Pharmacogenomics (PGx) offers the promise of utilizing genetic fingerprints to predict individual responses to drugs in terms of safety, efficacy and pharmacokinetics. Early-phase clinical trial PGx applications can identify human genome variations that are meaningful to study design, selection of participants, allocation of resources and clinical research ethics. Results can inform later-phase study design and pipeline developmental decisions. Nevertheless, our review of the clinicaltrials.gov database demonstrates that PGx is rarely used by drug developers. Of the total 323 trials that included PGx as an outcome, 80% have been conducted by academic institutions after initial regulatory approval. Barriers for the application of PGx are discussed. We propose a framework for the role of PGx in early-phase drug development and recommend PGx be universally considered in study design, result interpretation and hypothesis generation for later-phase studies, but PGx results from underpowered studies should not be used by themselves to terminate drug-development programs.

**Keywords:** clinical research, drug development, early-phase, PD, PGx, pharmacodynamics, pharmacogenetics, pharmacogenomics, pharmacokinetics, PK, POC, proof-of-concept

A genomic biomarker is defined as ‘a measurable DNA and/or RNA characteristic that is an indicator of normal biologic processes, pathogenic processes and/or response to therapeutic or other interventions’. Pharmacogenomics (PGx) is defined as ‘the study of variations of DNA and RNA characteristics as related to drug response’. Pharmacogenetics is a subset of PGx and is defined as ‘the study of variations in DNA sequence as related to drug response’ [1]. Another way of thinking about the difference between pharmacogenetics and PGx is that the former deals with single genes and the latter with the entire human genome [2,3]. PGx focuses on the predictive outcome of drug interventions as opposed to genomic predictors of the natural course of illness, diagnostics and prognostics (‘genomics’). PGx is the study of the interaction between the drug and the individual in terms of drug efficacy, safety and pharmacokinetics (PK). It incorporates information from across the translational spectrum, from gene–disease relationships through confirmatory clinical studies, into the development and eventual application of new drugs [4]. PGx offers the promise of delivering personalized and targeted drug therapy to those most likely to benefit from it [5–8]. PGx may also contribute to the design and interpretation of clinical trials and improve effectiveness, safety and overall benefit:risk ratio of drugs in development [9,10]. It has the potential for earlier arrival at informed developmental decisions – an alternative to the ‘trial and error’ approach. Both the US FDA and NIH have identified PGx as a key tool in the drug-development armamentarium [11,12]. This review will start with a general background of PGx, proceed with a discussion of PGx applications in early-phase research and conclude with recommendations for future development of the field. PGx stakeholders to whom this review is addressed include drug developers, clinical and preclinical investigators, biostatisticians, translational science experts and drug-development consultants, clinical research operators (early phase), analytics operators and experts, business developers, healthcare payers, regulators and policy makers.

**History**

PGx considerations, even before the term was in use, can be traced to ancient times [13]. The principle of personalized treatment tailored to someone’s physical and physiological constitution is long-held [14,15]. Pythagoras’ reluctance to pass through fava bean fields, possibly contributing to his death by captors, may have been due to his recognition of his own glucose-6-phosphate dehydrogenase (G6PD) deficiency and the hemolytic anemia associated with consumption of fava beans [16,17]. Treating someone based on their body type or other constitutional factors, as is the practice in ancient Ayurveda, Chinese, Tibetan and Iranian traditional medicine systems, incorporates genetic
features into treatment considerations [13–15,18,19]. Evidence-based medicine may have yet to validate some claims of traditional medicine, but these examples demonstrate that the principles of PGx and related personalized medicine were long part of medical approaches and practice. More recently, at the dawn of a modern trend, the advent of PGx as a science and development tool was first conceptualized by Motulsky in 1957 [20–22]. A twin study of dicumarol pharmacokinetics provided the first evidence of genetic-based pharmacokinetics [23]. Another example of a PGx-guided individualized intervention that may not be thought of as such but has been practiced for decades is anesthesia, where the doses of the anesthetics are continuously adapted to the individual’s signs and symptoms [24].

