BloodSpot

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BloodSpot: a database of gene expression profiles and transcriptional programs for healthy and malignant haematopoiesis

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ABSTRACT
Research on human and murine haematopoiesis has resulted in a vast number of gene-expression data sets that can potentially answer questions regarding normal and aberrant blood formation. To researchers and clinicians with limited bioinformatics experience, these data have remained available, yet largely inaccessible. Current databases provide information about gene-expression but fail to answer key questions regarding co-regulation, genetic programs or effect on patient survival. To address these shortcomings, we present BloodSpot (www.bloodspot.eu), which includes and greatly extends our previously released database HemaExplorer, a database of gene expression profiles from FACS sorted healthy and malignant haematopoietic cells. A revised interactive interface simultaneously provides a plot of gene expression along with a Kaplan–Meier analysis and a hierarchical tree depicting the relationship between different cell types in the database. The database now includes 23 high-quality curated data sets relevant to normal and malignant blood formation and, in addition, we have assembled and built a unique integrated data set, BloodPool. Bloodpool contains more than 2000 samples assembled from six independent studies on acute myeloid leukemia. Furthermore, we have devised a robust sample integration procedure that allows for sensitive comparison of user-supplied patient samples in a well-defined haematopoietic cellular space.

INTRODUCTION
A decade of intense studies of the genetic programs underlying normal and malignant haematopoiesis has resulted in a number of gene-expression data sets, which can potentially help answer questions concerning the molecular mechanisms governing normal haematopoiesis and how these are de-regulated in cancer. To researchers and clinicians with limited bioinformatics experience, these data have been available through online databases in the form of raw or semi-processed files but remained largely inaccessible for analysis, let alone comparison with user-supplied in-house data. Recently, a number of web interfaces have been generated to facilitate single gene queries of in-house data (ImmGen Gene Skyline (1), Gene-expression Atlas (2), Leukemia Gene Atlas (3) and Differentiation Map (2)) or curated, compiled and processed data sets (HemaExplorer (3), Gene Expression Commons (4), A HeamAtlas (5), BloodChIP (6), BloodExpress (7) and CODEX (8)). These tools provide information on the expression of single genes, but fail to answer the main questions as to whether these genes influence patient survival or if genes or pathways are regulated in similar or inverse patterns. We have previously published a comprehensive database of mRNA microarray samples from FACS sorted healthy and leukemic bone marrow samples (3) which has proven a useful and popular resource for researchers working within the areas of cellular differentiation, haematopoiesis and leukaemia.

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Here, we present a complete overhaul and significantly expanded version of the original database, with a new and interactive interface, all freely available online. The new database redefines current approaches to explorative data integration, presentation and visualisation of gene-expression in the haematopoietic system. Consequently, all these improvements called for a new name: BloodSpot.

The core function of BloodSpot is to provide an expression plot of genes in healthy and cancerous haematopoietic cells at specific differentiation stages. To present these haematopoietic gene profiles, we have developed a novel visualisation chart that simply integrates the benefits of strip charts and violin plots. The server accepts either a unique gene name (gene alias) or a gene signature name from the MSigDB database. Of note, an auto-complete mechanism helps finding the right names for genes and gene signatures. To contextualise the haematopoietic gene expression profile, two additional levels of visualisation are available: an interactive hierarchical tree that shows the relationship between the samples displayed and a Kaplan–Meier plot based on a high-quality Acute Myeloid Leukemia (AML) data set (9). Additionally, we added a large body of curated data sets to the database, which users can query seamlessly. Significantly, we provide a new integrated data set of samples from AML patients along with FACS sorted samples from healthy individuals. This new integrated data set provides the most detailed picture of the gene expression landscape in healthy and malignant haematopoiesis to date. Finally, the database provides the possibility of comparing user-supplied leukaemia samples to healthy cells.

