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Synergies between centralized and federated approaches to data quality: a report from the national COVID cohort collaborative

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Research and Applications

Objective: In response to COVID-19, the informatics community united to aggregate as much clinical data as possible to characterize this new disease and reduce its impact through collaborative analytics. The National COVID Cohort Collaborative (N3C) is now the largest publicly available HIPAA limited dataset in US history with over 6.4 million patients and is a testament to a partnership of over 100 organizations.

Materials and Methods: We developed a pipeline for ingesting, harmonizing, and centralizing data from 56 contributing data partners using 4 federated Common Data Models. N3C data quality (DQ) review involves both automated and manual procedures. In the process, several DQ heuristics were discovered in our centralized context, both within the pipeline and during downstream project-based analysis. Feedback to the sites led to many local and centralized DQ improvements.

Results: Beyond well-recognized DQ findings, we discovered 15 heuristics relating to source Common Data Model conformance, demographics, COVID tests, conditions, encounters, measurements, observations, coding completeness, and fitness for use. Of 56 sites, 37 sites (66%) demonstrated issues through these heuristics. These 37 sites demonstrated improvement after receiving feedback.

Discussion: We encountered site-to-site differences in DQ which would have been challenging to discover using...
federated checks alone. We have demonstrated that centralized DQ benchmarking reveals unique opportunities for DQ improvement that will support improved research analytics locally and in aggregate.

**Conclusion:** By combining rapid, continual assessment of DQ with a large volume of multisite data, it is possible to support more nuanced scientific questions with the scale and rigor that they require.

**Key words:** electronic health records, data accuracy, COVID-19

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**INTRODUCTION**

COVID-19 has precipitated a worldwide public health emergency requiring responsive action from all branches of medical science, including informatics. The National COVID Cohort Collaborative (N3C), sponsored by the NIH National Center for Advancing Translational Sciences (NCATS), is a data-driven response to this challenge. Through leading-edge technology, N3C uses harmonized electronic health record (EHR) data to support pioneering collaborative research that spans the full COVID disease cycle—from risk factors, to disease progression, to treatment decisions. The result of this collaboration is a research environment that addresses technical, legal, and policy barriers to rapid discovery and dissemination of actionable clinical findings to optimize the acute and long-term health outcomes of diverse populations nationwide.

At the heart of the N3C collaborative is a centralized enclave of EHR data assembled from Clinical and Translational Sciences Award (CTSA) hubs, Institutional Development Award (IDeA) Networks for Clinical and Translational Research (IDeA-CTR) hubs, and the OCHIN network. As of this writing, it is the largest ever assembly of harmonized EHR data for research in the United States, comprising 6.4 million patients from 56 sites including 7.2 billion rows of data. N3C ingests and harmonizes patient-level EHR data from participating sites for patients with positive COVID-19 tests or whose symptoms are consistent with COVID-19. Additional records collected include persons who have tested negative for COVID-19 (and have never tested positive) to support comparative studies.4 Data harmonization is made possible through the efforts of the National Center for Data to Health (CD2H) and subject matter experts from Observational Health Data Sciences and Informatics (OHDSI), the Patient-Centered Clinical Research Network (PCORnet), the Acruial to Clinical Trials (ACT) network, and TriNetX.

**Precursors to N3C: federated data networks**

Clinical data repositories from EHR sources have evolved over the decades. Ad hoc database designs from the early period of EHR adoption had limited generalization across sites. The earliest federated models in the 1990s ultimately gave rise to Sentinel and to Common Data Models (CDMs) such as PCORnet, the ACT network, TriNetX, and Observational Medical Outcomes Partnership (OMOP), which later became OHDSI. In a federated data network, each participating site’s data stay behind its institutional firewall, but are structured according to a CDM. This enables queries and results to be shared across sites rather than raw data.

