RESEARCH ARTICLE

Predicting ionizing radiation exposure using biochemically-inspired genomic machine learning [version 1; peer review: 3 approved]

Jonathan Z.L. Zhao¹,², Eliseos J. Mucaki¹, Peter K. Rogan¹-⁵

¹Department of Biochemistry, Schulich School of Medicine and Dentistry, Western University, London, ON, N6A 2C1, Canada
²Department of Computer Science, Faculty of Science, Western University, London, ON, N6A 2C1, Canada
³Department of Epidemiology & Biostatistics, Schulich School of Medicine and Dentistry, Western University, London, ON, N6A 2C1, Canada
⁴CytoGnomix Inc., London, ON, N5X 3X5, Canada
⁵Department of Oncology, Schulich School of Medicine and Dentistry, Western University, London, ON, N6A 2C1, Canada

Abstract

Background: Gene signatures derived from transcriptomic data using machine learning methods have shown promise for biodosimetry testing. These signatures may not be sufficiently robust for large scale testing, as their performance has not been adequately validated on external, independent datasets. The present study develops human and murine signatures with biochemically-inspired machine learning that are strictly validated using k-fold and traditional approaches.

Methods: Gene Expression Omnibus (GEO) datasets of exposed human and murine lymphocytes were preprocessed via nearest neighbor imputation and expression of genes implicated in the literature to be responsive to radiation exposure (n=998) were then ranked by Minimum Redundancy Maximum Relevance (mRMR). Optimal signatures were derived by backward, complete, and forward sequential feature selection using Support Vector Machines (SVM), and validated using k-fold or traditional validation on independent datasets.

Results: The best human signatures we derived exhibit k-fold validation accuracies of up to 98% (DDB2, PRKDC, TPP2, PTPRE, and GADD45A) when validated over 209 samples and traditional validation accuracies of up to 92% (DDB2, CD8A, TALDO1, PCNA, EIF4G2, LCN2, CDKN1A, PRKCH, ENO1, and PPM1D) when validated over 85 samples. Some human signatures are specific enough to differentiate between chemotherapy and radiotherapy. Certain multi-class murine signatures have sufficient granularity in dose estimation to inform eligibility for cytokine therapy (assuming these signatures could be translated to humans). We compiled a list of the most frequently appearing genes in the top 20 human and mouse signatures. More frequently appearing genes among an ensemble of signatures may indicate greater impact of these genes on the performance of individual signatures. Several genes in the signatures we derived are present in previously proposed signatures.

Open Peer Review

Reviewer Status  
Invited Reviewers

1 2 3


1 Daniel Oh, University of North Carolina at Chapel Hill, Chapel Hill, USA
2 Michael D. Story, University of Texas Southwestern Medical Center, Dallas, USA
3 Liang-hao Ding, University of Texas Southwestern Medical Center, Dallas, USA
4 Roel Quintens, Belgian Nuclear Research Centre, Mol, Belgium
5 Mohamed Mysara, Belgian Nuclear Research Centre, Mol, Belgium

Any reports and responses or comments on the article can be found at the end of the article.
Conclusions: Gene signatures for ionizing radiation exposure derived by machine learning have low error rates in externally validated, independent datasets, and exhibit high specificity and granularity for dose estimation.

Keywords
Ionizing Radiation Exposure, Machine Learning, Gene Signatures, Molecular Diagnostics, Validation, Biodosimetry, Support Vector Machine, Minimum Redundancy Maximum Relevance

Corresponding author: Peter K. Rogan (progan@uwo.ca)

Author roles: Zhao JZL: Conceptualization, Data Curation, Formal Analysis, Investigation, Methodology, Software, Validation, Writing – Original Draft Preparation, Writing – Review & Editing; Mucaki EJ: Conceptualization, Investigation, Methodology, Software, Supervision; Rogan PK: Conceptualization, Formal Analysis, Funding Acquisition, Investigation, Methodology, Project Administration, Supervision, Writing – Original Draft Preparation, Writing – Review & Editing


Grant information: Natural Sciences and Engineering Research Council of Canada (NSERC Discovery Grant RGPIN-2015-06290); the Canadian Foundation for Innovation; Canada Research Chairs, and CytoGnomix Inc. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Copyright: © 2018 Zhao JZL et al. This is an open access article distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

How to cite this article: Zhao JZL, Mucaki EJ and Rogan PK. Predicting ionizing radiation exposure using biochemically-inspired genomic machine learning [version 1; peer review: 3 approved] F1000Research 2018, 7:233 (https://doi.org/10.12688/f1000research.14048.1)

**Introduction**

Potential radiation exposures from industrial nuclear accidents, military incidents, or terrorism are threats to public health\(^1\). There is a need for large scale biodosimetry testing, which requires efficient screening techniques to differentiate exposed individuals from non-exposed individuals and to determine the severity of exposure\(^2\). Current diagnostic techniques, including the cytogenetic gold standard\(^3-6\), may require several days to provide accurate dose estimates\(^7-9\) of large cohorts. To address the need for faster diagnostic techniques that accurately measure radiation exposures, gene signatures based on transcriptomic data have been introduced\(^10\). Probit regression models of radiation response using 25 probes on peripheral blood samples achieved up to 90% accuracy for distinguishing between irradiated blood samples and unirradiated controls\(^7\). A 74-gene classifier based on nearest centroid expression levels was 98% accurate in distinguishing four levels of irradiation from controls\(^9\). This level of performance implies that samples exposed to different levels of radiation may be distinguishable based on mRNA expression levels of different genes. While this suggests the feasibility of transcriptional modeling of radiation responses, validation with external datasets is required to establish its reliability for rapid diagnostics. A caveat of these signatures is that they have not all been externally validated on datasets independent of the source data used for model development. A 29-gene signature modelled using a support vector machine (SVM) was externally validated on such a dataset, resulting in 80% accuracy in distinguishing higher (≥8Gy) from lower dose (<2Gy) radiation exposure in novel samples\(^7\). The present study derives signatures with improved performance on externally validated samples by employing a different selection of modelling techniques. The machine learning pipeline used here addresses some of the previous limitations through a more rigorous feature selection process and stricter validation procedures.

Previously, the Student’s t-test\(^7\), the F-test\(^10\), and correlation coefficients\(^7\) were used to identify potential radiation biomarker genes. Although statistical criteria can distinguish genes that are differentially expressed upon radiation exposure, they do not eliminate expressed genes with redundant responses to radiation exposure. Redundancy increases the possibility of overfitting, thereby reducing the generalizability of these models to predict responses in independent datasets. We address this limitation with the information theory-based criterion for gene selection known as minimum redundancy maximum relevance (mRMR)\(^11-13\), which ranks genes according to shared mutual information between expression levels and radiation dose (relevance), and by minimizing mutual information shared by expression values of these and other genes (redundancy)\(^11,12\). mRMR outperforms ranking criteria based solely on maximizing relevance\(^12\). In contrast with heuristic approaches like differential expression, we only consider genes with evidence of a relationship to radiation response, which significantly limits the number of model features. Biochemically-inspired genomic machine learning (ML) has been used to derive high performing gene signatures that predict chemotherapy and hormone therapy responses\(^13-15\). From an initial set of mRMR-derived biochemically relevant genes, wrapper approaches for feature selection\(^16\) are used to find an optimal set of genes that predict exposure to radiation.

It can be challenging to obtain highly accurate models that perform well on externally validated samples for several reasons. Aside from biases in training data, batch effects and lack of reproducibility may introduce systematic and random sources of variability into gene expression microarray data. Different source datasets can impact data normalization, reducing model performance. We utilize two validation procedures. The first is a signature-centric approach that mirrors external k-fold validation\(^7\). The limitation of signature-centric validation is that, while signatures allow for the identification of important genes associated with radiation response, a tangible model is required to generate actual diagnostic predictions. To address this limitation, we also use a second model-centric approach, which we term “traditional validation”. This procedure applies quantile normalization to training and test data before a model is fitted to the training data. This quantile method has been shown to be more effective than scaling, loess, contrast, and non-linear methods in reducing variation between microarray data\(^17\). Model validation was not expected to perform as well as signature validation, because quantile normalization is not always successful in eliminating variation between microarray datasets, whereas k-fold validation is independent of this source of variation. This study shows that robust model validation is a critical step in reproducibly predicting which individuals have been exposed to significant levels of radiation.

