DATA NOTE

A curated transcriptome dataset collection to investigate the immunobiology of HIV infection [version 1; referees: 3 approved]

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Abstract

Compendia of large-scale datasets available in public repositories provide an opportunity to identify and fill current gaps in biomedical knowledge. But first, these data need to be readily accessible to research investigators for interpretation. Here, we make available a collection of transcriptome datasets relevant to HIV infection. A total of 2717 unique transcriptional profiles distributed among 34 datasets were identified, retrieved from the NCBI Gene Expression Omnibus (GEO), and loaded in a custom web application, the Gene Expression Browser (GXB), designed for interactive query and visualization of integrated large-scale data. Multiple sample groupings and rank lists were created to facilitate dataset query and interpretation via this interface. Web links to customized graphical views can be generated by users and subsequently inserted in manuscripts reporting novel findings, such as discovery notes. The tool also enables browsing of a single gene across projects, which can provide new perspectives on the role of a given molecule across biological systems. This curated dataset collection is available at: http://hiv.gxbsidra.org/dm3/geneBrowser/list.

Keywords

Transcriptomics, Bioinformatics, Software, HIV, Immune Response, Big Data

This article is included in the Sidra Medicine gateway.

This article is included in the Data: Use and Reuse collection.

Open Peer Review

Referee Status: ✔ ✔ ✔

Invited Referees

version 1
published 11 Mar 2016

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2 Nicolas Chomont, Université de Montréal, Canada
3 José Alcamí Pertejo, Instituto de Salud Carlos III, Spain
Francisco Diez-Fuertes, Instituto de Salud Carlos III, Spain

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Comments (0)
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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Introduction

Uncovering the gene transcription signature associated with different outcomes of HIV infection is paramount to a deeper understanding of HIV pathogenesis and to identifying potential therapeutic targets for improving immunological response and for eradicating HIV infection. HIV has a complex life cycle during which it engages multiple host cellular components, including the immune cells in which it replicates, undermining immune functions. It also hijacks host transcription factors and enzymes to assure viral production and subsequent infections. HIV dysregulates host genes resulting in aberrant immune response, disease progression, and opportunistic infections. The ability to pool and analyze samples across various groups of HIV infected individuals with different disease outcomes and across various cell types or tissues, offers a unique opportunity to define common denominators of the immune control of HIV infection, the regulation of HIV replication, and/or the virus-host interaction. With this in mind, we make available, via an interactive web application, a curated collection of transcriptome datasets relevant to HIV infection.

With over 65,000 studies deposited in the NCBI Gene Expression Omnibus (GEO), a public repository of transcriptome profiles, the identification of datasets relevant to a particular research area is not straightforward. Furthermore, GEO is primarily designed as a repository for storing data, rather than for browsing and interacting with the data. Thus, we used a custom web application, the gene expression browser (GXBX), to host a collection of datasets that we identified as particularly relevant to the study of the immunobiology of HIV infection. This tool has been described in detail and the source code released as part of a recent publication. It allows seamless browsing and interactive visualization of large volumes of heterogeneous data. Users can easily customize data plots by adding multiple layers of information, modifying the sample order and generating links that capture these settings and can be inserted in email communications or in publications. Accessing the tool via these links also provides access to rich contextual information essential for data interpretation. This includes for instance access to gene information and relevant literature, study design, and detailed sample information.

Material and methods

Identification of relevant datasets

Potentially relevant datasets deposited in GEO were identified using an advanced query based on the Bioconductor package GEOmetadb, version 1.30.0, and on the SQLite database that captures detailed information on GEO data structure (https://www.bioconductor.org/packages/release/bioc/html/GEOmetadb.html). The search query was designed to retrieve entries where the title or summary contained the word HIV, and were generated from human samples using Illumina or Affymetrix commercial platforms.

