Implementing pharmacogenomics decision support across seven European countries: The Ubiquitous Pharmacogenomics (U-PGx) project

Kathrin Blagec,1 Rudolf Koopmann,2 Mandy Crommentuijn – van Rhenen,3 Inge Holsappel,3 Catheline H van der Wouden,4 Lidija Konta,5 Hong Xu,1 Daniela Steinberger,2,5,6 Enrico Just,2 Jesse J Swen,4 Henk-Jan Guchelaar,4 and Matthias Samwald1

1Section for Artificial Intelligence and Decision Support; Center for Medical Statistics, Informatics, and Intelligent Systems; Medical University of Vienna, Vienna, Austria, 2bio.logis Genetic Information Management GmbH, Frankfurt am Main, Germany, 3Medicines Information Centre; Royal Dutch Pharmacists Association (KNMP), The Hague, The Netherlands, 4Department of Clinical Pharmacy & Toxicology, Leiden University Medical Center, Leiden, The Netherlands, 5bio.logis Center for Human Genetics, Frankfurt am Main, Germany and 6Institute for Human Genetics, Justus Liebig University, Giessen, Germany

Corresponding Author: Dr. Matthias Samwald, CeMSIIS, BT88, Medical University of Vienna, Spitalgasse 23, 1090 Vienna, Austria; matthias.samwald@meduniwien.ac.at

Received 1 September 2017; Revised 23 November 2017; Editorial Decision 7 January 2018; Accepted 9 January 2018

ABSTRACT

Clinical pharmacogenomics (PGx) has the potential to make pharmacotherapy safer and more effective by utilizing genetic patient data for drug dosing and selection. However, widespread adoption of PGx depends on its successful integration into routine clinical care through clinical decision support tools, which is often hampered by insufficient or fragmented infrastructures. This paper describes the setup and implementation of a unique multimodal, multilingual clinical decision support intervention consisting of digital, paper-, and mobile-based tools that are deployed across implementation sites in seven European countries participating in the Ubiquitous PGx (U-PGx) project.

Key words: clinical decision support systems, pharmacogenomics

INTRODUCTION

Pharmacogenomics (PGx), i.e., using genetic data to guide drug-dosing and selection, emerged as a promising strategy for making pharmacotherapy safer and more effective.1–4 A successful implementation of PGx into clinical practice strongly depends on the availability of clinical decision support (CDS) tools that translate raw genetic test results into concise and clinically actionable therapeutic recommendations, and make those results available to healthcare providers at the point of care.5

Several projects utilizing different variants of PGx testing and CDS have been launched; these projects are described and compared in detail in Supplementary Material S1.6–23

A common factor of these successful PGx implementation projects is the delivery of CDS via the electronic health record (EHR), either as an interruptive alert at the time of prescribing, or as part of the patient’s digital record. Although the availability of EHRs in hospital settings significantly increased within the past decade and reached adoption rates of >50% in most developed countries, nationwide availability still cannot be expected in most regions.24,25 Moreover, a lack of interoperability between different existing EHR systems as well as their fragmented availability beyond hospital settings still constitute substantial barriers to the efficient and secure sharing of PGx data.25–28

In this paper, we describe the development and implementation of a unique, multi-modal, and multi-lingual PGx CDS strategy...
across 7 European countries in the context of the Ubiquitous PGx (U-PGx) project that enables the delivery of PGx CDS in the presence of diverse and fragmented healthcare infrastructures.

Implementing CDS in the U-PGx project

Project setting

Widespread adoption of PGx-guided prescribing in routine care will heavily depend on the availability of robust data from large-scale clinical studies that demonstrate improved clinical outcomes and cost-effectiveness of PGx testing when applied to broad patient populations. The Ubiquitous PGx (U-PGx) project was initiated to address this need by implementing PGx panel testing and CDS across 7 European countries and measuring patient outcomes and cost-effectiveness. The project started in January 2016 with a total duration of 3 years and a budget of 15 million Euros from the Horizon 2020 EU research program.

The project includes a clinical study which was initiated in early 2017. It is designed as a prospective, block-randomized, controlled study, and a total of 8100 patients are planned to be enrolled over the course of three years (Clinicaltrials.gov NCT03093818). Patient recruitment takes place at one or more healthcare institutions in each of the 7 participating countries.