**Background**

There are many interindividual differences in response to drug therapy, many may not matter, but some do and may pertain to the interaction of drug with genetically determined physiological and pathophysiological mechanisms. The differences that matter pertain to three main categories or domains of study in PGx: PK, beneficial pharmacodynamics (PD; beneficial, or efficacy), and adverse PD (adverse or toxicity). In every step of PK, or the processes that govern what the body does to the drug (principally absorption, distribution, metabolism and excretion), there could be genetic variability in the following: gastrointestinal tract environment and absorption, active transport across membranes, metabolism, protein binding. Similarly, genetic traits can impact a drug’s PD, or the mechanism(s) governing the drug’s actions at its targets (drug-related phenotypes [4]). The impact of interspecies genomic differences (between rhesus monkeys and humans) on a drug’s target binding and consequent dramatic clinical manifestations is exemplified by the first-in-man study of the anti-CD28 monoclonal antibody TGN1412 [25,26]. Genetic differences in response to drugs (e.g., due to CYP450 polymorphisms and G6PD deficiency) may be inter- or intra-ethnic [17,27–30]. PGx principles may inform drug developers’ marketing strategies and healthcare providers’ insurance policies [31–33].

Three main PGx approaches are applied in drug development:

* Candidate gene studies. Also called monogenic (or simply genetic rather than the below genomic approaches) or targeted pathway analysis [10], these are studies of single genes known or hypothesized to impact specific pathways relevant to drug response (e.g., CYP2D6 metabolism) [34]. These studies are hypothesis-driven with predetermined and limited numbers of specific genetic associations between the pharmacological intervention and its outcomes. The disadvantage of the approach is that it may overlook associations with unknown and unanticipated genetic candidates. The advantage is the small number of analyses and the often large effect sizes [34,35]. This has been the first approach to be used in PGx studies because it did not require access to the full human genome, available only since 2000 [23,34].

* Genome-wide association studies (GWAS) involve comparing two groups of individuals with differing characteristics (e.g., drug response) and identifying associations with many known genetic variants. Since the genetic variants are chosen without prior knowledge or suspicion of an association with the differing clinically relevant phenotypes (e.g., disease, responses to treatment), GWAS studies are considered an unbiased, ‘agnostic’ approach [36]. GWAS may assess associations of up to 1 million genetic variants with clinically relevant outcomes [37]. Associations found in GWAS do not necessarily imply causality of the identified genetic marker in the variation of the phenotype. Hence, follow-up studies on the impact of modifications in gene expression on the phenotype are required [4]. The multiple associations studied require adequate statistical corrections. With a few notable exceptions and notwithstanding the relatively small numbers tested so far using GWAS, it appears that PGx effect sizes using GWAS are usually small [34,35].

* Whole-exome and whole-genome sequencing – the most comprehensive approach that analyzes the entire human genetic material for variants relevant to drug response [38]. It may help identify novel or very rare variants that would not be discovered via GWAS, but nevertheless may need to be combined with GWAS to evaluate associations with drug response. Although interpreting such data would be difficult. With reduced cost of whole-genome sequencing (at ~US$3000 in 2012 [4]) this approach is expected to become the dominant PGx approach.

**Uses in clinical practice**

Personalized medicine and the related term, individualized therapy, have been defined as
the practice of adapting treatment to an individual’s medical condition, genetics, demographics, environment and lifestyle [38–40]. PGx addresses the genetic component of personalized medicine as it applies to drug therapy. Table 1 lists examples of successful applications of PGx in clinical care. The approach holds the promise of providing more effective, targeted treatments and quicker arrival at optimal care. This could translate into better public health benefits and eventually reduced hospitalizations and burden of illness, and increased productivity of affected individuals [33,41–43]. PGx has the potential to improve the health of vulnerable populations and neglected diseases [44]. Nevertheless, the application of PGx knowledge and principles in clinical practice has been slow [4,32,37,45–48]. PGx-driven clinical practice guidelines with decision-making algorithms informed by controlled-clinical trials may increase precision, accuracy and relevance of recommendations and subsequent applicability [28,49].

PGx use in drug development
Understanding the science of PGx: the genotype–phenotype interactions that are relevant to drug response [50] are:

* Disease genotypes and phenotypes – understanding the genetic origins, mechanisms and manifestations of illnesses is crucial to cure, prevention or mitigation efforts, the processes that influence restoration of health, and the design of drugs to carry out these effects.

* Drug – the therapeutic agent whose actions are determined by the availability of genomic-driven targets and processes [51].

* Drug-relevant genotypes (genetic and epigenetic) – these are the genomic components (DNA and RNA) coding and determining drug-relevant phenotypes (see below). The genetic material may be germline (heritable, part of host constitution) or acquired (e.g., tumor mutations), and in either case may be modified by epigenetic and/or nongenetic elements (e.g., infections and toxins) temporarily or permanently [52].

