ImmuCellDB: An Indicative Database of Immune Cell Composition From Different Tissues and Disease Conditions in Mouse and Human

Ziyi Chen1,2,3, Han Na1,2 and Aiping Wu1,2*

1 Institute of Systems Medicine, Chinese Academy of Medical Sciences & Peking Union Medical College, Beijing, China, 2 Suzhou Institute of Systems Medicine, Suzhou, Suzhou, China, 3 Department of Infectious Diseases, The Second Hospital of Nanjing, The Affiliated Hospital of Nanjing University of Chinese Medicine, Nanjing, China

Immune cell composition is highly divergent across different tissues and diseases. A comprehensive resource of tissue immune cells across different conditions in mouse and human will thus provide great understanding of the immune microenvironment of many diseases. Recently, computational methods for estimating immune cell abundance from tissue transcriptome data have been developed and are now widely used. Using these computational tools, large-scale estimation of immune cell composition across tissues and conditions should be possible using gene expression data collected from public databases. In total, 266 tissue types and 706 disease types in humans, as well as 143 tissue types and 61 disease types, and 206 genotypes in mouse had been included in a database we have named ImmuCellDB (http://wap-lab.org:3200/ImmuCellDB/). In ImmuCellDB, users can search and browse immune cell proportions based on tissues, disease or genotype in mouse or humans. Additionally, the variation and correlation of immune cell abundance and gene expression level between different conditions can be compared and viewed in this database. We believe that ImmuCellDB provides not only an indicative view of tissue-dependent or disease-dependent immune cell profiles, but also represents an easy way to pre-determine immune cell abundance and gene expression profiles for specific situations.

Keywords: immune cell, deconvolution, human, mouse, transcriptome, database

INTRODUCTION

Tissues infiltrating immune cells have long been recognized as important regulators in both healthy and disease conditions. In response to different stimuli, normal and abnormal immune reactions may be produced by the immune system. For instance, autoimmune diseases can occur when the immune reactions targeting our body are too strong, whereas tumors can be established when immune responses to malignant cells are too weak. When fighting external pathogens, inflammation or infection can occur depending on the magnitude and duration of immune responses (https://www.budandtender.com/blogs/bud-tender-blog/your-endocannabinoid-and-immune-system). In addition to local immune responses, systematic multi-organ immune responses frequently happen
in many diseases. Immune states in multiple irrespective areas can also be reshaped by some cytokines, metabolites, etc., that are transported by the circulatory system (1). Therefore, knowledge of the constitution of tissue immune cells under different conditions should greatly enhance our understanding of their roles.

Usually, tissue immune cell abundance is measured using well-known methods including flow cytometry (2), immunochemistry (3), etc. However, these experimental-based procedures are usually conducted in a laboratory and are time-consuming when batch processing many biological samples. Additionally, cell type-specific markers and corresponding antibodies are not readily available in many circumstances. Although some public databases of flow cytometry data like Immport (4) or FlowRepository (5) offer users access to download experimental data corresponding to a specific study, the number of tissue and disease categories is still small and may restrict researchers from querying tissue immune cell abundances they are interested in. Recently, with the advancement of high-throughput transcriptome measuring technologies, multiple computational tools have already been designed and used to study the abundance of tissue immune cells in terms of omics data, including DNA microarrays, RNA-seq, and DNA methylation, etc. (6, 7) The suitable performance of these computational-based methods has been validated in multiple studies. Compared to an experimental based strategy, tissue immune cell composition can be rapidly estimated from genomics data. Additionally, tissue transcriptome data from most tissue and disease types has already been deposited into some public database like Gene Expression Omnibus (GEO) (8). These represent a great resource for researchers for transcriptome data under different conditions (9, 10).