The relevance of each entry returned by this query was assessed individually. This process involved reading through the descriptions and examining the list of available samples and their annotations. Sometimes it was also necessary to review the original published report in which the design of the study and generation of the dataset are described in more details. We identified 87 datasets meeting the search criteria and containing HIV infected samples (some studies related to HIV problematics contained uninfected samples only). Out of the 87 datasets, 41 were generated from tissues or cells isolated from HIV infected individuals, 46 contained cell lines or primary cells infected in vitro. Since molecular, cellular and physiological processes involved in the context of in vivo and in vitro infections are dramatically different, we decided to create two separate collections. Here we describe the “in vivo collection” composed of 34 curated datasets (after filtering out datasets that did not meet quality control criteria, as described in “Dataset Validation” section, or datasets generated using an unsupported array platform). Of the 34 datasets, 7 are from whole blood, 7 from peripheral blood mononuclear cells (PBMCs), 8 from CD4+ and/or CD8+ T-cells, 4 from monocytes, 1 from dendritic cells (DCs), and 7 from tissues different from blood (Figure 1). Four datasets comprise samples from patients co-infected with tuberculosis (TB), one dataset comprises samples from AIDS related lymphomas, and four datasets addressed HIV infected patients with neurological disorders, such as HIV related fatigue syndrome, major depression disorder (MDD), or HIV-Associated Neurocognitive Disorder (HAND). Among the many noteworthy datasets, several stood out, such as the extensive

![Figure 1](https://www.bioconductor.org/packages/release/bioc/html/GEOmetadb.html)
Table 1. List of datasets constituting the collection, also available at http://hiv.gxbsidra.org/dm3/geneBrowser/list.

<table>
<thead>
<tr>
<th>Title</th>
<th>Platform</th>
<th>Number of samples</th>
<th>Sample source</th>
<th>Validation genes</th>
<th>GEO ID</th>
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<tr>
<td>Blood Transcriptional Signature of hyperinflammation in HIV-associated Tuberculosis</td>
<td>Illumina HG-U133A</td>
<td>187</td>
<td>Whole blood</td>
<td>N/A</td>
<td>GSE84111</td>
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<td>CD4+ T Cell Decline is Predicted by Differential Expression of Genes in HIV seropositive patients</td>
<td>Affymetrix HG-U133A</td>
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<td>CD4+ T cell gene expression in virologically suppressed HIV-infected patients during Maraviroc intensification therapy</td>
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<td>Chronic CD4+ T cell Activation and Depletion in HIV-1 Infection: Type I Interferon-Mediated Disruption of T Cell Dynamic</td>
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<td>Comparison of gene expression profiles of HIV-specific CD8 T cells from controllers and progressors</td>
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<td>CD8, CD4-neg</td>
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<td>Comparison of transcriptional profiles of CD4+ and CD8+ T cells from HIV-infected patients and uninfected control group</td>
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<td>CD4, CD8</td>
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<td>Differential Gene Expression in HIV-Infected Individuals Following ART</td>
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<td>Differential Gene Expression of Soluble CD8+ T-cell mediated suppression of HIV replication in three older children</td>
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<td>Expression data from HIV exposed and uninfected women</td>
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<td>Genome-wide mRNA expression correlates of viral control in CD4+ T cells from HIV-1 infected individuals</td>
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<td>GSE18233</td>
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<td>Genome-wide transcriptional profiling of HIV positive and negative children with active tuberculosis, latent TB infection and other diseases</td>
<td>Illumina HumanHT-12 v4</td>
<td>491</td>
<td>Whole blood</td>
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<td>GSE39941 (GSE39939 + GSE39940)</td>
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<td>Genome-wide analysis of gene expression in whole blood from HIV-1 progressors and non-progressors</td>
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<td>Whole blood</td>
<td>N/A</td>
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<td>Genome-wide transcriptional profiling of HIV positive and negative adults with active tuberculosis, latent TB infection and other diseases - GSE37250_famil</td>
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<td>537</td>
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<td>HIV-1 infection in human PBMCs in vivo</td>
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<td>Inflammation and macrophage activation in adipose tissue of HIV-infected patients under antiretroviral treatment</td>
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<td>Adipose tissue</td>
<td>ADIPOQ</td>
<td>GSE19811</td>
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<td>Longitudinal comparison of monocytes from an HIV viremic vs avirrmeic state</td>
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<td>16</td>
<td>Monocytes</td>
<td>CD14</td>
<td>GSE5220</td>
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<td>Microarray Analysis of Lymphatic Tissue Reveals Stage-Specific, Gene-Expression Signatures in HIV-1 Infection</td>
<td>Affymetrix HG-U133_Plus_2</td>
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<td>Lymph node</td>
<td>XIST</td>
<td>GSE16363</td>
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<td>Molecular Classification of AIDS-Related Lymphomas</td>
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<td>17</td>
<td>Tissues</td>
<td>XIST</td>
<td>GSE17189</td>
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<td>The National NeuroAIDS Tissue Consortium Brain Gene Array: Two types of HIV-associated neurocognitive impairment</td>
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<td>Brain</td>
<td>XIST</td>
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<td>The Relationship between Virus Replication and Host Gene Expression in Lymphatic Tissue during HIV-1 Infection</td>
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<td>Lymph node</td>
<td>XIST</td>
<td>GSE21589</td>
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<td>Transcriptional profiling of CD4 T-cells in HIV-1 infected patients</td>
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<td>CD4+ T cells</td>
<td>CD3, CD4</td>
<td>GSE23879</td>
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<td>Transcriptome analysis of HIV-infected peripheral blood monocytes</td>
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<td>Monocytes</td>
<td>CD14</td>
<td>GSE50011</td>
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<td>Transcriptome analysis of primary monocytes from HIV+ patients with differential responses to therapy</td>
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<td>Monocytes</td>
<td>CD14</td>
<td>GSE52900</td>
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<td>Whole Blood Transcriptional Response to Early Acute HIV -GPL10558</td>
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<td>Whole blood</td>
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<td>GSE29429</td>
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<td>Whole blood</td>
<td>XIST</td>
<td>GSE29429</td>
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</table>

