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Impact of SLCO1B1 pharmacogenetic testing on patient and healthcare outcomes: A systematic review

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Abstract
Demonstrated improvements in patient outcomes will facilitate the clinical implementation of pharmacogenetic testing. Using the association between solute carrier organic anion transporter family member 1B1 (SLCO1B1) and statin-associated muscle symptoms (SAMS) as a model, we conducted a systematic review of patient outcomes after delivery of SLCO1B1 results. Using PubMed and Embase searches through December 19, 2017, we identified 37 eligible records reporting preliminary or final outcomes, including 6 studies delivering only SLCO1B1 results and 5 large healthcare system-based implementation projects of multi-pharmacogene panels. Two small trials have demonstrated at least short-term improvements in low-density lipoprotein cholesterol after SLCO1B1 testing among previously statin intolerant patients. Evidence from large implementation projects suggests that SLCO1B1 results may change prescribing patterns for some high-risk patients. No study has reported improvements in SAMS or cardiovascular events or tracked the economic outcomes of SLCO1B1 testing. Ongoing studies should collect and report outcomes relevant to pharmacogenetics stakeholders.
Introduction

The field of pharmacogenetics is one beneficiary of the last decade’s accelerated pace of genomic discovery(1, 2). Hundreds of drug-gene associations have now been identified that have the potential to help prescribers and patients optimize the risk-benefit ratio of pharmacotherapy(3). These pharmacogenetic associations could be ideal candidates for the translation of genomic discovery into patient care, since test results might have ready actionability to inform drug selection and dose(4, 5) and might lend themselves to clinical decision support (CDS) interventions in the electronic health record (EHR)(6-9). Some healthcare systems in the United States and worldwide are making large investments to implement clinical pharmacogenetics programs into their healthcare delivery systems(10-14), but most health care is still delivered in settings without the routine use of pharmacogenetic testing. Innovation generally diffuses slowly throughout medical practice(15), and for pharmacogenetics, this lag is exacerbated in part by healthcare providers and insurers who remain unconvinced of its value in improving the health care and outcomes of patients(16-19). Evidence that pharmacogenetic testing improves patient outcomes is needed to break this impasse(16-19).

One well validated drug-gene association is the interaction between simvastatin and the solute carrier organic anion transporter family member 1B1 (SLCO1B1) gene for the risk of statin-associated muscle symptoms (SAMS). Statins, or 3-hydroxy-3-methylglutaryl-CoA reductase inhibitors, are cholesterol-lowering medications used by over 30 million Americans that have dramatically reduced the risk of cardiovascular disease (CVD) and death in the U.S.(20). Statins are generally well tolerated, but up to 20% of patients describe muscle aches or weakness(21, 22) and up to 1 in 10,000 experience life-threatening myopathy(23-25). In 2008, a high-profile genome-wide association study using data from the Study of the Effectiveness of Additional Reductions in Cholesterol and Homocysteine (SEARCH) trial identified a robust association
between a common genetic variant in \textit{SLCO1B1} and simvastatin-related myopathy (26). Each copy of the C minor allele at rs4149056 (contained within the \textit{SLCO1B1*5, *15, and *17} haplotypes) increased myopathy risk by a factor of 4.5, such that CC homozygotes had a 16.9-fold increased risk compared to TT homozygotes (26). Numerous other studies have gone on to replicate the association between \textit{SLCO1B1} and SAMS, particularly with simvastatin (27-30). In 2012, the Clinical Pharmacogenetics Implementation Consortium (CPIC) issued its first recommendations for simvastatin prescribing and dosing when a patient’s \textit{SLCO1B1} genotype is known (31), and some early-adopter healthcare systems are now incorporating \textit{SLCO1B1} genotyping into their clinical and research pharmacogenetics programs (10-14).

Although it makes intuitive sense that the use of \textit{SLCO1B1} testing in clinical care would help prescribers and patients avoid statin-related adverse effects, intuition might not be sufficiently persuasive to promote its widespread adoption among providers and payers. Several reviews have examined the validity of the \textit{SLCO1B1}-SAMS association (30, 32, 33), but none has reviewed the prospective outcomes of integrating \textit{SLCO1B1} genotype information into clinical care. Given the primacy of outcomes data in determining the clinical utility of pharmacogenetic testing, we performed a systematic review of studies reporting outcomes after the delivery of \textit{SLCO1B1} results.

Results

\textit{Search results}

The database searches described in the Methods section below yielded 5,374 unique records (Figure 1). Manual author and reference searches identified another 57 potentially eligible records, including one personal author communication providing unpublished results. After full-text review, we identified 37 records describing 16 eligible studies with completed or
preliminary outcomes, including 2 nonrandomized trials and 4 randomized controlled trials (RCT). Table 1 presents the characteristics of these studies. We identified 17 records describing another 10 studies with potentially eligible study designs that have not yet reported outcomes.