However, there are still no available web database search engine for users to query the differences in abundance of tissue immune cells between different tissue and disease types. With tissue expression data accumulated in GEO, an in-depth knowledge of the inner immune cell constitution allows easy prediction from tissue expression data. Therefore, predicting the composition of tissue immune cells from tissue transcriptome data should greatly accelerate our understanding of the dynamics of immune cells in different situations. Furthermore, a user-friendly database interface covering all possible tissue and disease combinations will be convenient for those seeking to view the landscape of tissue immune cell abundance under different situations.

Thus, by mining the transcriptome data profiles from microarray data from both humans and mice in GEO, transcriptome data including nearly 266 tissue types and 706 disease types in humans and 143 tissue types, 61 disease types, and 206 genotypes from mouse were collected. Next, using immune cell estimation tools developed for human and mouse, the composition of 21 immune cells was calculated. By integrating resources for immune cell composition and gene expression across different tissue and disease conditions into this database, users can browse, search, and view data in accordance with the selection items provided. This database represents an indicative resource for tissue immune cell compositions over tissue types, diseases, and genotypes, and can provide a convenient querying tool for other researchers.

MATERIALS AND METHODS

Data Collection and Preprocessing

ImmuCellDB aims to provide an indicative landscape of tissue immune cell composition across different tissue and disease conditions (Figure 1A). All human and mouse transcriptome data were mined from GEO with the R package “GEOmetadb” (11). The cell lines, Image result for flow cytometry sorted or magnetic bead enriched cell types, and some synthetic mixtures were filtered out based on their characteristics, title, and source name for each Sample accessions numbers (GSM). Finally, a summary table including GSM IDs, GSE IDs, Platform name, and sample description content (the title, source name and characteristics of GSM and the summary and the overall design part of GSE for each sample) were tabulated (Supplementary Figure 1).

Then, the characteristics for each sample including tissue type, disease type, and genotype were manually extracted from the sample description contents. As there were many different forms of names for some tissues, diseases, and genotypes, samples under the same conditions may have been misclassified into different groups. Therefore, a standardized disease name was helpful for avoiding confusions made by different disease nomenclature. Here, all disease types collected from GEO were manually retrieved from International Classification of Diseases version 11 (ICD-11) and unified according to the disease classification rules in this database.

With respect to all these selected samples, the raw CEL files were downloaded and samples profiled on the sample microarray platforms were processed with the same pipeline (Figure 1B). The probsets for each microarray platform were annotated with the help of some custom CDF packages (http://brainarray.mbnimed.umich.edu/Brainarray/Database/CustomCDF/). All scripts used to process raw data can be found on Github (https://github.com/wuaipinglab/ImmuCC/tree/master/ImmuCellDB/). Then, expression matrices for different microarray platforms were merged into one large matrix according to the common genes shared by them. To ensure that the expression levels were distributed in a similar range, a quantile normalization method was used for each dataset. In addition, gene IDs from mouse expression data were converted into the orthologous human gene IDs.

Immune Cell Proportion Estimation

Using processed gene expression data, tissue immune cell compositions across all different conditions could be calculated with some previously reported deconvolution tools designed specifically for microarray data (Figure 1B) (6). For human samples, the relative proportion of 22 immune cells was estimated with CIBERSORT (9). With respect to the mouse transcriptome data, the composition of 25 immune cells was predicted with our own model array_ImmuCC (10). To help the
users view the marker genes selected for each immune cell, a matrix for the relationship between each immune cell and all signature genes was available at the Supplementary Table 1. The signature genes for human immune cells were directly downloaded from the Supplementary Table 1 of CIBERSORT (https://static-content.springer.com/esm/art%3A10.1038%2Fnmeth.3337/MediaObjects/41592_2015_BFnmeth3337_MOESM207_ESM.xls) (9). The signature genes for mouse immune cells were derived from our previous research (10). Samples with P values > 0.05 were discarded as this may reflect a lower immune cell infiltration (12).