**Figure 2.** Thematic composition of the dataset collection. Word frequencies extracted from titles of the studies loaded into the GXB tool are depicted as a word cloud. The size of the word is proportional to its frequency.
Gene expression browser (GXB) – dataset upload and annotation

Once a final selection had been made, each dataset was downloaded from GEO as a Simple Omnibus Format in Text (SOFIT) file. It was in turn uploaded on a dedicated instance of the GXB, an interactive web application developed at the Benaroya Research Institute, hosted on the Amazon Web Services cloud. Available sample and study information were also uploaded. Samples were grouped according to possible interpretations of study results and gene rankings were computed based on different group comparisons (e.g. comparing samples form HIV negative vs HIV positive patients, with or without antiretroviral therapy, in different stages of disease progression, or with or without co-infection, depending on the focus of respective studies).

GXB – short tutorial

The GXB software has been described in detail in a recent publication. This custom software interface provides users with a means to easily navigate and filter the dataset collection available at http://hiv.gxbsidra.org/dm3/geneBrowser/list. A web tutorial is also available online: https://gxb.benaroyaresearch.org/dm3/tutorials.gsp#.gxbtut. Briefly, datasets of interest can be quickly identified either by filtering on criteria from pre-defined lists on the left side of the dataset navigation page, or by entering a query term in the search box at the top of the dataset navigation page. Clicking on one of the studies listed in the dataset navigation page opens a viewer designed to provide interactive browsing and graphic representations of large-scale data in an interpretable format. This interface is designed to present ranked gene lists and to display expression results graphically in a context-rich environment. Selecting a gene from the rank-ordered list on the left of the data-viewing interface will display its expression values graphically in the screen’s central panel. Directly above the graphical display, drop down menus give users the ability: a) To change the rank list by selecting different comparisons (in cases where the dataset is split in more than two groups), or to only include genes that are selected for specific biological interest. b) To change sample grouping (Group Set button); in some datasets, user can switch between interpretations where samples are grouped based on cell type or disease, for example. c) To sort individual samples within a group based on associated categorical or continuous variables (e.g. gender or age). d) To toggle between a bar plot view and a box plot view, with expression values represented as a single point for each sample. Samples are split into the same groups whether displayed as a bar plot or a box plot. e) To provide a color legend for the sample groups. f) To select categorical information to be overlaid at the bottom of the graph. For example, the user can display gender or smoking status in this manner. g) To provide a color legend for the categorical information overlaid at the bottom of the graph. h) To download the graph as a portable network graphics (png) image or the table with expression values as a comma separated values (csv) file. Measurements have no intrinsic utility in absence of contextual information. It is this contextual information that makes the results of a study or experiment interpretable. It is therefore important to capture, integrate and display information that will give users the ability to interpret data and gain new insights from it. We have organized this information under different tabs directly above the graphical display. The tabs can be hidden to make more room for displaying the data plots, or revealed by clicking on the blue “hide/show info panel” button on the top right corner of the display. Information about the gene selected from the list on the left side of the display is available under the “Gene” tab. Information about the study is available under the “Study” tab. Information available about individual samples is provided under the “Sample” tab. Rolling the mouse cursor over a bar plot, while displaying the “Sample” tab, lists any clinical, demographic, or laboratory information available for the selected sample. Finally, the “Downloads” tab allows advanced users to retrieve the original dataset for analysis outside this tool. It also provides all available sample annotation data for use alongside the expression data in third party analysis software. Other functionalities are provided under the “Tools” drop-down menu located in the top right corner of the user interface. These functionalities include notably: a) “Annotations”, which provides access to all the ancillary information about the study, samples and the dataset, organized across different tabs; b) “Cross Project View”, which provides the ability to browse across all available studies for a given gene; c) “Copy Link”, which generates a mini-URL encapsulating information about the display settings in use and that can be saved and shared with others (clicking on the envelope icon on the toolbar inserts the url in an email message via the local email client); and d) “Chart Options”, which gives user the option to customize chart labels.