As part of the intervention, patients who receive a first prescription of a drug, for which a pharmacogenomic guideline is available, are tested for a panel of more than 48 clinically relevant PGx variants across 13 important pharmacogenes relevant for dose optimization of 41 drugs. PGx testing is deployed in a mixed “reactive-preemptive” approach, meaning that a full PGx panel is ordered at the time of first prescription of a drug and dosage can be optimized based on PGx guidelines. PGx panel results are then readily available for optimizing prescription of other drugs prescribed in future interactions with the healthcare system.

While having to deal with heterogeneity (e.g., different types of EHRs) is not uncommon in larger CDS implementation projects, this challenge is aggravated within the U-PGx project, where technical framework conditions range from complete absence of any information technology (IT) infrastructure at some implementation sites to sophisticated IT systems with the ability to provide active CDS via automated alerts at others (see Table 1). Furthermore, differences do not only exist between different participating countries but also between participating sites within the same country. Ensuring a standardized intervention while still making optimal use of existing technical capabilities and meeting essential country-specific requirements is therefore a key requirement in the U-PGx project.

Another fundamental challenge lies in maximizing the accessibility of PGx results within and between different healthcare settings. Fragmented health IT infrastructures within countries and lacking interoperability, as encountered in this project, entail a significant risk of experiencing silo effects, meaning that information is trapped in one system or institution and sharing between different systems or institutions is impeded. Preemptive PGx testing is based on the rationale that testing patients for an entire panel of the most important PGx variants at once may be cost-effective, because they will likely profit several times from their PGx results in future interactions with the healthcare system. Avoiding silo effects by ensuring the accessibility and sharing of PGx results within and between different healthcare settings, e.g., in- and outpatient settings, and healthcare providers is therefore vital for a successful implementation of a preemptive PGx strategy.

Besides these technical requirements, the international setting of the project also poses unique challenges caused by the diversity of languages and regulatory frameworks.

Devising a multi-modal CDS implementation strategy

To overcome challenges associated with implementing PGx CDS on an international level and across multiple clinical sites with largely diverging IT infrastructures, the U-PGx consortium has devised a unique implementation strategy drawing from a spectrum of CDS delivery modes deployed inside and outside of EHRs that

Table 1. Characteristics of Existing IT Infrastructures at the U-PGx Implementation Sites

| Infrastructure characteristics | NL | GB | IT | ES | AT | SI | GR |
|-------------------------------|----|----|----|----|----|----|----|
| EHR inpatient setting         | Yes| Yes| Partially| Yes| Partially| No| No|
| EHR outpatient setting        | Yes| Partially| Yes| Yes| Yes| No| No|
| Text reports                  | Yes| Yes| Yes| Yes| Yes| No| No|
| Active CDS                    | Yes (for PGx, DDIs, contraindications, and drug dose) | Yes (for allergies and DDI) | No| Yes (for allergies)| No| No|
| Passive CDS                   | No| Yes| Yes| Yes| Yes| No| No|
| LIMS                          | Yes| Yes| Yes| Yes| Yes| No| No|
| Structured laboratory results | Yes| Yes| Yes| Yes| Yes| No| No|

Abbreviations: AT, Austria; DDI, Drug-drug interactions; EHR, Electronic health record; ES, Spain; GB, Great Britain; GR, Greece; IT, Italy; LIMS, Laboratory information management system; NL, The Netherlands; PGx, Pharmacogenomics; SI, Slovenia.
to the local languages of each participating country (English, German, Greek, Slovenian, Spanish, and Italian) by certified translators and validated by consortium members. Furthermore, representatives from clinical implementation sites curated a list of the most common local trade names for all drugs covered by the project. Phenotype designations (e.g., “CYP2D6 ultrarapid metabolizer”) remained in English to preserve a standardized designation across all implementation sites. For the initial phase of the project, the phenotype and genotype terminology of the G-Standaard was adopted unchanged (see Supplementary Material S3); efforts to standardize and harmonize existing PGx terminologies and therapeutic recommendations developed by different working groups are currently underway.38,39