The CDMs have been key to using EHR data for research; however, secondary analytic uses—particularly involving multiple contributing sites—require resource-intensive quality control to achieve the required uniformity and specificity to support open-ended research. Despite their importance, data quality (DQ) checks are inconsistently applied and implementation methodologies are largely not evaluated. Further, evaluations of DQ often fail to accurately determine the data’s “fitness for use,” which evaluates both its intrinsic and intentional aspects. DQ evaluations must take into account the initial purpose of the data in their source systems, as well as the intended use of these data once harmonization processes have transformed them for use in secondary research. Each of the aforementioned federated networks has methods to promote local DQ and adherence to data model conventions; these methods vary in maturity. As examples, OHDSI offers its DQ Dashboard tool for sites to run against their local CDM in order to evaluate adherence to OMOP CDM convention and diagnose common issues; PCORNet requires a quarterly data “curation” and quality check that uses prepackaged SAS scripts; and ACT has a “smoke test” to ensure federated network query response and has a DQ Dashboard of its own under development; and TriNetX employs a growing library of DQ metrics and visualizations with site benchmarking as the basis for evaluating the results. Because the data are federated, DQ evaluation and remediation are performed locally at the site-by-site personnel. These local checks ensure that data conform to the specifications of the chosen CDM, and may also check for data anomalies such as statistical outliers, invalid dates, biological implausibility, abundant missing data, and other common clinical data issues. Such approaches support the alignment of data; however, with the data remaining behind the institutional firewalls, it can be difficult to assess conformance or determine overall variability across sites, especially if the CDM does not have the capability of executing ad hoc DQ-related queries across the data network.

**N3C’s centralized approach**

In contrast to the federated approach, N3C pools data from each partner site in accordance with its signed Data Transfer Agreement and harmonizes all submitted data to the OMOP CDM. By the time, data are submitted to N3C, sites have already applied a layer of local DQ checks, and are submitting data that are “clean” by the local definition. Once data are merged across sites, additional opportunities for improvement may become apparent due to the ability to efficiently compare and benchmark among similar sites. N3C centralized DQ methods complement and augment the foundation accomplished by sites in the context of the federated networks.

**Data quality**

DQ emerged in the 2010s as an explicit subject of attention and research in informatics. Notably, the Patient-Centered Outcomes Research Institute (PCORI) funded an effort to define DQ, resulting in the “Harmonization” framework of Kahn and colleagues. The dimensions of DQ defined there (internal verification, external validation, conformance, completeness, and plausibility) are the basis of the PCORNet and OHDSI conception and framework for DQ checking. N3C continued to build on this foundation for our centralized DQ approach.
OBJECTIVE

To describe N3C’s approach to DQ curation and explore the value that a centralized data architecture and approach to DQ adds to what sites can accomplish locally, either alone or in the context of a federated data research network.

MATERIALS AND METHODS

N3C’s data harmonization pipeline

N3C has engaged in a precedent-setting endeavor to centrally harmonize the 4 major CDMs (OMOP, PCORnet, ACT, and TriNetX) into OMOP (Figure 1). This enables N3C’s EHR dataset to span 56 institutions at the time of this writing while putting minimal burden on sites themselves. The EHR data spans multiple data domains, including patient demographics, encounter details, diagnoses, procedures, medications, lab tests, and clinical observations; a detailed list of supported fields can be found in the OMOP 5.3.1 common data model specification. The pipeline evolved over 3 implementation environments and is presently running in Palantir Foundry.

Centralized DQ assessment

After transformation to OMOP, each site’s inaugural N3C submission is loaded into the N3C Data Quality Portal (DQP), which performs a series of automated DQ checks prior to manual evaluation by the Data Ingestion & Harmonization (DI&H) team (see “Acknowledgments” for members). The DQP is built on queries similar to OHDSI’s DQ Dashboard and provides a series of targeted visualizations that allow the team to assess each site’s DQ in areas of importance for COVID research (see Table 1). The DQP supports site data review both in the context of the site’s source CDM and in comparison to the other 3 CDMs.

N3C purposely uses a light touch during these DQ checks, placing a high value on including as much submitted data as possible, with the understanding that each site’s data likely contains local idiosyncrasies and inconsistencies that are acceptable so long as they are known. Such issues (e.g., a site is able to provide only outpatient data, only supports a subset of vital signs, or is frequently missing units of measure) can be reported in N3C’s release notes, but would not prevent a site from “passing” on to inclusion in that week’s release. Issues that can prevent sites from passing (“Must Pass”), and thus make up our minimum data standards, are detailed in Table 1.