Dataset validation

Quality control checks were performed by examination of profiles of relevant biological markers. Known leukocyte surface markers were used to verify consistency of the information provided by dataset depositors, and to identify instances where contamination of samples by other leukocyte populations may be confounding. The markers that were used include: CD3 (CD3D), a T-cell marker; CD4 and CD8 (CD8A), markers of CD4+ and CD8+ T cells respectively; CD11c (ITGAX), an mDC marker; CD14, expressed by monocytes and macrophages; or Adiponectin (ADIPOQ), expressed in adipose tissue. Expression of the XIST transcripts, which expression is gender-specific, was also examined in datasets containing relevant information, to determine its concordance with demographic information provided with the GEO submission (respective links in Table 1).

Data availability

All datasets included in our curated collection are also available publically via the NCBI GEO website: www.ncbi.gov/geo; and are referenced throughout the manuscript by their GEO accession numbers (e.g. GSE44228). Signal files and sample description files can also be downloaded from the GXB tool under the “downloads” tab.
Author contributions
JB and DC conceived the theme for this dataset collection. DC, SP and CQ designed the software. SP, CQ and SB installed and tested the software and programmed portions of the web application. SB uploaded datasets. JB curated and annotated datasets. JB and DC prepared the first draft of the manuscript. All authors were involved in the revision of the draft manuscript and have agreed to the final content.

Competition interests
No competing interests were disclosed.

Grant information
JB, SB and DC were supported by the Qatar Foundation.

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

Acknowledgments
We would like to thank all the investigators who decided to make their datasets publicly available by depositing them in GEO.

References


Data Source
Open Peer Review

Current Referee Status: ✔️ ✔️ ✔️

Version 1

Referee Report 20 April 2016
doi:10.5256/f1000research.8824.r12871

José Alcamí Pertejo 1, Francisco Diez-Fuertes 2
1 Centro Nacional de Microbiologia, Instituto de Salud Carlos III, Majadahonda, Madrid, Spain
2 Instituto de Salud Carlos III, Majadahonda, Madrid, Spain

Blazkova et al. describe an interactive web application that includes 34 different transcriptome datasets. This open tool facilitates access to transcriptome analysis in the HIV field allowing meta-analyses on transcriptomic changes in HIV infection.

As strengths of the article I will highlight:
- The application is friendly and easy to use and allowed us to compare our results with a large collection of databases in a comprehensive way.
- The software allows searches related with a particular gene and how its expression is modified in different scenarios (infected vs non-infected, long term non-progressors vs typical progressors, treated vs untreated).
- The cellular types in which dataset have been obtained are indicated.
- Datasets included have been selected according to their interest and high methodological standards. For example, when contamination with cell types different from those initially targeted are detected the studies are not considered for the final dataset thus enhancing the quality of the results.

I would propose some suggestions to improve this interesting tool:
- All the studies were performed with microarrays. It would be important to discuss if the inclusion of data using RNA-seq approaches and the current units used in these studies (FPKMs, RPKMs, TPMs) could be incorporated in the future.
- It should be clarified if the results among the different studies are normalized or just described with the units used in each study. If data normalization has been performed it would important to describe how it was done.

Overall it represents an important effort that can be useful for many researchers working in the field of HIV genetics and pathogenesis.

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 15 May 2016

Jana Blazkova, Sidra Medical and Research Center, Qatar

Thank you for your positive review and valuable feedback.

1. We are working on including RNA-seq data (see answer to a similar comment made by Amalio Telenti).

2. The data are not normalized among the different studies, but there is a normalization for individual studies. We are uploading raw or background subtracted data (based on the input in GEO), we then floor the data (give it a minimum value of 10, if it is below 10), and perform quantile normalization. In case that only normalized data are available, we present them as they are.