Five pilot studies and one RCT have studied the delivery of \textit{SLCO1B1} results specifically\cite{34-43}, all but one of which were conducted by investigators at Duke University\cite{34-40, 42, 43}. Three of the Duke pilot studies involved pharmacists in the identification of patients for \textit{SLCO1B1} genotyping and/or the formulation of medication recommendations based on \textit{SLCO1B1} results\cite{34-38, 40}. A fourth Duke study, a pilot nonrandomized trial, enrolled 58 patients with prior statin nonadherence and studied the impact of \textit{SLCO1B1} results delivery on statin prescriptions and medication adherence, compared to concurrent controls\cite{39}. This pilot informed the design of a larger trial, the only published RCT designed specifically to examine the clinical impact of \textit{SLCO1B1} testing. In this RCT by Voora and colleagues\cite{42, 43}, 167 patients with prior statin intolerance were randomly allocated to \textit{SLCO1B1} results delivery to patients and providers at baseline versus at study end. The study was powered to detect a 1-point difference in the primary outcome of a medication adherence scale; secondary outcomes included low-density lipoprotein cholesterol (LDL-C) at 3 and 8 months, the Brief Pain Inventory, and the Short Form Health Survey (SF-12) quality-of-life measure\cite{42, 43}.

In addition, five large healthcare system-based pharmacogenetics implementation projects of multigene panels including \textit{SLCO1B1} have reported preliminary or final results: four in the U.S. (1,200 Patients Project\cite{11, 44-49}; INGENIOUS\cite{13, 50, 51}; the Marshfield Clinic\cite{12, 52}; and the RIGHT protocol\cite{8, 11, 12, 53-56}) and one from La Paz University Hospital in Spain\cite{10}(Table 1). Most of these are using passive or active CDS in the EHR to support prescriber utilization of pharmacogenetic results in several different clinically actionable pharmacogenes. The remaining studies identified by our search included smaller single-arm
intervention studies of multigene panels (11, 49, 57-65) and one pilot RCT of genome sequencing (66, 67) (Table 1).

**Study quality and risk of bias**

The included studies generally had poor to moderate quality and risk of bias (Table S1). Most intervention studies suffered from poor comparability on the Newcastle-Ottawa Scale due to the absence of a comparator group.

**Reported outcomes**

Review of the outcomes reported in the eligible studies led to the conceptual model of patient and healthcare outcomes shown in Figure 2, which guided the creation of the 6 outcome categories in Table 2 and our presentation of the results below. In this conceptual model, \( SLCO1B1 \) results might act on provider and patient attitudes and behaviors to effect a change in clinical, economic, and other outcomes (Figure 2). As presented below, we categorized all reported outcomes as either *utility outcomes* (including clinical outcomes and healthcare utilization and economic outcomes) or *process outcomes* that might mediate the relationship between \( SLCO1B1 \) testing (including provider utilization and attitudes, prescribing behavior and prescriptions, medication adherence, and other patient-reported outcomes).

*Utility outcomes*

Clinical outcomes

Only two studies to date have quantitatively reported clinical outcomes after \( SLCO1B1 \) testing. The Duke nonrandomized pilot trial among previously statin-intolerant patients reported a non-
significantly greater reduction in LDL-C in the intervention group (-12 ± 45 mg/dL) compared to concurrent controls (+6 ± 38 mg/dL, p=0.06) after one year(39). In the subsequent RCT, LDL-C values were significantly lower in the intervention group compared to the control group at 3 months (132 ± 42 mg/dL vs. 144 ± 43 mg/dL, p=0.04) but not at 8 months (129 ± 38 mg/dL vs. 141 ± 44 mg/dL, p=0.07)(43). Improvements observed in total cholesterol, but not high-density lipoprotein cholesterol or triglycerides, were consistent with these LDL-C changes. In a follow-up analysis, when patients in the usual care arm received their SLC01B1 results at the end of the study, they had a greater decrease in LDL-C values compared to intervention patients during the same post-study period, such that the two arms ultimately achieved similar LDL-C reductions from baseline. This RCT found no significant differences in medication side effects between the intervention arms, as measured by pain and quality-of-life instruments(43). Other studies of pharmacogenetic testing have made general qualitative statements that no participants experienced medication side effects during the observation periods(41, 57, 60, 66). No study has reported creatinine kinase values, SAMS, or cardiovascular events after SLC01B1 testing.