**Methods**

**Datasets**

Murine gene expression datasets\(^18\) were obtained from peripheral blood (PB) mononuclear cell samples of ten-week old C57B16 mice that either received total body radiation at 50 cGy, 200 cGy, or 1000 cGy or were not exposed. Post-exposure, total RNA was isolated after 6 hours and expression was determined by microarray analysis using Operon Mouse V3.0.1 (Gene Expression Omnibus (GEO): GPL4783 from GSE10640[GPL4783])\(^19\) and Operon Mouse V4.0 arrays (GEO: GPL6524 from GSE10640[GPL6524])\(^19\). Similar analyses were performed with human expression microarrays\(^14\), including datasets GEO: GSE6874[GPL4782], GSE10640/GPL6522\(^19\), GSE1725\(^20\), and GSE701\(^13\). GSE6874 and GSE10640 consist of PB samples collected 6 hours post-exposure from healthy donors and patients undergoing total body irradiation at 150–200 cGy analyzed with Operon Human V3.0.2 (GEO: GPL4782) and Operon Human V4.0 (GEO: GPL6522) microarrays. GSE10640/GPL6522 consists of 32 patients treated with alkylator-based chemotherapy without radiation. GSE1725 contains lymphoblastoid cell line samples derived from 57 subjects treated with 500 cGy. RNA was extracted 4 hours after exposure. Expression was measured using Affymetrix Human Genome U95 Version 2 Array (GEO: GPL8300). GSE701 contains lymphoblastoid cell lines from Fondation Jean Dausset-CEPH which were irradiated at 300 cGy or 1000 cGy and extracted 1–24 hours after exposure. Expression was measured using the Affymetrix Human Genome U95A Array (GEO: GPL91).

**Preprocessing (Figure 1, panel i)**

Rows and columns of microarray data that are less than 95% complete were removed and any remaining missing values were imputed using the nearest-neighbor algorithm. Only genes that
are common across all datasets have been retained. Expression values of each probe were transformed to z-scores and the mean expression value of probes for the same gene have been assigned as the expression of each gene.

Biochemically-inspired gene selection\textsuperscript{13–15} (Figure 1, panel ii) A literature search has been conducted to identify genes implicated in radiation response using the search queries “radiation genes,” “radiation response genes,” and “radiation signatures” on PubMed. Cited genes comprise those differentially expressed after radiation exposure, genes present in DNA repair databases and other radiation signatures, and evolutionarily conserved genes that were highly expressed in radio-resistant species. A list of 998 genes was compiled\textsuperscript{22–35, Supplementary Table X}) for deriving signatures.

Minimum Redundancy Maximum Relevance (mRMR) gene ranking\textsuperscript{11,12} (Figure 1, panel iii) Rank is assigned by incremental selection of genes based on the mutual information difference (MID) criterion\textsuperscript{11,12}. Highly ranked genes have expression information that shares mutual information with radiation exposure and shares little information with expression of other genes. The MID criterion used to select the next ranked gene is \( \max_{i \in \Omega} \sum_{h \in S} I(i,h) - \sum_{j \in S} I(i,j) \), where \( i \) is a gene selected from \( \Omega \), the total gene space, \( S \) is the set of genes selected before \( i \), \( |S| \) is the number of genes selected before \( i \), \( I(i,h) \) is the mutual information between expression of gene \( i \) and radiation dose \( h \), and \( I(i,j) \) is the mutual information between expression of gene \( i \) and expression of gene \( j \).

Support Vector Machine (SVM) Learning SVM models are classifiers that use hyperplane boundaries to separate samples into exposure classes by maximizing the distance between the separating hyperplanes and samples of each class. The \texttt{fitcecoc} function of MATLAB 2017a’s Statistics and Machine Learning Toolbox\textsuperscript{36} with a SVM template was used to fit SVM models to training data. The \texttt{fitcecoc} function was used because it allows the fitting of multiclass models, which was required for analysis of murine samples that were irradiated at four different exposure levels. The SVM models use the Gaussian radial basis function kernel and a range of selected
box-constraint and kernel-scale parameters. The box-constraint, denoted by the variable $C$, determines how severely misclassifications are penalized during training. The kernel-scale, denoted $\sigma$, represents the width of the Gaussian radial basis function. These parameters collectively control the tradeoff between underfitting and overfitting\(^{37}\). After feature selection, a grid search is performed to determine the optimal ($C, \sigma$) combination for values of $C$ and $\sigma$ between 1 and 100000 (inclusive) by powers of 10 such that $C \geq \sigma$.

**Feature selection (FS) (Figure 1, panel iv)\(^ {16}\)**

Greedy feature selection was used to derive signatures. Complete sequential feature selection (CSFS) sequentially adds genes to an initially empty base set. The added gene is the highest mRMR-ranked gene that is not already included. This is repeated until all genes have been evaluated and the best performing subset of genes is identified. Forward sequential feature selection (FSFS) sequentially adds genes from the top 50 mRMR ranked genes to an initially empty base set. The added gene is the one whose addition improves the model by the greatest margin. Backward sequential feature selection (BSFS) sequentially removes genes from the top 30 mRMR ranked genes. The gene removed is the one whose removal causes the greatest improvement in the model. For BSFS and FSFS, we measure model improvement using misclassification or log loss during k-fold validation (see Performance metrics section below). Genes are added or removed until model performance plateaus. During feature selection, $C$ and $\sigma$ parameters need to be chosen for SVM learning (see SVM Learning section above). Thus, each signature is characterized by the feature selection algorithm used, the dataset used to derive it, and the $C-\sigma$ combination used for its SVM models during feature selection. This leads to a large number of possible signatures (see Supplementary Files Y1–Y7). Supplementary Files Y1–Y3 and Supplementary Files Y6–Y7 contain k-fold validation results from which the top 20 signatures (evaluated using average validation log loss), in particular, were analyzed (Figure 2, Figure 3, Figure 6, Figure 7).

**Figure 2.** Heat map depicting the gene compositions of the top 20 murine signatures derived from different datasets: GSE6874[GPL4783] and GSE10640[GPL6524]. Gene frequency values are first scaled within datasets and then scaled across datasets to ensure values between 0 and 1.
Validating signatures (Figure 1, panel v)
Stratified k-fold validation was used to validate signatures. Samples of the validation dataset were partitioned into k sets, comprised of an approximately equal distribution of radiation levels. For validation, each set was used to test a model trained on the remaining sets, resulting in predictions for all samples in the dataset. Advantages of this approach are that variation between datasets is not pertinent and that signatures can be validated on differently labeled datasets (with samples irradiated at different levels).

Validating models (Figure 1, panel vi)
Model validation requires separate training and test datasets (the training set is often used for FS). Genes from the signature are extracted from the training and test sets and their expression values are quantile normalized by sample. An important distinction between our approach and a previous study is that quantile normalization is applied immediately before validation, so expression of only the genes present in the signature being validated have been normalized. By contrast, previous approaches perform quantile normalization over entire datasets; while this reduces variability in expression values within datasets, it also suppresses the dynamic range, with potential consequential effects on the prognostic value of expression data. After normalization, an SVM model was fit to training datasets and used to generate predictions from the test dataset.