Categorization of Different Tissue Diseases and Immune Cells
To facilitate viewing of results, a tree browser for different tissue and disease types was constructed. First, all collected tissue

---

**FIGURE 1** | Schematics for ImmuCellDB. (A) A brief introduction for the relationship of tissues and diseases in the database. (B) The workflow for the construction of database. Modules for data collection, data preprocessing, immune cell estimation and database construction are shown. (C) A pie plot for the relative proportion of species and top 10 most popular tissues and diseases. (D) A screenshot for the homepage of ImmuCellDB. The home page was consisted of four parts that is the introduction for the database, statistics of sample data, the worldwide distribution for all users to this database and a body map of human and mouse organs.
names were unified according to their anatomical location. The tissues were then hierarchically organized into three levels. Next, to keep the consistency across all disease types, name unification was also manually performed for all disease names. The corrected disease names were then manually searched from ICD-11 (https://icd.who.int/browse11/l-m/en). Both the disease classification in ICD-11 and the GEO characteristics of data were taken into consideration, and all samples were then grouped into a corresponding disease tree. Firstly, since the disease names of samples provided in GEO were not the standard names, it is hard for us to directly put them into the correct positions in the ICD disease classification tree. At first, we had unified the disease names with reference to ICD-11 and prior knowledge. For a disease with multiple disease names, these disease names were replaced with the correspondent disease name listed in ICD-11. For instance, both Breast carcinoma, Mammary gland tumor and Breast tumor, Breast cancer were grouped into Breast cancer. For some disease names used in GEO while not existed in ICD-11, the original disease names from GEO were kept. For instance, the prostate cancer was classified as Adenocarcinoma of prostate (2C82.0), Other specified malignant neoplasms of prostate (2C82.Y) and Malignant neoplasms of prostate, unspecified (2C82.z) in ICD-11. In ImmuCellDB, the prostate cancer was grouped as Adenocarcinoma of prostate, prostate cancer and non-metastatistic prostate cancer. Except for Adenocarcinoma of prostate, the other two diseases were not described in ICD-11 which were kept in ImmuCellDB. Another question is that too many hierarchical branches of diseases are not convenient for searching. To address this issue, the disease classification tree provided in our database was only consisted of three levels. We classified all diseases into groups according to the top-level of the disease classification tree in ICD-11. For instance, for certain infectious or parasitic diseases, we put all diseases including HIV infection, Leprosy, Influenza infection into this group. Then, diseases were merged into the second level according to prior knowledge. With respect to HIV infection, the group was finally consisted of multiple disease names including HIV infection, HIV-1 infection, HIV-1 subtype C infection, etc. Finally, to ensure the comparability of the immune cell types predicted between these two species, immune cell types calculated with these two computational models were unified according to their location in our immune cell differentiation tree. Specifically, they were aggregated in the following ways. For human immune proportion data, Mast cells = Mast cells resting + Mast cells activated, and T cells CD4 activated = T cells CD4 memory activated. For mouse immune proportion data, T cells CD8 = T cells CD8+ + T cells CD8 + T cells CD8, T cells CD4 activated = Th1 Cells + Th2 Cells + Th17 Cells, and Dendritic cells resting = Dendritic cells immature. Finally, immune cells predicted by these two models were merged into 21 types including: Mast cells, Neutrophils, Eosinophils, Monocytes, Macrophages M0, Macrophages M1, Macrophages M2, Dendritic cells activated, Dendritic cells resting, B cells memory, B cells naive, Plasma cells, T cells CD4 naive, T cells CD4 memory, T cells follicular helper, Tregs, T cells CD4 activated, T cells CD8, Gamma delta T cells, NK cells resting, and NK cells activated.

Web Server Construction
ImmuCellDB was implemented using the MongoDB system. The metadata for species, tissues, diseases or genotypes from each sample were stored in the MongoDB database with a unique key ID. The web interface of this database was constructed based on the Node.js framework, which comes with an Application Programming Interface of the MongoDB built on top of JavaScript libraries including jQuery, Bootstrap, ECharts, Highcharts and additional plugins. All figures and tables were generated with R scripts.

