Pharmacogenomics: A road ahead for precision medicine in psychiatry

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ABSTRACT

Psychiatric genomics is providing insights into the nature of psychiatric conditions that in time should identify new drug targets and improve patient care. Less attention has been paid to psychiatric pharmacogenomics research, despite its potential to deliver more rapid change in clinical practice and patient outcomes. The pharmacogenomics of treatment response encapsulates both pharmacokinetic (“what the body does to a drug”) and pharmacodynamic (“what the drug does to the body”) effects. Despite early optimism and substantial research in both these areas, they have to date made little impact on clinical management in psychiatry. A number of bottlenecks have hampered progress including a lack of large-scale replication studies, inconsistencies in defining valid treatment outcomes across experiments, a failure to routinely incorporate adverse drug reactions and serum metabolite monitoring in study designs, and inadequate investment in the longitudinal data collections required to demonstrate clinical utility. Nonetheless advances in genomics and health informatics present distinct opportunities for psychiatric pharmacogenomics to enter a new and productive phase of research discovery and translation.

INTRODUCTION

Psychiatric disorders are major contributors to the global burden of disease, accounting for 19% of all years lived with disability worldwide (Rehm and Shield, 2019). The societal impact of these conditions arises from their high prevalence, with one in 5-6 adults affected annually (Baker, 2020), as well as their substantial effects on morbidity and premature mortality. Notably, the global burden of psychiatric disorders has remained high throughout recent decades (Jorm et al., 2017), in contrast to improvements seen in communicable, and other non-communicable diseases (Martinez et al., 2020; Murray et al., 2020). The reasons for the lack of progress in prevention and treatment are complex, but they include limited understanding of pathogenesis, pathophysiology and of the mode of action of existing treatments and why they are ineffective for many people. This is compounded by the inadequate provision of mental healthcare. Only 20% of individuals with common psychiatric disorders, such as depression and anxiety, have access to appropriate treatment in Europe and North America, and this “treatment gap” is likely to be even wider in low- and middle-income countries (Vigo et al., 2019).

Even among those who can access treatment, fewer than 50% will receive an “effective” therapy, broadly defined as one leading to a positive response with sustained improvement in therapeutic outcomes (De Silva et al., 2014; Wong et al., 2010). Established treatments for particular major mental health disorders also show broadly equivalent levels of effectiveness when assessed in large cohorts (Cipriani et al., 2018; Lieberman et al., 2005), and thus treatment decisions often centre around preferences based on generic adverse reaction profiles without knowledge of the patient’s actual risks (National Collaborating Centre for Mental Health, 2014). Adverse drug reactions and lack of effectiveness are common reasons for psychiatric medication being discontinued (Legge et al., 2016), with only a minority of patients remaining on prescribed treatments for the full therapeutic course (Jones et al., 2006; Lieberman et al., 2005). These issues are exacerbated by the requirement for long-term treatment of many patients (Demyttenaere, 2019), and together these factors result in a large
fraction of those diagnosed with major psychiatric disorders failing to benefit from current standards of care (Figure 1A). While this is a criticism levelled at psychopharmacological treatments (Muscatello et al., 2020), many of these same issues also apply to the evidence-based psychological treatment approaches used in psychiatry (Holmes et al., 2018).

The prescription of drugs that are ineffective or cause harmful side effects has substantial implications for individuals and the resources of health systems (Fava et al., 2017; Muscatello et al., 2020). In part, this is a reflection of the “efficacy-effectiveness gap”, the disconnect between the results of randomised controlled trials (“efficacy”) of interventions and their real-world performance (“effectiveness”; Eichler et al., 2011), due in large part to the limited degree to which trial participants and processes reflect clinical practise. This phenomenon is apparent in all areas of medicine, indeed meta-reviews have highlighted that drugs routinely used in psychiatry have equivalent efficacy to those used in other specialities (Huhn et al., 2014; Leucht et al., 2012). These analyses also found large individual variability in response, including frequent remission under placebo, which creates an upper bound to the efficacy of treatments when measured as means across groups of patients.

Many factors have been associated with individual differences in treatment response in psychiatric disorders (Perna et al., 2020; Stern et al., 2018) including demographics and lifestyle traits (sex, age, ancestry, body mass index, smoking habits, socioeconomic status), symptom profiles (frequency, severity and stage of illness, age at onset, comorbidities) and treatment provision (primary or secondary care, treatment adherence monitoring, concomitant prescription of other drugs or psychotherapy). Recent research has explored potential explanations behind some of these associations, such as physiological differences between sexes (LeGates et al., 2019), and variation in markers of neurobiological function affecting response to particular medications (McCutcheon et al., 2019). More specifically, genetic variation makes important contributions to variability in treatment response, as well as to the occurrence of adverse effects of medication. This raises the possibility of applying the many advances in human genomics to understand and predict individual variability in drug response, which is the aim of pharmacogenomics research.