Performance metrics (Figure 1, panel vii)
Performance was determined by comparing predicted radiation doses with actual radiation exposures of each sample. Metrics included misclassification error rate, goodness-of-fit, and multi-class log loss. Misclassification is the percentage of samples that were incorrectly classified, goodness-of-fit is the average absolute value difference between predicted radiation exposure and actual radiation exposure, and multi-class log loss is

$$\text{log loss} = \frac{1}{N} \sum_{i=1}^{N} \sum_{j=1}^{M} y_{ij} \log p_{ij},$$

where $N$ is the number of samples, $M$ is the number of class labels, $p_{ij}$ is the predicted probability that observation $i$ is in class $j$, and $y_{ij}$ is an indicator variable equal to 1 if sample $i$ is in class $j$ and 0 otherwise.

Results
We discovered radiation gene signatures using the microarray data of human and mouse peripheral blood samples and human lymphoblastoid cell lines, which were validated either according to signature (Figure 1, panel v) or with the respective model (Figure 1, panel vi). The murine data were obtained from a wider range of radiation exposure levels (0 cGy, 50 cGy, 200 cGy, 1000 cGy) than the human whole body radiation datasets, which were binary comparisons of radiation effects (0 cGy vs. 150-200 cGy, 0 cGy vs. 500 cGy, or 300 cGy vs. 700 cGy). This made possible the discovery of murine gene signatures with finer granularity for discriminating individuals exposed to different

Figure 3. Scatter plot depicting the mutual information each gene’s expression shares with radiation exposure (averaged over GSE6874[GPL4783] and GSE10640[GPL6524]). The size of each circle is proportional to the frequency at which the gene appears in the top 20 murine signatures ranked by log loss averaged over GSE6874[GPL4783] and GSE10640[GPL6524]. The genes presented match those of the Figure 2.
exposure levels, which is not currently feasible with the human samples.

Murine gene signatures

Table 1 displays the murine signatures derived using our pipeline which had the best performance metrics during k-fold validation on an independent dataset. In addition to the signature information, we report the feature selection algorithm (FS Algorithm) used to discover the signature, the internal validation performance metrics (FS Misclassification fraction and FS Log Loss function). Validation performance metrics on external dataset(s) are indicated by the Validation Misclassification fraction, Validation Log Loss function, and Validation goodness of fit (or GoF). In the FS Misclass. and FS Log Loss columns, one value is always N/A because signatures are derived by optimizing either misclassification or log loss, but never both. The remaining murine signatures are presented in Supplementary Files Y6 and Supplementary Files Y7.

A list of the most consistently appearing genes in the best performing signatures were obtained by pooling the top 20 murine signatures (assessed by validation log loss) from GSE6874[GPL4783] and GSE10640[GPL6524], and respectively collating the top 17 and 19 most frequent genes. The union of these two sets comprises 33 genes displayed within a heat map based on the frequencies of each gene (Figure 2). Surprisingly, the compositions of signatures derived from both datasets are not as similar as one may expect. The genes that appear more frequently in signatures derived from one dataset infrequently appear in the other even though both datasets consisted of the same types of samples irradiated at the same exposure levels.

The shared mutual information of these expressed genes with radiation dose (Figure 3) indicates whether only high mutual information genes appear in the best signatures or whether some lower mutual information genes may also be selected by our feature selection algorithms. The frequency of each gene among these signatures (represented by diameter of the circle) correlates with the mutual information between expression and radiation dose ($\rho = 0.8016$). However, it would be an oversimplification to create signatures based solely upon mutual information, since some genes in lower performing signatures exhibit higher mutual information content. Development of accurate signatures requires more than a collection of gene features whose individual expression values share information with radiation dose, since many of these genes may reveal similar information, and redundant machine learning model features. For instance, Bax and Blink are both common among the best murine signatures, even though Blink shares much more mutual information with radiation dose than Bax expression. Since Blink and Bax are involved in completely different pathways – Bax is an inducer of apoptosis whereas Blink is involved in a B-cell antigen receptor signaling pathway required for optimal B-cell development – they provide different types of information to the overall model. Conversely, we also observe that genes with high information content, such as Ms4a1, may appear less frequently than genes with lower information content, such as Glipr2.

Although mRMR prioritizes genes with non-redundant, complementary contributions, subsequent wrapper steps of forward and backward sequential feature selection occur independently of the mRMR ranking. mRMR reduces the list of features considered by these algorithms, but it is possible for only high mutual information genes to be selected for the final signature. Thus, the inclusion of lower mutual information genes, such as Ube2vl and Urod, reinforces the effectiveness of the mRMR method.

The cellular roles of these protein products (Figure 2 and Figure 3) demonstrate a variety of pathways and functions (Figure 4), some of which have previously discussed. These include DNA repair genes (Pold2 and Pold1), inducers of apoptosis (E2a4, Bax, and Phlda3), chaperonins (Cct3 and

---

**Table 1. Best murine signatures assessed by K-Fold validation.**

<table>
<thead>
<tr>
<th>Signature (C, $\sigma$)</th>
<th>FS(^1) Algo.</th>
<th>FS(^1) Misclass.</th>
<th>FS(^1) Log Loss</th>
<th>Validation Misclass.</th>
<th>Validation Log Loss</th>
<th>Validation GoF(^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>a) Derived from GSE6874[GPL4783] and 5-fold Validated on GSE10640[GPL6524] (n = 75)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phlda3 Blink Bax Cdkn1a Cct3 Pold1 Cd79b E124 Elf2ak4 Ccn1 Glipr2 Hexb Pou2af1 Swap70 Apex1 Ptpn1 Mdm2 Tpst1 Ly6e Sdcbp (10, 10)</td>
<td>BSFS</td>
<td>N/A</td>
<td>0.08</td>
<td>0.08 ± 0.00</td>
<td>0.29 ± 0.02</td>
<td>15 ± 0</td>
</tr>
<tr>
<td>Phlda3 Blink Bax Cdkn1a Cct3 Tlam Pold1 Cd72 Cd79b E124 Galt Elf2ak4 Ms4a1 Ccn1 Glipr2 Gga2 Sh3bp5 Hexb Gcdh Pou2af1 Swap70 Apex1 Ptpn1 Mdm2 Tpst1 Ly6e Sdcbp Lcn2 Suclg2 (100000, 100)</td>
<td>BSFS</td>
<td>0.04</td>
<td>N/A</td>
<td>0.10 ± 0.00</td>
<td>0.23 ± 0.01</td>
<td>26 ± 1</td>
</tr>
<tr>
<td>Cdkn1a Blink Phlda3 Sdcbp Ccn1 (1000, 100)</td>
<td>FSFS</td>
<td>N/A</td>
<td>0.13</td>
<td>0.17 ± 0.00</td>
<td>0.49 ± 0.01</td>
<td>12 ± 0</td>
</tr>
<tr>
<td><strong>b) Derived from GSE10640[GPL6524] and 6-fold Validated on GSE6874[GPL4783] (n = 103)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blink Ccn1 Tpst1 Pold4 Elf2ak4 Atp5i (100000, 100)</td>
<td>FSFS</td>
<td>N/A</td>
<td>0.12</td>
<td>0.11 ± 0.00</td>
<td>0.35 ± 0.01</td>
<td>25 ± 0</td>
</tr>
<tr>
<td>Blink Polk Sod3 Ube2v1 Elf2ak4 (100000, 100)</td>
<td>FSFS</td>
<td>N/A</td>
<td>0.22</td>
<td>0.20 ± 0.00</td>
<td>0.64 ± 0.01</td>
<td>18 ± 0</td>
</tr>
</tbody>
</table>

\(^1\)FS: Feature Selection. \(^2\)GoF: Goodness of Fit.
Cct7, cell cycle regulators (Ccng1 and Cdkn1a), B-cell development genes (Cd79b and Blnk), B-cell antigens (Cd72 and Ms4a1), and a stress-response kinase that inhibits protein synthesis globally (Eif2ak4).