Data Access
To enable quick access of the data deposited in this database, all transcriptome and immune cell composition data are also available at Figshare (https://figshare.com/articles/dataset/ImmuCellDB/13546529). All transcriptome data used in ImmuCellDB were collected from public database and will not be used for other commercial purpose.

RESULTS
Database Summary
ImmuCellDB is a database for tissue immune cell composition and gene expression in nearly 266 tissue types and 706 disease types in humans and 143 tissue types, 61 disease types, and 206 genotypes in mouse (Figure 1C). According to the expression data collected from GEO, the abundance of 21 different types of immune cells was inferred from the transcriptome data of tissues using deconvolution tools. To reduce the display space, tissue types and disease types were categorized into some major groups. ImmuCellDB contains five major modules: home, search, analysis, tools and tutorial (Figure 1D). From the home module, users can view the introduction for this database, the worldwide distribution of users and the sample number for each tissue. By clicking each tissue in the body of human and mouse, all samples belong to this tissue will be selected. The search module gives an access to tissue immune cell compositions over certain tissue types, disease types or genotypes. The analysis module is consisted of four tools: ImmDiff, ExpresDiff, Correlation and NetCor. The variation and correlation of immune cell abundance and gene expression level between different conditions can be compared and viewed with these tools. An overview for the functions and results of the search module and analysis module were indicated at Supplementary Figure 2. The tools module allows users to estimate the composition of tissue immune cell with three our previously reported tools: Array_ImmuCC (10), Seq_ImmuCC (13) and Tissue_ImmuCC (14). The tutorial module provides an introduction for search, analysis and tools. Also, a worked example of how to find a valuable information from this database was also available at the Supplementary Materials.

Search Module
ImmuCellDB provides a tree browser for users to query for tissue immune cell compositions over certain tissue types, disease types
or genotypes using the search module. As indicated in Figure 2A, by sequentially selecting the species, tissue type, disease type or genotype from the drop-down box, two figure types plus a table will be returned to the result page. From the image figure box, a box plot for the relative proportion of all 21 immune cells in each condition was generated (Figure 2B). Next, to illustrate the variation for each immune cell over different conditions, the proportional distribution of each immune cell across all selected conditions was displayed with a boxplot (Figure 2C). In some situations, users may want to integrate this retrieved information with other data. To meet these special requirements, the table consisted of the sample information including GSM IDs, tissue types, disease types or genotypes, and the abundance of 21 immune cells. Users can download all these figures and tables directly from this result page.

Analysis Module

ImmDiff Tool

Many factors including species, tissue types, disease types, or genotypes may have an impact on the context of tissue immune cells. To evaluate the differences in tissue immune cell levels, the ImmDiff tool in the Analysis function was designed to compare the immune cell proportion between two specified condition groups. Based on the two groups selected from the selection box covering species, tissue type, disease type, or genotype, a box plot was used to show the distribution of all immune cells between these two groups (Figure 3A). Moreover, a comparison between these two groups was carried out with a t-test, and the immune cells with significant differences in proportion were marked with asterisks. Furthermore, users can compare their own data with the samples collected in this database using the “Comparing with uploaded data” module. The same result format will be generated under this selection. Based on this analysis, immune cells with different abundance can be successfully identified.

ExpresDiff Tool

Gene expression levels also vary across different species, tissue types, disease types, and genotypes. Knowledge of the differentially expressed genes under different conditions is useful for understanding the divergence of transcriptome levels. The ExpresDiff tool developed here allows users to identify the gene expression changes between different condition groups. After setting two biological conditions in the selection box, the gene expression difference between these two groups could be obtained. The differential expression analysis was conducted using a confect ranking method with the “topconfects” package and ranked according to the confident effect size (15). Users can download the whole table directly from the result page. Then, a plot representing the fold change and averaged expression level of genes with the highest effect size was generated (Figure 3B). Additionally, to investigate the expression pattern of the most differentially expressed genes, the expression of the top 50 genes over all samples belonging to these two groups was exhibited using a heatmap. Similar to the ImmDiff tool, users can also compare their own data with the samples deposited in this database using the “Comparing with uploaded data” module. It should be noted that comparison of...
the gene expression differences between human and mouse is not suggested here.