* Drug-relevant phenotypes (biomarkers) – these are physiological and/or pathological phenotypes that are relevant to drug actions. These could be efficacy outcomes, adverse events, drug plasma concentrations or other PK/PD biomarkers. As the experience with gefitinib demonstrates, earlier identification of a predictive biomarker (e.g., the presence of EGFR mutation in the case of gefitinib; Table 1), understanding the connection between genetic markers and drug response can make drug-development programs more focused, effective and productive [29]. Membrane transporters and genetic variations in their expression will influence PK, efficacy and safety of drugs [53,54]. The drug may use normal physiological systems and processes to reach the target (e.g., chronotherapeutics – the impact of circadian rhythms on drug response [55]) and may impact normal physiological systems in a diseased individual.

Applications of PGx methods in clinical trials in early-phase clinical development
Application of PGx principles in drug development is a continuous process that starts with discovery and continues through the drug’s postmarketing period [3,31,56–68]. Most PGx discoveries to date have been made in postmarketing studies [10,37], but a consensus is being established over the utility and possibly the indispensable nature of PGx approaches as components of all stages of drug-development programs [4,35]. Some also recommend that all commonly used medications should undergo PGx investigation [35]. Early-phase development is an important landmark in the overall R&D process since it presents the first introduction of the drug to humans. It is a phase characterized and limited by small sample sizes and trial durations due to ethical and economic reasons, as well as by uncertainties regarding doses, outcome measures, target populations, efficacy and toxicity parameters, their scope and magnitude.

Mechanistic pathways relevant to drug PD (efficacy and safety) affected by genetic polymorphisms with relevance to drug targets, and actions, can be determined before Phase I studies in humans [69]. Similarly, genetic factors affecting drug metabolism, transporters and other PK parameters can be studied in vitro and in vivo prior to initiation of human studies [55,70]. Such findings can then be incorporated in early-phase human study design. For example, genetic variations in HLA detected by in vitro tests may allow prediction of allergic and other idiosyncratic adverse events in the near future [69,71]. Drug targets may also be identified through their adverse event profile, and what may be undesirable or unexpected effects in one development program may prove desirable and beneficial in another [51]. PGx offers the potential of excluding patients at
Figure 1. Total number of trials with pharmacogenomics outcomes (and percentage of total trials) per year in the clinicaltrials.gov database. Studies are divided into early phase (Phases 0, I and II) and late phase (Phases III and IV) 1999–2012. PGx: Pharmacogenomics.

Figure 2. The percentage of pharmacogenomics studies sponsored by industry and academia.

Figure 3. Number of pharmacogenomics studies per therapeutic area. CVS: Cardiovascular system; GI: Gastrointestinal; PGx: Pharmacogenomics.

Analysis of clinicaltrials.gov database for pharmacogenomics in clinical trials.

**Purpose:** To assess the scope of application of PGx principles and outcomes in clinical trials as reflected in trials registered on clinicaltrials.gov. **Methodology:** Clinicaltrials.gov database [201] was accessed on 1 January 2013. The ‘search for studies’ feature was used to conduct the search with ‘pharmacogenomics’ as the keyword (also returning entries for ‘pharmacogenomic’). Each study entry was reviewed by the two authors and information on phase, sponsor, therapeutic area, objectives, study start date, sample size and end points captured. **Results:** A total of 323 studies included PGx as outcome (primary or secondary) in the entire 14 years (1999 to 2012) available in the clinicaltrials.gov database. This represents 0.23% of the total 138,416 studies registered during that period (mean: 341.4 participants per study; standard deviation: 878.2). 164 were early-phase (Phase 0, I and II) studies comprising 0.32% of the total 50,994 early-phase studies. PGx study numbers increased gradually from 2003 but have plateaued since 2008. Out of the total, 258 (80%) have been sponsored by academic institutions, 27% by industry and 7% were industry–academic collaborations. Oncology, at 193 studies (59.7%), was the therapeutic area with most PGx, followed by CNS (45 studies; 13.9%). See Figures 1–3. **Conclusion:** Over the 14-year period of the clinicaltrial.gov database, PGx applications in clinical trials (both general and early phase) remained limited with minimal growth.

