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Authors

Sirota, Marina Thomas, Cristel G Liu, Rebecca <u>et al.</u>

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OPEN Data Descriptor: Enabling precision medicine in neonatology, an integrated repository for preterm birth research

Marina Sirota^{1,2}, Cristel G. Thomas³, Rebecca Liu⁴, Maya Zuhl⁵, Payal Banerjee⁵, Ronald J. Wong⁶, Cecele C. Quaintance⁶, Rite Leite⁷, Jessica Chubiz⁸, Rebecca Anderson⁹, Joanne Chappell¹⁰, Mara Kim^{11,12}, William Grobman¹³, Ge Zhang¹⁰, Antonis Rokas^{11,12}, Sarah K. England⁸, Samuel Parry⁷, Gary M. Shaw⁶, Joe Leigh Simpson⁴, Elizabeth Thomson³, Atul J. Butte^{1,2} & March of Dimes Prematurity Research Centers*

Preterm birth, or the delivery of an infant prior to 37 weeks of gestation, is a significant cause of infant morbidity and mortality. In the last decade, the advent and continued development of molecular profiling technologies has enabled researchers to generate vast amount of 'omics' data, which together with integrative computational approaches, can help refine the current knowledge about disease mechanisms, diagnostics, and therapeutics. Here we describe the March of Dimes' Database for Preterm Birth Research (http://www.immport.org/resources/mod), a unique resource that contains a variety of 'omics' datasets related to preterm birth. The database is open publicly, and as of January 2018, links 13 molecular studies with data across tens of thousands of patients from 6 measurement modalities. The data in the repository are highly diverse and include genomic, transcriptomic, immunological, and microbiome data. Relevant datasets are augmented with additional molecular characterizations of almost 25,000 biological samples from public databases. We believe our data-sharing efforts will lead to enhanced research collaborations and coordination accelerating the overall pace of discovery in preterm birth research.

¹Institute for Computational Health Sciences, University of California, San Francisco, CA 94158, USA. ²Department of Pediatrics, University of California, San Francisco, CA 94158, USA. ³Northrop Grumman Health Solutions, Rockville, MD 20850, USA. ⁴Enterprise Science And Computing, Inc., Rockville, MD 20850, USA. ⁵March of Dimes, White Plains, NY 10605, USA. ⁶March of Dimes Prematurity Research Center at Stanford, Department of Pediatrics, Stanford University School of Medicine Stanford, CA 94305, USA. ⁷Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA 19104, USA. ⁸Department of Obstetrics and Gynecology, Washington University in St Louis, St. Louis, MO 63110, USA. ⁹Department of Human Genetics, University of Chicago, Chicago, IL 60637, USA. ¹⁰Cincinnati Children's Hospital Medical Center and University of Cincinati College of Medicine, Cincinati, OH 45267, USA. ¹¹Department of Biological Sciences, Vanderbilt University, Nashville, TN 37235, USA. ¹²Department of Biomedical Informatics, Vanderbilt University Medical Center, Nashville, TN 37212, USA. ¹³Department of Obstetrics and Gynecology, Feinberg School of Medicine, Northwestern University, Chicago, IL 60637, USA. Correspondence and requests for materials should be addressed to M.S. (email: marina.sirota@ucsf.edu) *A full list of members appears in the Author Contributions.

| Design Type(s) | data integration objective • database creation objective | | | |
|--------------------------|--|--|--|--|
| Measurement Type(s) | premature birth | | | |
| Technology Type(s) | digital curation | | | |
| Factor Type(s) | Data Type • research organization | | | |
| Sample Characteristic(s) | | | | |

Background & Summary

In the last decade, the advent and continued development of genotyping and next-generation sequencing technologies has enabled researchers to generate a vast amount of molecular data. The current and evergrowing availability of public 'omics' databases, including Gene Expression Omnibus^{1,2}, Array Express³, dbGAP^{4,5}, and numerous other repositories, along with computational tools to reveal molecular drivers of disease at a network level, present a unique and new opportunity to refine current knowledge about disease mechanisms, diagnostics, and therapeutics. In addition, novel technologies are allowing highthroughput measurements of the human genome, epigenome, transcriptome, and immunome at the population level⁶. There have been several disease- and phenotype-specific efforts to aggregate datasets and to make them available in the public domain, with exemplary strides by the cancer community including The Cancer Genome Atlas (TCGA)⁷, which captures 'omics' profiling of tens of thousands of cancer samples; Cancer Cell Line Encyclopedia (CCLE), which focuses on molecular profiling of cancer cell lines; the Library of Integrated Network-Based Cellular Signatures (LINCS)⁸, which focuses on transcriptomic profiling of small molecules and other perturbations in cancer lines; and many others. However, other fields are lagging behind in the data-sharing realm. To this end, we aim to enable open data-sharing efforts in the fields of obstetrics and gynecology. Here, we present our March of Dimes (MOD) Database for Preterm Birth Research (http://www.immport.org/resources/mod), a unique resource that captures a variety of 'omics' datasets related to preterm birth (PTB).

