The effects of electronic medical record phenotyping details on genetic association studies: HDL-C as a case study

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Abstract

Background: Biorepositories linked to de-identified electronic medical records (EMRs) have the potential to complement traditional epidemiologic studies in genotype-phenotype studies of complex human diseases and traits. A major challenge in meeting this potential is the use of EMR-derived data to extract phenotypes and covariates for genetic association studies. Unlike traditional epidemiologic data, EMR-derived data are collected for clinical care and are therefore highly variable across patients. The variability of clinical data coupled with the challenges associated with searching unstructured clinical notes requires the development of algorithms to extract phenotypes for analysis. Given the number of possible algorithms that could be developed for any one EMR-derived phenotype, we explored here the impact algorithm decision logic has on genetic association study results for a single quantitative trait, high density lipoprotein cholesterol (HDL-C).

Results: We used five different algorithms to extract HDL-C from African American subjects genotyped on the Illumina Metabochip \((n = 11,519)\) as part of Epidemiologic Architecture for Genes Linked to Environment (EAGLE). Tests of association between HDL-C and genetic risk scores for HDL-C associated variants suggest that the genetic effect size does not vary substantially across the five HDL-C definitions.

Conclusions: These data collectively suggest that, at least for this quantitative trait, algorithm decision logic and phenotyping details do not appreciably impact genetic association study test statistics.

Keywords: Electronic medical record, Genetic risk score, HDL-C, eMERGE network, PAGE I study

Background

Biorepositories linked to de-identified electronic medical records (EMR) are an emerging resource for genetic association studies [1]. Compared with traditional epidemiologic studies, EMR-based studies offer multiple advantages including relative ease of ascertainment, rapid accrual of samples and associated data, longitudinal measures, and the potential for lengthy follow-up. Another major advantage of EMR-based or clinic-based studies is their potential for pharmacoepidemiology and other applications associated with personalized medicine.
While clinic-based studies linked to EMRs offer multiple advantages, they also offer multiple challenges when accessed for research such as genetic association studies. A major challenge of the EMR is that the data are not collected for research purposes; that is, the data are collected as part of routine clinical care. Therefore, unlike traditional epidemiologic studies, there is no “baseline” measurement or examination of all study participants, and the number of overall measurements and exams can vary widely by patient. This variability is in stark contrast to longitudinal epidemiologic studies where participants are surveyed and examined uniformly every few years.

Because of the variable and somewhat erratic nature of the EMR data, investigators accessing these data for genetic association studies must make specific decisions in developing phenotype algorithms designed to extract outcomes and covariates for analysis. For example, for a commonly studied measurement such as body mass index, the investigator has multiple options including the first height and weight mentioned, the last height and weight mentioned, an average of all heights and weights mentioned for all clinic visits, the height and weight mentioned closest to another clinical diagnosis (such as type 2 diabetes), and so on.

Many of the challenges associated with EMR-based phenotyping are being addressed by collaborative consortiums such as the electronic MÉdical Records and GÉnomics (eMERGE) network, a cooperative group of several DNA biorepositories in the United States linked to EMRs funded by the National Human Genome Research Institute [2,3]. A major goal of the eMERGE network is the development of portable algorithms designed to define disease outcomes for use in genetic association studies [4]. Algorithms developed under eMERGE have been used successfully for single study site [5,6] and as eMERGE-wide studies [7-13], the latter of which demonstrate the portability of these algorithms despite possible variations in clinical practice. A portion of the eMERGE EMR-derived phenotypes have also been mapped back to PhenX variables using the PhenX Toolkit [14], suggesting that EMR-derived phenotypes are comparable to epidemiologic collected phenotypes [15].