**PHARMACOLOGICAL PROCESSES AND GENOMIC VARIATION**

The relationship between the dose of a drug and its effect, broadly speaking, depends upon the combined action of two sets of processes, pharmacokinetics and pharmacodynamics (Hefner et al., 2013). Pharmacokinetics (“what the body does to a drug”) refers to the set of biotransformations that a drug undergoes in the body, through absorption, distribution, metabolism, and excretion (ADME) processes. Whereas, pharmacodynamics (“what the drug does to the body”) refers to the physiological and biological responses caused by that drug, mostly reflecting ligand-receptor occupancy and downstream effects. Pharmacogenomics seeks to identify the genetic basis of these processes, often with research targeted to specific drugs or drug-gene pairs (Krebs and Milani, 2019; Roden et al., 2019). This discipline was pioneered by the work of Friedrich Vogel, Arno Motulsky and Werner Kalow in the late 1950s and early 1960s, with important tenets, such as the tranethnic variability of drug metabolising enzymes, becoming fully established by the 1980s (Müller and Rizhanovsky, 2020).

Early pharmacogenomics research was based upon the “candidate gene” analysis framework consolidated in the 1990s, which rapidly generated many drug-gene associations (Evans and
Johnson, 2001). Several of these advanced into Phase 1 clinical trials, sparking the interest of a pharmaceutical industry that saw pharmacogenomics as a potential solution to the “innovation deficit” of traditional drug development pipelines (Norton, 2001). However, as in other areas of genomic research, there were difficulties in replicating the results of candidate gene studies, a consequence of their limitations in establishing causal variants and controlling for genomic confounders (Peters et al., 2010; Walgren et al., 2005). Thus, few of the initial discoveries fulfilled the criteria necessary to demonstrate clinical utility, prompting the creation of international consortia to explicitly assess the actionability and therapeutic potential of pharmacogenomic evidence, such as the Dutch Pharmacogenetics Working Group (DPWG; Swen et al., 2008) and the Clinical Pharmacogenomics Implementation Consortium (CPIC; Relling and Klein, 2011). Despite criticisms of the field at that time (Ioannidis, 2013), the work of these consortia, together with the move towards genome-wide association studies (GWAS) as standard hypothesis-generating experimental designs, contributed to the consolidation of pharmacogenomics as one of the pillars of the “precision medicine” model of healthcare in the last decade (Hicks et al., 2019).

Currently, more than 1,000 basic research studies in pharmacogenomics are published every year (Müller and Rizhanovsky, 2020), and 165 guidelines on the use of pharmacogenomic information for specific drug-gene pairs are available in the PharmGKB repository as of May 2021, including 33 relevant for psychiatry (Bousman et al., 2021). Recommendations in these guidelines are derived from a thorough process of literature curation and standardised variant annotation (Barbarino et al., 2018), partially aligned with guideline standards from the US National Academy of Medicine (Caudle et al., 2014). They are increasingly recognised by industry and regulators in drug development and labelling processes (Schuck and Grillo, 2016; Yin and Vandell, 2019). PharmGKB guidelines also assess the quality of supporting evidence and, when indicated, make genotype-based prescribing recommendations (Relling et al., 2020).

However, these guidelines have not been widely adopted by clinical services, including those providing psychiatric care, as in most cases the trials needed to demonstrate their clinical utility to the standards required by public healthcare settings have yet to be carried out (Caudle et al., 2014; Russell et al., 2021). Additionally, important financial and regulatory constraints are still being explored, particularly those regarding the circumstances in which to implement genetic testing and who should cover its cost (Mitropoulou et al., 2020).

**GENOMICS OF DRUG METABOLISM**

The majority of the established drug-gene pairs and associated dosing guidelines relate to pharmacokinetic effects (particularly drug metabolism). Due to the difficulty of directly studying molecular interactions *in vivo*, pharmacological studies of large cohorts typically rely on measuring markers of drug metabolism and response. Treatment outcomes are sometimes used as proxies, particularly if the aim of the study concerns specific treatment effects (Grady and Ritchie, 2011; Roden et al., 2019), but the plasma concentration of an active drug metabolite offers advantages over other proxy measures since it’s a relatively close marker to biochemical pathways and a potential correlate of drug efficacy (Hefner et al., 2013). Metabolite concentrations can be also be analysed as precise quantitative phenotypes, and thus are well-suited to the statistical methods optimised for genetics and genomics studies (Grady and Ritchie, 2011; Suhre et al., 2011). This perhaps underlies the productivity of this line of research, which is responsible for most of the current drug-gene associations with potential
clinical applicability in psychiatric settings (Table 1; Bousman et al., 2021) and in general (Tornio and Backman, 2018).