PTB, or the delivery of an infant prior to 37 weeks of gestation, is a significant cause of infant morbidity and mortality. Globally, approximately 11% of infants are born prematurely every year, totaling nearly 15 million births. Infants born preterm are at risk for a variety of adverse outcomes, such as respiratory illness, blindness, and cerebral palsy, with associated complications resulting in nearly one million deaths each year^{9–11}. Despite many attempts for preterm birth prevention, there is still an acute problem with prevalence rising according to the World Health Organization. Spontaneous preterm birth, accounting for two thirds of all preterm births, is considered a complex phenotype with no single known cause¹² or biological basis^{13,14}. One mechanism that has been associated with spontaneous preterm birth is chorioamnionitis, a condition associated with microbial infection of the amniotic fluids. Other suggested causes of PTB are progesterone deficiency, cervical insufficiency, disruption of the immune tolerance of the mother towards the fetus and disruption of the vaginal microbial balance, causing an inflammatory process^{15–19}. Because of the complexity of the phenotype, a comprehensive integrative approach is needed to better understand the etiology of preterm birth and inform new diagnostic and therapeutic strategies.

The March of Dimes is dedicated to not only helping affected infants and families; but also preventing PTB. Only modest progress has been made in identifying the underlying causes of PTB, so the MOD has made this a top research priority. To foster a new model of collaboration with the hope of leading to transformative discoveries, six MOD Prematurity Research Centers have been launched. The goals are to integrate scientists from individual disciplines and to form innovative collaborations that can accelerate research discoveries²⁰. The first Center was launched in 2011 at Stanford University School of Medicine to study infection and inflammation, transcriptomics and lead bioinformatics effort in PTB research. The second center, the MOD Prematurity Research Center - Ohio Collaborative, was opened in 2013 with a focus on evolutionary biology and genetics in PTB as well as molecular development of pregnancy, progesterone signaling and racial disparities in PTB. In 2014, two more centers were launched. On November 10, 2014 the MOD Prematurity Research Center at Washington University was established in St. Louis to apply bioengineering to study cervical remodeling and uterine contractility, as well as determining if chronodisruption is a risk factor for preterm birth. On November 17, 2014 the MOD Prematurity Research Center at the University of Pennsylvania was opened to address questions of cervical remodeling, placental development and bioenergetics in the context of PTB. The fifth center, MOD Prematurity Research Center University of Chicago-Northwestern-Duke was launched in 2015 with a focus on what causes premature birth, and specifically address six interrelated transdisciplinary research themes around gene regulation. The most recent center was launched in February 2018 at the Imperial College London with a focus on how the body recognizes and interacts with bacteria and other microbes in the birth canal that may increase the risk of premature birth.

There have been several previous efforts to aggregate and integrate genomics data for PTB research²¹. dbPTB²² is a resource that collects genomic data across a large number of studies focusing on linking

information from published literature with data from expression databases, linkage studies, and pathway analyses to identify biologically relevant genes for testing in an association study of genetic variants and PTB. However, the database is limited to disease-gene and pathway associations. GEneSTATION²³ is a comprehensive database that integrates cross-species genomic, transcriptomic, and evolutionary data to advance the understanding of the genetic basis of gestation - and pregnancy-associated phenotypes and to accelerate the translation of discoveries from model organisms to humans. This resource specifically focuses on evolutionary and genomic data and is not designed for capturing other 'omics' technologies that cannot be directly mapped to human genes or genetic elements (e.g., microbiome data). In addition, dbPTB does not allow for bioinformatics re-analyses of the processed datasets they aggregate and only summarizes the data at the results level; whereas, GEneSTATION is designed for exploratory data analyses and not as a tool for large-scale re-analyses or meta-analyses of pregnancy 'omics' data in general. A comprehensive resource that can capture more diverse types of data and enable data re-use and re-analyses is needed.

The MOD Database for Preterm Birth Research aims to organize scientific and clinical research data across the six MOD-Funded Prematurity Research Centers with the goal of enhancing research collaborations and accelerating the overall pace of discoveries in this field. Data from the Centers includes a diverse set of processed 'omics' data and results files as well as data generation protocols to support reanalyses and meta-analyses of the datasets. As of January 2018, the database references 9 studies across over 350 patients and with individual level molecular data on nearly 8,000 samples from 6 measurement modalities. Four additional large-scale GWAS studies across tens of thousands of patients are also included, for which only summary-level data are available. The repository includes genomic, transcriptomic, immunonological, and microbiome data that are available freely to the scientific community. Having all the data aggregated as part of the same resource, substantially extends the value the data allowing researchers to integrate the information and ask novel research questions.

Methods

The database development was undertaken in collaboration with Northrop Grumman (NG) – Health Solutions, partner of the National Institute of Allergy and Infectious Diseases (NIAID) Division of Allergy, Immunology, and Transplantation (DAIT). Since 2004, NG has been the prime contractor for the ImmPort²⁴ database and data-sharing portal, working with researchers at UCSF and Stanford to ensure that NIAID-funded discoveries serve as the foundation of future research. In order to host the MOD Database for Preterm Birth Research, we used the existing infrastructure, data model, and repository schema, known as ImmPort (www.immport.org)^{24,25}. ImmPort encompasses immunological data across a wide range of diseases and conditions including several pregnancy-related studies. Study data includes over 50 diverse types of data including arrays, mass cytometry (CyTOF), enzyme-linked immunosorbent assays (ELISA), flow cytometry, and gene expression as well as others.