Decision support tools
U-PGx GIMS does not only act as a centralized knowledge base in the project but also serves as the main portal for the upload of genetic data obtained from the U-PGx genotyping platform, and the retrieval of patient-specific PGx reports in various formats. For this purpose, GIMS offers a wide range of secure data transfer capabilities, ranging from simple file (.csv) imports and uploads to modern web-based application programming interface technologies, including Hyper Text Transfer Protocol Secure (HTTPs)-based representation state transfer (RESTful) services and common standards like Health Level Seven International (HL7) and Fast Healthcare Interoperability Resources (FHIR).40,41

As illustrated in Figure 2, U-PGx GIMS enables PGx CDS by the following means: (1) the secure transfer of PGx test results and patient-specific dosing recommendations in a structured format for incorporation into local EHRs for use in passive or active CDS; (2) the generation of a PGx report in Portable Document Format or Open Document (ODT) format that can be filed either in the patient’s digital or paper-based health record; and (3) the generation of a “Safety-Code” card that enables mobile-based PGx CDS independent of existing IT infrastructures (see Supplementary Material S2).

To conform to privacy and data security regulations, all data that are exchanged between the implementation site and the centralized GIMS (i.e., PGx test results, PGx reports) are done with pseudonyms. Matching of PGx reports with identifying patient information occurs locally at each implementation site.

PGx report
Delivering PGx CDS for an entire panel of PGx variants in a paper-based form requires a careful report design to avoid overwhelming clinicians. The U-PGx report was therefore structured to provide information most relevant at the point of care – such as for which drugs a dosage adjustment is recommended for the respective patient – right at the beginning of the report, whereas additional information – such as the patients detailed PGx results – are provided on the following pages (see Supplementary Material S2).

The report is generated by the GIMS Diagnostic Report Module, which is certified as a medical device and holds the Conformité Européene (CE) mark in accordance with European legislation (EEC 93/42, EC 2007/47).

Safety-code card
To complement paper-based CDS solutions at clinical sites that lack an EHR infrastructure and to maximize the accessibility and sharing of PGx results within and between different healthcare settings and healthcare professionals, the “Safety-Code” card system is deployed at all participating institutions.42,43

---

Figure 1. Decision support solutions in U-PGx. The U-PGx CDS strategy combines several complementary modes of delivering patient-specific PGx therapeutic recommendations to healthcare providers at the point of care, with or without integration into local EHRs. Active, interpretive CDS alerts clinicians of relevant gene-drug interactions via a pop-up message in the EHR or e-prescription system at the time of prescribing. Passive CDS is delivered either inside the EHR system as a digital report, or outside the EHR system via mobile- and paper-based solutions. The different decision support solutions deployed in the U-PGx project including the underlying knowledgebase are described in detail below.

U-PGx knowledge base
Technical realization
For U-PGx, the curation of the knowledge base and the automated translation of genetic data to associated phenotypes and recommendations are handled by the Genetic Information Management Suite (GIMS), a Drupal-based content management system developed and operated by the U-PGx partner bio.logis Genetic Information Management GmbH.32,33 Using a web-based content management system for knowledge base maintenance offers several advantages compared to using local or static solutions, such as a central workflow for editing, translating, reviewing, and validating content and a transparent change history across all participating sites (see Supplementary Material S2).

Content curation
The Dutch Pharmacogenetics Working Group (DPWG) is an ongoing effort to develop concise and clinically actionable recommendations for risk-phenotypes based on systematic literature review, and is formally associated with U-PGx. Up to the time of this writing, the DPWG has authored guidelines for 92 gene-drug pairs across 17 genes, all of which are incorporated into the G-Standaard, a comprehensive Dutch drug database, and regularly updated.29,34 This subset of the G-Standaard, containing the PGx-based therapeutic recommendations, including the data structure that links genotype-predicted phenotypes with active ingredients and therapeutic recommendations was adopted unchanged for the U-PGx knowledge base and is also available to interested parties via an open-source license (see Supplementary Material S3).

Chemical substances and active ingredients are identified by their Chemical Abstracts Service (CAS) number and the Anatomical Therapeutic Chemical (ATC) Classification System code within the knowledge base, respectively; a comprehensive systematic vocabulary such as Systematized Nomenclature of Medicine – Clinical Terms (SNOMED CT) is currently not utilized.35–37 An overview and further description of the knowledge base data model is provided in Supplementary Material S4.