Correlation Tool
The correlation between immune cell abundance and gene expression values may reflect a possible direct or indirect relationship between genes and immune cells. For each cell type, the significantly correlated genes may relate to their character, such as differentiation stage and functional roles. For instance, a positive correlation between T cells CD8 and PD1 was associated with the dysfunction of CD8 T cells (16). The Correlation tool designed here offers access for users to easily view the correlation between genes and immune cells. With samples in the selected species, tissue types, disease types, cell types, and gene symbols, the Spearman correlation between gene expression level and immune cell abundance was calculated. A dot plot was used to illustrate the correlation between each immune cell and gene pairs across all selected samples (Figure 3C). A linear fitting was also added to the dot plot to help interpret the tendency across all data sets.

NetCor Tool
With the Correlation tool described above, analysis of the correlation between a small amount of genes and immune cell pairs is also possible. However, a comprehensive picture of the correlation between immune cell abundance and gene expression values can provide an important resource for their inner relationship in distinct tissue microenvironments. The NetCor tool makes this analysis possible by integrating all correlated immune cell and gene pairs into a table. For all selected data sets, the Spearman correlation between the expression level of each gene and the abundance of each immune cell was calculated. Finally, all immune cell types and gene pairs whose correlation efficiency exceeds 0.7 were returned to the result page. The results are displayed in a tabular format, with each row representing a correlated immune cell type and gene pairs, as well as a correlation coefficient.

DISCUSSION
We report here the ImmuCellDB database, which provides access to the immune cell proportion and gene expression data from over 266 tissue types and 706 disease types in human and 143 tissue types, 61 disease types, and 206 genotypes in mouse. Utilizing the results estimated from tissue transcriptome data, ImmuCellDB provides a collective resource for tissue immune cell abundances and gene expression profiles from different conditions.

Using ImmuCellDB, the distribution of the relative proportion of 21 immune cells in each tissue and disease combinations, and the variation in the distribution of each immune cell over different tissue and disease groups can be viewed with the Search module. Limited by the size of the web page, only the first 10 selected tissue and disease groups are used to generate a plot when the number of tissue and disease combinations is larger than 10. In addition to searching from the database, tissue immune cell abundance and gene expression file differences between different tissues and disease conditions can be calculated. The immune cells with different abundance and the genes with significantly different expression levels between these two groups can be calculated with ImmDiff and ExpresDiff. This information will support an understanding of the different immune reaction pathways and gene transcription profiles between any two selected conditions. Besides an analysis of immune cells and genes separately, the correlation between any immune cell and gene pairs can also be defined according to the Spearman correlation between cell abundance and gene expression level. With the Correlation and NetCor tools, users can obtain the correlation profile of immune cells and genes under any specified conditions. It should
be noted that a high correlation coefficient was not equal to a participation of a gene in a cell’s function.

However, all datasets used in this database were derived from DNA microarray-based transcriptome platforms because more immune cell types were included in computational tools designed specifically for microarray data. Additionally, multiple methods have been developed to get the abundance of tissue immune cells from other omics data, including bulk RNA-seq (13), bisulfite sequencing (17), and ATAC-seq (18). Leveraging these sequencing-based data resources, we can estimate the immune cell counts from other bulk tissue omics data. In addition to bulk studies, some single cell detecting methods such as single cell RNA-seq (19) and mass cytometry (20) can also be used to determine cell amounts. By assigning each cell to the correspondent cell type, the proportion of each cell population can be successfully enumerated (21). Therefore, further integrating the immune cell abundance estimated from multi-level omics data into this database should expand the application of ImmuCellDB to additional tissues and disease categories.