The platform is freely available, and requires no login, at: www.bloodspot.eu

DATA CONTENT UPDATES

Available data sets

BloodSpot is a database of mRNA expression in healthy and malignant haematopoiesis and includes data from both humans and mice. The database is subdivided into several data sets that are each accessible for browsing through the new interface. Data sets are organised by organism of origin and disease status. The data sets are organised as follows: first, human healthy haematopoietic cells, then human leukaemia and finally healthy mouse haematopoietic cells. BloodSpot contains the data sets from our previous HemaExplorer (3) as well as new published data sets, all manually processed as described in Rapin et al. (10). All data sets available in BloodSpot were generated using oligonucleotide microarray chips, except for one mouse data set that was generated using RNA sequencing technology. For completeness, the database also includes the content of other online databases that we deem relevant for the study of haematopoiesis in the framework of BloodSpot. These external databases include the Differentiation Map (DMAP) (2) and the ImmunoLogical Genome project (ImmGen) (1).

In total the platform encompasses more than 5000 samples (see Tables 1–3). All data sets were controlled for quality, appropriately normalised and adjusted for batch effects when necessary (11,12).

BloodPool

One new feature of BloodSpot is BloodPool, an aggregated and integrated data set grouping the results of multiple studies focusing on AML. By means of our batch correction methods this data set can be used to study gene expression (programs) in AML in comparison with healthy corresponding cells (see Figure 1). Using the computational method developed in Rapin et al. (10), we have also computed gene expression fold changes relative to their nearest normal counterparts for all AML profiles in BloodPool. BloodPool is available for browsing within BloodSpot and can be selected as any of the other available data sets.

MSigDB and CMAP gene signatures integration

We collected all gene signatures available from the Molecular Signatures Database (MSigDB) (13) (version 4.0) (http://www.broadinstitute.org/gsea/msigdb/) and computed, for each signature, the mean expression values for all samples in all data sets. These mean values summarise the expression of a signature for each sample. Connectivity map (CMAP) (13) signatures were generated with the rank matrix provided by
Table 1. Data sets for normal hematopoiesis

<table>
<thead>
<tr>
<th>Data set</th>
<th>Organism</th>
<th>Source</th>
<th>Sample numbers</th>
<th>Cell types</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal hematopoiesis with AMLs</td>
<td>Human</td>
<td>GSE42519</td>
<td>34</td>
<td>HSC, MPP, CMP, MEP, GMP, early PM, late PM, MY, MM, BC, PMN</td>
<td>Rapin et al. (20)</td>
</tr>
<tr>
<td>Normal hematopoiesis (HemaExplorer)</td>
<td>Human</td>
<td>GSE17054</td>
<td>2</td>
<td></td>
<td>Majeti et al. (21)</td>
</tr>
<tr>
<td>Normal hematopoiesis (HemaExplorer)</td>
<td>Human</td>
<td>GSE19599</td>
<td>4</td>
<td>GMP, MEP</td>
<td>Andersson et al. (22)</td>
</tr>
<tr>
<td>Normal hematopoiesis (HemaExplorer)</td>
<td>Human</td>
<td>GSE11864</td>
<td>2</td>
<td>Monocytes</td>
<td>Hu et al. (23)</td>
</tr>
<tr>
<td>Normal hematopoiesis (HemaExplorer)</td>
<td>Human</td>
<td>E-MEXP-1242</td>
<td>2</td>
<td>Monocytes</td>
<td>Wildenberg et al. (24)</td>
</tr>
<tr>
<td>Normal hematopoiesis (DMAP)</td>
<td>Human</td>
<td>GSE24759</td>
<td>211</td>
<td>Normal Hematopoiesis</td>
<td>Novershtern et al. (2)</td>
</tr>
<tr>
<td>Mouse normal hematopoietic system</td>
<td>Mouse</td>
<td>GSE14833, GSE6506, GSE15907</td>
<td>&gt;700</td>
<td>Normal Hematopoiesis</td>
<td>Di Tullio et al. (25), Chambers et al. (26)</td>
</tr>
</tbody>
</table>