One of the best murine signatures derived from GSE10640 [GPL4783]: Phlda3, Blnk, Bax, Cdkn1a, Cct3, Pold1, Cd79b, Ei24, Eif2ak4, Ccng1, Glipr2, Hexb, Pou2af1, Swap70, Apex1, Pigm1, Mdm2, Tpx1, Ly6e, Sdcbp consistently achieved <10% misclassification error with SVM parameters $C = 10$, $\sigma = 10$. However, for samples that are incorrectly classified according to this signature, the misclassification percentage does not reveal the actual deviation from the correct dose. The confusion matrix visualizes the prediction accuracy of this signature on GEO: GSE10640 [GPL6524] (Figure 5). Indeed, the performance of the matrix shows that the predicted errors for a small fraction of samples deviate from the actual exposures by no more than a single adjacent exposure level. Although the predictions presented in the confusion matrix come from a single iteration of k-fold validation, the standard error associated with misclassification for this signature is extremely low (0.0013) so this confusion matrix is representative of nearly all possible iterations of k-fold validation.

### Human gene signatures

The best performing signatures obtained from each human dataset, assessed by k-fold validation, are presented in Table 2. Although four human radiation datasets were available, GSE701 contained only 10 samples, which was insufficient for derivation of a unique gene signature. While k-fold validation removes the requirement for inter-dataset normalization, it assesses the ability of signatures (genes) to predict radiation exposure without tying the signatures to corresponding models. Each signature is characterized by the feature selection algorithm and its validation statistics, which have been averaged over the 3 independent datasets that were excluded from the original data used to derive the signature. Since traditional validation typically requires separate training and test sets that feature samples irradiated at the same exposure levels, only signatures derived from GEO: GSE6874 [GPL4782] and GEO: GSE10640 [GPL6522] could be analyzed. Table 3 presents the best human signatures according to this validation approach.
Figure 5. Confusion matrix for murine signature: Phlda3, Bax, Cdkn1a, Cct3, Tlam, Poid1, Cd72, Cd79b, El24, Galt, Elf2ak4, Ms4a1, Ccng1, Glipr2, Gga2, Sh3bp5, Hexb, Gcdh, Pou2af1, Swap70, Apex1, Ptpn1, Mdm2, Tpst1, Ly6e, Sdcbp, Lcn2, Sucig2. Numerators represent the number of samples in each category while denominators represent the total number of samples that were irradiated at a given exposure level (i.e. is the sum of the number of samples in each row).

Table 2. Best human signatures assessed by K-Fold validation.

<table>
<thead>
<tr>
<th>Signature (C, σ)</th>
<th>FS Algo.</th>
<th>Average Misclass.</th>
<th>Average Log Loss</th>
<th>Average GoF</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>a) Derived from GSE1725 and K-Fold Validated on GSE10640[GPL6522] (n = 85), GSE6874[GPL4782] (n = 78), and GSE701 (n = 10)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GADD45A DDB2 (1, 1)</td>
<td>FSFS</td>
<td>0.07</td>
<td>0.40</td>
<td>24</td>
</tr>
<tr>
<td>PPM1D DDB2 CCNF Cdkn1a PCNA GADD45A PRKAB1 TOB1 TNFRSF10B MYC Ccnb2 PTP4A1 BAX CcnA2 ATF3 LG1 CCNG1 FHL2 Ppp1r2 MBd4 Rasgrp2 Ubc Ninj1 Trim22 Il2rb Tp53bp1 Ptprcap Eef1d Ptpre Rad23b Eif2b4 Stx11 Ptn6 Stk10 Psmd1 Btg3 Mlh1 Rnpep Hspd1 Ung Ptprc Ptpra Bcl2 Gss Sh3bp5 Tpp2 Idh3b Ccnh Stk1 Hsp27 Eif4ebp2 Hpaa4 Fads2 Rpa3 Gzmk Anxa4 Icam1 Ppid Lm02 Ppido Nudt1 Fuc Polor2a L19 Rpa1 Pts Tnfrsf4 Rpa2 Psmd8 Gcdh Man2c1 Ptntr2 Ruvbl1 Atph Gk Cd79b Mapk4 Pold1 Ptk2h Akt2 Moap1 Cong2 Aldoa Srd5a1 Hat1 Xrcc1 Eif2s3 Rad1 Ube2a Zfp36l1 Cdb8a Taldo1 Gpx4 Ssbp2 Ercc3 Atps0 Pepd Eif4g2 Aco2 Hexb Ubex3a Arpc1a Psmd10 Prcp Ppib Znf337 Cetn2 Rpl29 (10000, 10000)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>b) Derived from GSE10640[GPL6522] and K-Fold Validated on GSE1725 (n = 114), GSE6874[GPL4782] (n = 78), and GSE701 (n = 10)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ddb2 Rad17 Psmd9 Ly9 Ppih Pcna Mdhd2 Moap1 Tp53bp1 Ppm1d Atsg1 Bcl2l2 Eno2 Ptp4a1 Psmd8 Lgi1 Fdps Ogdh Ccng1 Psmd1 (100, 100)</td>
<td>BSFS</td>
<td>0.05</td>
<td>0.39</td>
<td>15</td>
</tr>
<tr>
<td>Ddb2 Hspd1 Icam1 Ptp4a1 Gtf3a Ly9 (100000, 10000)</td>
<td>FSFS</td>
<td>0.08</td>
<td>0.16</td>
<td>43</td>
</tr>
<tr>
<td>Rad17 Tnfrsf10b Psmd9 Ly9 Ppih Pcna Znf337 Mdhd2 Tp53bp1 Ppm1d Zfp36l1 Atps0 Gadd45a Ptp4a1 Psmd8 Lgi1 Atps0 Fdps Ogdh Psmd1 (100000, 10000)</td>
<td>BSFS</td>
<td>0.05</td>
<td>0.22</td>
<td>11</td>
</tr>
<tr>
<td><strong>c) Derived from GSE6874[GPL4782] and K-Fold Validated on GSE1725 (n = 114), GSE10640[GPL6522] (n = 85), and GSE701 (n = 10)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ddb2 Prkdc Prkch Igj (100000, 10000)</td>
<td>FSFS</td>
<td>0.02</td>
<td>0.27</td>
<td>7</td>
</tr>
<tr>
<td>Ddb2 Prkdc Tpp2 Ptpre Gadd45a (10000, 100)</td>
<td>FSFS</td>
<td>0.02</td>
<td>0.07</td>
<td>5</td>
</tr>
</tbody>
</table>
This type of external validation is the most challenging due to the variability associated with different microarray experiments and batch effects of different platforms. This potentially explains the lower performance obtained by traditional validation (Table 3) compared with k-fold validation on the same datasets (Table 2). The remaining human signatures are described in Supplementary Files Y1–Y5.

To determine which human genes are most consistently selected, the most frequently appearing genes (11 or 12 depending on number of equally prevalent genes in different signatures) were compiled from the top 20 human signatures (assessed by lowest average log loss during k-fold validation) from GSE10640[GPL6522], GSE6874[GPL4782], and GSE1725. The union of these three lists indicates the relative frequencies of each gene (Figure 6). Figure 7 visualizes the mutual information of gene expression (Figure 6) shared with radiation dose.

While most genes have similar representation in signatures derived from different datasets, GADD45A and DDB2, in particular, are significantly more frequent in those derived from GSE1725 and GSE10640[GPL6522]. GADD45A and DDB2 are present in signatures derived from samples irradiated at different exposures (GADD45A – 500 cGy, DDB2 – 150-200 cGy). This raises questions as to whether these genes have a larger influence on the accuracy of individual signatures and whether their expression is calibrated to radiation exposure levels. Removal of these gene features was performed to address their impact. Genes of interest have been removed from each of the top 20 human signatures derived from various datasets and then the signatures were revalidated excluding these features (Table 4). The difference between the validation metrics preceding and following removal of a gene represents the changes in misclassification, log loss, and goodness of fit, respectively. GADD45A appears in 14 of the top 20 signatures derived from GSE1725. Of the 14 signatures, 10 were single gene signatures, as GADD45A alone was expected to sufficiently distinguish irradiated from unirradiated samples. In these cases, it was assumed that a null signature would perform as well as a predictor that randomly draws predictions from a uniform distribution of doses. Removal of GADD45A from these 14 signatures, results in an average increase in misclassification, log loss, and goodness of fit by 0.319, 0.368, and 109 cGy, respectively (see Table 4a). In contrast, elimination of BAX, which only appears in 2 of the top 20 signatures derived from GSE1725 and results in an average increase in misclassification, log loss, and goodness of fit by 0.018, 0.147, and 2.95 cGy respectively (Table 4c). Comparing the effects of removing DDB2 (Table 4c) and PRKAB1 (Table 4f) from the top 20 GSE10640[GPL6522] signatures confirms the impact of genes that frequently occur within the most accurate gene signatures.