Healthcare utilization and economic outcomes

Some studies have tracked the costs and resources required to conduct their pharmacogenetics projects or those incurred as a result of that implementation(10, 38, 50, 66). The clinical pharmacogenetic service in Spain cost the national health system €202,140 over 3 years, with the cost of each consultation averaging €216(10). Preliminary results from the INGENIOUS RCT of pharmacogenetic panel testing show that genotyping the first 106 participants generated 25 actionable genotypes and prompted 10 (9%) consult requests by physicians(50). Two studies have reported patient willingness-to-pay for multigene pharmacogenetic testing that included SLC01B1. In one, participants in the OSU-Coriell Personalized Medicine Collaborative RCT of genomic counseling, 28% of whom had an actionable SLC01B1 result, reported a mean (SD) willingness to pay of $56 ($81) for a clinical pharmacogenetics service(59). In the second, the
RIGHT Protocol at the Mayo Clinic, a survey of 869 participants who had undergone panel pharmacogenotyping found that 42% were not willing to incur out-of-pocket costs for pharmacogenetic tests; 58% of the rest reported a maximum willingness-to-pay of $100 (54). No study has reported downstream healthcare costs after receipt of SLC01B1 results specifically, although the MedSeq Project pilot RCT found no differences in 6-month healthcare costs between participants receiving genome sequencing including SLC01B1 genotyping versus no genome sequencing (mean $1490 vs. $1142, excluding the costs of sequencing and interpretation) (66).

Process outcomes

Provider utilization and attitudes

Studies in which results were delivered to prescribers enabled an examination of how frequently they interfaced with the information and their attitudes about its value. Studies in which providers initiated SLC01B1 testing generally reported low testing uptake (10, 34-36). Studies delivering SLC01B1 results to providers through the EHR without provider initiation have reported providers’ EHR transactional data. For example, the 1,200 Patients Project of more than a dozen drug-gene pairs reported that 69% of 2,279 patient visits over 3 years were associated with a provider log-in to the pharmacogenetics CDS system within 72 hours (48). About one-third of patients' active medications had associated pharmacogenetic alerts (0.5% red, 13% yellow, and 21% green) (48); a 10-month analysis in the first 608 patients reported that providers clicked on 100%, 72%, and 20% of red, yellow, and green alerts, respectively (47). However, SLC01B1 transactions were not specifically reported. During the first 14 months that SLC01B1 CDS was in production in the RIGHT Protocol, there were 0.7 interruptive alerts per month for simvastatin orders attempted for rs4149056 TC or CC patients among 3,788 patients seen by 1,247 unique providers (55). Studies surveying providers about their experiences have reported overall positive attitudes about pharmacogenetics, including its
clinical relevance and impact on management(10, 47, 60), although no study reported provider
attitudes about SLC01B1 testing specifically.

Prescribing behavior and prescriptions

Most studies have reported or are actively collecting data on the impact of SLC01B1 results on
medication prescriptions, measured either from the EHR or provider or patient report. Some
studies of pharmacogenetic panel testing have reported only composite medication changes(10,
60, 61, 63), while small pilot studies have reported specific cases where SLC01B1 results guided
therapy(34, 35, 38, 40, 57, 66). Large studies with SLC01B1 CDS alerts have reported counts of
medication changes attributed to pharmacogenetic results. The 1,200 Patients Project reported
that 25% of 2,279 visits over 3 years had medication changes; simvastatin was the drug with the
highest percentage of changes influenced by CDS (69%), although this represented only 8
simvastatin discontinuations in SLC01B1 C carriers among 868 patients(48). In the first 3 years
of the Marshfield Clinic project, there have been 5 CDS alert recommendations triggered by
simvastatin prescriptions, only one of which was followed, prompting the provider to prescribe
atorvastatin instead (personal communication, Terrie Kitchner, December 6, 2017). Similarly,
pharmacists in a Duke pilot study did not recommend any simvastatin prescription changes to
the providers caring for 6 patients with carrier or homozygous SLC01B1*5 results(40). Two
controlled studies have examined prescribing behavior. In the Duke nonrandomized pilot trial,
55% of patients with a history of statin non-adherence had statin prescriptions 4 months after
receiving SLC01B1 results, compared to 20% of concurrent controls with statin prescriptions
after one year (p<0.001)(39). In the subsequent RCT, more participants receiving SLC01B1
results were on statin therapy at 3 months compared to usual care (55% vs. 38%, p=0.04), but
this difference was not statistically significant after 8 months (54% vs. 37%, p=0.07)(43).
Medication adherence

While prescriptions largely reflect provider behavior, medication adherence is a patient behavior. Small pilot studies have either found no impact of \textit{SLCO1B1} testing on statin adherence or did not collect data to enable pre-post or between-group comparisons\cite{34, 35, 38, 41}. The Duke nonrandomized pilot study among patients with prior statin discontinuation found that 47% of intervention patients reported taking a statin after 4 months compared to 15% of concurrent controls after one year ($p<0.001$)\cite{39}. The subsequent RCT, however, found no differences in adherence or the Medication Possession Ratio after 3 or 8 months between the subsets of patients in both arms reinitiated on statin therapy\cite{43}. The authors reported that intervention patients perceived higher necessity of their medications than control patients at 3 months, but not at 8 months\cite{43}, consistent with observations from the pilot study\cite{39}.