Altogether, ImmuCellDB is a valuable resource for researchers to study the immune cell abundance and gene expression levels of tissues in different conditions.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

ZC and AW conceived and designed the study. ZC and AW analyzed the data and results. HN contributed to the discussion and analysis of the studies. ZC and AW wrote the paper. All authors contributed to the article and approved the submitted version.

REFERENCES

1. Tisoncik-Jo GJ, Gasper DJ, Kyle JE, Eifeld AJ, Selinger C, Hatta M, et al. Integrated Omics Analysis of Pathogenic Host Responses During Pandemic H1N1 Influenza Virus Infection: The Crucial Role of Lipid Metabolism. Cell Host Microbe (2016) 19(2):254–66. doi: 10.1016/j.chom.2016.01.002
2. Cabral-Marques O, Schimke LF, de Oliveira EB Jr, Al Khawanky N, Ramos RN, Al-Ramadi BK, et al. Flow Cytometry Contributions for the Diagnosis and Immunopathological Characterization of Primary Immunodeficiency Diseases With Immune Dysregulation. Front Immunol (2019) 10:2742. doi: 10.3389/fimmu.2019.02742
3. Duraiyan J, Govindarajan R, Kaliyappan K, Palanisamy M. Applications of Immunohistochemistry. J Pharm Bioallied Sci (2012) 4(Suppl 2):S307–9. doi: 10.4103/0975-7406.100281
4. Bhattacharya S, Dunn P, Thomas CG, Smith B, Schaefer H, Chen J, et al. ImmPort: Toward Repurposing of Open Access Immunological Assay Data for Translational and Clinical Research. Sci Data (2018) 5:180015. doi: 10.1038/sdata.2018.15
5. Spillen J, Breuer K, Rosenberg C, Kotecha N, Brinkman RR. FlowRepository: A Resource of Annotated Flow Cytometry Datasets Associated With Peer-Reviewed Publications. Cytometry A (2012) 81(9):727–31. doi: 10.1002/cyto.a.22106
6. Chen Z, Wu A. Progress and Challenge for Computational Quantification of Tissue Immune Cells. Brief Bioinform (2021). doi: 10.1093/bib/bbaa358
7. Avila Cobos F, Vandesompele J, Mestdagh P, De Preter K. Computational Deconvolution of Transcriptomics Data From Mixed Cell Populations. Bioinformatics (2018) 34(11):1969–79. doi: 10.1093/bioinformatics/bty019
8. Barrett T, Wilhite SE, Lefoud P, Evangelista C, Kim IF, Tomashevsky M, et al. NCBI GEO: Archive for Functional Genomics Data Sets–Update. Nucleic Acids Res (2013) 41(Database issue):D991–5. doi: 10.1093/nar/gks1193
9. Newman AM, Liu CL, Green MR, Gentles AJ, Feng W, Xu Y, et al. Robust Enumeration of Cell Subsets From Tissue Expression Profiles. Nat Methods (2015) 12(5):453–7. doi: 10.1038/nmeth.3337
10. Chen Z, Huang A, Sun J, Jiang T, Qiu FX, Wu A. Inference of Immune Cell Composition on the Expression Profiles of Mouse Tissue. Sci Rep (2017) 7:40508. doi: 10.1038/s41598-017-11773-y
11. Zhu Y, Davis S, Stephens R, Meltzer PS, Chen Y. GEOmetadb: Powerful Alternative Search Engine for the Gene Expression Omnibus. Bioinformatics (2008) 24(23):2798–800. doi: 10.1093/bioinformatics/btn529
12. Ali HR, Chilton L, Pharoah PD, Markowitz F, Calses P. Patterns of Immune Infiltration in Breast Cancer and Their Clinical Implications: A Gene-Expression-Based Retrospective Study. PLoS Med (2016) 13(12):e1002194. doi: 10.1371/journal.pmed.1002194
13. Chen Z, Quan L, Huang A, Zhao Q, Yuan Y, Yuan X, et al. Seq-ImmuCC: Cell-Centric View of Tissue Transcriptome Measuring Cellular Compositions