However, the diagnostic contributions of GADD45A and DDB2 expression to the radiation levels at which samples were exposed (500 cGy and 150-200 cGy respectively) are confounding. The effects on model performance resulting from removal of GADD45A from the GSE10640[GPL6522] signatures (Table 4b) versus the GSE1725 signatures (Table 4a) are discordant. ΔMC is higher when GADD45A is removed from GSE1725, but ΔLL is higher when GADD45A is removed from GSE10640[GPL6522]. ΔLL is large when GADD45A is removed from both datasets, which is consistent with the importance of GADD45A at both radiation doses. Indeed, GADD45A expression has been demonstrated to be rapidly induced by radiation levels as low as 2 Gy[23]. Similar discordance was observed in the feature removal experiments of DDB2 (Table 4c, 4d).

As was the case with murine signatures, genes appearing in the best human signatures do not necessarily share high mutual information with radiation dose. However, the compositions of the human signatures are dominated by four genes, DDB2, GADD45A, PCNA, and PPM1D, which all share a lot of information with radiation dose (DDB2: 0.55, GADD45A: 0.39, PCNA: 0.51, PPM1D: 0.46). The functions associated with these and less frequently appearing genes are depicted in Figure 8. The pathways and functions represented include keratinocyte differentiation.

### Table 3. Best performing human signatures assessed by traditional validation.

<table>
<thead>
<tr>
<th>Signature (C, T)</th>
<th>FS Algo.</th>
<th>FS Log Loss</th>
<th>Validation Log Loss</th>
<th>Validation GoF</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) Derived from GSE10640[GPL6522] and Validated on GSE6874[GPL4782] (n = 78)</td>
<td>FSFS</td>
<td>2.0E-14</td>
<td>0.70 ± 0.03</td>
<td>25 ± 0</td>
</tr>
<tr>
<td>DDB2 H5PD1 MAP4K4 GTF3A PCNA MDH2 (1000, 10)</td>
<td>FSFS</td>
<td>0.07</td>
<td>0.20 ± 0.03</td>
<td>35 ± 5</td>
</tr>
<tr>
<td>DDB2 GTF3A TNFRSF10B (1, 1)</td>
<td>BSFS</td>
<td>N/A</td>
<td>0.08 ± 0.00</td>
<td>14 ± 0</td>
</tr>
<tr>
<td>DDB2 CDA8 TALDO1 PCNA EIF4G2 LCN2 CDKN1A PRKCH E01 PPM1D (10000, 1)</td>
<td>BSFS</td>
<td>N/A</td>
<td>0.12 ± 0.00</td>
<td>21 ± 0</td>
</tr>
</tbody>
</table>

DDB2 H5PD1 MAP4K4 GTF3A PCNA MDH2 (1000, 10)
DDB2 GTF3A TNFRSF10B (1, 1)
DDB2 CDA8 TALDO1 PCNA EIF4G2 LCN2 CDKN1A PRKCH E01 PPM1D (10000, 1)

This table lists indicates the relative frequencies of each gene (Figure 6) shared with radiation dose.

### Table 4. Feature removal from the top 20 human signatures.

<table>
<thead>
<tr>
<th>Feature</th>
<th>FS Log Loss</th>
<th>Validation Log Loss</th>
<th>Validation GoF</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) Derived from GSE10640[GPL6522] and Validated on GSE6874[GPL4782] (n = 78)</td>
<td>FSFS</td>
<td>2.0E-14</td>
<td>0.70 ± 0.03</td>
</tr>
<tr>
<td>DDB2 H5PD1 MAP4K4 GTF3A PCNA MDH2 (1000, 10)</td>
<td>FSFS</td>
<td>0.07</td>
<td>0.20 ± 0.03</td>
</tr>
<tr>
<td>DDB2 GTF3A TNFRSF10B (1, 1)</td>
<td>BSFS</td>
<td>N/A</td>
<td>0.08 ± 0.00</td>
</tr>
<tr>
<td>DDB2 CDA8 TALDO1 PCNA EIF4G2 LCN2 CDKN1A PRKCH E01 PPM1D (10000, 1)</td>
<td>BSFS</td>
<td>N/A</td>
<td>0.12 ± 0.00</td>
</tr>
</tbody>
</table>

DDB2 H5PD1 MAP4K4 GTF3A PCNA MDH2 (1000, 10)
DDB2 GTF3A TNFRSF10B (1, 1)
DDB2 CDA8 TALDO1 PCNA EIF4G2 LCN2 CDKN1A PRKCH E01 PPM1D (10000, 1)

This table lists indicates the relative frequencies of each gene (Figure 6) shared with radiation dose.
Figure 7. Scatter plot depicting the mutual information each gene’s expression shares with radiation exposure (averaged over GSE10640[GPL6522], GSE6874[GPL4782], and GSE1725). The size of each circle is proportional to the frequency at which the gene appears in the top 20 human signatures ranked by average validation log loss from GSE10640[GPL6522], GSE6874[GPL4782], and GSE1725. The genes shown are also the same as those indicated in Figure 6.

Figure 6. Heat map depicting the gene compositions of the top 20 human signatures derived at different radiation doses: 150-200 cGy (GSE10640[GPL6522], GSE6874[GPL4782]) and 500 cGy (GSE1725). Frequencies are first scaled within and then between datasets to ensure values between 0 and 1.
Table 4. Effect of removing genes from signatures of different datasets.

<table>
<thead>
<tr>
<th>GSE1725 Validation (0 vs 500 cGy)</th>
<th>GSE10640 Validation (0 vs 150-200 cGy)</th>
<th>GSE6874 Validation (0 vs 150-200 cGy)</th>
<th>GSE701 Validation (300 vs 700 cGy)</th>
<th>Average</th>
</tr>
</thead>
<tbody>
<tr>
<td>ΔMC</td>
<td>ΔLL</td>
<td>ΔGoF</td>
<td>ΔMC</td>
<td>ΔLL</td>
</tr>
<tr>
<td>a) Removal of GADD45A from signatures derived from GSE1725</td>
<td>0.446</td>
<td>0.008</td>
<td>N/A*</td>
<td>0.367</td>
</tr>
<tr>
<td>b) Removal of GADD45A from signatures derived from GSE10640[GPL6522]</td>
<td>0.001</td>
<td>0.011</td>
<td>0.658</td>
<td>0.001</td>
</tr>
<tr>
<td>c) Removal of DDB2 from signatures derived from GSE10640[GPL6522]</td>
<td>0.128</td>
<td>0.166</td>
<td>64.2</td>
<td>0.078</td>
</tr>
<tr>
<td>d) Removal of DDB2 from signatures derived from GSE1725</td>
<td>0.012</td>
<td>0.044</td>
<td>N/A*</td>
<td>0.069</td>
</tr>
<tr>
<td>e) Removal of BAX (control for GADD45A) from signatures derived from GSE1725</td>
<td>N/A**</td>
<td>0.08</td>
<td>N/A*</td>
<td>0.024</td>
</tr>
<tr>
<td>f) Removal of PRKAB1 (control for DDB2) from signatures derived from GSE10640[GPL6522]</td>
<td>N/A**</td>
<td>0.001</td>
<td>N/A*</td>
<td>0.011</td>
</tr>
</tbody>
</table>

*ΔGoF is always N/A for the dataset used to derive signatures because GoF is never used as the optimized metric during signature development (see Feature Selection Algorithms section under Methods).
**Unavailable because the top 20 human signatures derived from GSE1725 were all obtained by optimizing log loss rather than misclassification.

