FAQ – CRDW/UIC CIRCLE: Data Request Services

What is the CRDW?
The Center for Clinical Translational Sciences (CCTS) Biomedical Informatics (BI) Core continually develops a Clinical Research Data Warehouse (CRDW) to support the University of Illinois research enterprise and collaborator community. The purpose of the CRDW is to provide a single, secure, managed release point for human subjects’ data for use in research. The CRDW has an optimized Research Patient Repository (RPR) that stores and integrates human subjects’ data, including protected health information (PHI) and personally identifiable information (PII), from various sources that can be used to derive data-marts for clinical, translational and basic research purposes.

What is UIC CIRCLE?
The UIC Clinical Information Repository for Cohort Learning and Exploration (UIC CIRCLE) refers to the optimized Research Patient Repository (RPR), along with the related consultation, data request, and data delivery service provided to UIC researchers. This process entails working closely with researchers and utilizing segments of the CRDW’s RPR in order to fulfill requests for clinical data extracts and to ensure the secure transfer and storage of sensitive data.

Where does the CRDW & UIC CIRCLE data come from?
The data warehouse and repository are currently built with extracts from the UI Health hospital electronic medical record and billing systems.

What types of data are available?
*Date range of optimized data: January 1, 2010 – Present*
- Patient demographic information (i.e. age, sex, race/ethnicity)
- General visit description (i.e. visit type, length of stay, discharge disposition, medical service)
- Scheduling information (i.e. appointments, appointment type, appointment status)
- Insurance (i.e. Primary/Secondary Insurance, Financial Class)
- Medications (Cerner Multum & RxNorm)
- Vitals (blood pressure, temperature, etc.)
- General patient information (i.e. height, weight, BMI)
- Lab test results (name of lab test and/or codes [Sunquest & LOINC])
- Diagnoses (diagnosis names and/or codes [ICD-9-CM & ICD-10-CM])
- Procedures (procedure names and/or codes [ICD-9-CM Vol 3, ICD-10-PCS, CPT, & HCPCS])
- Maternity (MaRS & Cerner Maternity)

What are some examples of data NOT readily available?
- Clinical notes (work in progress – availability TBD)
- Dentistry data (work in progress – availability TBD)
- Radiology/Pathology Imaging Data
- Transplant data
- Pharmacy fill data (e.g. retail pharmacy)
Time estimates: How long does it take for an investigator to receive data?
This depends on the complexity of the request – some requests require calculations not easily obtained from UIC CIRCLE, and others may require crossing systems to match patient data for a complete dataset. Typically, a request can be completed in 4-6 weeks. However, some requests can be fulfilled more quickly. Investigators will receive an individualized time estimate after their request is evaluated and approved.

What is the general process of requesting data extracts or informatics support?
1) Investigators or a designated delegate are required to submit a service request form through the CCTS Service Request Site. A confirmation email will be received when the request is received and assigned to the CCTS Biomedical Informatics (BI) Core.
2) The BI Core Manager will follow up with requester when the request is received – Requests are processed in the order they are received, but investigators should inform the CCTS BI Core team of any critical deadlines.
3) A Data Request Authorization (DRA) form is sent to the investigator for completion. The BI Core Manager will review this form and send back a PDF copy with additional instructions.
   a. Requests are reviewed for technical and regulatory feasibility by the CCTS BI Core Team.
   b. The investigator will be contacted to gather additional requirements, as needed.*
4) A technical consultation may be required - The CCTS (BI) Core Manager will contact the PI to set up a meeting time.
5) If the project is feasible, the CCTS BI Core team will review and agree on a project plan, timeline, and estimated costs with the investigator prior to data extraction.
6) The CCTS BI Core team will contact the investigator with any updates during the time of project execution and send a notice when the project is complete. A signed IDUA may be required prior to receiving results.*

*IRB approvals, an approved DRA form, and a signed Internal Data Use Agreement (IDUA) may be required for completion of data requests – See Summary of Required Documentation for Each Dataset Type on page 4.

Are there any costs associated with this process?
The first hour of consultation and requirements gathering, as well as requests for counts that require 1 hour of work or less, are free of charge. Subsequent hours of service and data extraction will be charged at a cost of $85/hour. Estimated costs will be established and approved by the investigator prior to data extraction.