PGx: Pharmacogenomics.
<table>
<thead>
<tr>
<th>Drug</th>
<th>Disease</th>
<th>Phenotype</th>
<th>Type of impact (PK, toxicity, efficacy)</th>
<th>Percentage of patients that could benefit</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiopurine therapy</td>
<td>Examples include acute</td>
<td>TPMT intermediate activity</td>
<td>Toxicity (myelosuppression)</td>
<td>3–14% – those homozygous or heterozygous TPMT [28]</td>
<td>Homozygous TPMT-deficient patients (less than 0.5% of individuals) usually experience life-threatening myelosuppression</td>
</tr>
<tr>
<td></td>
<td>lymphoblastic leukemia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>and ulcerative colitis</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Codeine (but</td>
<td>Pain</td>
<td>CYP2D6 conversion of codeine to</td>
<td>PK (metabolism) and consequent efficacy, toxicity</td>
<td>0–10% poor metabolizers; 1–2% ultrarapid metabolizers [83]</td>
<td>Poor metabolizers have no analgesia; ultrarapid metabolizers risk morphine toxicity</td>
</tr>
<tr>
<td></td>
<td>(but applies to many</td>
<td>morphine, the active metabolite</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>drugs)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Warfarin</td>
<td>Hypercoagulable states</td>
<td>CYP2C9 metabolism of S-warfarin,</td>
<td>PK (metabolism) and consequent efficacy, toxicity,</td>
<td>Asian (86%) and Caucasian (55%) incidence of ‘low dose’ individuals from either VKORC1 or CYP2C9 variants (from at least one VKORC1 haplotype A or CYP2C9 variant allele) [91]</td>
<td>Narrow therapeutic range Major bleeding in 10–16% [10] PGx findings led to US label updates in 2007 and 2010</td>
</tr>
<tr>
<td></td>
<td>(e.g., thromboembolism)</td>
<td>the potent enantiomer [10]</td>
<td>(anticoagulation), as well as toxicity (lower chance of major hemorrhage) [10,33]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>VKORC1 inhibition by warfarin (VKORC1 gene encodes the target enzyme of warfarin)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>Hypercoagulable states</td>
<td>LOF allele of CYP2C19</td>
<td>PK (CYP2C19 metabolism to active drug)</td>
<td>Up to 30% of study population [92]; 2–14% of the general population are poor CYP2C19 metabolizers [85]</td>
<td>PGx findings led to US label updates in 2009 and 2010</td>
</tr>
<tr>
<td></td>
<td>(e.g., thromboembolism)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gefitinib</td>
<td>NSCLC</td>
<td>EGFR</td>
<td>Efficacy</td>
<td>12–18% [29]</td>
<td>PGx findings made during drug development; first EGFR-targeted category approved for NSCLC [29]</td>
</tr>
<tr>
<td>Trastuzumab</td>
<td>Breast cancer</td>
<td>HER2-positive tumors</td>
<td>Efficacy (overall survival and disease-free survival)</td>
<td>20% of breast cancer patients</td>
<td>Cardiac toxicity an important consideration in benefit:risk analysis [93]</td>
</tr>
<tr>
<td>Abacavir</td>
<td>HIV</td>
<td>HLA-B*5701</td>
<td>Safety (identifying serious adverse event: hypersensitivity reaction)</td>
<td>5–8% of clinical trial patients [10,94]</td>
<td>Postmarketing – leading to US label update in 2008 100% negative-predictive value for immunologically confirmed hypersensitivity reaction</td>
</tr>
<tr>
<td>Ivacaftor (Kalydeco)</td>
<td>Cystic fibrosis</td>
<td>G551D mutation in the CFTR gene</td>
<td>Efficacy [95]</td>
<td>The mutation exists in 4% of cystic fibrosis patients</td>
<td>Development of the drug guided by PGx principles [37]</td>
</tr>
<tr>
<td>Irinotecan</td>
<td>Colorectal cancer</td>
<td>Lower capacity to metabolize SN-38</td>
<td>Toxicity [96]</td>
<td>UGT1A1*28 mutation in 26–31% of Caucasians, 42–56% of Africans and 9–16% of Asians, but may be higher in Indians [97]</td>
<td>One of the first drugs to receive pharmagenomically guided labeling requirements by the US FDA in 2005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(active metabolite of irinotecan)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>patients homozygous for the UGT1A1*28 allele</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

LOF: Loss of function; NSCLC: Non-small-cell lung cancer; PGx: Pharmacogenomics; PK: Pharmacokinetics; SN-38: 7-ethyl-10-hydroxy-camptothecin (the active metabolite of irinotecan).
risk for adverse events from clinical trials and eventual clinical care. PGx identification and exclusion of those most likely to experience an adverse event, as in the case of abacavir hypersensitivity, have resulted in increased utilization and sales of this drug [72].