Figure 8. Depiction of major cellular functions of most frequently appearing genes of the best human signatures (see Figure 5 and Figure 6).
Validation Performance

N/A 99%

TP53BP1).

In the future, a similar analysis could be performed in clinical studies of human subjects exposed to different radiation levels, which might prove useful for determining treatment eligibility after exposure to high levels of myelosuppressive radiation.

A comparison of the most frequently appearing genes in the optimal human (Figure 6) and mouse signatures (Figure 2) with signatures previously derived in other studies reveals little overlap (Table 5). The compositional differences can be attributed to types of samples used for model training, microarray platforms used, and feature selection techniques used in deriving signatures. However, genes consistently selected in optimized signatures in at least three independent studies include BAX, DDB2, GADD45A, LY9, and TRIM22. Expression of these genes is indeed predictive of radiation dose and not just a result of noise in individual datasets. An ensemble signature consisting of these genes achieves up to 92.3% accuracy in k-fold validation over 277 samples and up to 81.2% accuracy in traditional validation over 78 samples. The quality of the gene signature is largely determined by the quality and amount of training data used to fit the SVM model. Thus, this level of accuracy is not the upper bound on the performance of an SVM of the ensemble signature. Additional data at exposures with fixed levels of radiation in matched training and testing samples could improve model performance.

Ensemble models should be considered which combine genes discovered in different well-performing signatures. Although the most frequently represented human and murine genes were

### Discussion

Biochemically inspired genomic signatures of human and murine radiation response exhibit high accuracies in validating independent datasets (98% in k-fold validation, 92% by traditional methods). Some of the human signatures exhibit among the highest specificities reported (e.g. the signature DDB2, CD8A, TALDO1, PRKCH, EIF4G2, LCN2, CDKN1A, PRKCH, ENO1, PPM1D) and exhibited 92% accuracy when validated on GSE10640[GPL6522]. This dataset contains both radiation therapy patients (150-200 cGy) and controls (0 cGy) which include healthy donors and chemotherapy patients treated with alkylators. Thus, the signature distinguished radiation-induced and chemotherapy-associated DNA damage.

Some of the best performing signatures consisted of one to three gene features. The first signature in Table 2 contains GADD45A and DDB2, and exhibits a misclassification error rate of 7%. These relatively short signatures have advantages over longer signatures with similar performance. It is more likely that the model can be generalized to a wider spectrum of data, when fewer features are required, and from a practical standpoint, diagnostic tests based on fewer gene expression measurements are less susceptible to experimental error.

BAX, an inducer of apoptosis, was the single gene shared among those frequently appearing in both murine and human signatures. One possible explanation for this is that the mouse datasets featured samples irradiated at four levels while human datasets contained samples irradiated at two levels. Genes selected by multi-class model algorithms may better discriminate radiation dose. Nonetheless, the radiation response pathways of mice are not necessarily similar to those of humans. In fact, Lucas et al. have shown that the murine signatures they developed are not translatable to human samples.

Furthermore, only two genes, including BAX, are shared by the human and murine signatures derived by Dressman et al. (human)

None of the samples exposed to ≥200 cGy are misclassified below this radiation dose based on the multi-class murine signatures (Figure 5). In the future, a similar analysis could be performed in clinical studies of human subjects exposed to different radiation levels, which might prove useful for determining treatment eligibility after exposure to high levels of myelosuppressive radiation.

### Table 5. Genes found in best performing signatures and previously derived signatures.

<table>
<thead>
<tr>
<th>Prior Studies</th>
<th>Validation Performance</th>
<th>Shared Genes in Signatures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dressman et al. (human)</td>
<td>K-Fold (internal) 90%, K-Fold (external) N/A, Traditional (external) N/A</td>
<td>BAX, DDB2, PRKCH</td>
</tr>
<tr>
<td>Dressman et al. (mouse)</td>
<td>N/A, N/A, N/A</td>
<td>Bax, Cd72, Cd79b, Cdkn1a, Ei24, Galt, Glipr2, Ly6d, Ms4a1, Tlam</td>
</tr>
<tr>
<td>Paul et al. (human)</td>
<td>98%, N/A, N/A</td>
<td>BAX, DDB2, GADD45A, LY9, PCNA, PPM1D, PTP4A1, RASGRP2, TRIM22</td>
</tr>
<tr>
<td>Lu et al. (human)</td>
<td>~90%, 86%, N/A</td>
<td>DDB2, FHL2, GADD45A, LY9, TRIM22</td>
</tr>
<tr>
<td>This study (human)</td>
<td>100%, 98%, 92%</td>
<td>N/A</td>
</tr>
<tr>
<td>This study (mouse)</td>
<td>99%, 92%, N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>
compiled, genes common to one dataset did not appear equally frequently in signatures from the other. This discordance may possibly result of noise in the different datasets, or perhaps to intrinsic differences between them. Compilation of frequently appearing genes in different datasets may be useful for discovery of consistently represented genes that are incorporated into high-performance signatures.

The types of data available for this study and the analytical approaches we used potentially limited the interpretation of these gene signatures. Blood samples of mouse and human datasets were all collected within 24 hours of exposure. Thus, signatures derived on these datasets may only be valid in white blood cells with a limited time window (<24 hours). Additionally, one of the datasets we used to derive signatures, GSE6874, appears to have been a particularly noisy dataset, based on the average misclassification rates on GSE10640, GSE1725, and GSE6874 of 0.03, 0.02, and 0.11, respectively. Assuming that it is possible to differentiate samples irradiated at different levels of exposure using expression data, the feature selection misclassification metric estimates the theoretical limit of how well differentially irradiated samples can be separated based on expression. The surprisingly high feature selection misclassification values obtained from GSE6874 may therefore be indicative of greater levels of noise in the data. Lastly, the greedy feature selection algorithms used to derive signatures cannot guarantee optimal results, that is, we cannot confirm that we have found the best possible signatures from each dataset for predicting radiation exposure. This potentially explains the discordance in gene composition between murine datasets (Figure 2).

Nevertheless, the validation performance of radiation signatures is significantly improved (Table 5). The signatures that were externally k-fold validated achieved nearly 100% accuracy. Some of our human signature models are also externally validated in the traditional sense (i.e. using a single model). This validation method, which is representative of an actual scenario, achieves >90% accuracy, and is directly relevant to creating a routine, efficient and highly accurate expression-based radiation prognostic assay.

Data availability
All data underlying the results are available as part of the article and no additional source data are required.


Code is available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

Competing interests

Grant information
Natural Sciences and Engineering Research Council of Canada (NSERC Discovery Grant RGPIN-2015-06290); the Canadian Foundation for Innovation; Canada Research Chairs, and CytoGnomix Inc.

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Supplementary material
Supplementary File X: This spreadsheet lists all the genes found from our literature search (see Methods) that were considered during feature selection. For each gene, we report the reason for inclusion and a link to the paper containing the supporting evidence.

Click here to access the data.

Supplementary Files Y1-Y7: These files contain information concerning all the signatures derived for this paper. Each file contains the validation results of signatures derived from a particular dataset. Files Y1-Y3 contain the k-fold validation results of human signatures derived from GSE1725, GSE6874, and GSE10640, respectively, while Y4-Y5 contain the traditional validation results of human signatures derived from GSE6874 and GSE10640, respectively. Files Y6-Y7 contain the k-fold validation results of mouse signatures derived from GSE10640[GPL4783] and GSE10640[GPL6524], respectively. Each supplementary file contains the following columns: Signature, FS Algorithm, C, sigma, FS Misclassification, FS Log Loss, K, Misclassification, Misclassification Error, Log Loss, Log Loss Error, Goodness of Fit, and Goodness of Fit Error. These headings are described in the tab titled “Legend” in Files Y1-Y7. In addition, Files Y1-Y3 have three extra columns: Average Misclassification, Average Log Loss, and Average Goodness of Fit, which represent the misclassification, log loss, and goodness of fit, respectively, averaged over all validation sets.