Other patient-reported outcomes

Uncontrolled studies have reported that some patients had difficulty recalling their specific \textit{SLCO1B1} results\cite{35, 38} and had variable understanding of them\cite{10, 40, 59}. Nonetheless, patients generally perceived the information as useful to their providers\cite{35, 38, 54, 59}. Pilot studies have also reported that patients generally had no concerns or distress after receiving pharmacogenetic results\cite{35, 40, 66}.

Discussion

Ten years after the publication of the association between \textit{SLCO1B1} and SAMS\cite{26}, we found few high-quality studies reporting patient outcomes after the delivery of \textit{SLCO1B1} results. Most notably, a pilot trial and subsequent small RCT among previously statin intolerant patients observed at least short-term improvements in LDL-C after \textit{SLCO1B1} testing. Although these
findings require replication, the 10-mg/dL reduction in LDL-C the investigators observed, if sustained, would result in a 5% lower 5-year risk of major CVD event(68). Apart from this, while the proposed benefit of SLC01B1 testing is the avoidance of SAMS, it is worth noting that no study has empirically demonstrated this outcome or the impact of SLC01B1 testing on CVD events. Evidence from small pilot studies and large healthcare system implementation projects does suggest that SLC01B1 results may change providers’ prescribing patterns for some, but not all, high-risk patients receiving simvastatin, but to date, the number of potential opportunities to observe prescription changes in large healthcare systems with SLC01B1 CDS in the EHR has been small. Receipt of SLC01B1 test results seems generally well tolerated by providers and patients. No study has specifically tracked the economic impact of SLC01B1 testing and its downstream outcomes.

The clinical validity of the SLC01B1-SAMS association has been well established; that is, the observed association between rs4149056 in SLC01B1 and statin-associated myotoxicity of varying severity has been replicated in numerous studies, particularly for simvastatin(30). The PharmGKB knowledge resource rates the genotype-phenotype association between SLC01B1 and simvastatin myopathy as having the highest level of evidence (Level 1A)(69). On the other hand, the clinical utility of SLC01B1 testing, or its ability to inform a change in clinical management that demonstrably improves patient outcomes, is less certain. A 2013 review by Stewart found no studies comparing clinical outcomes between patients whose statin prescriptions were guided or not guided by SLC01B1 results(32), and Sorich found no studies of the cost-effectiveness of SLC01B1 genotyping(70). A more recent review examined 89 studies purporting to address either the clinical validity or clinical utility of pharmacogenetic testing for statin use and found almost all claims of clinical utility to be lacking when examined against benchmarks such as number needed to genotype, the effect and risks of the intervention, and costs per quality adjusted life year(33). Many of the studies we identified in the present review

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are collecting data on prescription changes, a measure of the actionability of pharmacogenetic results. Still, the absence of prospective outcomes data for pharmacogenetic testing will continue to make health insurers reluctant to cover the costs of testing(71) and many clinicians reluctant to incorporate pharmacogenetics into their practices(72).

Our review of patient outcomes after \textit{SLCO1B1} testing prompts the following recommendations for future research. First, although RCT evidence may not be necessary to justify every clinical application of pharmacogenetic testing(73-75), prospectively collected outcomes data might be, ideally from studies with suitable control groups. Investigators are encouraged to identify and collect data from concurrent matched controls for the participants in ongoing and planned pharmacogenetic projects, to enable a less biased determination of the impact of testing.

Second, these outcomes should include those of interest to patients, providers, and payers, including clinical outcomes, quality of life, and costs. Third, even with the multigene pharmacogenetic panels that some implementation projects are using, it is important to report outcomes specific to individual pharmacogenetic tests, such as \textit{SLCO1B1} for statins and \textit{cytochrome P450 family 2 subfamily C member 19 (CYP2C19)} for clopidogrel(76). Very few studies using panels in our review reported outcomes pertaining to \textit{SLCO1B1} results specifically. Although panels enable efficiencies of scale in genotyping, additional costs such as the development and implementation of CDS for each drug-gene pair are not trivial(77). Locus-specific outcomes data will enable a determination of the returns on those investments. Fourth, as more outcomes data accrue, the effect of context on those outcomes should be examined, including the degree of patient and provider engagement in the process, the type of CDS used in results delivery, and the characteristics of patients most likely to benefit. To date, the strongest evidence supporting the use of \textit{SLCO1B1} testing derives from an RCT among previously statin-intolerant patients(43). This finding is consistent with the French National Network of Pharmacogenetics recommendation that \textit{SLCO1B1} testing is potentially useful for patients.
experiencing SAMS after statin initiation or with at least one SAMS risk factor; it does not recommend routine preemptive \textit{SLCO1B1} testing before general simvastatin initiation\cite{78}. Further research should examine the clinical utility of \textit{SLCO1B1} testing among statin-naïve patients and among patients already tolerating statin therapy.

