**Plan Overview**

*A Data Management Plan created using DMPTool*

**DMP ID:** https://doi.org/10.48321/D15S4Z

**Title:** Improving the quality of stem-cell derived insulin-producing cells

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**Data Manager:** Xing Jian

**Project Administrator:** Ruthsabel O'Lexy

**Funder:** National Institutes of Health (nih.gov)

**Funding opportunity number:** RFA-DK-21-030

**Template:** NIH-GDS: Genomic Data Sharing

**Project abstract:**

Type 1 diabetes (T1D) is a chronic autoimmune disease characterized by the loss of pancreatic \(\beta\)-cells. It accounts for up to 10% of all current cases of diabetes in the United States, and its overall incidence has significantly increased since 2002. The insulin-producing cells (SC-\(\beta\)-like cells) derived from human induced pluripotent stem cells (iPSCs) are promising renewable resources for both \(\beta\)-cell replacement T1D therapies and biomimetic devices to study human pancreatic islets’ function ex vivo. However, SC-\(\beta\)-like cells generated from current protocols do not meet all the critical properties of mature \(\beta\)-cells and are therefore not fully functional. The cells can reach maturation stage after transplantation into mouse models; however, this procedure is slow and costly, and it may not predict responses in humans accurately. The proposed project attempts to develop a superior protocol that would generate better functional cells with less time, expense, and variability. The proposed research has significant potential to achieve its long-term goal, namely, generating highly functional \(\beta\)-cells from iPSCs to create a much-needed model system for biomimetic devices and \(\beta\)-cell replacement therapies.

This R03 application’s rationale is based on published data from SC-\(\beta\)-like cells and mouse models. Recent studies of SC-\(\beta\)-like cells indicate that the key mature \(\beta\)-cell-specific transcription factor MAFA is only transiently expressed in the differentiation process, and its loss in the final differentiation stage leads to reduced...
β-cell functionality. Previous studies also showed that inducing MAFA expression in these cells would improve cell properties. Mouse animal model and cell line studies identified two biological pathways that are used in β-cells to upregulate MAFA levels. In the first pathway, m6A methylation on MAFA mRNA increases its stability and abundance. In the other pathway, high concentration of glucose increases MAFA expression by the hexosamine biosynthetic pathway. The objective of this project is to use small molecules related to these pathways to upregulate MAFA levels so that the critical functional properties of these cells can improve. To achieve its objective, this project will pursue two specific aims: aim 1 - using small molecule modulators of the m6A RNA methylation pathway to increase MAFA expression levels in SC-β-like cells; aim 2 - using small molecule modulators of hexosamine biosynthetic pathway to increase MAFA expression levels in SC-β-like cells. The six critical properties of human β-cells outlined by the Human Islet Research Network (HIRN) will be used as guidelines to characterize SC-β-like cells. Project data will be generated from the Coriell Institute’s facilities, with the exception that the next-generation sequencing will be carried out at an external high-throughput sequencing core facility, and analyzed with help from Coriell’s Genomics & Epigenomics Core and its Bioinformatics Core.

Start date: 04-30-2023

End date: 04-29-2025

Last modified: 10-13-2022

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Improving the quality of stem-cell derived insulin-producing cells

Data type

- Explain whether the proposed research involves human data, non-human data, or both.
- List the type(s) of genomic data that will be shared (e.g., sequence, transcriptomic, epigenomic, and/or gene expression data) and whether it is individual-level data, aggregate-level data, or both.
- List any other information such as relevant associated data (e.g., phenotype or exposure data) and information necessary to interpret the data (e.g., study protocols, data collection instruments, survey tools) that the investigator anticipates sharing.

This study will use de-identified human cell lines from the National Institute of General Medical Sciences (NIGMS) Human Genetic Cell Repository (https://www.coriell.org/1/NIGMS) without unique identify information. High throughput transcriptomic sequencing data generated in this study will be generated at single cell level. Single-cell transcriptome libraries will be prepared by a 10X Genomics Controller, and be sequenced by Illumina sequencers in a core facility. The raw data will be processed by Cell Ranger pipeline (proprietary software by 10X Genomics) and the Seurat workflow (open-source software). The raw data (FASTQ and TSV format) and the processed data (TSV format), as well as the associated metadata (TXT format) will be deposited to an NIH-designated data repository (see the next section), and the total size of the data is estimated to be 5 Gb. The phenotype data for the the cell line donors are publicly available at Coriell Institute website (www.coriell.org). The methods section of publications from this study will provide technique details of the experimental procedures. Other researchers should be able to download and independently perform analysis on the data, and validate the results of publications from this study. No custom-created code or in-house software will be used for data analysis.