Click here to access the data.
References


40. Myeloid Cytokines for Acute Exposure to Myelosuppressive Doses of Radiation (Hematopoietic Subsyndrome of ARS), Cytokine - Radiation Emergency Medical Management [Internet]. [cited 2018 Jan 12]. Reference Source

Roel Quintens
Radiobiology Unit, Institute for Environment, Health and Safety, Belgian Nuclear Research Centre, Mol, Belgium

Mohamed Mysara
Expert Group Interdisciplinary Biosciences, Institute for Environment, Health and Safety, Belgian Nuclear Research Centre, Mol, Belgium

Summary
In this study, the authors have identified gene signatures for radiation dose prediction using machine learning methodologies based on publically available microarray results from human and murine samples (mostly lymphocytes) exposed to ionizing radiation. Their signatures have been independently validated showing a high specificity for dose estimation. The authors have used a novel method, based on the concept of minimum redundancy maximum relevance. This generated signatures which often contained genes that had not previously been identified as potential radiation biomarkers. In all, this is a well-conducted study with relevance to the field. However, we do have some comments/questions/remarks, as outlined below.

Introduction

- Several other studies have applied machine learning methodologies to identify predictive radiation exposure biomarkers. Some of these have been reviewed in Hall et al., Mut Res 2017, Supplementary Table 3.5.1.1.

- Another important aspect of gene expression is alternative splicing, which also occurs in response to ionizing radiation (e.g. Sprung et al., PLoS ONE 2011; Forrester et al., PLoS ONE 2012; Macaeva et al., Sci Rep 2016). The latter study also showed for the first time the suitability of exon signatures as sensitive radiation biomarkers, and highlights the importance of prior knowledge at the exon level for subsequent primer- or probe-based assays (e.g. qRT-PCR). This may be discussed.

Methods
In the data Pre-processing “Rows and columns of microarray data that are less than 95% complete were removed and any remaining missing values were imputed using the nearest-neighbor algorithm” How many rows and columns were removed, and on which basis was the 95% threshold selected. Also, what is the effect of the nearest-neighbor algorithm on the data “over-fitting”. Is it possible to perform PCA on the data after removing any row/columns with less than 100% completeness and compare to the currently presented approach (95% removal and filling the remainder of the missing data)? This would allow the visualization of the effect of the proposed methodology on the segregation between the various records.

Only genes common to all datasets were retained. Does that mean common between mouse and human datasets? How were aliases identified?

The second step in the process is the selection of genes based on a non-exhaustive list of publications. Why was this necessary if the mRMR method for feature selection was applied?

I particularly like the idea of performing quantile normalization after feature selection. Is this something that has been published before? Can the authors speculate (or maybe even compare) about the performance of their method on pre-normalized datasets?

Concerning the method used for the “Validation of models”, I would think this approach would be more vulnerable towards the test/training dataset. What would occur to the accuracy when doing the normalization over-all of the data? Would the accuracy change drastically? Is it possible to extend the testing to cover additional data?

Many datasets exist on human PBMCs/whole blood irradiated with a range of doses. Why were these datasets not considered for this study while lymphoblastoid cell lines were?

It would be helpful to have a comparison of model performance with that of “traditional” machine learning methods, as used in some of the indicated references.

Results

“We discovered radiation gene signatures using the microarray data of human and mouse peripheral blood samples and human lymphoblastoid cell lines, which were validated either according to signature.” Were the human lymphoblastoid cell line and prepheral blood samples grouped together in one model? If so, would it be possible to visualize how the expression data of the shortlisted genes for each data type separately (using PCA for example)?

Can the authors comment on their observation that signatures derived from both murine datasets are not very similar? Apart from “noise, or intrinsic differences in the datasets”, could it possibly also be a consequence of the method used, i.e. mRMR in which low mutual information genes are selected? Based on Fig. 2 and 6 it seems that genes with higher mutual information in general have higher frequencies. Which seems logical.

The authors state that Ms4a1 appears less frequently than Glipr2. However, from the sizes of the circles, Ms4a1 seems to appear more frequently than Glipr2. Please verify this statement.

Are genes in Tables 1 and 2 ranked according to their frequency, importance,…?
How do the authors explain the low frequencies of human signature genes (Fig. 6), compared with murine (Fig. 2)?

Likewise, can the authors explain the large number of genes with low mutual information in the human signature (23 out of 26 <0.4), compared with the murine signature (4 out of 33 <0.4).

Although I like the idea of mRMR, it is somewhat counterintuitive to have genes with little mutual information to be important for dose prediction. This seems to be confirmed by the fact that the compositions of human signatures are dominated by genes with high mutual information (in fact, these are all well known p53-dependent genes which appear in a high number of published radiation signatures).

Discussion

I understand the advantage of small signatures in terms of practicality. However, in case of a real emergency, in which individuals have been irradiated without good knowledge about the exact time since exposure, larger gene signatures may provide the additional benefit of having different dynamics per gene. This may help to also predict not only the dose, but also the time since exposure. Furthermore, one-gene signatures may suffer from higher variability among the population compared to larger gene signatures.

I believe results from other, similar studies may be briefly situated in the introduction/discussion.

References


Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Partly
Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Radiobiology; Biodosimetry; Molecular Biology; Developmental Neuroscience

We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

---

**Author Response 09 Jun 2018**

**Peter Rogan, Western University, London, Canada**

- Regarding other studies that identify predictive radiation exposure biomarkers, we have now added text citing these and other studies in paragraph 1 of the introduction of version 2 of our paper.

- We investigated whether alternative splicing in response to ionizing radiation might affect the expression values that were used in training or validation of machine learning models we derived. These values could theoretically be distorted if the hybridization probes used to quantify gene expression in these microarray data were predominantly located in cassette exons. We carefully analyzed the probes on one of the most prominent genes in our signatures: DDB2, which consists of 10 exons in total. Out of the three human datasets used for deriving signatures, GSE6874[GPL4782] lacked information about the location of its DDB2 probe. However, the DDB2 probes in GSE10640[GPL6522] and GSE1725 were located in exons 8-10. According to UCSC Genome browser, the only transcript variants (NM_001300734.1, mRNA AB107039, and mRNA BC050455) involved skipping or fusing exons 3-7. Thus, in this case, alternative splicing does not affect our results, since the DDB2 probes avoid alternatively spliced exons. In general, probes seem to be designed to avoid alternatively spliced regions. Although we have not verified this for all the genes in our signatures, we speculate that taking an average over multiple probes reduces any potential effect of alternative splicing.

- The nearest neighbor analysis was performed to avoid inclusion of genes or individuals with sparse data in the analysis. We conservatively selected a threshold of 95% completeness to ensure that the original source data were reliable. At this threshold, none of the rows or columns or each of the datasets we used for deriving signatures (GSE1725, GSE6874[GPL4782], GSE10640[GPL6522], GSE10640[GPL4783], GSE10640[GPL6524]) were removed (based on genes common among all datasets). The effect of nearest neighbours on overfitting was minimal. Upon restriction to the set of genes available for feature selection, there were no missing values in GSE1725 and GSE6874[GPL4782], a single missing value in GSE10640[GPL6524] and GSE10640[GPL6522], and 62 missing values in GSE10640[GPL4783]. However, in GSE10640[GPL4783], of the genes available for feature selection, only four genes contained at least one missing expression value: Rad51, Ptpre, Gadd45a, and Pola1. None of these genes are among the 33 most frequently appearing genes of our murine signatures (see Figure 2). (Recall that GSE10640[GPL4783] is a murine dataset). Additionally, overfitting is mitigated by validation using independent
datasets. The nearest neighbor source data occurs before model development, so the model is not fit at that point.