With the recommendations above, ongoing projects using \textit{SLCO1B1} genotyping in research or clinical care in the U.S. and internationally have a tremendous opportunity to contribute to the lack of evidence for its clinical utility. We identified 10 institutions with ongoing studies whose designs and planned outcomes would have been eligible for this review\cite{11, 12, 14, 79-82}. Most of these represent multi-institutional efforts such as the Electronic Medical Records and Genomics (eMERGE)-PGx Consortium\cite{12}, the Pharmacogenomics Research Network Translational Pharmacogenetics Program (TPP)\cite{11}, the Implementing Genomics in Practice (IGNITE) Consortium\cite{13}, and the seven-country Ubiquitous Pharmacogenomics (U-PGx) Consortium\cite{14}. These projects are collecting a range of prescription, clinical, and economic outcomes, and their large scale will enable more precise estimates of the clinical utility of pharmacogenetic testing. For example, while preliminary reports suggest that individual healthcare systems may observe few instances where \textit{SLCO1B1} results change medication prescriptions, a recent update from the TPP reported that 14,508 \textit{SLCO1B1} results have been reported in the EHR of five participating institutions, of which 3,513 (24\%) were actionable\cite{11}. In addition to these large projects of pharmacogenetics panels, we are conducting the Integrating Pharmacogenetics in Clinical Care Study, an RCT specifically examining the impact of \textit{SLCO1B1} genotyping on LDL-C and concordance with statin therapy guidelines among statin-naïve patients (ClinicalTrials.gov Identifier: NCT02871934).
This review has a few limitations to note. The paucity of published clinical utility outcomes and the heterogeneity in other outcomes reported after \textit{SLCO1B1} testing precluded meta-analysis or between-study comparisons. We examined outcomes data from only a single specific gene-drug pair as an in-depth case study, paradigmatic of the state of the evidence for most other pharmacogenetic tests. It is unknown how the use of multigene pharmacogenetic panels would change the impact of \textit{SLCO1B1} information alone, since multiple genetic test results can interact in unpredictable ways on patient outcomes(83). While \textit{SLCO1B1} testing might be increasingly common in medical practice outside academic centers, we were only able to examine outcomes published in the biomedical literature.

In conclusion, despite advances bringing pharmacogenetic testing to clinical care, we found few patient outcomes reported after the delivery of \textit{SLCO1B1} results, a well validated pharmacogenetic locus. Outcomes data are needed to accelerate the pace of this clinical translation.

\textbf{Methods}

\textit{Protocol and registration}

The Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) Guidelines were followed for this systematic review (see PRISMA checklist, Table S2, in the Supplemental Information). We performed initial scoping searches of PubMed and Embase on June 22, 2017 before registering the review protocol on the PROSPERO register of systematic reviews on August 28, 2017 (CRD42017074795). After our initial searches and record review, we updated our searches on December 19, 2017 to identify new records.
Search strategy

We searched the PubMed and Embase databases for published reviews, meta-analyses, and primary studies published in or after 2005 using combinations of the following search strategy concepts: pharmacogenetics/pharmacogenomics, precision medicine, SLC01B1, statins, cardiovascular disease. The full search strategies are included in the Supplemental Information.

Scope and eligibility criteria

The primary aim of the systematic review was to review the evidence for the clinical utility of SLC01B1 testing, as determined by patient outcomes observed after SLC01B1 genotyping. We included intervention studies in which 1) participants were directly or indirectly (e.g. via their providers) given their SLC01B1 genotype results as a part of a research study or clinical care and 2) subsequent outcomes were prospectively collected and reported. Our scoping search enabled us to determine the appropriate breadth of eligibility criteria for both study designs and outcomes. Identifying few eligible randomized trials, we chose to additionally include pilot studies, implementation projects, and nonrandomized trials. We excluded case reports. We excluded studies reporting the association between SLC01B1 genotype and statin effects (i.e. the clinical validity of the SLC01B1 genotyping), as this has been reviewed in detail elsewhere, particularly by CPIC(30, 32, 33). We excluded retrospective observations among cohorts who had undergone direct-to-consumer pharmacogenetic testing (reviewed in (84)). We excluded records reporting only the frequency of SLC01B1 genotypes among participants or in which SLC01B1 genotype results were used to estimate hypothetical recommendations for medication changes (e.g. (5)). We included records in any language. After our scoping search identified few studies with eligible designs that reported clinical outcomes such as biomarker changes, morbidity, or mortality, we defined an eligible outcome broadly as any provider- or patient-reported outcome, EHR-derived outcome, or other study outcome measured after an intervention that involved the reporting of SLC01B1 results to providers and/or patients. Our
approach to categorizing these clinical utility and process outcomes is described above in the
Results. Any record describing an ongoing study whose design and planned outcomes would be
eligible for inclusion was noted so that authors could be contacted for more information.