Datasets archived in ImmPort cover a broad spectrum of study-specific data including protocol designs, assay protocols, treatment and sampling time points, and subject demographics. Datasets are curated and organized by the ImmPort team prior to sharing with the scientific public. Curation efforts include adoption of community standards and controlled vocabularies across a broad spectrum of variables to improve data consistency within and across studies facilitating data re-use. Data upload templates instantiate standards and ontologies and are the result of ongoing outreach with domain experts, standards working groups, and research consortia. Data are de-identified. ImmPort has adopted best-practices in human study participant de-identification such that Health Insurance Portability and Accountability Act of 1996 (HIPAA)-restricted data are not captured in ImmPort. In addition, ImmPort references genetics data stored within the National Center for Biotechnology Information (NCBI) Database of Genotypes and Phenotypes (dbGaP), ensuring ImmPort data safeguards potentially identifying information.

For each study (Fig. 1), a summary page is created, which contains the description, principal investigators (PIs), links to the publication, and external repositories with raw data. Only processed 'omics' data are stored in the repository to avoid duplication of effort. The information is drawn from the publication when available and confirmed by the PIs. Study design, types of assessments, and mechanistic assays with links to the protocols as well as the processed files are listed as separate tabs for each study. Finally, demographics are aggregated across all the samples for each study cohort where individual level data are available. Each subject is given a unique ID and when data from multiple assays is available for the same subject, the data can be accessed by the subject ID. All the human subjects work has been approved by the relevant IRB boards at each institution.

Data Records

As of January 2018, the resource indexed 13 studies (Data Citations1–13), with individual level molecular measurements on nearly 8,000 samples from more than 350 patients, capturing data from 6 measurement modalities (Fig. 2). The resource is further divided into three sections: 1) "curated datasets"; 2) "research highlights"; and 3) "other relevant publicly-available resources relevant to PTB research". The resource is updated at least on a quarterly basis as new studies get added.

The studies referenced by the database (Table 1) cover a diverse set of modalities including microbiome, CyTOF, RNA-Seq, cell-free DNA and RNA sequencing, and genotyping. The protocols for

SDY1157: Immune response throughout human pregnancy

| Study data available for download for Registered Users. Please read the terms and conditions of this User Agreement. | | | | | |
|--|---|--|--|--|--|
| Summary Design | Assessment Demographics Mechanistic Assays Study Files | | | | |
| Accession: | SDY1157 | | | | |
| DOI: | 10.21430/M3OV4WX72N | | | | |
| Title: | Immune response throughout human pregnancy | | | | |
| PI: | Brice Gaudilliere - Stanford University School of Medicine | | | | |
| Туре: | Observational | | | | |
| Condition Studied: | Immune system during Pregnancy | | | | |
| Brief Description: | This study combined the high-parameter functional profiling of peripheral immune cells with a previously unknown cell signaling?based Elastic Net (csEN) algorithm to infer a model of interrelated immune features accurately predicting the timing of immunological adaptations over the entire course of a term pregnancy. | | | | |
| Start Date: | 2015-01-01 | | | | |
| Detailed Description: | The phenotype and intracellular signaling activities of all major innate and adaptive immune cells were simultaneously quantified in serial whole-blood samples throughout pregnancy. Three sets of data were generated by quantifying the abundances of peripheral immune cell sub-sets, capturing endogenous intracellular signaling activities, and determining the capacity of immune cell subsets to respond to stimulation with receptor-specific ligands. The csEN algorithm was adapted from the Elastic Net (EN) regularized regression method and accounts for the influence of previous biological knowledge of receptor-specific signal transduction on the generation of single-cell mass cytometry data. This algorithm allowed to infer a model of interrelated immune features that accurately predicts the timing of immunological adaptations over the entire course of a term pregnancy. | | | | |
| Objectives: | Determine whether a precise chronology of pregnancy-related immunological adaptations is detectable from the mass cytometry analysis of peripheral immune ce phenotype and functional changes during pregnancy. | | | | |
| Endpoints: | CyTOF | | | | |
| Gender Included: | Female | | | | |
| Subjects Number: | 28 | | | | |
| Download Packages: | Study Download Packages | | | | |
| Contract/Grant: | | | | | |
| Data Completeness: | 2 - Complete set of descriptive data and results, as ascertained by ImmPort. | | | | |

Figure 1. Screenshot of an example study indexed in the resource (SDY1157).