The eMERGE network has been successful in designing and implementing EMR-based algorithms for multiple phenotypes; however, it is unclear if the decision logic underlying each algorithm for phenotypes with repeated measures impacts downstream analyses for genetic association studies. To explore this possible impact, we as the Epidemiologic Architecture for Genes Linked to Environment (EAGLE) as part of the larger Population Architecture using Genomics and Epidemiology (PAGE) I study [16] conducted a genetic association study for the commonly measured and studied high density lipoprotein cholesterol (HDL-C). We created five HDL-C algorithms to extract this quantitative trait from African American subjects genotyped with the Illumina Metabochip [17] and available in BioVU, the Vanderbilt biorepository linked to de-identified electronic medical records [18]. Overall, we demonstrate that the genetic effect size estimates and levels of significance are similar across all HDL-C extraction methods attempted suggesting that commonly used decision logic for repeated measures in the EMR may not have appreciable impacts on downstream analyses conducted for genetic association studies.
Methods

Study population

All study subjects are drawn from BioVU, Vanderbilt University Medical Center’s biorepository linked to de-identified electronic medical records. A description of BioVU, including its oversight and ethics, has been previously published [18,19]. In brief, DNA is extracted from discarded blood samples drawn for routine clinical care from Vanderbilt University affiliated outpatient clinics. The DNA sample is linked to the patient’s de-identified EMR known as the Synthetic Derivative (SD). The SD contains billing (ICD-9) codes, procedure codes and labs. Prescription medication, including dose, is available in the SD through MedEx [20], an algorithm that extracts medications and their signature mentions from free-text entries available in the EMR. The SD also contains all clinical notes.

As EAGLE, a study site of PAGE I, we genotyped mostly non-European descent DNA samples available in BioVU as of 2011 on the Illumina Metabochip (described below), hereto referred as “EAGLE BioVU” (n = 15,863) [21]. The present study is limited to African Americans within EAGLE BioVU (n = 11,519).

HDL-C definitions

HDL-C measurements were extracted from a de-identified EMR using five different methods. First, for each subject, the median HDL-C value of all documented HDL-C measurements was collected (“All HDL-C”). Next, both first and last reported HDL-C were mined from the subject’s laboratory data (“First HDL-C” and “Last HDL-C”). Lastly, HDL-C values were extracted for subjects both prior to (“pre-medication HDL-C”) or following (“post-medication HDL-C”) evidence of lipid-lowering medications and the median value was reported. EAGLE BioVU clinical notes were searched for evidence of lipid-lowering drugs for each subject using medication class as well as medication generic and brand names (Table 1). For each mention of lipid-lowering drug use, we extracted the date of medication mention to compare against date of HDL-C lab to determine if that measurement of HDL-C was “pre-medication” or

| Table 1 Lipid-lowering medication class and list of drugs |
|--------------------------------------------------------|
| **Fibrates** | **Resins** | **Statins** |
|---------------------------------|-----------------|-------------------------------|
| Gemfibrozil (Lopid®) | Cholestyramine (Questran®, Questran® Light, Prevalite®, Locholest®, Locholest® Light) | Atorvastatin (Lipitor®) |
| Fenofibrate (Antara®, Lofibra®, Tricor®, Triglide™) | Colestipol (Colestid®) | Fluvastatin (Lescol®) |
| Clofibrate (Atromid-S) | Colesevelam HCl (WelChol®) | Lovastatin (Mevacor® and Altoprev™) |
| | | Pravastatin (Pravachol®) |
| | | Rosuvastatin Calcium (Crestor®) |
| | | Simvastatin (Zocor®) |
| | | Lovastatin + niacin (Advicor®) |
| | | Atorvastatin + amlodipine (Caduet®) |
| | | Simvastatin + ezetimibe (Vytorin™) |

Four major medication classes containing lipid-lowering medications were used to search the clinical notes: fibrates, niacin, resins, selective cholesterol absorption inhibitors (Ezetimibe or Zetia®), and statins (also known as an HMG CoA reductase inhibitors). For each of the medication classes included in the search, we have listed the specific drugs considered, including both the generic and brand names.
“post-medication.” Subjects with no evidence of lipid-lowering medication prescriptions were considered “pre-medication HDL-C.” All HDL-C values used in this analysis were collected when the subject was 18 years or older.

Genotyping and SNP selection
A total of 15,863 DNA samples from mostly non-European descent subjects were genotyped on the Illumina Metabochip, including 11,519 African Americans, by Vanderbilt University Center for Human Genetics Research DNA Resources Core. The Illumina Metabochip is a custom array of approximately 200,000 variants chosen as GWAS-identified index variants or GWAS-identified regions for fine-mapping based on data from the first iteration of the 1000 Genomes Project [17]. Quality control of the Illumina Metabochip data for EAGLE BioVU followed the quality control procedures outlined in Buyske et al. [22].