Utilization of PGx principles may be especially favorable in Phase II studies [32]. But even though O’Donnell et al. report only 19% positive PGx findings (defined as ‘worthy of additional follow-up and validation’) in Phase I compared with 70% in Phase II, such findings may still return the investment by contributing to more effective Phase II and Phase III designs and shorter time to critical developmental decisions (i.e., decisions influencing patent-life utilization of the drug under study or back-up candidates). A comprehensive review of PGx study designs is available elsewhere [69]. Randomized controlled trials in early-phase may be preferred for PGx studies as they provide more definitive information regarding predictive properties of biomarkers and subgroups than open-labeled or single-armed trials. Even though randomized controlled trials are more expensive, they may return the investment by providing more precise information about efficacy and safety of the drug in subpopulations of interest and facilitate faster and more focused later-phase development [29]. However, if clear association is demonstrated between genetic polymorphism and drug response in in vitro studies such randomization may be unethical (Box 1, point 3). Adaptive designs based on PGx data collected in the early part of the study may improve the efficiency and cost–effectiveness of early-phase PGx trials.

Genetic profiling of subgroups at risk for adverse events may allow reintroduction of effective drugs that were withdrawn by regulators owing to rare but serious adverse events in the postmarketing period. Such ‘resuscitation’, ‘resuscitation’ and ‘repurposing’ of drugs has been proposed by the NIH as a key initiative in the fight against stagnation in drug development [11,69]. Finally, PGx technologies may facilitate development of drugs for vulnerable populations and rare diseases, reduce ethnic disparities in applicability, success and generalizability of PGx and disease-specific genomic data (e.g., from target validation, in vitro and in vivo preclinical studies), consultations with subject-matter experts and regulators, and validation of assays, methods and models involved [73,74]. It is further recommended that all drugs in development be considered for application of PGx principles in their early-phase clinical development, and especially those meeting the criteria outlined in Box 4. Toxicity and PK PGx biomarkers may have advantage over efficacy biomarkers in Phase I, even in oncology studies where efficacy Phase I studies are routine [32].

Methodological PGx considerations should take into account the small, short and exploratory nature of early-phase trials. PGx findings in such studies are likely to be hypothesis-generating rather than hypothesis testing, but nevertheless capable of providing mechanistic support of PGx effects relevant to drug response. If identified, these effects will need to be confirmed in definitive, larger studies in later stages of development [10]. PGx study designs including participant selection criteria and outcomes should be tailored to the drug and disease under consideration [10,69]. Genetic material should be collected from all participants if possible to increase power in these usually small studies, reduce selection bias and optimize the opportunity to identify and study outliers. Samples may be analyzed at a later stage of drug development when they can be combined with other sources of information to increase the likelihood of detecting meaningful signals. Potential relevance of active metabolites to efficacy and toxicity (as in the case of codeine, tamoxifen and clopidogrel) should be considered when designing PGx studies. PGx-specific statistical methods should be used in study design and analyses of the data [75].

Operational standardization is crucial for the applicability, success and generalizability of PGx studies. Prospective and routine collection of biological samples (germline: e.g., blood, buccal cells; acquired mutations: e.g., tumor tissue and metastases) from healthy volunteers or patients in all clinical trials should be practiced [10,76]. This should be accompanied by proper, long-term storage of DNA, RNA and tissues relevant to drug response. Routine PGx language should

---

**Challenges & recommendations for PGx applications in early-phase trials**

Box 2 outlines the challenges to PGx applications in early-phase development and Box 3 outlines recommendations aimed at increasing the likelihood of success of such applications. To take full advantage of the potential contribution of PGx approaches, study designs and data to early-phase drug development, PGx strategies should be considered well before entry into human trials, preferably at least 2 years before anticipated first-in-man studies. This will allow sufficient time to gather available drug-specific and disease-specific genomic data (e.g., from target validation, in vitro and in vivo preclinical studies), consultations with subject-matter experts and regulators, and validation of assays, methods and models involved [73,74].
Box 1. Benefits of pharmacogenomics applications in drug development.