The CCTS BI Core will also work with investigators who are awaiting funding or seeking informatics support on a grant. Requests for provisional cost waivers will be evaluated on a per-project basis.

Should I cite the CCTS in any publications or presentations?
Yes, we ask that you please acknowledge CCTS CRDW/UIC CIRCLE support by referencing grant number UL1TR002003.
What are the data request and data set types? (See corresponding tables on page 4)

**Data Counts** - Assistance in obtaining general counts of clinical data (i.e. the number of patients seen for a specified condition or set of conditions).

*Investigators can also access, QUICKSet/i2b2, to obtain some counts data on their own. Training is required to access this system - [http://training.ccts.uic.edu/](http://training.ccts.uic.edu/).*

**Data Extracts** - Assistance in obtaining complex sets of clinical data for a study, such as identifiable patient information or detailed queries for specific data elements.

**Data Set Types:**

1) **Requests that are preparatory for research/aggregate counts – Cohort Identification:**

   Provides the number of patients within stipulated parameters. No PHI is given - only a count of patients or events. This type of data is typically requested in preparation for a research proposal/grant submission to determine study feasibility.

2) **Requests for De-identified Data Sets:**

   This data set is a result of a positive aggregate count (mentioned in #1 above). Health data is provided without any of the 18 direct identifiers (as defined by HIPAA) and is non-coded. This type of data is typically requested in preparation for a research proposal/grant when more specific information about a potential cohort is needed but individual patients do not need to be identifiable.

3) **Requests for Coded Data Sets:**

   This data set is the same as type 2 (also stripped of any direct identifiers), but is provided to the researcher in a uniquely coded rather than a de-identified manner. This allows CRDW staff to re-identify the data and provide an updated data set on a continuous basis. The researcher cannot request re-identification of the dataset. The key to enable linkage of a coded dataset with PHI will remain only with the CRDW staff. This type of data can be either a) retrospective that includes all existing patient data at the time data is requested or b) prospective that includes all new incoming patient data. This can include updates on patient’s data that has already been collected.

4) **Requests for a Limited Data Set:**

   Similar to dataset type 3 above, this data set contains health data that does not include direct identifiers (such as name and street address). However, limited data sets MAY contain the following indirect identifiers:
   a. Town or City, state, zip code;
   b. Ages in years up to 90 years (must aggregate all ages 90 or older)
   c. Dates directly related to an individual – such as birth date, date of death, admission date, discharge date, visit date, diagnosis date, etc. (Month/year is preferred).

   Like data set type 3, the data request could be a snapshot of what is available at the time of the request (retrospective) or updated data could be provided on an ongoing basis (prospective). The researcher may not request additional subject identifiers.

5) **Requests for Information for Recruitment Purposes/Identified Data Sets –**

   This type of data set includes contact information of patients within a limited data set. The goal is to promote best practices for recruiting and consenting potential research subjects. This dataset is provided such that potential subjects can be approached to determine their interest in participating in research projects. This data set is typically provided for the purpose of pre-trial screening or possible recruitment for a clinical trial, interventional, or observational study. Data provided may include patient names, medical record number, telephone number, address, clinic schedule (time and location), primary care physician name, and provider.

6) **Requests for Defined Identified Cohort Data Sets –**

   This type of dataset contains identified patient data that includes health data for a research study. The researcher defines a group of human subjects for whom a specific set of data is requested. The identified cohort could be individuals that have provided consent in the context of an approved IRB protocol, individuals identified through one of the above processes, or individuals identified through the researcher’s clinical activities. The data request could be a snapshot of what is available at the time of request or updated data that could be provided on an ongoing basis.
### Example uses of requested dataset types

<table>
<thead>
<tr>
<th>Dataset Request Type</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aggregate Counts</td>
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<tr>
<td>De-identified Data Set</td>
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<tr>
<td>Coded Data Set</td>
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<tr>
<td>Limited Data Set</td>
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<tr>
<td>Information for Recruitment Purposes</td>
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<tr>
<td>Identified Data Set</td>
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<table>
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<tr>
<th>Uses of requested dataset</th>
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<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
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</thead>
<tbody>
<tr>
<td>Study feasibility; Hypothesis testing</td>
<td>Chart review, retrospective only (no updates or supplements)</td>
<td>Chart review, retrospective and prospective</td>
<td>Chart reviews that require certain identifier such as zip code or dates of service</td>
<td>Recruitment for clinical studies, clinical registries</td>
<td>Studies on an identified cohort, clinical trials</td>
<td></td>
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</tbody>
</table>

