. and Jemal,A. (2012) Cancer statistics, 2012. *CA Cancer J. Clin.*, **62**, 10–29.
- Gentleman,R.C., Carey,V.J., Bates,D.M. et al. (2004) Bioconductor: open software development for computational biology and bioinformatics. *Genome Biol.*, **5**, R80.
- Seal,R.L., Gordon,S.M., Lush,M.J. et al. (2011) genenames.org: the HGNC resources in 2011. *Nucleic Acids Res.*, **39**, D514–D519.
- Popple,A., Durrant,L.G., Spendlove,I. et al. (2012) The chemokine, CXCL12, is an independent predictor of poor survival in ovarian cancer. *Br. J. Cancer*, **106**, 1306–1313.
- Bentink,S., Haibe-Kains,B., Risch,T. et al. (2012) Angiogenic mRNA and microRNA gene expression signature predicts a novel subtype of serous ovarian cancer. *PLoS One*, **7**, e30269.
- Partheen,K., Levan,K., Osterberg,L. et al. (2006) Expression analysis of stage III serous ovarian adenocarcinoma distinguishes a sub-group of survivors. *Eur. J. Cancer*, **42**, 2846–2854.
- Yoshida,S., Furukawa,N., Haruta,S. et al. (2009) Expression profiles of genes involved in poor prognosis of epithelial ovarian carcinoma: a review. *Int. J. Gynecol. Cancer*, **19**, 992–997.
- Crijns,A., Fehrmann,R., de Jong,S. et al. (2009) Survival-related profile, pathways, and transcription factors in ovarian cancer. *PLoS Med.*, **6**, e24.
- Denkert,C., Budczies,J., Darb-Esfahani,S. et al. (2009) A prognostic gene expression index in ovarian cancer - validation across different independent data sets. *J. Pathol.*, **218**, 273–280.
- Yoshihara,K., Tajima,A., Yahata,T. et al. (2010) Gene expression profile for predicting survival in advanced-stage serous ovarian cancer across two independent datasets. *PLoS One*, **5**, e9615.
- Mok,S.C., Bonome,T., Vathipadiekal,V. et al. (2009) A gene signature predictive for outcome in advanced ovarian cancer identifies a survival factor: microfibril-associated Glycoprotein 2. *Cancer Cell*, **16**, 521–532.
- Konstantinopoulos,P.A., Spentzos,D., Karlan,B.Y. et al. (2010) Gene expression profile of BRCAness that correlates with responsiveness to chemotherapy and with outcome in patients with epithelial ovarian cancer. *J. Clin. Oncol.*, **28**, 3555–3561.
- Meyniel,J.P., Cottu,P.H., Decraene,C. et al. (2010) A genomic and transcriptomic approach for a differential diagnosis between primary and secondary ovarian carcinomas in patients with a previous history of breast cancer. *BMC Cancer*, **10**, 222.
- Bonome,T., Levine,D., Shih,J. et al. (2008) A gene signature predicting for survival in suboptimally debulked patients with ovarian cancer. *Cancer Res.*, **68**, 5478–5486.
- Gillet,J.P., Calcagno,A., Varma,S. et al. (2012) Multidrug resistance-linked gene signature predicts overall survival of patients with primary ovarian serous carcinoma. *Clin. Cancer Res.*, **18**, 3197–3206.
- Ferriss,J.S., Kim,Y., Duska,L. et al. (2012) Multi-gene expression predictors of single drug responses to adjuvant chemotherapy in ovarian carcinoma: predicting platinum resistance. *PLoS One*, **7**, e30550.
- Yoshihara,K., Tsunoda,T., Shigemizu,D. et al. (2012) High-risk ovarian cancer based on 126-gene expression signature is uniquely characterized by downregulation of antigen presentation pathway. *Clin. Cancer Res.*, **18**, 1374–1385.
- Murph,M., Liu,W., Yu,S. et al. (2009) Lysophosphatidic acid-induced transcriptional profile represents serous epithelial ovarian carcinoma and worsened prognosis. *PLoS One*, **4**, e5583.
- Ouellet,V., Provencher,D.M., Maugard,C.M. et al. (2005) Discrimination between serous low malignant potential and invasive epithelial ovarian tumors using molecular profiling. *Oncogene*, **24**, 4672–4687.
- Tohill,R.W., Tinker,A.V., George,J. et al. (2008) Novel molecular subtypes of serous and endometrioid ovarian cancer linked to clinical outcome. *Clin. Cancer Res.*, **14**, 5198–5208.