1. Identifying drug targets – those relevant to the drug’s efficacy, safety, pharmacokinetics
2. Study design [98]:
   a. Outcomes that could be impacted by genotype variability:
      i. Pharmacokinetics – area under the curve, $C_{\text{max}}$, $T_{\text{max}}$, clearance, volume of distribution, half-life, trough drug concentrations [10]
      ii. Pharmacodynamics – biomarkers related to efficacy and/or toxicity [99]
      iii. Drug–drug interactions
   b. Dose selection – influenced by existing population and subpopulation information on dose–response and concentration–response relationships of genes relevant to the drug or disease under study [10,100]
   c. Therapeutic window for the drug: identifying drug plasma concentrations that are between toxic levels (upper limit) and non-effective levels (lower limit)
   d. Selection of participants (i.e., contributing to more meaningful inclusion/exclusion criteria) – for example: inclusion of ethnic or other subpopulations with known genetic variants relevant to drug response (e.g., apoE [101])
   e. Prospective selection of covariates to understand gene-covariate impact on variability [10]
   f. Statistics – exploratory and hypothesis-generating approach is the rule in early-phase studies
3. Ethics: adhering to pharmacogenomic principles would enable more ethical study designs by limiting the testing of new medications to those most likely to benefit and least likely to experience adverse outcomes
4. Acquisition of information and expertise required to design future studies (i.e., Phase III and Phase IV)
5. Potential for genotype-specific regulatory approvals and product labeling [29,69,102]
6. Drug ‘rescue’ and ‘repurposing’
7. Vulnerable population and rare disease drug development [44]

be incorporated into informed consents and obtained, preferably, from all trial participants at screening to avoid introduction of selection bias by those who discontinue the study. Technological platforms (e.g., analytic and diagnostic) suited to the drug, disease and design under consideration should be identified and validated prior to study initiation.

Business development and pipeline decision-making and considerations may drive PGx approaches. PGx data may drive and contribute to drug labeling and related business

Box 2. Challenges to methodology and applications of pharmacogenomics in early-phase clinical trials.

1. Limited knowledge of genotypes and phenotypes relevant to drug response (efficacy/safety/pharmacokinetics). This may be particularly applicable to the development of drugs with new targets and/or mechanisms of action (applies to items next two points below as well)
2. Limited knowledge regarding disease phenotypes and genotypes
3. Limited knowledge regarding target populations and ethnic subpopulations relevant to drug response
4. Limited availability of technology and expertise to design and interpret PGx studies
5. Methodological hurdles – PGx signals may be too weak for the limited power of early-phase studies. Frueh argued ‘you cannot be personalized if you randomize’ [103] implying the goals of personalized medicine and the current gold standard of clinical research might be conceptually and methodologically in conflict
6. Regulatory hurdles to the standardization of PGx study methodologies and use of PGx data in drug labels and clinical practice [86,88,104]
7. Ethical hurdles – PGx studies require a definition of meaningful efficacy and toxicity thresholds. However, such thresholds are challenging to define [105]. Equity of access to expensive diagnostics and treatments is another ethical issue relevant to PGx applications [106]. Comprehension and confidentiality hurdles can limit healthy volunteer and patient participation in PGx studies [107]
8. Business and financial considerations:
   a. Limited knowledge, at earlier phases of development, about a drug’s eventual market value, extent and scope of developmental program
   b. Limited adoption of PGx principles by drug developers. Only 27% of studies that include PGx outcomes are carried out by industry according to our analysis of the clinicaltrials.gov database
   c. Healthcare payers – although enthusiastic about PGx, are still struggling to translate study findings into effective policies [32,33]
   d. Cost – the cost of whole-genome sequencing was approximately US$3000 in 2012 [4]. PGx cost is trending down while speed and efficiency are increasing. Whole-genome sequencing has the advantage that it only needs to be done once per lifetime

PGx. Pharmacogenomics.
development and marketing strategies. PGx findings may play an important part in decisions to rescue or repurpose drugs based on promising PGx signals in subgroup analyses.

PGx in Phase 0 (microdosing) studies can be used to study PK parameters (absorption, distribution, metabolism and excretion), transporter polymorphisms and imaging biomarkers \[76-79\]. An example is the utilization of PET to image receptor binding of the drug labeled with positron-emitting nuclides. Doses needed to image receptor binding are typically in the microdose levels, thus benefiting from regulatory leniency at the pre-investigational new drug phase

---

**Box 3. Recommendations for the application of pharmacogenomics in early-phase clinical trials.**

**A. General (including PGx applications during preclinical development)**

1. **Logistics:**
   - i. Early planning – 2 years before human trials
   - ii. PGx considered in all POC clinical trials
   - iii. Drug/disease genomic data gathered
   - iv. Experts identified and consulted
   - v. Regulators engaged
   - vi. Validation of genotyping and phenotyping