Providing feedback to sites

As previously described, N3C’s signature “white glove,” or one-on-one DQ support, provides feedback and individualized source model-specific help to sites. Members of N3C’s Phenotype and Data Acquisition (P&D&A) team (see “Acknowledgments” for members) serve as liaisons between sites and the DI&H team. The P&D&A team is composed of subject matter experts in each of N3C’s 4 supported data models. Each site is assigned an expert in their source model as their P&D&A point of contact. After a site’s initial payload is evaluated using the DQP, the site’s P&D&A contact compiles a list of data issues in the “must pass” and “heads up” categories and emails that list to the site. Corrections are generally iterative in nature, and correction cycles will continue until the site passes all “must pass” checks. Some corrections are simple, while others require individual troubleshooting meetings with the P&D&A team, attendance at P&D&A office hours, or sharing code snippets.

More recently, we have started an initiative to provide some of the benchmarking data generated by the DQP directly to sites, in visual format. Our centralized architecture gives us the unique ability to provide this type of benchmarking data and may reveal opportunities for DQ improvement of which sites were previously unaware. One of the visualizations we sent, a heatmap illustrating “coverage” of different vital signs for COVID inpatients across a variety of anonymized sites, is shown in Figure 2. These visualizations allow sites to compare their DQ to that of other sites that are using the same CDM. Hierarchical clustering was used to bring together sites with similar profiles of vital reporting.

Assessing N3C’s DQ impact

To assess the impact of N3C DQ feedback on sites’ local DQ, we reviewed all submitted DQ issues filed on site data in the N3C Data Enclave and performed a qualitative analysis to extract DQ heuristics, as well as the number of sites to which each heuristic applied. Only “released” sites (i.e., sites whose data are available for research in the Enclave) were included in the analysis; a denominator of 56 sites. Issue instances were counted if they were in the “Must Pass” category. Sites that are still working through data issues (and are thus not yet released) are not included in the denominator, and “Heads Up”-type issues or simple formatting errors (e.g., incorrect delimiters, missing headers, etc.) were not included in the count of issues.

RESULTS

Table 2 provides an accounting of the DQ issues found and improvements made by N3C-participating sites based on our feed-
Table 1. Data quality issue types

<table>
<thead>
<tr>
<th>Check type</th>
<th>Data checks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source CDM conformance</td>
<td><strong>Must Pass</strong>: All tables required by the native CDM specs are present, with all CDM-required fields populated; fields that use a controlled value set (eg, “M” for male, “F” for female, etc.) are populated with valid values</td>
</tr>
<tr>
<td>Demographics</td>
<td><strong>Must Pass</strong>: Count of patients qualifying for COVID phenotype is reasonable when compared with sites of similar size; sex, race, and ethnicity distributions reasonable for the site’s population; month of birth evenly distributed throughout the calendar year <strong>Heads Up</strong>: &gt;20% of race or ethnicity is missing or “No Matching Concept”</td>
</tr>
<tr>
<td>COVID tests</td>
<td><strong>Must Pass</strong>: All COVID tests must be coded with an OMOP standard concept (or, for non-OMOP source data, the LOINC equivalent); all COVID test results must be coded with an OMOP standard concept (or, for non-OMOP source data, the equivalent controlled vocabulary term); numbers of negative and positive COVID tests are reasonable when compared with sites of similar size <strong>Heads Up</strong>: High numbers of COVID tests with null results</td>
</tr>
<tr>
<td>Conditions</td>
<td><strong>Must Pass</strong>: Clinical encounters are present that are coded with U07.1 (ICD-10 code for COVID), and those encounters are distributed across various visit types (eg, outpatient, inpatient, emergency)</td>
</tr>
<tr>
<td>Encounters</td>
<td><strong>Must Pass</strong>: Clinical encounters are distributed across a variety of standard visit types (eg, outpatient, inpatient, emergency); the distribution of visit types is reasonable when compared with similar sites; the majority of inpatient visits have valid end dates; the mean duration of visits of various types is reasonable for that type of visit; the vast majority of visit end dates are later than or equal to the visit start date</td>
</tr>
<tr>
<td>Measurements/observations</td>
<td><strong>Heads Up</strong>: The site supports only a small number (eg, 5–10) of unique measurement or observation types</td>
</tr>
<tr>
<td>Coding completeness</td>
<td><strong>Must Pass</strong>: No more than 20% of records in any domain are coded with nonstandard OMOP concept IDs without further explanation (OMOP sites only); no more than 20% of records in any domain are coded with “0—No Matching Concept” without further explanation (affects OMOP sites only); the PERSON_ID attached to all records in domain tables must exist in the PERSON table; primary keys are valid (ie, no duplicate rows in any table); if applied by the site, date shifting is consistent within each patient across all domains</td>
</tr>
<tr>
<td>Fitness for use</td>
<td>Use of the data by researchers often reveals additional DQ issues for one or more sites (eg, sparsely populated body mass index data, in the context of a study of obesity and COVID). In these cases, we report the findings to sites so that they can take action in their local data if they wish to have their site’s data included in the study</td>
</tr>
</tbody>
</table>