Based on the genotype variants included in the U-PGx panel, rules for translating from genotypes to haplotypes and phenotypes were curated by PGx experts in the project.

DPWG guidelines for clinically actionable phenotype-drug pairs covered by the U-PGx genotyping panel were translated from Dutch
This card is part of a mobile clinical decision support (CDSS) called the Medication Safety Code system which enables quick retrieval of patient-relevant PGx drug dosing guidelines even in the absence of a local EHR infrastructure.

Designed as a credit card-sized plastic card, the “Safety-Code” card contains a quick response (QR) code that can be decoded and interpreted by common smartphones and other devices (Figure 3). After scanning the QR code, the medical professional is led to a website that provides drug dosing recommendations customized to the PGx profile of the patient.

Furthermore, the “Safety-Code” card contains an overview of the patients’ most important PGx test results including a list of drugs for which PGx-based dosing adjustments are recommended. Patients participating in the PRE emptive Pharmacogenomic testing for prevention of Adverse drug Reactions (PREPARE) study are asked to carry their “Safety-Code” cards with them and display them to medical professionals when pharmacotherapy is initiated or altered, which has the additional benefit of promoting patient engagement.

Card contents are generated through GIMS (see Supplementary Material S2), and physical cards are printed locally at implementation sites with dedicated card printers.

The basic architecture of the Medication Safety Code system is publicly available via an open-source license (see Supplementary Material S3).

CDS implementation at clinical sites

Depending on their existing IT infrastructure and associated technical capabilities, each of the participating countries uses at least 2 complementary CDS methods for providing health care providers with patient-specific PGx-based therapeutic recommendations (Table 2). While main methods differ per country, all sites deploy the “Safety-Code” card as an adjunct method to optimize information diffusion within and between different healthcare providers.

Lessons learned

While a final reflection and assessment of our implementation efforts can only be conducted after completion of the project, we nevertheless want to share the most important experiences collected over the course of this initial project phase.

Firstly, sufficient time should be ensured for the curation of the knowledge base, and in particular, the curation of the mapping between the raw data output of the genotyping platform. In addition, it is advisable to establish a workflow for dealing with rare variants that may not be covered by the knowledge base. For U-PGx, such variants are reported to a dedicated mailing list by implementation site representatives, reviewed by experts in the consortium, and added to the knowledge base.
Table 2. Utilization of CDS tools in Each Participating Country

| Planned CDS intervention | NL | UK | I | E | A | SLO | GR |
|--------------------------|----|----|---|---|---|-----|-----|
| Automatic alerts via her | xx | xx | xx | xx | xx | xx  | xx  |
| Paper-based PGx reports  | x  | x  | x  | x  | x  | x   | x   |
| Digital PGx reports via EHR | xx | xx | xx | xx | xx | xx  | xx  |
| Safety-Code card | x  | x  | x  | x  | x  | x   | x   |

Each participating country uses one of the listed CDS interventions as their main method for providing physicians and pharmacists with patient-specific drug dosing recommendations, depending on their existing infrastructure and IT capabilities. Furthermore, each country deploys additional methods to facilitate the transfer of results between different healthcare settings (e.g., inpatient and outpatient setting). Main methods are marked with xx, adjunct methods are marked with x.

Furthermore, the ability to quickly interpret and return partial genotyping results should be considered. At early implementation stages, some sites had problems with assays for 1 or 2 genes in the panel, and requested the generation of PGx reports based solely on the results of the remaining genes. This was not anticipated in the initial design of the reporting software for pre-emptive PGx.

Finally, in case an integration into local EHR infrastructure is envisaged, sufficient lead time should be scheduled for establishing communication and collaboration with the local IT department and dealing with often encountered bureaucratic obstacles, such as obtaining necessary permits. This observation resonates with the experiences reported earlier by Herr et al.30

DISCUSSION AND CONCLUSION

While automated alerts displayed via the EHR at the time of prescribing are commonly viewed as the gold standard for delivering CDS, their implementation is tied to the availability of an adequate technical infrastructure which is currently still insufficient in many healthcare settings.44,45 By developing and implementing a multimodal CDS concept, we demonstrate the feasibility of implementing PGx CDS at multiple clinical sites across 7 countries in the presence of largely fragmented and diverse health care infrastructures. We use several complementary methods, including digital, paper-based, and mobile CDS solutions that allow each participating institution to choose their preferred combination of CDS tools that best fit their institutional preferences and technical requirements.