Based on a previous fine-mapping study of HDL-C using Metabochip [23], seven of the 22 fine-mapped HDL-C loci exhibited evidence of association at \( p < 1 \times 10^{-4} \) in African Americans. The seven index SNPs from these seven associated HDL-C loci were selected for use in calculating the genetic risk score (GRS, Table 2).

Statistical methods
Both a weighted and unweighted GRS were calculated in PLINK [24]. In general, the GRS is calculated for each subject by counting the number of effect alleles (0, 1, or 2) across each SNP, multiplying that number by the known effect size (for the unweighted GRS, effect sizes were set equal to one), summing those values, and dividing by the number of non-missing SNPs, thus providing the average score per SNP. Effect estimates for the weighted GRS were based on the meta-analysis of PAGE African Americans [23]. Linear regression, adjusted for sex, with GRS as the independent variable and HDL-C measurement as the dependent variable was used to determine the beta coefficient.

Results
Approximately 43% of the 11,519 African American subjects genotyped on the Illumina Metabochip as part of EAGLE had at least one HDL-C measurement available in the EMR (Table 3). The median number of clinic visits and medical records lengths in years was three each while the median ICD-9 code mentions (for unique codes) was 54. The median value for HDL-C ranged from 48–51 across the five different HDL-C definitions explored here (Table 3).

| SNP     | Gene of interest | Effect Allele | Effect on HDL-C \( (mg/dl) \) |
|---------|------------------|---------------|-------------------------------|
| rs247617 | CETP             | C             | -0.111                        |
| rs1077834 | LIPC           | A             | -0.033                        |
| rs10096633 | LPL            | G             | -0.042                        |
| rs189069311 | APOAS         | A             | -0.080                        |
| rs255054 | LCAT            | A             | -0.042                        |
| rs6601299 | PPP1R3B        | A             | -0.063                        |
| rs4810479 | PLTP            | G             | -0.029                        |

\(^{1}\)Beta coefficients were drawn from meta-analysis results of PAGE African Americans [23].
We first calculated the unweighted GRS using seven HDL-C associated variants (Table 2) for each African American in EAGLE BioVU with at least one HDL-C measurement. The number of HDL-C risk alleles ranged from 3 to 12, which the majority of subjects having 8 risk alleles (Figure 1).

We then performed tests of association for each of the five HDL-C definitions using the unweighted GRS as the independent variable. The unweighted GRS was significantly associated with each of the five HDL-C definitions, and the levels of significance ranged from $4.06 \times 10^{-86}$ (post-medication HDL-C; $n = 2,085$) to $3.73 \times 10^{-197}$ (first HDL-C; $n = 4,910$). Because level of significance is influenced by sample size, we then plotted each resulting beta and 95% confidence intervals to compare the effect sizes of the unweighted GRS across the five different HDL-C definitions (Figure 2). The unweighted GRS effect size was similar across the five different HDL-C definitions (Figure 2). Results from the weighted GRS do not appreciably differ from the unweighted results (data not shown).

### Discussion

We demonstrate here that for HDL-C, a commonly studied quantitative trait for cardiovascular disease risk, algorithm decision logic and phenotyping details applied to repeated measures available in the EMR do not appreciably affect downstream

| Variable                          | No. Obs. | Median | IQR |
|-----------------------------------|----------|--------|-----|
| Medical record length (years)     | 4,912    | 3      | 8   |
| Clinic visits (N)                 | 4,912    | 3      | 6   |
| ICD-9 codes (N)                   | 4,897    | 54     | 85  |
| All HDL-C (mg/dl)                 | 4,890    | 51     | 23  |
| First HDL-C (mg/dl)               | 4,912    | 50     | 23  |
| Last HDL-C (mg/dl)                | 4,912    | 49     | 22  |
| pre-medication HDL-C (mg/dl)      | 4,074    | 51     | 23  |
| post-medication HDL-C (mg/dl)     | 2,086    | 48     | 22  |

Number of observations (No. Obs.) as well as medians and interquartile ranges (IQR) are given for each variable. Includes only unique ICD-9 codes per individual.
genetic association test statistics or overall study conclusions. These data, along with on-going algorithm development within the eMERGE network [5,25], suggest that phenotypes derived from EMR-based repositories are robust to the underlying variability inherent in clinical collections. Although not explicitly tested here, the similarities of genetic effect sizes observed here for the five HDL-C definitions in the same sample suggest that any one of these EMR-derived test statistics robust to algorithm decision logic can be included in meta-analyses with traditional epidemiologic studies.