Table 2. Data sets for leukemic patients

<table>
<thead>
<tr>
<th>Data set</th>
<th>Organism</th>
<th>Source</th>
<th>Patient numbers</th>
<th>Cell types</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>AML Normal Karyotype data sets</td>
<td>Human AML</td>
<td>GSE15434</td>
<td>251</td>
<td>NK-AML, WBM</td>
<td>Kohlman et al. (28)</td>
</tr>
<tr>
<td>AML TCGA data sets</td>
<td>Human AML</td>
<td>TCGA</td>
<td>183</td>
<td>Various genetic aberrations, including t(8;21), inv(16), t(15;17), t(11q23), complex karyotype, WBM AML, ALL and preleukemic stages.</td>
<td>TCGA (9)</td>
</tr>
<tr>
<td>Leukemia MILE study</td>
<td>Human AML</td>
<td>GSE13159</td>
<td>2096</td>
<td>AML, ALL and preleukemic stages.</td>
<td>Haferlach et al. (29,30)</td>
</tr>
<tr>
<td>AML versus normal</td>
<td>Human AML</td>
<td>GSE6891, GSE13159</td>
<td>91</td>
<td>NK-AML, WBM</td>
<td>de Jonge et al. (31,32)</td>
</tr>
<tr>
<td>Bloodpool</td>
<td>Human AML</td>
<td>GSE13159, GSE15434, TCGA, GSE16804, GSE14468</td>
<td>251/2076</td>
<td>Mainly AML, ALL and preleukemic stages.</td>
<td>all references above</td>
</tr>
</tbody>
</table>

Data normalisation

All data were normalised and batch corrected to eliminate potential lab batch effects. For this we performed Robust Multi-array Average (RMA) (14) normalisation of all microarray .CEL data files partitioned by origin, and next applied ComBat (http://jlab.byu.edu/ComBat/) (12) an empirical Bayes method implemented in the R language. The batches were defined to be the study name/number, while the covariates was assigned to the relevant cell type. The resulting integrated gene expression databases can be visualised directly or compared to external samples provided by the user. See Tables 1–3 for an overview of the data presented in BloodSpot and the normalisation procedure used. All AML data sets available in BloodSpot are normalised according to Rapin et al. (10) and further batch corrected using ComBat when necessary. This processing schema ensures that the samples are normalised in the context of normal haematopoiesis and according to state of the art batch correction methods, regardless of the origin of the data.

For RNA-seq data, we used the Blue Collar Bioinformatics RNA-seq pipeline (mapping on mm10 mouse genome with TopHat version 2 (15), (https://bcbio-nextgen.readthedocs.org/)) to obtain normalised count data from raw fastq files from Lara-Astiaso et al. (16). We report count data processed using the variance stabilising transformation method from the DESeq2 package (17).

Abbreviations and sample annotations

Abbreviations for all cell types can be found below the plot by clicking the ‘Abbreviations’ link. Typically, the user can find more detailed information about each cell type such as a longer, more informative name, and for healthy cells data sets the immunophenotype, when available. Links to the raw unprocessed data can also be found here.

Available genes

The server is restricted to genes found in our database of Affymetrix Human 133U plus 2, Affymetrix Human...
Table 3. Data set overview