FUNDING

This work was supported by: 1 The National Key Plan for Scientific Research and Development of China (2016YFD0500301), 2 The CAMS Initiative for Innovative Medicine (2016-I2M-1-005), 3 The Six-talent Peaks Project in the Jiangsu Province (SWYY-169), 4 The Open Project Program of the National Laboratory of Pattern Recognition (NLPR) (201900004), 5 The Non-profit Central Research Institute Fund of Chinese Academy of Medical Sciences (2018RC310022), 6 Central Public-Interest Scientific Institution Basal Research Fund (2016ZX310195, 2017PT31026 and 2018PT31016).

ACKNOWLEDGMENTS

We would like to thank all members in Wu lab to their suggestions for the optimization of web interface.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.670070/full#supplementary-material

Supplementary Figure 1 | An overview for transcriptome datasets selection and filtering process.

Supplementary Figure 2 | An introduction for the functions and results of Search module and the Analysis module.

Supplementary Table 1 | Matrix for the relationship between each immune cell and all signature genes. In this table, the marker genes were indicated as ‘1’ and the remaining genes were indicated as ‘0’.

Supplementary Materials | Three simple work example on how to find a valuable information from this database.
of Immune Microenvironment From Mouse RNA-Seq Data. Front Immunol (2018) 9:1286. doi: 10.3389/fimmu.2018.01286
14. Chen Z, Li C, Shen Q, Liu W, Qin FX, Wu A. Tissue-Specific Deconvolution of Immune Cell Composition by Integrating Bulk and Single-Cell Transcriptomes. Bioinformatics (2020) 36(3):819–27. doi: 10.1093/bioinformatics/btz672
15. Harrison PF, Pattison AD, Powell DR, Beilharz TH. Topconfects: A Package for Confident Effect Sizes in Differential Expression Analysis Provides a More Biologically Useful Ranked Gene List. Genome Biol (2019) 20(1):67. doi: 10.1186/s13059-019-1674-7
16. McClanahan F, Riches JC, Miller S, Day WP, Kotsiou E, Neuberg D, et al. Mechanisms of PD-L1/PD-1-Mediated CD8 T-Cell Dysfunction in the Context of Aging-Related Immune Defects in the Emicro-TCL1 CLL Mouse Model. Blood (2015) 126(2):212–21. doi: 10.1182/blood-2015-02-626754
17. Onuchic V, Hartmaier RJ, Boone DN, Samuels ML, Patel RY, White WM, et al. Epigenomic Deconvolution of Breast Tumors Reveals Metabolic Coupling Between Constituent Cell Types. Cell Rep (2016) 17(8):2075–86. doi: 10.1016/j.celrep.2016.10.057
18. Li H, Sharma A, Luo K, Qin ZS, Sun X, Liu H. DeconPeaker, A Deconvolution Model to Identify Cell Types Based on Chromatin Accessibility in ATAC-Seq Data of Mixture Samples. Front Genet (2020) 11:392. doi: 10.3389/fgene.2020.00392
19. Gomes T, Teichmann SA, Talavera-Lopez C. Immunology Driven by Large-Scale Single-Cell Sequencing. Trends Immunol (2019) 40(11):1011–21. doi: 10.1016/j.it.2019.09.004
20. Hartmann FJ, Bendall SC. Immune Monitoring Using Mass Cytometry and Related High-Dimensional Imaging Approaches. Nat Rev Rheumatol (2020) 16(2):87–99. doi: 10.1038/s41584-019-0338-z
21. Abdelaal T, Michielsen L, Cats D, Hoogduin D, Mei H, Reinders MJT, et al. A Comparison of Automatic Cell Identification Methods for Single-Cell RNA Sequencing Data. Genome Biol (2019) 20(1):194. doi: 10.1186/s13059-019-1795-2

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher’s Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2021 Chen, Na and Wu. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.