“Must Pass” and “Heads Up” data check for release into the N3C Data Enclave.

DQ: data quality; N3C: National COVID Cohort Collaborative; OMOP: Observational Medical Outcomes Partnership.

back. These improvements map to the checks we perform against each site’s first payload, detailed earlier in Table 1.

These heuristics revealed DQ issues in 37 (66%) of the 56 sites, all of which demonstrated improvement after receiving feedback from N3C. Selected examples are detailed here.

Example of heuristic #1: not using (or improperly using) source CDM’s controlled vocabulary in one or more fields

There are numerous examples of fields that require controlled vocabularies among the 4 source CDMs. Nearly a quarter of all N3C sites have violated the use of these vocabularies on one or more occasions within their source CDM. Examples of this issue include nonuse or improper use of the ACT race and ethnicity vocabulary (eg, using local codes rather than the controlled vocabulary’s value sets), incorrect DX and DX_TYPE agreement (eg, labeling an ICD-10-CM diagnosis code as type “Other”) in PCORnet, or using standard concepts in an inappropriate domain (eg, filing conditions wrongly in the OBSERVATION table) in OMOP. These types of errors often create the illusion of missing data. Having centralized access to sites’ source data gives the N3C team the ability to diagnose the issue in detail and offer sites ways to remedy the problem.

Example of heuristic #2: COVID test result values not standardized or null

Eleven sites submitted nonstandard, null, or otherwise unusable COVID test result values in their initial submissions. Quality issues included:

- Nonharmonizable COVID test results (eg, submitting a free-text result rather than the source CDM’s controlled vocabulary equivalent).
- Null COVID test results in excess of a reasonable number of pending results.
- COVID test results that used the source CDM’s controlled vocabulary, but mapped to an unusual concept (eg, one site mapped results to OMOP concept ID 45877980, “Not,” presumably for a negative test).

Even where it is not a source CDM requirement to map every qualitative test result to the CDM’s controlled vocabulary, for N3C’s use case, harmonized COVID tests are essential. We worked with sites to prioritize these mappings, even as new COVID test codes continued to emerge over the course of the pandemic. Figure 3 shows the improvements made by these 11 sites over time.

Example of heuristic #4: implausible distribution of visit types

Each source CDM has its version of an encounter or visit table, as well as a controlled vocabulary to assign a “type” to visits, such as inpatient, outpatient, or emergency. Different models’ vocabularies have different levels of specificity for visit types; the N3C DQ process accounts for this by aggregating multiple valid visit type codes to a higher-level category (eg, “Inpatient Hospital” and “Inpatient Visit” in the OMOP vocabulary can roll up to an overall category of Inpatient for the purposes of quality analysis). Even when these roll-ups are taken into account, however, 7 N3C sites had implausible
distributions of visit types, where implausibility was defined by the overall distribution across sites. Such implausibility is illustrated here for 5 of the 7 sites that specifically had an implausible proportion of in-patient visits. One of the 5 sites started with inpatient visits significantly above the mean (Figure 4A), while the other 4 had significantly fewer (Figure 4B). As shown in Figure 4, after feedback from N3C, all 5 sites improved the quality of their visit type mappings to bring their in-patient visit proportion closer to the mean. Of note, the proportions at times worsened with subsequent payloads, pointing out the need for recurring vigilance: Fixing a problem once does not mean it stays fixed.