<table>
<thead>
<tr>
<th>Data set</th>
<th>Features</th>
<th>Samples</th>
<th>Normalisation method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leukemia MILE study</td>
<td>67191</td>
<td>2095</td>
<td>1</td>
</tr>
<tr>
<td>Normal human hematopoiesis with AMLs</td>
<td>67191</td>
<td>296</td>
<td>1,7</td>
</tr>
<tr>
<td>Immgen Key populations</td>
<td>47273</td>
<td>256</td>
<td>2</td>
</tr>
<tr>
<td>AML versus normal</td>
<td>67191</td>
<td>252</td>
<td>3</td>
</tr>
<tr>
<td>AML TCGA data set</td>
<td>67191</td>
<td>244</td>
<td>1</td>
</tr>
<tr>
<td>AML TCGA data set versus normal</td>
<td>67191</td>
<td>244</td>
<td>3</td>
</tr>
<tr>
<td>AML Normal Karyotype</td>
<td>54675</td>
<td>234</td>
<td>1</td>
</tr>
<tr>
<td>AML Normal Karyotype versus normal</td>
<td>67191</td>
<td>234</td>
<td>3</td>
</tr>
<tr>
<td>Normal human hematopoiesis (DMAP)</td>
<td>35459</td>
<td>211</td>
<td>4</td>
</tr>
<tr>
<td>Immgen abT cells</td>
<td>47273</td>
<td>190</td>
<td>2</td>
</tr>
<tr>
<td>Immgen Dendritic cells</td>
<td>47273</td>
<td>151</td>
<td>2</td>
</tr>
<tr>
<td>Immgen MFs Monocytes Neutrophils</td>
<td>47273</td>
<td>114</td>
<td>2</td>
</tr>
<tr>
<td>Immgen B cells</td>
<td>47273</td>
<td>103</td>
<td>2</td>
</tr>
<tr>
<td>Normal human hematopoiesis (HemaExplorer)</td>
<td>57270</td>
<td>77</td>
<td>5</td>
</tr>
<tr>
<td>Immgen gdT cells</td>
<td>47273</td>
<td>76</td>
<td>2</td>
</tr>
<tr>
<td>Immgen Stem and progenitor cells</td>
<td>47273</td>
<td>76</td>
<td>2</td>
</tr>
<tr>
<td>Mouse normal hematopoietic system</td>
<td>57613</td>
<td>67</td>
<td>4</td>
</tr>
<tr>
<td>Immgen Activated T cells</td>
<td>47273</td>
<td>55</td>
<td>2</td>
</tr>
<tr>
<td>Immgen NK cells</td>
<td>47273</td>
<td>47</td>
<td>2</td>
</tr>
<tr>
<td>Immgen Stromal cells</td>
<td>47273</td>
<td>39</td>
<td>2</td>
</tr>
<tr>
<td>Mouse normal (RNA seq)</td>
<td>45426</td>
<td>52</td>
<td>6</td>
</tr>
<tr>
<td>BloodPool</td>
<td>67191</td>
<td>2120</td>
<td>1,7</td>
</tr>
<tr>
<td>BloodPool versus normal</td>
<td>67191</td>
<td>2076</td>
<td>3,7</td>
</tr>
</tbody>
</table>

Normalisation method legend:
1. Each cancer sample is normalised together with a set of samples from sorted normal myeloid populations. All samples where normalised using RMA. Comparison of gene expression values is not possible with other data sets in Bloodspot.
2. All samples from the ImmGen data sets were normalised together with RMA. Samples were subsequently attributed to the different data sets in Bloodspot. This means that comparison of gene expression values is possible across all ImmGen data sets.
3. The data are normalised according to Rapin et al. Briefly, each cancer sample is normalised together with a set of samples from sorted normal myeloid populations. Next, using a PCA-based method, the 5 closest normal samples from the cancer sample are averaged and this computed normal sample are next compared to the cancer sample allowing for computation of gene expression fold changes. See Supplementary Methods and Rapin et al. (10).
4. All samples were normalised using RMA. Comparison of gene expression values is not possible with other datasets in Bloodspot.
5. See our previous work (Bagger et al. (3)).
6. The data were processed using the bcbio nextgen RNA-seq pipeline. Count data were subsequently processed with DESeq2’s variance stabilising transformation method.
7. The data was batch corrected using ComBat, taking study number as batch.

133UA and Affymetrix Human 133UB chips for human, and GeneChip Mouse Genome 430 2.0 and Affymetrix Mouse Gene 1.0 ST Arrays for mouse. For the RNA-seq data set UCSC annotation for the mm10 genome was used.

In order to handle gene aliases, a dictionary of gene aliases was constructed from NCBI ftp://ftp.ncbi.nlm.nih.gov/gene/DATA/ and The HUGO Gene Nomenclature Committee (HGNC) www.genenames.org. Ambiguous gene aliases were not included when constructing the dictionary. The alias conversion is only used when the query is not an official gene symbol or probe name. The end result allows for greater flexibility regarding gene names input and faster browsing.