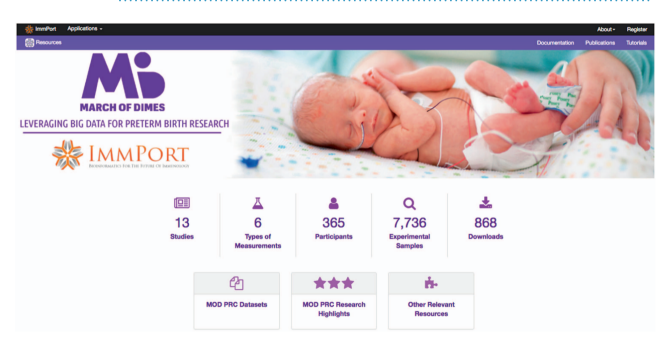


Figure 2. The MOD Database for Preterm Birth Research. A screenshot of the data repository resource as of January 2018. The landing page contains repository statistics, including the number of studies, participants, experimental samples, and study downloads. There are three sections including: 1) a table of the studies with corresponding links; 2) research highlights listing a selection of recent publications with links to the studies; and 3) a compilation of other relevant publicly-available datasets.

| ID | Type of Data | Patients | Samples | Principal Investigator | Center | Unprocessed Data | Processed Data | PubMed ID |
|-----------------------------------|------------------------------|----------|---------|--|--|---|-----------------------------|---------------------------|
| SDY465 (Data Citation 1) | Microbiome 16S | 47 | 4,122 | David Relman | Stanford University | Raw reads Link to SRA | .BIOM format | 26283357 ²⁶ |
| SDY475 (Data Citation 2 | CYTOF | 23 | 95 | Martin Angst | Stanford University | 3,000,000 cells per sample | Cell Clusters | 26190063 ²⁷ |
| SDY775 (Data Citation 3) | Microbiome 16S | 7 | 69 | Samuel Parry and Frederic Bushman | University of Pennsylvania | Raw reads | .BIOM format | 27338728 ²⁸ |
| SDY1155 (Data Citation 4) | RNA-Seq | 15 | 75 | Catalin Buhimschi | Ohio Collaborative | Sequencing | Counts matrix | 27452435 ²⁹ |
| SDY1164 (Data Citation 6) | Microbiome 16s | 136 | 2,177 | David Relman | Stanford University | Raw reads | .BIOM | 28847941 ³⁰ |
| SDY1157 (Data Citation 7) | CYTOF | 28 | 112 | Nima Aghaeepour and Brice Gaudilliere | Stanford University | 3,000,000 cells per sample | Cell clusters | 28864494 ³¹ |
| SDY673 (Data Citation d8) | Cell Free DNA and RNA Seq | 50 | 188 | Stephen Quake | Stanford University | ~20,000,000 reads per sample | Counts matrix | 28904056 ³² |
| SDY1175 (Data Citation 8) | Cell Free DNA Seq | 16 | 58 | Stephen Quake | Stanford University | ~20,000,000 reads per sample | Abundance matrix | 28830999 ³³ |
| SDY776* (Data Citation 9) | Genotyping | 1,705 | 1,705 | Louis Muglia | Ohio Collaborative | Genotypes Link to dbGaP | Association test results | 26284790 ^{34,35} |
| SDY1173* (Data Citation 10) | Genotyping | ~40k | ~40k | Louis Muglia | Ohio Collaborative | Genotypes Link to dbGaP | Association test results | 28877031 ³⁵ |
| SDY1206* (Data Citation 11) | Microbiome | 77 | 149 | Molly Stout | Washington University in St. Luis | Raw reads link to Short Read Archive (SRA) | Abundance matrix | 28549981 ³⁶ |
| SDY1215* (Data Citation 12) | Genotyping (Mitochondria) | ~15k | ~15k | Neal Sondheimer | University of Pennsylvanya in collab with Stanford | Genotypes Link to dbGaP | Association test results | 29249523 ³⁷ |
| SDY1205* (Data Citation 13) | Genotyping | ~15k | ~15k | Marina Sirota and Atul Butte | Stanford in collab with Ohio Collaborative | Genotypes Link to dbGaP | Association test results | 29317701 ³⁸ |

Table 1. MOD-funded studies in the MOD database for preterm birth research as of January 2018. *No individual level genomic data is available, only summary statistics and results.

| Study ID | Database | Туре | Number of Samples | Tissue |
|-----------------|----------|------------------|--------------------------|---|
| GSE46510 | GEO | Transcriptomics | 154 samples | Maternal Blood |
| GSE59491 | GEO | Transcriptomics | 326 samples | Maternal Blood |
| GSE73685 | GEO | Transcriptomics | 183 samples | Amnion, Cord Blood, Decidua, Maternal Blood |
| phs001320.v1.p1 | dbGaP | Transcriptomics | 58 samples | Chorionic villus sampling |
| PRJNA242473 | SRA | Microbiome | 349 samples | Vaginal Microbiome |
| phs000735.v1.p1 | dbGaP | Microbiome | 48 samples | Placenta |
| phs000256.v3.p2 | dbGaP | Microbiome | 3,474 samples | Vaginal Microbiome |
| phs000332.v2.p2 | dbGaP | Genetics | 1,779 samples | Blood |
| phs000714.v1.p1 | dbGaP | Genetics | 2,928 samples | Blood |
| phs000353.v1.p1 | dbGaP | Genetics | 3,478 samples | Blood |
| phs000103.v1.p1 | dbGaP | Genetics | 2,000 mother-child pairs | Blood |
| phs001055.v1.p1 | dbGaP | Genetics | 2,397 samples | Blood |
| phs000276.v2.p1 | dbGaP | Genetics | 5,415 samples | Blood |
| SDY37 | ImmPort | ELISA, Cytokines | 55 samples | Maternal Blood |
| SDY36 | ImmPort | Vaccine Response | 300 samples | Maternal Blood |