Review process

The titles and abstracts of all search records were screened for potential eligibility by two
independent reviewers; discrepancies were resolved by discussion and consensus among the
study team. Potentially eligible records progressed to full record review for determination of
eligibility. The references of review papers and eligible studies were manually searched for
additional eligible studies.

Data abstraction

A Microsoft Excel database was used to abstract the following from each eligible study: country;
study design; patient population and number; the genotyping intervention, including any
genotypes other than SLC1B1 reported and any associated decision support; any control
group; any quantitative or qualitative outcome reported, including the method of collection and
results; and the current status of the study. We categorized any intervention study with a
control group as a nonrandomized trial if historical or concurrent controls were used or as a
randomized controlled trial if participants were randomly allocated to the study arms(85). All
other eligible studies were categorized as intervention studies, which included
pharmacogenetics interventions delivered through pilot studies or operational clinical
innovation programs.
**Author communication**

For each ongoing study with a potentially eligible study design (typically identified through a manuscript describing the study design and rationale), we emailed the corresponding author(s) a link to a brief survey requesting any published or unpublished results from the study referenced in the records we identified (Supplemental Information). Each author was sent up to three requests, each separated by at least 7 days. We also performed targeted author searches to identify any additional records from these ongoing studies.

**Assessing study quality and risk of bias**

Bias and study quality were systemically assessed using the Newcastle-Ottawa Scale (NOS) for the quality of intervention studies and nonrandomized studies, with greater scores indicative of higher study quality(86). For the randomized controlled trials, the Jadad scale(87) was used for the study quality assessment.

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Figure Legends

**Figure 1:** PRISMA flow diagram of search results

**Figure 2:** Conceptual model of patient and healthcare outcomes after delivery of *SLCO1B1* pharmacogenetic test results

Supplemental Information
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<th>Population (n)</th>
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<th>Experimental intervention</th>
<th>Decision support</th>
<th>Control group (n)</th>
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<td>Intervention study (U.S.A.)</td>
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<td>• Genotyping of prospectively enrolled cohort</td>
<td>• CDS system outside EHR including green/yellow/red alerts with clinical summaries and interpretation</td>
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<td>Intervention study (U.S.A.)</td>
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<td>Duke University 1 (39)</td>
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<td>Duke University 2 (40)</td>
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<td>Test interpretation included CPIC guidelines</td>
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<td>Intervention study (U.S.A.)</td>
<td>Patients at 2 cardiology clinics (30)</td>
<td>CYP2C9, CYP2C19, CYP2D6, SLC01B1, VKORC1</td>
<td>Pre- and post-test MTM sessions with pharmacist for patients</td>
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<td>MTM also included recommendations for lifestyle modification and OTC medications</td>
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<td>Patients and referring cardiologists received PGx results</td>
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<td>Pharmacist discussed recommendations based on FDA and/or CPIC guidelines with cardiologist before sharing action plan with patients</td>
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</tr>
<tr>
<td>Duke University 4 (34-36)</td>
<td>Non-randomized trial (U.S.A.)</td>
<td>Primary care patients at 2 internal medicine practices (63)</td>
<td>CYP2C9, CYP2C19, CYP2D6, HLA-B*1502, SLC01B1, VKORC1</td>
<td>Site 1: Pharmacist on call: physician consulted pharmacist about PGx testing; pharmacists screened patients and notified physicians about eligibility</td>
<td>All participating physicians first attended a 1-hour CME session about PGx.</td>
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<td>Site 2: Pharmacist in-house: pharmacist screened patients and alerted physicians to availability of PGx testing for relevant medications</td>
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<tr>
<td>Duke University 5 (42, 43)</td>
<td>Randomized control trial (U.S.A.)</td>
<td>Patients not currently taking statins due to history of adverse effect, ineligible if prior rhabdomyolysis or CK &gt;10xULN (167)</td>
<td>SLC01B1</td>
<td>Genotyping at a research visit.</td>
<td>Physician reports:</td>
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<td>Patients and their physicians received PGx results by email</td>
<td>SLC01B1 results</td>
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<td>Risk of rhabdomyolysis</td>
<td>Risk of rhabdomyolysis</td>
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<td>General expectations of LDL-C reduction from different statin types &amp; doses</td>
<td>General expectations of LDL-C reductions from different statins</td>
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<tr>
<td>First Moscow State Medical University (41)</td>
<td>Intervention study (Russia)</td>
<td>Patients with hyperlipidemia already on statin therapy (35)</td>
<td>SLC01B1</td>
<td>Patients received results</td>
<td>Patient reports:</td>
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<td></td>
<td>SLC01B1 results</td>
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<td></td>
<td>Reassurance about which statins should be tolerable and lower CVD risk</td>
<td>Patient reports:</td>
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<td>General information about statins and CVD risk</td>
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</tbody>
</table>