- Berchuck,A., Iversen,E.S., Lancaster,J.M. et al. (2005) Patterns of gene expression that characterize long-term survival in advanced stage serous ovarian cancers. *Clin. Cancer Res.*, **11**, 3686–3696.
- Dressman,H., Berchuck,A., Chan,G. et al. (2007) An integrated genomic-based approach to individualized treatment of patients with advanced-stage ovarian cancer. *J. Clin. Oncol.*, **25**, 517–525.
- Berchuck,A., Iversen,E.S., Luo,J. et al. (2009) Microarray analysis of early stage serous ovarian cancers shows profiles predictive of favorable outcome. *Clin. Cancer Res.*, **15**, 2448–2455.
- Cancer Genome Atlas Research Network. (2011) Integrated genomic analyses of ovarian carcinoma. *Nature*, **474**, 609–615.
- Dressman,H.K., Berchuck,A., Chan,G. et al. (2012) Retraction. An integrated genomic-based approach to individualized treatment of patients with advanced-stage ovarian cancer. *J. Clin. Oncol.*, **30**, 678.

31. Sean,D. and Meltzer,P.S. (2007) GEOquery: a bridge between the Gene Expression Omnibus (GEO) and BioConductor. *Bioinformatics*, **23**, 1846–1847.
32. McCall,M.N., Bolstad,B.M. and Irizarry,R.A. (2010) Frozen robust multiarray analysis (fRMA). *Biostatistics*, **11**, 242–253.
33. Bolstad,B.M., Irizarry,R.A., Astrand,M. et al. (2003) A comparison of normalization methods for high density oligonucleotide array data based on variance and bias. *Bioinformatics*, **19**, 185–193.
34. Durinck,S., Moreau,Y., Kasprzyk,A. et al. (2005) BioMart and Bioconductor: a powerful link between biological databases and microarray data analysis. *Bioinformatics*, **21**, 3439–3440.
35. Altschul,S.F., Madden,T.L., Schaffer,A.A. et al. (1997) Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. *Nucleic Acids Res.*, **25**, 3389–3402.
36. Miller,J.A., Cai,C., Langfelder,P. et al. (2011) Strategies for aggregating gene expression data: the collapseRows R function. *BMC Bioinformatics*, **12**, 322.
37. Huttenhower,C., Schroeder,M., Chikina,M.D. et al. (2008) The sleipnir library for computational functional genomics. *Bioinformatics*, **24**, 1559–1561.
38. Bild,A.H., Yao,G., Chang,J.T. et al. (2006) Oncogenic pathway signatures in human cancers as a guide to targeted therapies. *Nature*, **439**, 353–357.
39. Kauffmann,A., Rayner,T.F., Parkinson,H. et al. (2009) Importing Array-Express datasets into R/Bioconductor. *Bioinformatics*, **25**, 2092–2094.
40. Shah,N.H., Jonquet,C., Chiang,A.P. et al. (2009) Ontology-driven indexing of public datasets for translational bioinformatics. *BMC Bioinformatics*, **10**(Suppl. 2), S1.
41. Kajiyama,H., Shibata,K., Terauchi,M. et al. (2008) Involvement of SDF-1alpha/CXCR4 axis in the enhanced peritoneal metastasis of epithelial ovarian carcinoma. *Int. J. Cancer*, **122**, 91–99.
42. Kulbe,H., Chakravarty,P., Leinster,D.A. et al. (2012) A dynamic inflammatory cytokine network in the human ovarian cancer microenvironment. *Cancer Res.*, **72**, 66–75.
43. Johnson,W.E., Li,C. and Rabinovic,A. (2007) Adjusting batch effects in microarray expression data using empirical Bayes methods. *Biostatistics*, **8**, 118–127.
44. Dai,M., Wang,P., Boyd,A.D. et al. (2005) Evolving gene/transcript definitions significantly alter the interpretation of GeneChip data. *Nucleic Acids Res.*, **33**, e175.
45. Li,Q., Birkbak,N.J., Gyorffy,B. et al. (2011) Jetset: selecting the optimal microarray probe set to represent a gene. *BMC Bioinformatics*, **12**, 474.
46. Langfelder,P. and Horvath,S. (2008) WGCNA: an R package for weighted correlation network analysis. *BMC Bioinformatics*, **9**, 559.