Table 2. Other publicly available studies relevant to preterm birth as of January 2018. *No individual level genomic data is available, only linking out to the relevant resources.

data generation are included in each study (Fig. 1), as are subject demographics and relevant clinical variables as well as processed 'omics' datasets. When possible, raw data stored in other public repositories such as SRA, dbGaP and GEO are linked. Publications arising from the studies are referenced on individual study pages, and the most recent papers are also listed on the "research highlights" section of the resource.

Publicly-available data are displayed in a table format, which currently lists 15 additional studies from GEO, dbGaP, SRA, and ImmPort (Table 2). Various types of transcriptomic, microbiome, and genomic

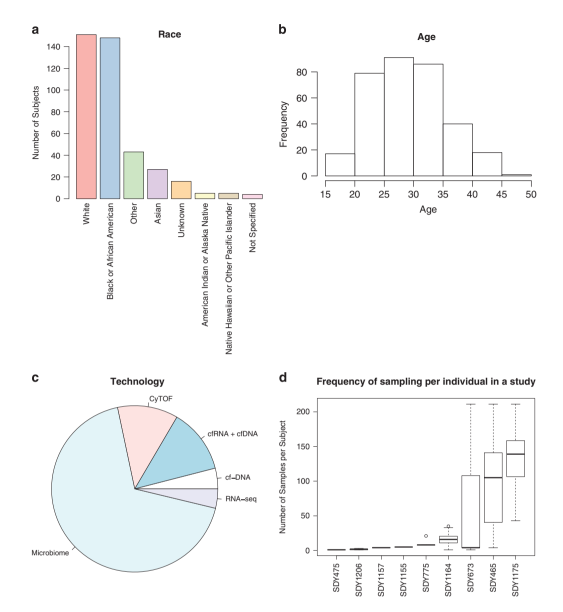


Figure 3. Overall Database Statistics on Individual Measurements (9/13 studies). (a) Race distribution across the cohort. (b) Age distribution across the cohort. (c) Percentage of samples profiled by each technology from the individual measurements. (d) Frequency of sampling per individual in each study.

data across a variety of relevant reproductive tissues are included in the database with molecular profiles for more than 25,000 samples. The studies are linked from the resource page to their respective study detail pages in their respective repositories.

Technical Validation

In order to technically validate the resource, we computed some overall statistics on the repository (Fig. 3) focusing on the studies where we had individual level data (as opposed to aggregated summary statistics). As of January 2018, the database cohort consisted primarily of samples from White and African-American individuals with smaller proportions of samples from Asian and other racial and ethnic groups (Fig. 3a). The age of participants that we have individual level data on ranged from 16 to 46 years of age with a mean of 29.7 ± 6.1 years (Fig. 3b). The majority of individuals (67.9%) in the resource have been profiled with microbiome 16s technology (Fig. 3c). There is a significant proportion of cell-free DNA and cell-free RNA measurements, as well as CyTOF (12.5 and 11.8% respectively). While the majority of the studies are cross-sectional and have very few samples per individual, there are several longitudinal studies (SDY465, SDY763 and SDY1175) with extensive sampling frequency (Fig. 3d).

We have also been tracking downloads through ImmPort for individual studies as well as bulk downloads as part of the ALLSTUDIES package, which contains the entirety of publicly-available ImmPort studies (Table 3). Individual study downloads counts range from 0 to 61 downloads with an

| Study | First Data Release | Date of First Data Release | Number of downloads as of 01/22/2018 | |
|---------------|--------------------|----------------------------|--------------------------------------|--|
| SDY465 | DR18 | March 18, 2016 | 41 | |
| SDY475 | DR16 | December 14, 2015 | 54 | |
| SDY673 | DR24 | November 7, 2017 | 5 | |
| SDY775 | DR20 | January 31, 2017 | 27 | |
| SDY776 | DR24 | November 7, 2017 | 1 | |
| SDY1155 | DR23 | September 29, 2017 | 8 | |
| SDY1157 | DR23 | September 29, 2017 | 13 | |
| SDY1164 | DR23 | September 29, 2017 | 40 | |
| SDY1173 | DR24 | November 7, 2017 | 5 | |
| SDY1175 | DR24 | November 7, 2017 | 8 | |
| SDY1205 | DR25 | January 4, 2018 | 61 | |
| SDY1206 | DR25 | January 4, 2018 | 0 | |
| SDY1215 | DR25 | January 4, 2018 | 0 | |
| Total | | | 263 | |
| Bulk Download | 666 | | | |

Table 3. Download statistics for each study as of January 2018.

average of 20 downloads per study in addition to a total of 666 downloads of the whole repository as of January 2018.