2. **Methodology:**
   - i. Exploratory and hypothesis-generating
   - ii. Study types: pharmacokinetics/pharmacodynamics in healthy volunteers and/or patients; dose-response
   - iii. Outcomes: phenotypes, genotypes of drug response, subpopulations, high-risk, outliers
   - iv. PGx data collected from all participants
   - v. Active metabolites considered as targets
   - vi. Validation of genotyping and phenotyping assays
   - vii. Statistics: PGx-specific statistical methods

3. **Operational:**
   - i. Routine collection of biological samples
   - ii. Long-term storage of DNA, RNA and tissues
   - iii. Identify technological platforms (e.g., diagnostics)
   - iv. Incorporation of PGx language into informed consents

4. **Business development & pipeline:**
   - i. Drug labeling
   - ii. Drug ‘rescue’ and ‘repurposing’

**B. Phase 0 (microdosing): PGx can be used to study:**

1. Pharmacokinetics (ADME – absorption, distribution, metabolism and excretion)
2. Transporter polymorphisms
3. Imaging biomarkers

**C. Phase I: In addition to the above, PGx can be used to study:**

1. Pharmacokinetics in MAD/SAD studies
2. Active metabolites
3. Toxicity
4. Drug–drug interactions
5. Efficacy (e.g., oncology) or proxy-efficacy biomarkers
6. Healthy volunteers (unless too toxic)
7. Validity and utility of preclinical data
8. Inform Phase II and later-phase study design

**D. Phase II: In addition to the above, PGx can be used to study:**

1. Focus on efficacy biomarkers and associated genomic parameters
2. Follow-up on any signals generated in earlier studies
3. Dose range and ‘therapeutic window’

**E. Validation & follow-up**

1. PGx signals validated in adequately powered studies
2. Low-frequency genotypes may preclude follow-up studies
3. PGx relevance to regulatory submissions/labeling
4. New indications, guidelines, pharmacovigilance

MAD: Multiple ascending dose; PGx: Pharmacogenomics; POC: Proof-of-concept; SAD: Single ascending dose.
of development [80,81]. Such imaging of receptor binding could help identify subpopulations exhibiting variability in binding to receptors of interest.

During Phase I, PGx can be used to study, in addition to the above, toxicity parameters in single and multiple ascending dose studies, drug–drug interactions, conversion to metabolically active compounds and, in some cases (e.g., oncology), drug efficacy or proxy-efﬁcacy biomarkers (i.e., biomarkers suggesting efﬁcacy even in healthy volunteers [e.g., reduction in blood pressure with antihypertensive drugs]). Phase I PGx studies can be used to explore magnitude and relevance of preclinical PGx signals pertaining to drug targets and mechanism of action and PK parameters [69]. Phase I PGx ﬁndings can inform Phase II and later-phase study design, including stratification of subpopulations, choice of doses, biomarkers and diagnostics [69].

In addition to the above (Phase 0 and Phase I) Phase II studies should focus on efﬁcacy biomarkers, follow-up on any signals generated in earlier studies (including preclinical) and help identify the range and therapeutic window of dosing regimens (especially useful for drugs with a narrow therapeutic index) [82]. These in turn should be followed by conﬁrmation in large, adequately-powered and long-term studies where possible, and results incorporated, after regulatory approval, into product labeling, practice guidelines, repurposing of approved drugs (new indications), healthcare policies and pharmacovigilance programs [11,28,83,84].

Economics of PGx applications in early-phase drug development

Will PGx applications improve the economics of drug development? Utilization of PGx principles in drug development may lower drug costs by reducing size and length of clinical trials, permit earlier arrival at developmental decisions and increase the postapproval patent-protection period [89]. Hence, the impact of expensive PGx methodology and smaller eventual market may be mitigated by smaller and shorter studies, more efﬁcient and cost-effective drug-development programs. Importantly, drug-development programs may ground to a halt and effective drugs may not reach regulatory approval if subpopulations experience prohibitive levels of adverse events or efﬁcacy signals are diluted by inclusion of non-responsive subpopulations. In addition, an ethical challenge presents itself

---

**Box 4. Criteria favoring inclusion of pharmacogenomics design in early-phase clinical trials.**

1. Outcomes suggesting clear and meaningful differences among subgroups: exceptional efﬁcacy, severe toxicity and outlier pharmacokinetics
2. Known drug-response phenotypes
3. Drug-response phenotypes are associated with known genomic markers
4. Existing assays for the phenotypic biomarkers and genomic markers
Pharmacogenomics (PGx) is the study of variations of DNA and RNA characteristics as related to drug response. PGx offers the promise of matching the person and the illness with the optimal treatment. There is a growing number of clinical applications of PGx in routine healthcare and a large number of potential future applications.