The genes robustly identified by pharmacogenomics research, in both psychiatry and general medicine, include about 40% of the 266 human protein-coding genes that are known to be implicated in ADME processes (Tilleman et al., 2019), though this definition can be expanded to around 800 if known drug targets are also considered (Schärfe et al., 2017). ADME genes appear consistently in candidate gene studies of pharmacogenomic traits, although it is reassuring that they also often emerge through data-driven approaches, such as GWAS. Indeed, genome-wide assays are often enriched for these “usual suspects”, and signals can be as narrow as to implicate individual alleles or SNPs within these genes (Loukola et al., 2015; Pardiñas et al., 2019). These pharmacogenomic studies also benefit from decades of in vitro and model organism studies, which have already clarified the role of many of the causal genes and pathways in ADME phenotypes (Neavin et al., 2016). They have also allowed the definition, for a restricted set of drug metabolising enzymes, of standardised metrics of protein function (“activity scores”). Such scores allow individuals to be classified into the rapid-poor metaboliser spectrum for specific drugs (Caudle et al., 2017) and have formed the basis of many pharmacogenetic prescribing guidelines including those for psychiatry (Table 1).

The fact that a small number of genes have been robustly implicated from analyses of relatively small samples in pharmacogenomic studies sits in contrast to large-scale genomic studies of complex traits and disorders which identify an ever-expanding number of associations reflecting polygenic genetic architectures (Visscher et al., 2017). This can be explained by the underlying relationship between genetic variation and pharmacokinetics, particularly drug metabolism. In general, the genomic studies of endogenous and xenobiotic metabolism have revealed fewer genetic determinants than equivalent studies in complex traits, with most metabolites showing oligogenic architectures characterised by a few common variants of large effect size (Timpson et al., 2018). Heritability estimates, however, are similarly large (Lauschke and Ingelman-Sundberg, 2019), which raises the possibility that genetic predictors of drug metabolism could be created using relatively small sets of genomic markers as recently shown possible by research on other oligogenic traits (Zhang et al., 2020).

The apparent simplicity of the genomic architecture of these pharmacogenomic traits is complicated by challenges in fully characterising the genomic variation within these genes. Several important ADME genes, of which the best known is probably CYP2D6, harbour structural variation that is not well-captured by standard genotyping protocols (Lauschke et al., 2017). As a consequence of this, insights and results from much of the early literature on these genes have to be cautiously interpreted (Yang et al., 2017), which compromises applications and trials in clinical settings (Cavallari et al., 2019). It should be noted that complex genomic regions within ADME genes are not uncommon, and originated from the evolution of gene clusters through duplication and homologous recombination (Meech et al., 2012; Thomas, 2007), favouring a genomic environment with relaxed natural selection where novel mutations tended to drift (Hovelson et al., 2017). Thus, despite successes in identifying potential causal variants from the analysis of metabolite concentrations and related metrics, pharmacogenomic traits are likely to still have substantial unexplained (“hidden” or “missing”) heritability, which might require more complete sequencing and genotype-environment studies to be resolved (López-Cortegano and Caballero, 2019).
PHARMACOGENOMICS OF ADVERSE DRUG REACTIONS

Responses to drugs can include unwanted side effects. The preferred term for these is “adverse drug reactions” (ADRs) when they can be confidently assigned to a specific medication. ADRs are major causes of morbidity and mortality, and genetic variation makes important contributions to the risk of developing them (Carr and Pirmohamed, 2017). Many of the drug-gene pairings within PharmGKB resources highlight ADRs that are dependent on genotype (Relling et al., 2020), and these include treatments for psychiatric disorders.

In their simplest form, ADRs can be classified as “Type A” (pharmacological) or “Type B” (idiosyncratic). Pharmacological reactions arise from an adverse response to the known mechanism of action of a drug, occur in a dose-dependent fashion, and can be understood and potentially predicted from the drug’s known targets. Thus, genetic variation contributing to pharmacokinetic mechanisms can be relevant for these ADRs, and there are already several examples of highly penetrant risk alleles within genes encoding drug-metabolizing enzymes, such as CYP2C9 and bleeding on warfarin treatment, or CYP2D6 and opiate-induced respiratory depression (Carr and Pirmohamed, 2017). On the other hand, idiosyncratic ADRs are not predictable from the known pharmacological profile of the drug, are rarer (20% of all ADRs), but can be life-threatening and cause severe organ damage. Immunological processes have been implicated in many of these ADRs and the majority of guidelines relate them to polymorphisms within immune response genes, particularly the HLA system (Manson et al., 2020).