Usage Notes

The overarching goal of our data-sharing effort is to enable new scientific discoveries from the rich molecular resources that have been funded by the MOD to advance the research in PTB. By making all the data publicly available, we aim to engage the larger research community in tackling this important problem. We are also working on incorporating other types of data into this repository including activity monitoring, sociobehavioral data, imaging and others. In addition to the traditional 'omics', investigators are studying non-coding regulatory regions, role of mitochondria, patient-specific glycans and sleep-wake cycles in the context of birth timing. We hope our efforts will entice industry leaders in the areas of machine learning and artificial intelligence to develop and apply computational methodologies to PTB research by leveraging the data that we aggregate. Our group has had tremendous success in mining publicly-available datasets to enable computational drug discoveries in the areas of autoimmunity and cancer^{39–45}. More recently, we have carried out an integrative ancestry-specific GWAS analysis in PTB by integrating internal- and publicly-available data and identified several variants associated with early PTB³⁸ as well as a transcriptomics meta-analysis of maternal and fetal signals elucidating the role of the immune system in parturition timing⁴⁶. We will continue similar efforts in the area of PTB, and we hope to inspire others to follow suit. We hope this work enabling intersection of data layers related in the context of preterm birth will lead to discovery of novel diagnostic biomarkers and ultimately aid in formulating more effective interventional strategies for the management and prevention of PTB.

References

- 1. Barrett, T. et al. NCBI GEO: archive for functional genomics data sets--update. Nucleic Acids Res 41, D991–D995, doi:10.1093/nar/gks1193 (2013).
- 2. Edgar, R., Domrachev, M. & Lash, A. E. Gene Expression Omnibus: NCBI gene expression and hybridization array data repository. *Nucleic Acids Res* **30**, 207–210 (2002).
- Kolesnikov, N. et al. ArrayExpress update--simplifying data submissions. Nucleic Acids Res 43, D1113–D1116, doi:10.1093/nar/gku1057 (2015).
- Mailman, M. D. et al. The NCBI dbGaP database of genotypes and phenotypes. Nat Genet 39, 1181–1186, doi:10.1038/ng1007-1181 (2007).
- Tryka, K. A. et al. NCBI's Database of Genotypes and Phenotypes: dbGaP. Nucleic Acids Res 42, D975–D979, doi:10.1093/nar/gkt1211 (2014).
- 6. Consortium, G. T. Human genomics. The Genotype-Tissue Expression (GTEx) pilot analysis: multitissue gene regulation in humans. *Science* **348**, 648–660, doi:10.1126/science.1262110 (2015).
- Grossman, R. L. et al. Toward a Shared Vision for Cancer Genomic Data. N Engl J Med 375, 1109–1112, doi:10.1056/NEIMp1607591 (2016).
- 8. Subramanian, A. *et al.* A Next Generation Connectivity Map: L1000 Platform and the First 1,000,000 Profiles. *Cell* **171**, 1437–1452 e1417, doi:10.1016/j.cell.2017.10.049 (2017).
- Morken, N. H., Kallen, K. & Jacobsson, B. Outcomes of preterm children according to type of delivery onset: a nationwide population-based study. *Paediatric and perinatal epidemiology* 21, 458–464, doi:10.1111/j.1365-3016.2007.00823.x (2007).
- Schaaf, J. M., Mol, B. W., Abu-Hanna, A. & Ravelli, A. C. Ethnic disparities in the risk of adverse neonatal outcome after spontaneous preterm birth. Acta obstetricia et gynecologica Scandinavica 91, 1402–1408, doi:10.1111/aogs.12013 (2012).
- Bastek, J. A., Srinivas, S. K., Sammel, M. D. & Elovitz, M. A. Do neonatal outcomes differ depending on the cause of preterm birth? A comparison between spontaneous birth and iatrogenic delivery for preeclampsia. *American journal of perinatology* 27, 163–169, doi:10.1055/s-0029-1234036 (2010).