Data repositories

Identify the repository or repositories where the investigator plans to submit genomic data.

- **Human data:** Studies generating human genomic and any associated phenotypic data must use an NIH-designated data repository for submission. Plans must include whether the data will be available through unrestricted or controlled-access repositories. If data cannot be submitted to an NIH-designated repository, see Request for an Alternative Data Sharing Plan.

- **Non-human data:** Studies generating non-human genomic data may use any widely available repository as appropriate for the data.

High throughput transcriptomic sequencing data generated in this study will be deposited into Gene Expression Omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/), which is an NIH-funded repository, and freely distributes these data to the broad scientific community. In order to meet the MIBBI (Minimum Information for Biological and Biomedical Investigations) standard, the metadata (TSV format) will be prepared from the GEO metadata spreadsheet (https://www.ncbi.nlm.nih.gov/geo/info/examples/seq_template.xlsx). To facilitated file transfer and save storage space, the raw and processed data will be stored in GEO as compressed files (GZ format or TAR.GZ).
Submitted data will conform to relevant data and terminology standards in the single-cell transcriptome research field. The accession number (GSE) assigned to the data by GEO will be used as a persistent unique identifier, so that the data are findable and identifiable to the general public. No unusual data formats are planning to be used.

**Data Submission and Release Timeline**

Provide a timeline for how genomic data will be shared in a timely manner.

I will comply with the broad and rapid data sharing as outlined in the NIH Genomic Data Sharing Policy and will adhere to the changes to NIH-wide data sharing policies, such as the expected 2023 NIH Data Management and Sharing Policy. I agree to deposit high throughput transcriptomic sequencing data into GEO repository as soon as possible but no later than the date of the completion of the funded project period for the parent award or upon acceptance of the data for publication, or public disclosure of a submitted patent application, whichever is earlier. After the initial release of data, I will update the deposited data bi-annually. Data will be available at GEO to the general public as long as possible.

**Institutional Review Board (IRB) Review of Institutional Certification**

**Human data only:** IRB review of the investigator’s proposal for data submission is an element of the Institutional Certification, which assures that the proposal for data submission and sharing is appropriate.

Since this study will use de-identified human cell lines from the NIGMS Human Genetic Cell Repository without unique identify information, the IRB at Coriell Institute determines that this study is not a human-subject study, and no approval from IRB at Coriell Institute is required. I agree that I will not use any data generated from this study for identification purpose.

**Appropriate Uses of the Data**

Describe any limitations on the use of the data. These limitations should be decided by the submitting investigator and their institution, in consultation with the IRB or equivalent body. They should be based on the language in the informed consent form or the recommendations of an IRB or equivalent body.

The donors of the cell lines used in this study are participants of the Personal Genome Project (https://www.personalgenomes.org/). These samples have been consented for commercial use and for public posting of Personally Identifying Genetic Information (PIGI), which allows open-access, public posting of extensive genetic data. Participants the Personal Genome Project are consented to publicly share their genomic and trait data in an integrated, publicly-accessible format using a CC0 waiver or equivalent public domain license. Other than purposes of reasonable cost recovery, the participates of the Personal Genome Project are consented not to sell or license participant data or tissues. Since the data are intended to have open-access to the general public, data security will be controlled by GEO and NIH.
Statement of Designation of Genomic Summary Results (GSR)

Investigators should indicate if a study should be designated as “sensitive” or “not sensitive” for the purposes of access to GSR. This designation should be confirmed in the Institutional Certification Form.

Because of the broad consent of the donors of the cell lines used in this study, this study is designated as not sensitive for the purposes of access to GSR. Principle Investigator of this project, Xing Jian will serve as the data manager, and will be responsible for performing all the required data management and sharing activities and the execution the data management and sharing plan. The Director of Research Office at Coriell Institute, Ruthsabel O'Lexy will serve as the project administrator, and oversee the data management and sharing plan, and monitor the compliance with the plan quarterly.
Planned Research Outputs

Dataset - "Single Cell Transcriptome Analysis of Stem Cell Derived Insulin-Producing Cells"

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Planned research output details

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