Example of heuristic #15: data utility challenges
A number of N3C DQ findings have come from analysts using the data, spotting inconsistencies, and submitting issue tickets. One such example involves data on mortality, the most commonly investigated endpoint in N3C. Generally, sites document patient deaths in their source CDM with pairs of patient IDs and death dates. However, because not all CDMs require a death date to be present to note a patient as deceased, numerous sites provide patients IDs, but no dates. In one instance, a site’s Death data table included all of their patient IDs (for both living and deceased patients); in their definition, a null date denoted a living patient, and a populated date denoted a death. Taken individually, each of these structures makes sense. However, aggregate analyses that either ignore missing dates or require dates to be present would come to drastically different conclusions, either over- or underestimating mortality. In these cases, we worked individually with sites to standardize where possible, and in other cases, provided user education and analytical workarounds. Other researcher-identified site-level issues include those tied to a particular type of measurement, such as a misrepresented unit of measure for a specific type of value (discovered when comparing height/weight calculated body mass index [BMI] to reported BMI) or mismapped measurements (discovered when reviewing mean SpO2 by site). In each case, the DQ concern was referred for further “upstream” remediation in the pipeline beyond the project that discovered it and the site was informed.

Table 2. Data quality heuristics

<table>
<thead>
<tr>
<th>No.</th>
<th>Heuristic</th>
<th>Type</th>
<th>No. of sites</th>
<th>sites (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Not using (or improperly using) source CDM’s controlled vocabulary in one or more fields</td>
<td>Source CDM conformance</td>
<td>13</td>
<td>23.2</td>
</tr>
<tr>
<td>2</td>
<td>COVID test result values not standardized or null</td>
<td>COVID tests</td>
<td>11</td>
<td>19.6</td>
</tr>
<tr>
<td>3</td>
<td>Lacking/incorrectly populating field(s) required by source CDM</td>
<td>Source CDM conformance</td>
<td>9</td>
<td>16.1</td>
</tr>
<tr>
<td>4</td>
<td>Implausible distribution of visit types (eg, 75% inpatient)</td>
<td>Encounters</td>
<td>7</td>
<td>12.5</td>
</tr>
<tr>
<td>5</td>
<td>Large number of “No Matching Concept” records (OMOP source only)</td>
<td>Coding completeness</td>
<td>6</td>
<td>10.7</td>
</tr>
<tr>
<td>6</td>
<td>Lacking table(s) required by source CDM</td>
<td>Source CDM conformance</td>
<td>5</td>
<td>9.0</td>
</tr>
<tr>
<td>7</td>
<td>Many or all inpatient visits lacking valid end dates</td>
<td>Encounters</td>
<td>5</td>
<td>9.0</td>
</tr>
<tr>
<td>8</td>
<td>Few or no clinical encounters coded with U07.1</td>
<td>Conditions</td>
<td>5</td>
<td>9.0</td>
</tr>
<tr>
<td>9</td>
<td>Implausible count of patients qualifying for phenotype</td>
<td>Demographics</td>
<td>3</td>
<td>5.4</td>
</tr>
<tr>
<td>10</td>
<td>Small number of unique measurement/observation types</td>
<td>Measurement/observation</td>
<td>2</td>
<td>3.6</td>
</tr>
<tr>
<td>11</td>
<td>PERSON IDs in fact tables that are not in the PERSON table</td>
<td>Coding completeness</td>
<td>2</td>
<td>3.6</td>
</tr>
<tr>
<td>12</td>
<td>Primary keys are not unique</td>
<td>Coding completeness</td>
<td>2</td>
<td>3.6</td>
</tr>
<tr>
<td>13</td>
<td>Inconsistent local date shifting causing implausible timelines</td>
<td>Coding completeness</td>
<td>2</td>
<td>3.6</td>
</tr>
<tr>
<td>14</td>
<td>Implausible demographics (eg, 100% male patients)</td>
<td>Demographics</td>
<td>2</td>
<td>3.6</td>
</tr>
<tr>
<td>15</td>
<td>Data utility challenges (eg, missing mortality data)</td>
<td>Fitness for use</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Items compiled here are from a qualitative analysis of the “Must Pass” data issues filed on any one of the 56 currently released N3C sites that resulted in a fix by the site. Fitness for Use is an additional heuristic that applies to all sites and is thus also included here. Simple formatting errors (eg, incorrect delimiters) and noncritical “Heads Up” issues are excluded from this analysis.

aDenominator: 56 sites; 37 unique sites are represented across these categories.