As of June 2017, all CDS tools had been finalized and rolled out in the countries that were randomized to start with the study arm, i.e., Greece, Slovenia, and Spain. Roll-out of CDS at sites that are currently recruiting patients for the control arm (i.e., UK, the Netherlands, Austria, and Italy) will be commenced in summer 2018. The adoption and usability of the different CDS tools deployed in the U-PGx project will be evaluated at several time points throughout the study period; intermediary results will be used for continuous improvement of the tools.

We hope that the project will successfully address the remaining barriers to widespread adoption of PGx-guided prescribing and preemptive testing strategies.

COMPETING INTERESTS

DS has developed concepts of genetic information management (GIM) that were realized by the company founded as, the bio.logis Genetic Information Management GmbH. For this company, she is the CEO. She is also medical director of bio.logis Center of Human Genetics, a diagnostic institution that is applying and testing the GIM-systems in the context of medical care. Enrico Just is the CEO of the bio.logis Genetic Information Management GmbH. RK is an employee of bio.logis Genetic Information Management GmbH. The other authors have no competing interests to declare.

CONTRIBUTORS

KB wrote major parts of the manuscript. JJS, HJG, and MS were involved in the initial conception of the U-PGx systems. KB, RK, HX, and MS were directly involved in software engineering of components of the U-PGx systems. All authors were involved in shaping some aspects of the U-PGx systems. All authors assisted in revising the manuscript and gave final approval of the version to be published.

FUNDING

This work was supported by European Community's Horizon 2020 Programme grant number 668335 (U-PGx).

SUPPLEMENTARY MATERIAL

Supplementary material is available at Journal of the American Medical Informatics Association online.

REFERENCES

1. Pirmohamed M, Burnside G, Eriksson N, et al. A randomized trial of genotype-guided dosing of warfarin. New Engl J Med 2013;369:2294–2303.
2. Verhoef TI, Ragai G, de Boer A, et al. A randomized trial of genotype-guided dosing of acenocoumarol and phenprocoumon. N Engl J Med 2013;369:2304–2312.
3. Mallal S, Phillips E, Carosi G, et al. HLA-B*5701 screening for hypersensitivity to abacavir. N Engl J Med 2008;359:568–579.
4. Coenen MJH, de Jong DJ, van Marrewijk CJ, et al. Identification of patients with variants in TPMT and dose reduction reduces hematologic events during thiopurine treatment of inflammatory bowel disease. Gastroenterology 2015;149:907–17.e7.
5. Swen JJ, Huizinga TW, Gelderblom H, et al. Translating pharmacogenomics: challenges on the road to the clinic. PLoS Med 2007;4:e209.
6. O’Donnell PH, Bush A, Spatz J, et al. The 1200 patients project: creating a new medical model system for clinical implementation of pharmacogenomics. Clin Pharmacol Ther 2012;92:446–449.
7. O’Donnell PH, Danahey K, Jacobs M, et al. Adoption of a clinical pharmacogenomics implementation program during outpatient care–initial results of the University of Chicago “1,200 Patients Project.” Am J Med Genet C Semin Med Genet 2014;166C:68–73.
8. Gottesman O, Scott SA, Ellis SB, et al. The CLIPMERGE PGx Program: clinical implementation of personalized medicine through electronic health records and genomics-pharmacogenomics. Clin Pharmacol Ther 2013;94:214–217.
9. Scott SA, Owusu Obeng A, Botton MR, et al. Institutional profile: translational pharmacogenomics at the Icahn School of Medicine at Mount Sinai. Pharmacogenomics 2017;18:1381–1386.
10. Johnson JA, Elsey AR, Clare-Salzler MJ, et al. Institutional profile: University of Florida and Shands Hospital Personalized Medicine Program: clinical implementation of pharmacogenetics. Pharmacogenomics 2013;14:723–726.
11. Weitzel KW, Alexander M, Bernhardt BA, et al. The IGNITE network: a model for genomic medicine implementation and research. BMC Med Genomics 2016;9:1.