- Green, N. S. et al. Research agenda for preterm birth: recommendations from the March of Dimes. American journal of obstetrics and gynecology 193, 626–635, doi:10.1016/j.ajog.2005.02.106 (2005).
- Ferrero, D. M. et al. Cross-Country Individual Participant Analysis of 4.1 Million Singleton Births in 5 Countries with Very High Human Development Index Confirms Known Associations but Provides No Biologic Explanation for 2/3 of All Preterm Births. PLoS One 11, e0162506, doi:10.1371/journal.pone.0162506 (2016).
- 14. Chang, H. H. et al. Preventing preterm births: analysis of trends and potential reductions with interventions in 39 countries with very high human development index. Lancet **381**, 223–234, doi:10.1016/S0140-6736(12)61856-X (2013).
- Macones, G. A. *et al.* A polymorphism in the promoter region of TNF and bacterial vaginosis: preliminary evidence of geneenvironment interaction in the etiology of spontaneous preterm birth. *American journal of obstetrics and gynecology* 190, 1504–1508, discussion 1503A, doi:10.1016/j.ajog.2004.01.001 (2004).
- Roberts, A. K. et al. Association of polymorphism within the promoter of the tumor necrosis factor alpha gene with increased risk of preterm premature rupture of the fetal membranes. American journal of obstetrics and gynecology 180, 1297–1302 (1999).
- 17. Annells, M. F. *et al.* Interleukins-1, -4, -6, -10, tumor necrosis factor, transforming growth factor-beta, FAS, and mannose-binding protein C gene polymorphisms in Australian women: Risk of preterm birth. *American journal of obstetrics and gynecology* **191**, 2056–2067, doi:10.1016/j.ajog.2004.04.021 (2004).
- 18. Engel, S. A. *et al.* Risk of spontaneous preterm birth is associated with common proinflammatory cytokine polymorphisms. *Epidemiology* **16**, 469–477 (2005).
- Kalish, R. B., Vardhana, S., Gupta, M., Perni, S. C. & Witkin, S. S. Interleukin-4 and -10 gene polymorphisms and spontaneous preterm birth in multifetal gestations. *American journal of obstetrics and gynecology* 190, 702–706, doi:10.1016/j.ajog.2003.09.066 (2004).
- 20. Wise, P. H et al. Risky Business: Meeting the Structural Needs of Transdisciplinary Science. J Pediatr 191, 255–258, doi:10.1016/j.jpeds.2017.08.072 (2017).
- Eidem, H. R., McGary, K. L., Capra, J. A., Abbot, P. & Rokas, A. The transformative potential of an integrative approach to pregnancy. *Placenta* 57, 204–215, doi:10.1016/j.placenta.2017.07.010 (2017).
- 22. Uzun, A et al. dbPTB: a database for preterm birth. Database (Oxford) 2012, bar069, doi:10.1093/database/bar069 (2012).
- Kim, M. et al. GEneSTATION 1.0: a synthetic resource of diverse evolutionary and functional genomic data for studying the evolution of pregnancy-associated tissues and phenotypes. Nucleic Acids Res 44, D908–D916, doi:10.1093/nar/gkv1137 (2016).
- 24. Bhattacharya, S. et al. ImmPort: disseminating data to the public for the future of immunology. Immunol Res 58, 234–239, doi:10.1007/s12026-014-8516-1 (2014).
- 25. Bhattacharya, S. *et al.* ImmPort, toward repurposing of open access immunological assay data for translational and clinical research. *Sci Data* **5**, 180015, doi:10.1038/sdata.2018.15 (2018).
- DiGiulio, D. B. et al. Temporal and spatial variation of the human microbiota during pregnancy. Proc Natl Acad Sci USA 112, 11060–11065, doi:10.1073/pnas.1502875112 (2015).
- 27. Gaudilliere, B. *et al.* Implementing Mass Cytometry at the Bedside to Study the Immunological Basis of Human Diseases: Distinctive Immune Features in Patients with a History of Term or Preterm Birth. *Cytometry A* **87**, 817–829, doi:10.1002/cyto.a.22720 (2015).
- Lauder, A. P. et al. Comparison of placenta samples with contamination controls does not provide evidence for a distinct placenta microbiota. Microbiome 4, 29, doi:10.1186/s40168-016-0172-3 (2016).
- Ackerman, W. E. t. et al. Comprehensive RNA profiling of villous trophoblast and decidua basalis in pregnancies complicated by preterm birth following intra-amniotic infection. Placenta 44, 23–33, doi:10.1016/j.placenta.2016.05.010 (2016).
- Callahan, B. J. et al. Replication and refinement of a vaginal microbial signature of preterm birth in two racially distinct cohorts of US women. Proc Natl Acad Sci U S A 114, 9966–9971, doi:10.1073/pnas.1705899114 (2017).
- 31. Aghaeepour, N. et al. An immune clock of human pregnancy. Sci Immunol 2, doi:10.1126/sciimmunol.aan2946 (2017).
- Pan, W. et al. Simultaneously Monitoring Immune Response and Microbial Infections during Pregnancy through Plasma cfRNA Sequencing. Clin Chem 63, 1695–1704, doi:10.1373/clinchem.2017.273888 (2017).
- Kowarsky, M. et al. Numerous uncharacterized and highly divergent microbes which colonize humans are revealed by circulating cell-free DNA. Proc Natl Acad Sci U S A 114, 9623–9628, doi:10.1073/pnas.1707009114 (2017).
- 34. Plunkett, J. et al. An evolutionary genomic approach to identify genes involved in human birth timing. PLoS Genet 7,
- e1001365, doi:10.1371/journal.pgen.1001365 (2011).
- 35. Zhang, G. et al. Genetic Associations with Gestational Duration and Spontaneous Preterm Birth. N Engl J Med 377, 1156–1167, doi:10.1056/NEJMoa1612665 (2017).
- Stout, M. J. et al. Early pregnancy vaginal microbiome trends and preterm birth. Am J Obstet Gynecol 217, 356 e351–356 e318, doi:10.1016/j.ajog.2017.05.030 (2017).
- Crawford, N. et al. Divergent Patterns of Mitochondrial and Nuclear Ancestry Are Associated with the Risk for Preterm Birth. J Pediatr, doi:10.1016/j.jpeds.2017.10.052 (2017).
- 38. Rappoport, N. *et al.* A genome-wide association study identifies only two ancestry specific variants associated with spontaneous preterm birth. *Sci Rep* **8**, 226, doi:10.1038/s41598-017-18246-5 (2018).
- Aran, D. et al. Comprehensive analysis of normal adjacent to tumor transcriptomes. Nat Commun 8, 1077, doi:10.1038/s41467-017-01027-z (2017).
- Aran, D., Sirota, M. & Butte, A. J. Systematic pan-cancer analysis of tumour purity. Nat Commun 6, 8971, doi:10.1038/ncomms9971 (2015).
- Bagley, S. C., Sirota, M., Chen, R., Butte, A. J. & Altman, R. B. Constraints on Biological Mechanism from Disease Comorbidity Using Electronic Medical Records and Database of Genetic Variants. *PLoS Comput Biol* 12, e1004885, doi:10.1371/journal.pcbi.1004885 (2016).
- 42. Dudley, J. T. *et al.* Computational repositioning of the anticonvulsant topiramate for inflammatory bowel disease. *Sci Transl Med* 3, 96ra76, doi:10.1126/scitranslmed.3002648 (2011).
- Kosti, I., Jain, N., Aran, D., Butte, A. J. & Sirota, M. Cross-tissue Analysis of Gene and Protein Expression in Normal and Cancer Tissues. Sci Rep 6, 24799, doi:10.1038/srep24799 (2016).
- 44. Sirota, M. *et al.* Discovery and preclinical validation of drug indications using compendia of public gene expression data. *Sci Transl Med* **3**, 96ra77, doi:10.1126/scitranslmed.3001318 (2011).
- Sirota, M., Schaub, M. A., Batzoglou, S., Robinson, W. H. & Butte, A. J. Autoimmune disease classification by inverse association with SNP alleles. *PLoS Genet* 5, e1000792, doi:10.1371/journal.pgen.1000792 (2009).
- 46. Vora, B. et al. Meta-Analysis of Maternal and Fetal Transcriptomic Data Elucidates the Role of Adaptive and Innate Immunity in Preterm Birth. Front Immunol 9, 993, doi:10.3389/fimmu.2018.00993 (2018).