N3C: National COVID Cohort Collaborative; OMOP: Observational Medical Outcomes Partnership.
DISCUSSION

We have demonstrated that centralized DQ assessment reveals unique opportunities for iterative quality improvement for submitting sites. N3C’s DQ checks take into account various dimensions introduced in prior DQ work, such as conformance to source data models, density and completeness (eg, of COVID test results), and plausibility (eg, of percentage of inpatient visits). By participating in a consortium like N3C, sites receive routine feedback on their overall quality with tactical information on ways to address local issues. Moreover, N3C’s dedicated team of analysts with protected time to concentrate on DQ, deep subject matter expertise, and access to powerful visualization tools enable efficient support for participating sites in making rapid improvements in high-priority areas. This process can be transformative; by combining efficient, continual assessment of DQ with a large volume of multisite data, it is possible to support more nuanced scientific questions with the scale and rigor that they require.

Centralized data enable site-to-site comparisons

When examining data across N3C partner sites, the centralized approach revealed significant site-to-site DQ differences that would have been challenging to discover in isolation. Many of these “data issues” are indeed errors, but others arise from differences in interpretation or adoption of CDM components. Sites’ use of the encounter data domain is an example, where the definition of “one visit” can vary widely depending on the site’s EHR, organizational structure, or billing practices (see Figure 5). The site’s definitions of various visit types are generally not erroneous and likely comply with the rules of their source CDM. Problems surface, however, when data are combined across sites for multisite projects, leading to a need for data users to (1) understand that the issue exists and (2) develop consistent analytic workarounds and harmonization strategies. Benchmarking, or the ability to compare sites with their peers, is an efficient way to catch such issues.

Assessing the quantity and variety of data available per patient is another use case for benchmarking. Because instantiation and maintenance of CDMs are resource-intensive, it is common for sites to take a minimalistic approach to CDM data curation, particularly in the early days of implementing a new CDM. This may entail purposely
Choosing to support only a small number of lab tests, vitals, or other observations (eg, the top 50 most common labs; only blood pressure, weight, and height out of all possible vital measurements, etc.). However, this sparsity can have a big impact on what research questions are possible to answer using the data. Because N3C aims to be a multipurpose resource for COVID research, we use benchmarking to spot sites supporting fewer facts per patient (or a limited variety of clinical concepts) and encourage them to gradually add on as they can.

Benchmarking data can also serve as a clear and persuasive DQ communication tool. As shown in Figure 2, our ability to compare sites’ coverage of lab and vital concepts across sites gives sites a “report card” to see which concepts have more or less coverage at their peer sites. This information can help sites prioritize bringing in new types of data, or may spur an investigation of a data issue of which the site was unaware. It should be noted that federated networks (particularly PCORnet and TriNetX) are also capable of site-to-site benchmarking, but at an aggregated rather than detailed level. To use Figure 5 as an example, aggregate checks could produce the inpatient visit counts reported on the left side, but would not enable the row-level deep dive shown on the right. Though row-level checks enable more detailed benchmarking, both aggregate and row-level comparisons are highly valuable.

Centralized data enable crowd sourced DQ evaluation

The N3C repository not only pools data but also brings together multidisciplinary teams of clinicians, researchers, and statisticians from across the network of organizations with N3C Data Use Agreements. N3C Enclave researchers are invited to join any of more than 25 Clinical Domain Teams, where they can share domain-specific knowledge with a diverse set of peers. Researchers who join a Domain Team have access to a shared workspace where they can create sets of derived variables specific to their research question and conduct-related analyses. As detailed dataset reviews are an essential step in this process, these teams may reveal resolvable data issues such as hard-to-harmonize variations among contributing CDMs or incorrect mappings at the individual record level. Sometimes these quality errors, inconsistencies, or omissions are remedied within the Enclave by N3C, such as unit of measure harmonization, whereas for other issues these discoveries are referred back to the contributing site for remediation.