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<table>
<thead>
<tr>
<th>Study Title</th>
<th>Study Design</th>
<th>Eligibility</th>
<th>Outcome Measures</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>INGENIOUS** (13, 50, 51)</td>
<td>Randomized control trial</td>
<td>Safety-net hospital system of 10 clinics with common EHR (2,000 planned)</td>
<td>• Enrolling prescriber receives PGx reports &lt;br&gt; • PGx results uploaded to EHR and viewable by all providers &lt;br&gt; • Reports include alternative prescribing recommendations for index medication based on CPIC guidelines &lt;br&gt; • Subsequent prescription prompts an alert notifying provider a genotype report is available and gives dosing recommendations &lt;br&gt; • Providers may consult PGx service that documents recommendations in EHR</td>
<td>Patient does not undergo genotyping (4000 planned)</td>
</tr>
<tr>
<td>La Paz University Hospital (10)</td>
<td>Intervention study (Spain)</td>
<td>Patients receiving specialty care at a tertiary care teaching hospital (600)</td>
<td>Providers request testing from PGx unit, which generates report according to prespecified drug-gene protocol or after PGx consultation</td>
<td>Recommendation from PGx unit, based on CPIC and DPWG reviews</td>
</tr>
<tr>
<td>Marshfield Clinic** (12, 52)</td>
<td>Intervention study (U.S.A.)</td>
<td>Adults aged ≥50 years with healthcare system primary care physician and no prior use of simvastatin, warfarin, or clopidogrel (750 planned)</td>
<td>• Providers receive PGx results &lt;br&gt; • Patients have access to website with information about their PGx results</td>
<td>Active CDS alerts with CPIC dosing recommendation triggered by prescription in EHR</td>
</tr>
<tr>
<td>MedSeq Project (66, 67)</td>
<td>Randomized control trial</td>
<td>Generally healthy adult primary care patients (100)</td>
<td>Patients discussed interpreted genome report and family history pedigree with physician</td>
<td>Report included statement about simvastatin-associated myopathy risk</td>
</tr>
<tr>
<td>OSU-Coriell Personalized Medicine Collaborative* (11, 49)</td>
<td>Intervention study (U.S.A.)</td>
<td>Participants with heart failure and hypertension enrolled in RCT of genomic counseling for</td>
<td>• Patients received genetic reports by mail and by patient web portal &lt;br&gt; • Reports also uploaded to EHR</td>
<td>Reports to patients and in EHR included CPIC recommendations</td>
</tr>
<tr>
<td>Study</td>
<td>Intervention</td>
<td>Patients</td>
<td>Genes Assessed</td>
<td>Providers Ordered</td>
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<tr>
<td>PRIMER (60)</td>
<td>Intervention study (U.S.A.)</td>
<td>Patients of 27 providers with likelihood of exposure to a relevant medication (705)</td>
<td>COMT, CYP2D6, CYP2C19, CYP2C9, CYP1A2, CYP3A4, CYP3A5, F2, F5, MTHFR, OPRM1, SLCO1B1, SLC6A4, VKORC1</td>
<td>Providers ordered PGx panel testing and received report</td>
</tr>
<tr>
<td>RIGHT Protocol*** (8, 11, 12, 53-56)</td>
<td>Intervention study (U.S.A.)</td>
<td>Health system patients, including biobank participants, likely to initiate statin treatment within 3 years (3,788)</td>
<td>CYP2C9, CYP2C19, CYP2D6, HLA-B<em>1502, HLA-B</em>5701, SLCO1B1, TPM7, VKORC1</td>
<td>Preemptive genotyping with results available to provider in EHR and to patients through patient portal</td>
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<tr>
<td>Yale University (57)</td>
<td>Intervention study (U.S.A.)</td>
<td>Series of consecutive high-risk cardiovascular patients (32)</td>
<td>CYP2C19, CYP2C9, CYP2D6, CYP3A4, CYP3A5, F2, F5, MTHFR, SLCO1B1, VKORC1</td>
<td>Results reported to clinicians</td>
</tr>
</tbody>
</table>

*Part of the Pharmacogenomics Research Network Translational Pharmacogenetics Program (11)

**Part of the Implementing Genomics in Practice (IGNITE) Consortium (13)

***Part of the eMERGE-PGx Consortium (12)

**Abbreviations:** CDS (clinical decision support), CK (creatine kinase), CME (continuing medical education), CPIC (Clinical Pharmacogenetics Implementation Consortium), CVD (cardiovascular disease), DPWG (Dutch Pharmacogenetics Working Group), EHR (electronic health record), FDA (Food & Drug Administration), LDL-C (low-density lipoprotein cholesterol), MTM (medication therapy management), OSU (The Ohio State University), OTC (over-the-counter), PGx (pharmacogenetics), ULN (upper limit of normal)
Table 2. Utility and process outcomes after SLC01B1 genotyping in eligible studies