Competing Interests: No competing interests were disclosed.

Referee Report 15 April 2016

doi:10.5256/f1000research.8824.r13356

Nicolas Chomont
Department of Microbiology, Infectiology and Immunology, Université de Montréal, Montreal, QC, Canada

In this interesting article, Blazkova and colleagues describe the development of an interactive web application that allows HIV researchers to access a collection of transcriptome datasets relevant to HIV infection. The collection includes 34 datasets generated with human samples that have been carefully selected based on their relevance and quality control checks.

This is a very useful tool that can be easily used by non-experts in transcriptomics analyses. I have used it and I am convinced that it potentially represents an important contribution to the work performed by HIV researchers. I tested the accuracy of the tool (not in a formal way) by examining differences in the expression for several genes that are well-known to be modulated by HIV infection. The results are clearly presented and can be easily exported to be included in presentations/publications.

A few suggestions: I anticipate that the database will be updated on a regular basis. Therefore, it would be great to specify the date of the last update of the data set available online. Also, the possibility of adding RNAseq data would be important in the future. Maybe a brief description of each dataset would be useful too.

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 15 May 2016
**Jana Blazkova**, Sidra Medical and Research Center, Qatar

Thank you for your positive review and helpful suggestions.

1. That is a very good point, thank you for bringing it up. We will include the date of the last update on the webpage. F1000Research editors also encourage us to take advantage of the fact that their platform supports versioning. Also we will be able to update the table and list of studies as they become available. It is not clear whether this would require an additional round of review but we do not anticipate updates to be made more than once or twice yearly.

2. We are working on including RNA-seq data (see answer to a similar comment made by Amalio Telenti).

3. A brief description is included under the “Study” tab of each dataset (e.g. [https://www.youtube.com/watch?v=Te0lggbXjIY](https://www.youtube.com/watch?v=Te0lggbXjIY)). We envisage to set this view as a default, when opening individual studies.

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

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The article by Blazkova and colleagues constitutes an important contribution to the HIV field. It crystallizes the efforts of multiple groups that characterized the host transcriptional response to infection by providing a viewer of data that are not immediately accessible in a structured interface. I have assessed the performance of the tool, and found it intuitive and user-friendly.

It extends efforts of my group to provide facilitated access to genomic data in HIV disease ([http://www.guavah.org/](http://www.guavah.org/)).

I would bring two aspects up for discussion. First, that this tool should evolve to display RNAseq data = new generation sequencing data are increasingly available, effectively displacing microarrays. RNAseq is also easier for standardization across studies. Second, that users should be attentive to the subtleties of analysis: covariates such as gender, age, cellularity, analytical platforms and batch effects can influence expression profiles significantly. In-depth analysis may thus require downloading of original expression data.

**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.
Thank you for your positive review and valuable comments.

1. We are actually working on extending the supported platforms to high-throughput RNA sequencing. For now, a trial RNA-seq dataset concerning gene expression profiling of immune cell subsets across several diseases has been uploaded (GSE60424, https://gxb.benaroyaresearch.org/dm3/geneBrowser/show/396, http://journals.plos.org/plosone/article?id=10.1371/journal.pone.0109760). Concerning HIV immunobiology, we identified 13 datasets generated by high-throughput RNA sequencing, and only 2 of them (65 transcriptional profiles in total) were generated using cells from HIV infected individuals. Thus, not including the RNA-seq generated data doesn't have a major impact on comprehensiveness of the collection so far, nevertheless, we will definitely include this platform in the future.

2. We agree with the reviewer, that the covariates may have an influence on data interpretation; unfortunately, it should be noted that relevant information is not always available. In the case of batch information we estimate this number to be 5-10% of submissions. This is probably a point worth opening for discussion between GEO and community stakeholders. An advantage of relying on multiple studies however lies in the fact that independent validation can be obtained readily and outlier studies can be identified and further analyze for potential effect of another variable. It should also be noted that from within the GXB, the original expression data can be downloaded under the “Downloads” tab, as a SOFT format or series matrix data file (e.g. https://www.youtube.com/watch?v=TbMTht2Z2NU). The same dataset can also be directly accessed from GEO, in various formats (e.g. https://www.youtube.com/watch?v=O7RgeYD4SPs).

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