

United States Renal Data System (USRDS) Agreement for Release of Data

Project Title	
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In this agreement, "Requester Organization" means	

- A. The National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK), through the United States Renal Data System (USRDS) Coordinating Center, will provide the Requester data extracted from the USRDS research database (the "Data"), via download or on CDs, DVDs, or other media type, which constitutes a Limited Dataset within the meaning of the HIPAA privacy regulations.
- B. The sole purpose of providing the Data is the conduct of legitimate and approved biomedical, cost-effectiveness, and/or other economic research by the Requester.
- C. The Requester shall not use the Data for purposes that are not related to biomedical research, cost-effectiveness, economic and/or other epidemiological research. Purposes for which the Data may not be used include, but are not limited to,
 - the identification and targeting of under- or over-served health service markets primarily for commercial benefit
 - the obtaining of information about providers or facilities for commercial benefit
 - insurance purposes such as redlining areas deemed to offer bad health insurance risks
 - adverse selection (e.g., identifying patients with high-risk diagnoses).

Any use of the Data for research not specified in the original proposal must be approved by the USRDS Project Officer (PO).

- D. The Requester shall not use the Data to identify individual beneficiaries or individual providers on the files.
- E. The Requester shall not combine or link the Data provided with any other collection or source of information that may contain information specific to individuals on the files, except where a waiver of authorization has been approved by the Requester's IRB/Privacy Board and NIDDK.
- F. The Requester shall not publish or otherwise disclose the Data in the files to any person or organization unless the Data have been aggregated (that is, combined into groupings of Data such that the Data are no longer specific to any individuals within each grouping), and no cells (aggregates of Data) contain information on fewer than ten individuals or fewer than five providers or facilities. The Requester shall not publish or otherwise disclose Data that identify individual providers or facilities, or from which such identities could be inferred.

However, the Requester may release Data to a contractor for purposes of data processing or storage if (1) the Requester specified in the research plan submitted to the USRDS Project Officer that Data would be released to the particular contractor, or the Requester has obtained written authorization from the PO to release the Data to such contractor, and (2) the contractor has signed a data release agreement with the PO.

- G. A copy of any aggregation of Data intended for publication shall be submitted to the PO for review for compliance with the confidentiality provisions of this agreement prior to submission for publication and, if not approved, shall not be published until compliance is achieved. The PO must respond within 30 days.
 - The Approval Request Checklist may be found at:
 - https://www.usrds.org/2016/appx/3 1 USRDS Manuscript Approval Request Checklist 16.pdf
- H. Appropriate administrative, technical, procedural, and physical safeguards shall be established by the Requester to protect the confidentiality of the Data and to prevent unauthorized access to it. The safeguards shall provide a level of security outlined in OMB Circular No. A-130, Appendix III Security of Federal Automated Information Resources, which sets forth guidelines for security plans for automated information systems in Federal agencies.
- I. No copies or derivatives shall be made of the Data in these files except as necessary for the purpose authorized in this agreement. The Requester shall keep an accurate written account of all such copies and derivative files, which will be furnished upon request to the PO. The USRDS Data files covered in this data use agreement (DUA) may be retained by the Requester until the date specified by the PO in the approval letter, at which time Requester may request renewal of this data use agreement to extend the retention period to comply with legal or institutional recordkeeping requirements or to maintain the integrity of the research or research publications. If at any time during the data retention period the DUA between USRDS and CMS is canceled, the Requester will be contacted to destroy the files in their possession. At the completion of the activities in the research plan, the file(s) and any derivative files and copies shall be destroyed. At that time, the Requester will inform the USRDS and the PO in writing that the files have been destroyed.
- J. For the purpose of inspecting security procedures and arrangements, authorized representatives of the NIDDK and/or of CMS will, upon request, be granted access to premises where the Data are kept.

USRDS AGREEMENT FOR RELEASE OF DATA

K. The following USRDS Data file(s) is/are covered under this Agreement.

Standard Analysis Files (SAFs) requested	d:
□ Core	\square Dialysis Morbidity and Mortality Study (DMMS)
☐ Transplant	\square Comprehensive Dialysis Study (CDS)
☐ Hospital	\square Clinical Performance Measures
☐ CKD 5% Cohort Core	☐ Case Mix Adequacy (CMA)
\square CKD 5% Cohort Hospital	\square Active-Adipose Study (AAS)
☐ CROWNWeb Clinical Data	
For the following SAFs, indicate the clair	n year(s) requested as well:
\square Institutional Claims (pre-1989 through 201	4 available)
☐ Physician/Supplier Claims (1991–2014 av	vailable)
☐ Part D (2006–2014 available)	
\square Pre-ESRD Institutional Claims (incident y	rears 1995-2014)
\square Pre-ESRD Physician/Supplier Claims (in	ncident years 1995-2014)
☐ Pre-ESRD Part D (incident years 2008-2014	
☐ CKD 5% Institutional Claims (1992–2014	available)
☐ CKD 5% Physician/Supplier Claims (199	92–2014 available)
☐ CKD 5% Part D (2006–2014 available)	
Other:	
☐ Provider Crosswalk	☐ Physician Crosswalk
Requester Signature (for the Institutional C	Official for Data Assurance)
Authorized Signatory (name, title & date)	
Requester Address	
Requester Telephone Number	

USRDS AGREEMENT FOR RELEASE OF DATA

Investigator / Analyst signature	Name	Date
Investigator / Analyst signature	Name	Date
Investigator / Analyst signature	Name	Date
Investigator / Analyst signature	Name	Date
(attach additional signature pages as n	ecessary)	
USRDS Project Officer Signature		Date
Checklist:		Date
Checklist: DID YOU REMEMBER TO INCLUDE		Date
Checklist: DID YOU REMEMBER TO INCLUDE □ Signed copy of your institutions □ Copy of your project proposal in	al IRB approval memo	
Checklist: DID YOU REMEMBER TO INCLUDE □ Signed copy of your institutions □ Copy of your project proposal is http://www.usrds.org/2016/a	al IRB approval memo n recommended format at	ls using USRDS data.pdf
DID YOU REMEMBER TO INCLUDE □ Signed copy of your institutions □ Copy of your project proposal is http://www.usrds.org/2016/a □ Copy of this Data Use Agreement participants. Please note that any MODIFICATION additional files, require a new IRB	al IRB approval memo n recommended format at ppx/3/1 Outline for research proposal	ls using USRDS data.pdf PI, and all active whether they require original project proposal

06/14/2017 revision