**Conclusion**
The process of developing a new treatment involves progressive reduction of uncertainties regarding efficacy, safety, and, in the case of drugs, PK. PGx is potentially a powerful tool in reducing such uncertainties. Currently, PGx application in early-phase clinical research is minimal (at less than 1% of studies), stationary, and facing a host of challenges. We believe the promise of safe, effective, specific and personal treatments will serve as a powerful driver of PGx utilization in clinical development in general and early-phase research in particular and help overcome the challenges facing the PGx field. Familiarity with PGx methodologies and technologies, wider application and eventual reduced costs will lead to more efficient, cost-effective and productive drug-development programs and translation of genomic findings into meaningful improvements in patient care. We proposed a framework for the role of PGx in early-phase drug development and recommend PGx be universally considered in study design, result interpretation and hypothesis generation for later-phase studies, but PGx results from underpowered studies should not be used by themselves to terminate drug-development programs.

**Future perspective**
Notwithstanding the low and stationary rate of PGx utilization in early-phase clinical trials in the past decade, we predict a growing and steady increase in utilization for the upcoming 5–10 year period (2013–2023) and beyond. The main reason for this prediction is the belief that an approach that has the potential to provide valuable information for the development of effective, safe, specific and personal treatments will drive the approach past its challenges. Each of the challenges we have identified (limited knowledge about targets, genes and outcomes, limited familiarity with PGx, regulatory and ethical hurdles, methodological challenges and high costs) is manageable and improving [38,47]. In addition, current regulatory guidelines and efforts [10], pressures on industry to make drug development a more efficient and informed process [86,89], pressures from insurers to develop specific and cost-effective treatments [32] and public appeal for personalized therapeutics [90] will all serve as incentives to grow and establish PGx as an indispensable component of the clinical research and development process.

**Acknowledgements**
The authors would like to thank J Sundy and D Voora for their review and valuable comments made to the manuscript.

**Financial & competing interests disclosure**
The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties. No writing assistance was utilized in the production of this manuscript.

---

**Executive summary**

**Background & uses in clinical practice**
- Pharmacogenomics (PGx) is the study of variations of DNA and RNA characteristics as related to drug response. PGx offers the promise of matching the person and the illness with the optimal treatment. There is a growing number of clinical applications of PGx in routine healthcare and a large number of potential future applications.

**Clinicaltrials.gov analysis**
- Less than 1% of clinical trials registered at clinicaltrials.gov included PGx outcomes. The rate has been stationary for the past 6 years and was similar for early- and late-phase trials. Oncology accounted for approximately 60% and academia for 80% of PGx trials.

**Application in early-phase clinical development**
- PGx can substantially contribute to early-phase study design, selection of outcomes, stratification of participants, allocation of resources and improvement of clinical research ethics. Results can inform later-phase study design and pipeline developmental decisions.

**Challenges & recommendations for PGx application in early-phase trials**
- Challenges include: limited knowledge about targets, genes and outcomes, limited familiarity with PGx, regulatory and ethical hurdles, methodological challenges and high costs. We recommend PGx be universally considered in study design, result interpretation and hypothesis generation for later-phase studies, but PGx results from underpowered studies should not be used by themselves to terminate drug-development programs.
References


The document contains multiple references to pharmacogenomics and its application in drug development and personalized medicine. It highlights the importance of understanding genetic variations in drug metabolism and pharmacokinetics to optimize drug efficacy and reduce adverse effects. The text references various studies that explore the role of pharmacogenomics in early-phase clinical development, drug discovery, and in the context of breast cancer and other diseases. It also discusses the integration of pharmacogenomics into clinical trials and the role of pharmacogenomics in anticoagulant therapy. The document emphasizes the need for a deeper understanding of pharmacogenomics to improve patient care and develop more effective treatments.


103 Frueh FW. Back to the future: why randomized controlled trials cannot be the answer to pharmacogenomics and personalized medicine. Pharmacogenomics 10(7), 1077–1081 (2009).


Website
Clinicaltrials.gov database. www.clinicaltrials.gov