While our data suggest that EMR-derived phenotypes may be robust to certain aspects of the algorithm decision logic and phenotyping details, these data do not imply that genetic association studies are not impacted by poor phenotyping. Substantial literature has documented the need for rigorous case/control phenotyping as misclassification of either can lead to loss of power [26,27]. Careful phenotyping can also lead to insights into biological mechanisms or disease processes [28,29]. Finally, careful phenotyping is also essential for creative study design and genetic discovery [30].

The present study focuses on examining the impact algorithm decision logic has on genetic associations related to a single quantitative trait, HDL-C. As such, the conclusions offered here may be limited to HDL-C or to quantitative traits defined from repeated measures available in the EMR. Further study is needed to more fully explore the limitations and impact algorithm decision logic may have on genetic association studies for binary clinical outcomes such as myocardial infarction or pharmacogenomic studies for traits such as warfarin dosing. For the HDL-C data included here, additional limitations of the present study include limitations associated with extracting HDL-C from the EMR. For example, we searched clinic notes for mentions of lipid-lowering medication classes and drugs (genetic and brand names), but we did not include any common misspellings of these search terms. It is possible, therefore, that the “pre-medication” HDL-C definition contains HDL-C measurements while the subject was on lipid-lowering medication. Another limitation of the EMR is that, unlike most epidemiologic studies, fasting status or time to last meal is not available as a structured field. Here, we assumed that the HDL-C measured in EAGLE BioVU was measured for subjects who fasted for at least eight hours. This assumption is most likely incorrect, but its violation is unlikely to impact HDL-C levels substantially.

Figure 2 Additive effects of HDL-C risk alleles on various HDL-C measurements. Effect sizes (betas) from the linear regression analysis with the unweighted GRS, adjusted for sex, are shown as expected HDL-C levels (in mg/dl; black diamonds), along with their 95% confidence intervals.
Another limitation of the present study is related to sample size and power. We present here tests of association between various HDL-C derived variables and an unweighted GRS. The unweighted GRS, by design, is calculated by the number of risk alleles at loci known to be significantly associated with HDL-C levels. Therefore, with only a few thousand samples, we were able to statistically replicate the expected association between the unweighted GRS and the various HDL-C variables to further examine the genetic effect sizes estimated from these tests of associations. While the sample size of the present study was large enough for replicating known associations such as the loci represented in the unweighted GRS, the sample size is not large enough to perform discovery studies with the entire Metabochip dataset, even when limited to common variation (minor allele frequency >5%). Indeed, tests of association between the various HDL-C variables and common variants on the Metabochip failed to identify a statistically significant association after correction for multiple testing (data not shown). Furthermore, neither significance rankings nor genetic effect sizes could be reliably compared across HDL-C variables given the chance findings of non-significant tests of associations. Larger sample sizes are needed to make comprehensive comparisons of genetic effect sizes and significance rankings for EMR-derived phenotypes susceptible to algorithm decision logic and phenotyping details.

Despite the limitations, this study had multiple strengths including the depth of the clinical data and the diversity of EAGLE BioVU. EMR-derived datasets such as EAGLE BioVU coupled with genotype and sequence data promise to enrich existing and complimentary datasets for future genetic association studies for complex human diseases and traits.

**Conclusions**

These data collectively suggest that, at least for HDL-C, algorithm decision logic and phenotyping details do not appreciably impact genetics association study tests statistics.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

DCC and LD designed the experiment. RG, YB, EF-E, and JB performed the data extraction, performed the genotyping quality control, and assisted in the analysis. LD performed the main analyses and drafted the manuscript. DCC secured the major funding for the project. All authors read and approved the final manuscript.

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