- Regarding the questions about genes common to all datasets being retained: 1) Mouse and human signatures were derived separately. So only genes common to all human datasets were available for selection in deriving human signatures and only genes common to all mice datasets were available for selection in deriving mouse signatures. 2) Our scripts did not take gene name aliases into account. It is therefore possible that genes that were left out of the analysis because they were indicated by different names in different datasets.

- The initial list of publications we consulted may not have been complete, but it was the result of an extensive search. We were concerned that mRMR method applied to all genes without independent experimental support would result in Type II errors. We wanted to identify the key genes from the large volume of peer-reviewed work implicating various genes in radiation response. This hypothesis based study was not designed to discover novel genes, whose putative role in radiation response was unproven or unknown. We have automated the initial feature selection procedures using expression of all (including non-coding) genes, but this is beyond the scope of our efforts. The discovery strategy would require statistical correction for the likelihood of incorrectly rejecting a null hypothesis due to multiple comparison testing.

- Regarding quantile normalization after feature selection, this is the first time that we have used this approach. We attempted this because our initial efforts to derive signatures from pre-normalized data were unsuccessful (poor performance). Since we do not use the expression values for the majority of genes, there was no compelling reason to normalize across the entire set of expressed genes. We are not aware of other previous efforts, but cannot exclude this possible. We did not perform an exhaustive literature search for the method that we employed, since the universe of potential applications is nearly limitless. We speculate that if quantile normalization is done before feature selection, then the reduction in dynamic range in the transformed data may lead to the derivation of poorer signatures. To give a more quantitative answer, we took the top performing signature in the Y5 Supplementary file with respect to log loss (DD2 GTF3A TNFRSF10B) and re-validated with normalization over all data instead of just over genes in the signature; misclassification error was 2% higher and the goodness of fit was 4 cGy higher (log loss remained approximately the same). Interestingly, normalization performed over all of the data does not appear to have significant effects on the performance of our signatures. To “extend the testing to cover additional data,” it would be necessary to renormalize the initial set of genes to include these “additional genes”. If no other genes are to be included, only additional samples, then the expression values of the additional samples would need to be renormalized.

- Regarding other human datasets of irradiated PBMCs, in the feature selection stage, we specifically required datasets with large numbers of samples. The three human datasets we used for feature selection (GSE6874, GSE10640, GSE1725) were the largest ones we found deposited in GEO. In particular, GSE1725, the dataset with cell lines derived from patients, was the largest dataset available to us, containing 110 samples (171 samples in total, but 61 are UV irradiated). At the earliest stage of our project, we specifically chose datasets that maximized the number of common genes represented among them. This requirement enabled us to validate our signatures by either traditional or k-fold approaches. Genes available for feature selection must be present in all datasets in order for validation to work. We were surprised to find many datasets where key genes in our models were missing from expression data (see the last section on partial body irradiation data analysis in the results section of the revised version 2 of this paper).
Regarding comparison of model performance with other traditional machine learning methods, we have used these methods (eg. random forest, SVMs, decision trees) in previous gene expression studies (see reference 13 of version 1 which is reference 18 of version 2). The improved performance we describe here cannot be attributed to the specific model building approaches.

Regarding grouping of lymphoblastoid and PBL samples, models were always trained on one dataset at a time; they were not combined.

The dissimilarity of the murine datasets may be related to their use of different microarray platforms and were collected at different times. It is difficult to tease out the intrinsic differences of the datasets from the performance of the methods.

Regarding Ms4a1 and Glipr2, we verify the reviewers' observation. We corrected the density plot during preparation of the paper, but inadvertently neglected to make the corresponding change to the text. We have replaced Glipr2 with Ccngr1 or Eif2ak4.

Regarding ranking of genes in Tables 1 and 2, signatures derived using BSFS and CSFS list the genes according to mRMR rank. For signatures derived using FSFS, the genes would be listed according to the order in which they were selected. So in this sense, they are indeed ranked according to importance.

Regarding the low frequencies of human vs murine signature genes, in the descriptions of these figures, we mention that “Frequencies are first scaled within and then between datasets to ensure values between 0 and 1.” Human signature gene frequencies appear suppressed because DDB2 and GADD45A in particular were represented more frequently than any other gene by a large margin.

Regarding the numbers of genes with low mutual information in the human vs murine signatures, it may be relevant that expression data are not adjusted for either white blood cell count or body mass. We speculate that the gene expression response in the human samples reflects lower numbers of radiation exposed cells. This could dampen the signals and mutual information with radiation dose compared to the murine response.

Regarding the idea that it would be counterintuitive to have genes with low mutual information as important for dose prediction, it is true that signatures are dominated by genes whose expression values share high mutual information with dose. The purpose of mRMR is to make sure that we do not overlook the genes whose expression values may encode information that is not present in the genes whose expression values have high mutual information with dose, which is why sometimes genes with lower mutual information may appear as frequently or even a bit more frequently than genes with higher mutual information.

Regarding the single gene signature results, we agree that single gene signatures are more susceptible to extrinsic sources of variation unrelated to radiation exposure. However, simpler signatures may be necessary under laboratory conditions that limit the amount or complexity of testing, e.g. space radiation assays performed by astronauts.

**Competing Interests:** No competing interests were disclosed.
The authors are absolutely correct that in the event of a radiological accident blood is likely to be the most likely source for analytical materials that would reflect a radiation exposure. Given the analytical approach taken we tested this using two sets of data from normal human lung epithelial cells (HBECs) irradiated at multiple doses by γ-rays but also with Fe particles such as those found in the deep space environment. (Reference 8 in this manuscript)

We were interested in two things: 1) Would the results from lymphoid cells translate to epithelial cells? There is sufficient evidence in the literature to suggest signatures created with lymphoid cells are poor at predicting radio response in cells from tissue. 2) Would the results from γ-ray exposures translate to something more exotic like Fe particles which have discriminating gene sets that are both common to γ-ray and Fe particles as well as unique to the radiation type.

We chose the 10 gene signature derived from GSE6874 and validated against GSE10640. Given the limited size of our sample set, we were surprised to see Prediction Accuracies of 73% for γ-rays and 83% for Fe particle irradiation when examining the normal HBEC cells; and 85 and 76%, respectively, for a genetically manipulated HBEC (p53 knockdown, KRAS mutant over-expressing) cell line.

We look forward to testing this approach in genetically diverse tumor cells.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.
We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 09 Jun 2018

Peter Rogan, Western University, London, Canada

Thank you for your efforts to review our article and evaluate our software. We were excited about the significant results you obtained using the human signature on expression data from irradiated lung epithelial cells, and using the models to detect evidence of Fe particle radiation.

Competing Interests: No competing interests were disclosed.

Reviewer Report 14 May 2018

https://doi.org/10.5256/f1000research.15278.r33291

© 2018 Oh D. This is an open access peer review report distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Daniel Oh
Department of Radiation Oncology, East Carolina University, Greenville, NC, USA

Well written article. Good analysis of available gene expression data to create gene signatures for ionizing radiation exposure. Good discussion of the identified genes’ functions and roles in radiation response. Conclusions are well supported by the results. Hopefully the analysis and genes identified in this study will be incorporated and/or validated in future studies examining prediction of radiation exposure.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes
**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

---

**Author Response 09 Jun 2018**

**Peter Rogan**, Western University, London, Canada

Thank you for your kind comments. We agree that the approach and software should be useful for future studies of human radiation exposures. We are particularly motivated to apply multiclass SVMs, which were highly accurate in the murine dataset, to the analysis of a large set of radiation oncology patients exposed to different radiation doses.

**Competing Interests:** No competing interests were disclosed.

---

The benefits of publishing with F1000Research:

- Your article is published within days, with no editorial bias
- You can publish traditional articles, null/negative results, case reports, data notes and more
- The peer review process is transparent and collaborative
- Your article is indexed in PubMed after passing peer review
- Dedicated customer support at every stage

For pre-submission enquiries, contact research@f1000.com