**FUNCTIONALITY UPDATES**

Both the back-end and the front-end have been completely redesigned for interactive usage and speed of execution. The interface is built with a range of new functionalities, with a focus on simplicity of use (see Figure 2).

**Unified input**

BloodSpot takes a single gene name (or unambiguous gene alias) or gene signature name as query. Users can search for keywords such as ‘carcinomas’ or ‘cell cycle’ and will be provided with a list of matching gene signature names. When relevant, it is possible to select which probe-set to display from the list in the upper right corner of the main plot. By default, the probe with the overall highest intensity is at the top of the list. The option ‘Max probe’ will use the one probe with the highest intensity within each population.

**Default plot**

When visiting the interface the plot at the centre of the screen in the default view. This representation is a novel improved jitter strip chart of gene expression, a swift novel visualisation plot that draws from bar plots and violin plots where the jitter is controlled by the density of samples and normalised overall the columns in the chart. Thus the width of the data cloud shows how many samples have similar values (see Figure 3A and a comparison to existing data plot types in Supplementary Figure S1). For more details on this visualisation method please see (Sidiropoulos, N.,
Figure 2. BloodSpot interface details. After a gene alias is submitted to display its expression pattern, any of the top three panels can be clicked to magnify content. The three panels show, from left to right, a survival plot based on a high-quality AML data set displaying a full Kaplan–Meier analysis for any query gene or signature, an improved jitter strip chart of gene-expression plot that draws from bar plots and violin plots and an interactive hierarchical tree that shows the relationship between the samples displayed and allows changing the focus of the display. The Select Population button allows the user to select which populations to display. The Gene Correlations button shows in a table how much other genes or gene signatures correlate with the displayed gene. It is possible to click on the genes in the table to display their expression profile. The Print as PDF button allows the user to export the current plot in PDF format. The T-Test button allows you to perform significance test between pairs of populations (legend is as follows: NS: non significant; *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$). The Export Data as Text button allows you to export the raw data as text (CSV format). The Upload your own sample button allows for the upload of an Affymetrix HU133 plus 2.0 CEL file and for viewing it in the context of normal haematopoiesis. The dropdown menu in the upper right corner of the main plot can be used to select a probe representing the gene of interest; by default, the probe with the highest intensity is chosen. At the bottom of the main plot, a list of abbreviations is available that includes immunophenotypes when applicable.

Sohi, S.H., Rapin, N. and Bagger, F.O. (2015) SinaPlot: an enhanced chart for simple and truthful representation of single observations over multiple classes. bioRxiv, http://dx.doi.org/10.1101/028191). Both an R-package and a web-server have been implemented for those interested in make use of this plot type that we have named SinaPlot.

Survival plot

The chart shown to the left of the BloodSpot interface is a survival plot based on a high-quality AML data set from The Cancer Genome Atlas (TCGA). It displays a full Kaplan–Meier analysis of survival. The survival plots are only available for human data sets, sharing probes with the microarray platform used by the TCGA (Affymetrix U133 Plus 2) (see Figure 3B).

Tree plot

The chart shown to the right of the BloodSpot interface is an interactive hierarchical tree that shows the relationship between the samples displayed and allows changing the focus of the display. It is possible to mouse over the nodes to get the full name for long names. Nodes can be clicked to collapse a branch of the tree—this will also update the default plot in the middle and remove the same populations there (see Figure 3C).

Correlation of genes and gene signatures

For each gene and signature in every data set, we report the top correlating genes or signatures. Taking the haematopoietic fingerprint (e.g. the expression value of one gene over all haematopoietic cells) of all probe-sets and gene signatures in a given data set, we calculated the correlation matrix (Pearson) and present the highest positive and negative correlating genes/signatures. This feature allows for investigation of new associations between putative co-regulated genes or signatures that exhibit similar or inverse expression patterns over the course of haematopoiesis (see Figure 3D).