ADRs have a significant impact on health systems given their frequency and potential severity; 6–7% of hospital admissions in the UK have been attributed to ADRs and 19% of hospital inpatients experience significant ADRs (Pirmohamed et al., 2004), a rate that is likely higher in children and older people (Laatikainen et al., 2017; Sutherland et al., 2019). Their relevance in psychiatry is related to the fact that polypharmacy, the prescription of multiple medications, is common in psychiatric care and can lead to increased rates of ADRs through drug-drug interactions (Hefner et al., 2020). In addition, many psychiatric medications are so-called “dirty drugs”, having actions at multiple receptor targets with associated broad adverse effect profiles (Agid et al., 2007; Caraci et al., 2017). This is a counterpart to the classic idea of “magic bullet” drug development, whereby single-target drugs would improve treatment by minimising adverse effects. This approach has not yet led to advances in psychiatric treatment (Roth et al., 2004), which raises the question of whether the multi-target action of many medications may be central to their effectiveness (Ramsay et al., 2018).

These observations highlight the potential for developing prediction algorithms for ADRs to benefit patient safety and resource utilisation. Medications with pharmacogenomic indications, in which genomic data could be used to prevent or minimise ADRs, are slightly overrepresented among those prescribed in routine care (Barbarino et al., 2018), and it has been estimated that at least 65% of primary care patients are exposed to them over a 5-year period (Kimpton et al., 2019). The utility of this information is now being examined at scale, with the prospect of standardised pharmacogenomic variation being stored on routine health records and incorporated to alert and monitoring systems to guide prescribing decisions (Denny and Collins, 2021; Erika et al., 2017; Relling et al., 2020).

RISE, FALL AND PLATEAU OF PSYCHIATRIC PHARMACOGENOMICS

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There is a long history of pharmacogenomic studies of medications for mental health conditions, and the specific drug-gene pairs that have been identified from this work have been comprehensively reviewed elsewhere (Cacabelos, 2020; Lam and Someya, 2019). Early findings established effects of genetic variation in ADME enzymes on psychotropic medications (Coutts et al., 1999), prompting a large body of subsequent work which, despite its potential, has largely failed to be translated to clinical impact. In 2008, the field seemed to be “at the forefront of the use of pharmacogenomics in medicine” together with oncology, and five genetic tests were being marketed in the USA to support psychiatric treatment management (de Leon, 2009). However, these tests were soon discontinued or became outdated due to uncertain clinical utility and low consumer uptake, and have not been adopted by healthcare providers (de Leon, 2016). This experience reflects the paucity of research to support the clinical use of pharmacogenetic testing and reinforces the need to further develop the evidence base for the practical relevance of pharmacogenomic testing in common psychiatric conditions, including major depression and schizophrenia, and the limitations of commercial tests in providing the levels of reliable guidance demanded by clinicians, patients and regulators (Rakesh et al., 2020).

A major barrier to the implementation of pharmacogenomic testing in psychiatric care has been the lack of primary data of therapeutic utility for the vast majority of drug-gene pairs (Bousman et al., 2021). This is a prerequisite for the commissioning of new tests or interventions in many healthcare settings, as is the need to demonstrate health economic benefits. Advances are being made in this area with the advent of psychiatric clinical trials assessing the impact of testing for specific drug-gene pairs on psychiatric outcomes, although to date these have failed to demonstrate major therapeutic advantages. For example, there have been substantial efforts to assess the benefits of genomic testing for the treatment of major depression (Bousman et al., 2019), a necessary first step towards the clinical adoption of these procedures (Figure 1B). The largest trial of this kind, GUIDED, did not show a beneficial effect of pharmacogenomic guided prescribing for its primary outcome of symptom improvement against antidepressant standard-care (Greden et al., 2019). Results on treatment response and remission outcomes were more positive, although these were secondary analyses that have been questioned on conceptual and statistical grounds (Goldberg, 2019; Smith and Nemeroff, 2020). This lack of supportive evidence, coupled with the negative results of other studies (Perlis et al., 2020; Zeier et al., 2018), underlies the lack of support from the FDA, CPIC or DPWG for all current claims of genetic markers as potential predictors of antidepressant response or indeed any other pharmacodynamic outcome (Bousman et al., 2021). Nonetheless, all these organisations currently recognise that pharmacogenomic variation could be of relevance for antidepressant therapeutic management mainly due to its effects on pharmacokinetics (Bousman et al., 2021). This avenue, explored but not found significant in GUIDED, has however shown some promising results in recent prospective trials conducted within small-scale clinical settings (Papastergiou et al., 2021).