### Summary of Required Documentation for Each Dataset Type:

<table>
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</thead>
<tbody>
<tr>
<td><strong>1. Aggregate Counts</strong></td>
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</tr>
<tr>
<td>1a. Self Service</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
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<tr>
<td>1b. Consult Service</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td><strong>2. De-identified Data</strong></td>
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<td>Yes</td>
<td>No</td>
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</tr>
<tr>
<td><strong>3. Coded Data</strong></td>
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<tr>
<td>3a. Retrospective</td>
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<td>No</td>
</tr>
<tr>
<td>3b. Prospective</td>
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<td>Yes</td>
<td>As determined by OPRS</td>
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<td><strong>4. Limited Data Set</strong></td>
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<td></td>
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<tr>
<td>4a. Retrospective</td>
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<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>4b. Prospective</td>
<td>Yes</td>
<td>N/A</td>
<td>Yes</td>
<td>As determined by OPRS</td>
</tr>
<tr>
<td><strong>5. Identified data for Recruitment Purposes</strong></td>
<td>Yes</td>
<td>N/A</td>
<td>Yes</td>
<td>No*</td>
</tr>
<tr>
<td><strong>6. Patient Identified Data</strong></td>
<td>Yes</td>
<td>N/A</td>
<td>Yes</td>
<td>Yes/As determined by OPRS</td>
</tr>
</tbody>
</table>

*Data set provided only for recruiting and obtaining consent from potential subjects.
How do investigators retrieve their data extracts?
In most cases, completed datasets will be loaded into a secure REDCap (Research Electronic Data Capture) database. Access can only be granted to the REDCap projects after all proper approvals and steps are completed. [Establish a REDCap account]

What are the IRB Requirements? (See the table on page 4 to determine if your project requires IRB approval – Note: After your completed DRA form is reviewed, you will receive a PDF copy with a personalized summary of your requirements and next steps.)

For projects without approved protocol:

1. The PDF copy of your Data Request Authorization (DRA) form should be submitted to IRB with your protocol.
   1. The CCTS Biomedical Informatics Core Manager will email you a PDF copy of the DRA form after it is reviewed.
   2. Submit the DRA form in OPRS Live under other documents
   3. Update protocol to indicate the source of data requested (i.e. CCTS - CRDW – Clinical Research Data Warehouse/UIC CIRCLE)
   4. Ensure your protocol document lists all data elements requested – this should match your DRA form
2. Once the IRB application is approved, submit the approved DRA form, updated protocol document, and a copy of IRB approval letter to the CCTS Biomedical Informatics Core Manager

For projects with prior approved protocol:

1. Complete the Data Request Authorization (DRA) form and submit IRB protocol documents to CCTS Biomedical Informatics Core Manager.
   1. If not already stated, update the protocol to indicate the source of data requested (i.e. CCTS - CRDW/UIC CIRCLE) and ensure your protocol document lists all data elements requested (this should match your DRA form)
   2. Submit IRB amendment with completed DRA form – you will retrieve a PDF copy of the DRA form that can be uploaded to OPRS Live.
   3. Once the IRB application is approved, submit the approved DRA form, updated protocol document, and a copy of IRB approval letter to the CCTS Biomedical Informatics Core Manager

OR

1. If the protocol document already lists all requested data elements and source of data, the CCTS Biomedical Informatics Core Manager will review these items to determine if further IRB approval is required.
   1. If all requirements are met, no additional IRB approval will be required.
   2. If it is determined that additional IRB approval is required:
      1. Submit IRB amendment with the completed DRA form – you will retrieve a PDF copy that can be uploaded to OPRS Live.
      2. Once the IRB application is approved, submit the approved DRA form, updated protocol document, and a copy of IRB approval letter to the CCTS Biomedical Informatics Core Manager.