<table>
<thead>
<tr>
<th>Study/Institution</th>
<th>Utility outcomes</th>
<th>Process outcomes</th>
<th>Key findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Study/Institution</strong></td>
<td><strong>Clinical outcomes</strong></td>
<td><strong>Healthcare utilization and economic outcomes</strong></td>
<td><strong>Provider utilization and attitudes</strong></td>
</tr>
<tr>
<td>1200 Patients Project* (11, 44-49)</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>AltheaDx (61, 63)</td>
<td></td>
<td>X</td>
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</tbody>
</table>
| Duke University 1 (39) | X |  | X | X |  |  | • 47% vs. 15% of intervention vs. control patients taking statin after 4 months (p<0.001)  
• 1-year LDL-C reduction -12 ± 45 mg/dL in invention group vs. +6 ± 38 mg/dL in controls (p=0.06) |
| Duke University 2 (40) |  | X | X | X | X |  | • No prescription changes observed |
| Duke University 3 (37, 38) | X |  | X | X | X |  | • Pharmacist medication management and PGx visit lasted mean 16 (range 8-29) minutes  
• 3 PGX-based medication changes were made among 28 patients |
| Duke University 4 (34-36) | X |  | X | X | X | X | • Providers consulted pharmacist for 15 cases, averaging 5.7 minutes per consult.  
• 1/63 patients tested had simvastatin dose halved due to SLC01B1*5 result.  
• LDL-C lower in the intervention group vs. controls at 3 months (132 ± 42 mg/dL vs. 144 ± 43 mg/dL, p=0.04) but not 8 months (129 ± 38 mg/dL vs. 141 ± 44 mg/dL, p=0.07)  
• No between-group differences in pain or quality of life |
<p>| Duke University 5 (42, 43) | X |  | X |  | X |  |  |
| First Moscow State Medical University (41) | X |  |  |  |  |  | • No medication side effects reported |
| INGENIOUS** (13, 50, 51) |  | X |  |  |  |  | • Genotyping prompted PGX consults 10/106 (9%) patients |
| La Paz University Hospital (10) | X | X |  | X |  |  | • Clinical PGX service cost €202,140 over 3 years, with each consult averaging €216 |</p>
<table>
<thead>
<tr>
<th>Study</th>
<th>Simvastatin Prescriptions</th>
<th>Medication Changes</th>
<th>CDS Alerts</th>
<th>Mean Patient WILLINGNESS-TO-PAY</th>
<th>Medication Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Marshfield Clinic***</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>No medication side effects reported</td>
</tr>
<tr>
<td>MedSeq Project(66, 67)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>No between-group differences in 6-month healthcare costs (mean $1490 vs. $1142).</td>
</tr>
<tr>
<td>OSU-Coriell Personalized Medicine Collaborative* (11, 49, 58, 59, 62, 64, 65)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>Mean patient willingness-to-pay for clinical PGx service was $56 ($81)</td>
<td></td>
</tr>
<tr>
<td>PRIMER(60)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>42% of patients not willing to pay out-of-pocket costs for PGx tests; 58% of remainder reported willingness-to-pay of $100</td>
<td></td>
</tr>
<tr>
<td>RIGHT Protocol*** (8, 11, 12, 53-56)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>No medication side effects reported</td>
<td></td>
</tr>
<tr>
<td>Yale University(57)</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>No medication side effects reported</td>
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Outcomes are categorized as utility outcomes from SLCO1B1 testing or the process outcomes that might mediate the relationship between SLCO1B1 testing and utility outcomes. Clinical outcomes include biomarker changes, morbidity, and mortality, while healthcare utilization and economic outcomes include willingness-to-pay and the healthcare costs or other resources required to implement the intervention or resulting from its implementation. Provider utilization and attitudes include frequency of test ordering by providers, their use of the information, and their attitudes about its value. Prescribing behavior includes medication prescriptions, while medication adherence measures patient use of prescribed medications. Other patient-reported outcomes include patient recall of test results, concern or distress about results, and patient perceived utility of the information. Bolded text in the Key Findings column refer to utility outcomes (either clinical or healthcare utilization/economic outcomes). Abbreviations: CDS, clinical decision support; LDL-C, low-density lipoprotein cholesterol; PGx, pharmacogenetic.

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Records identified through database searching, 2005-2017 (n = 5,374)

Additional records identified through manual author and reference search and author contact (n = 57)

Records after duplicates removed (n = 4,863)

Records excluded by title and abstract (n = 4,474)

Records screened (n = 189)

Full-text articles excluded (n = 135)

Full-text articles assessed for eligibility (n = 54)

Ongoing potentially eligible studies without published outcomes (n = 17 records describing 10 studies)

Records included in qualitative synthesis (n = 37 records describing 16 studies)