Other built-in tools

Cell populations may be removed from the graphs using the ‘Select population’ button. The current plot displayed can be exported as PDF in publication-ready quality using the ‘Print as PDF’ button. The ‘T-Test’ button can be used to add the results from a students t-test for significance between pairs of populations to the plot. The legend is as following: NS: non-significant; *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$. The significance marks relies on t statistics for unequal sample sizes but assuming equal variance and the critical values are compared with a two-tailed probability. Finally, raw data can be exported as CSV using the ‘Export Data as Text’ button.
Figure 3. Main plots from BloodSpot for MEIS1. (A) Default view in BloodSpot. The plot is a novel improved jitter strip chart of gene expression that draws from bar plots and violin plots where the jitter is controlled by the density of samples and normalised over all the columns in the chart. (B) Survival plot based on a high-quality AML data set from The Cancer Genome Atlas (TCGA). It displays a full Kaplan–Meier analysis of survival. The survival plots are only available for human data sets, sharing probes with the microarray platform used by the TCGA. (C) Interactive hierarchical tree that shows the relationship between the samples displayed. Hovering over the nodes provides the full names of cell populations. Nodes can be clicked to collapse a branch of the tree—this will also update the default plot in the middle and remove the same populations there. The colour in the nodes represents the median expression of the queried gene. To accentuate the display in the trees, node size is also proportional to gene expression. Trees are based on literature (hierarchical differentiation), or overall sample correlation (correlation of samples). (D) Example table of genes and gene signatures correlating with MEIS1 expression in the default data set. This table appears when the user clicks on the ‘correlation’ button.

Upload sample

By clicking the ‘Upload sample’ button it is possible to analyse user-supplied samples produced on the Affymetrix U133 plus 2 platform. Significantly, doing so allows for the comparison of any myeloid microarray data to normal haematopoiesis. The resulting analysis is then displayed in a private session in the framework of BloodSpot along with a principal component analysis that shows the location of the uploaded sample in the hematopoietic sample space. The analysis is anonymous and requires no login. The resulting data set, including the uploaded sample, can then be queried along with the default data sets in a private session. All names and array information are stripped from the uploaded file before creating the database for the user session. Hence, the uploaded sample in the private session will appear simply as S,1 in all charts. The private sessions and uploaded data are deleted every day at GMT 1.30 pm.

EXAMPLES OF USE OF BLOODSPOT

To demonstrate the use of BloodSpot, we provide in the following section an example relying on data and analysis provided by the database.

MEIS1 is part of a transcriptional program required for the maintenance of MLL-rearranged AML (18). The expression of this gene is therefore often up-regulated in MLL leukaemias. Using Bloodspot, we investigated the expression pattern of MEIS1, and found it to be expressed at high levels in stem cells with decreasing expression as the cells differentiate (Figure 3A and C). Using the correlation function, we find that MEIS1 expression also correlates with the expression patterns of a number of Homeobox genes, including HOXA3, HOXA9 and HOXA10 which are also typically expressed early during haematopoiesis (19) (Figure 3D). Switching to the BloodPool data set, MEIS1 is found to be up-regulated in MLL leukaemias (Figure 4). Although the P-value in the survival plot does not reach statistical significance (0.055; see Figure 3B), the influence
patterns of single genes impact on patient survival, and addresses two key questions, namely, how gene-expression in developmental systems.

Haematopoiesis, leukaemia, basic immunology and gene expression in differentiated human blood cells. This will allow assessment of the origin of the blast population in AML patients as well as assessment of well known and novel genetic markers in the context of normal haematopoiesis, both of which could be important for stratification of difficult patient cases.

We have also integrated the largest pool of AML patient microarray samples to date and have computed gene expression fold changes for these profiles, thanks to our capability in the haematopoietic system. Thus the platform will help broaden the basis on which to generate hypotheses about potential therapeutic targets and expand the understanding of co-regulated genes and pathways, to support experimental findings from animal model systems.

**AVAILABILITY**

Bloodspot is accessible at [www.bloodspot.eu](http://www.bloodspot.eu)

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

**ACKNOWLEDGEMENTS**

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**REFERENCES**


![Figure 4. MEIS1 expression relative to the nearest normal counterpart in different AML subtypes, including MLL-rearranged AML.](image)