The ability for research teams to review row-level data in the N3C Enclave also helps end-users write more accurate analytic code by avoiding blind assumptions related to how representation of clinical facts varies by site or CDM. For example, when creating a flag to indicate the co-occurrence of a positive COVID test and a diagnosis code representing a comorbidity, date logic is applied in the code. Review of row-level data, which can be filtered to one or more sites whose variable distribution does not match other sites, may reveal that some measurement dates are representative of test result date instead of test order date—either of which may be acceptable in the source CDM. Though the data are not “wrong” in this case and do not need to be corrected by the contributing site, this ambiguity is a DQ issue nonetheless and requires resolution by data users during analysis. In general, identification of DQ issues that are highly dependent on the context of use is, by design, left for analysts pursuing specific questions to discover. In such cases, our centralized DQ team serves as a liaison to the sites and passes feedback from analysts to sites for local discussion and remediation, if deemed necessary and high value by the site. Issues of this type are extremely challenging or impractical to identify without access to the row-level data, in addition to the ability to compare across sites.

Centralization increases DQ efficiency

While centralization alone is not a recipe for improved DQ, it does present opportunities to implement generalized solutions at scale. N3C adopted 2 postprocessing DQ processes that illustrate this: (1) interrogating information “loss” in standardized terminology mappings and (2) performing post hoc evaluation of harmonized clinical information to ensure analytical utility. These solutions are more practical to achieve in a centralized environment where economy of scale provides the ability to easily see strings or terms that could be supported in value sets, or logic that could be modified to bring clinical information to the right analytical domain. Moreover, a centralized data ingestion pipeline allows for bidirectional improvement as the target model (OMOP, in this case) can evolve to include conventions that capture the heterogeneity of source system data.

Finally, centralized review simply puts additional trained eyes on a site’s data. The advantages to this are demonstrated by the fact that...
that N3C’s centralized review process found errors and room for improvement even in data that had passed either local review or one of the federated networks’ required checks. Indeed, this combination of local checks, by personnel who intimately understand their site’s data, and centralized checking, which can take advantage of economies of scale, may represent an ideal model for DQ assessment.

Limitations

N3C is devoted to a single disease, so some of our particular DQ checks derive from that focus. Though many of the issues identified here (e.g., encounter definitions, unit harmonization, plausibility) are not limited to COVID-related data, the emphasis of our suite of DQ checks is on variables required for COVID research. As an example, we ensure careful harmonization of qualitative COVID lab test results, but do not perform checks at the same level of detail on non-COVID labs. Still, many of our checks listed in Table 1 can apply to clinical data more generically.

Centralization does mean that data are further from the EHR source, and we rely on local staff to be final arbiters of their own DQ. In addition, as sites may have already executed one of the federated networks’ DQ checking protocols prior to submitting to N3C, the data that N3C receive may have already undergone a prior round DQ improvements. Yet, based on our results, we feel that the insights that sites receive from the composite experience of the entire Enclave add value even to previously improved data.

Despite the efficiency that N3C’s centralized Enclave enables for the assessment of DQ issues, much of the heavy lift of definitive DQ remediation lies with the submitting sites. We acknowledge that consortial data resources such as N3C are only possible because of the efforts of local teams and believe that centralized DQ supports collaboration and knowledge exchange that also helps improve local DQ. In informatics research, there is generally limited funding available to specifically support local DQ, which makes N3C sites’ engagement with our DQ process all the more impressive. This may be an indicator of demand for more centralized DQ in the future, given ongoing funding to do so. The ability to take some of the DQ workload off of local sites may incentivize site participation in future centralized repositories.

CONCLUSION

Federated data repositories, where the data remain at the generating site, offer the advantages of local curation by personnel deeply familiar with the data. Central repositories enable efficient DQ benchmarking at scale, and the generation of derivative, harmonized variables and units of measure for comparable and consistent analytics. Together, these advantages can synergize to a best of both worlds approach for DQ improvement and enhancement in clinical data repositories. Cooperation and communication between these complementary environments, as illustrated by the common data model communities and N3C, promise mutual advantage and maximal DQ.

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REFERENCES


