ImmPort Contract
SUBJECT DE-IDENTIFICATION PROCESS

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National Institutes of Health (NIH)
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Division of Allergy, Immunology, and Transplantation (DAIT)

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**SUBJECT DE-IDENTIFICATION PROCESS**

**SUBJECT DE-IDENTIFICATION SUMMARY**

The ImmPort team follows the Safe Harbor Method principles ([http://www.hhs.gov/ocr/privacy/hipaa/understanding/coveredentities/De-identification/guidance.html](http://www.hhs.gov/ocr/privacy/hipaa/understanding/coveredentities/De-identification/guidance.html)) in data capture templates and in data submission by:

- Rounding any “ages” submitted > 89 to 90 and add a comment indicating that this was done
- Not recording any dates within ImmPort templates; all time points and schedules are referenced to study day 0 (or a designated day in the study as determined by the data provider);
- Not recording any location information in ImmPort templates
- Not capturing any identifying information in ImmPort templates itemized in the Safe Harbor Method (Name, accounts, phone or fax numbers, email addresses, etc)

The ImmPort team severs the connection to subject identifiers in the source data management system by:

- If possible, surveying the data provider to determine whether local subject identifiers were used in the submission that would point back to a local system
- Hiding subject source identifiers, sample source identifiers, or experiment sample source identifiers in the ImmPort application to cut linkage with the source system
- Not exporting the subject source identifiers, sample source identifiers, or experiment sample source identifiers in any ImmPort generated data packages delivered online, at sFTP sites, or in the future, via Amazon AWS S3 storage
- Modifying result filenames to ImmPort accession numbers for individual result files (such as FCS, CEL, etc) if the filename may link back to subject source identifiers
- Not providing original, unstructured result files submitted by sites if they (1) haven’t stated concurrence with the disconnect with source identifiers; and/or (2) if the data can be parsed into standard ImmPort templates and source identifiers hidden

The ImmPort team does not share GWAS level SNP genotyping data and Next Gen sequencing data within ImmPort, and a formal request to NIAID similar to dbGAP is required for the data to be obtained.

For data provided in more “native” format (i.e., not in ImmPort template format), such as result files from closed clinical trials (assessments, lab tests, adverse events, etc):

- If the data source is ITN TrialShare, the dates are already obfuscated and are not identifying as are the subject identifiers; no special treatment is required
- For other data sources with closed clinical trial data (or the equivalent), data files are sent to our Stanford team’s HIPAA server and evaluated by HIPAA trained staff to (1) Identify any potentially identifying information, such as names, dates, locations, etc… (2) Remove any identifying information and normalize dates to study day 0; (3) Comment and Description fields are removed, unless determined to be completely devoid of identifying information or identifiers; (4) Return the data to NG staff for submission and processing