Data Citations

- 1. DiGiulio, D. B. et al. ImmPort https://doi.org/10.21430/M3D491LGDT (2016).
- 2. Gaudilliere, B. et al. ImmPort https://doi.org/10.21430/M3D8CS7ILY (2015).
- 3. Lauder, A. P. et al. ImmPort https://doi.org/10.21430/M3PZM1ERD2 (2017).

- 4. Ackerman, W. E. et al. ImmPort https://doi.org/10.21430/M34I5YT3K9 (2017).
- 5. Callahan, B. J. et al. ImmPort https://doi.org/10.21430/M37W3869AH (2017).
- 6. Aghaeepour, N. et al. ImmPort https://doi.org/10.21430/M3OV4WX72N (2017).
- 7. Pan, W. et al. ImmPort https://doi.org/10.21430/M3OARGGSY0 (2017).
- 8. Kowarsky, M. et al. ImmPort https://doi.org/10.21430/M33PSZ2FHV (2017).
- 9. Plunkett, J. et al. ImmPort https://doi.org/10.21430/M3AM8G2I2Q (2017).
- 10. Zhang, G. et al. ImmPort https://doi.org/10.21430/M3F345ZL81 (2017).
- 11. Stout, M. J. et al. ImmPort https://doi.org/10.21430/M3H1U3KJMZ (2018).
- 12. Crawford, N. et al. ImmPort https://doi.org/10.21430/M3VCNPMD4B (2018).
- 13. Rappoport, N. et al. ImmPort https://doi.org/10.21430/M37N6PJEQT (2018).

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Author Contributions

M.S., A.B., J.LS. and R.L. have conceptualized the resource. M.S. designed the resource and is overseeing the project., C.T., M.Z., P.B. and L.T. have carried out the implementation of the resource. R.W., C.Q., R.L., J.C., R.A., J.C., M.K., W.G., G.Z. and A.R. have helped with data deposition and sharing, as well as the design of the resource. L.M., C.O., S.E., G.M., S.P., G.S., D.S., A.B. and D.D. have provided oversight for resource design and data deposition. R.L. and J.L.S. have been involved in resource design and oversight. All authors have read and edited the manuscript.

March of Dimes Prematurity Research Centers

Deborah Driscoll⁷, George Macones⁸, Louis J Muglia¹⁰, Carole Ober⁹ & David K. Stevenson⁶

Additional Information

Competing interests: R.L. and J.L.S. are employees of the March of Dimes, which is the funding organization of this project. Since they have contributed to the resource design and inception, they are co-authors on this work.

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