In terms of healthcare impact, the most effective pharmacogenomic intervention in psychiatry is currently the assessment of rare HLA-B alleles for prevention of Stevens-Johnson syndrome and toxic epidermal necrolysis caused by carbamazepine, an anticonvulsant now most commonly used in psychiatry as a second-line treatment for bipolar disorder (Goodwin, 2003). With 77.4% sensitivity and 94.4% specificity in Han Chinese individuals, prescription changes
and screening protocols based on this finding have substantially reduced the incidence of this ADR across East Asia (Nguyen et al., 2019a; Pan et al., 2017). Similar procedures, again for carbamazepine, have been implemented outside of East Asia for genotyping HLA-A alleles to prevent hypersensitivity reactions (Phillips et al., 2018). Reflecting this evidence, in 2020 the FDA included carbamazepine and its structural analogue oxcarbazepine in its official review table of pharmacogenomic associations. To date, these are the only psychiatric drug-gene pairs labelled with explicit recommendations of pharmacovigilance informed, though not substituted, by preventative genotyping (U.S. Food and Drug Administration, 2021).

Whilst no other findings in psychiatric pharmacogenomics have achieved this level of clinical impact, cautious support is emerging for the potential utility of testing more widely. The International Society of Psychiatric Genetics (ISPG; McMahon et al., 2019) has recognised the potential benefits of the CPIC/DPWG guidelines in the prescription of certain antidepressants and antipsychotics (Table 1), a stance supported by a recent systematic review of CYP2C19 and CYP2D6 enzyme activity for these classes of medication (Milosavljević et al., 2020).

Although these two genes are implicated to a degree in the metabolism of 80% of psychiatric medications and are increasingly recognised in regulator-approved drug labels (Müller et al., 2013; van Schaik et al., 2020), it is worth noting that many psychiatric drugs do not yet have replicated pharmacogenomic associations with these genes or indeed any other ADME gene despite the large body of basic research on this topic (Kose and Cetin, 2018). Additionally, even in well-established associations between genetic markers and drug metabolism, downstream effects on drug response or ADRs need to be explicitly evaluated, not just inferred. Difficulties in both identifying primary associations and carrying out therapeutic evaluations are related to the broad challenges of psychiatric pharmacogenomic research that we outline below, particularly around underpowered “candidate gene” designs (Kose and Cetin, 2018; Kranzler et al., 2017). These challenges are reinforced by the lack of funding for large longitudinal studies that have proven beneficial in other areas of medicine (Pirmohamed et al., 2013; The SEARCH Collaborative Group, 2008). Consequently, recent comparisons between psychiatry and general medicine, particularly oncology, have found the latter to have a clear lead in terms of immediate healthcare impact, with psychiatric pharmacogenomics remaining a promise with unrealised potential (de Leon, 2016; Kranzler et al., 2017).

In summary, while most psychiatric care is supported by the prescription of medications, psychiatric drugs are not currently effective for large numbers of those living with mental health disorders, and treatment changes and discontinuation are commonplace. A genetic basis for individual differences in both therapeutic and adverse drug responses is well established, but has not yet been translated into real-world therapeutic improvements in psychiatry. The hope that this translation is possible comes from progress in other areas of medicine, the fact that psychiatric drugs are not less efficacious than non-psychiatric medications, and the repeated successes of genomic approaches in finding robust and replicable associations for drug metabolism and ADRs. It is thus worth exploring the reasons behind this slow advance, and whether specific improvements can be made to accelerate the pace and robustness of discoveries in psychiatric pharmacogenomics. This research might improve psychiatric drug efficacy by contributing to better drug design, but it is perhaps more likely to deliver its benefits through the offering of tailored treatments within the “precision medicine” paradigm. Here, it will be important to demonstrate valid stratification of groups of patients in terms of their...
response to a specific treatment (Figure 1B). However, treatment response can be a particularly hard concept to define in psychiatry, a limitation that has hampered research progress.

**MEASURING TREATMENT OUTCOMES AND RESPONSE**

Arguably the biggest challenge for psychiatric pharmacogenomics is the lack of widespread use of consistent, operationalised definitions of response to treatment. There are no diagnostic biomarkers in psychiatry, which forces diagnostic procedures to rely upon clinician assessment of subjective and observed clinical symptoms and their impact on patients’ functioning. Similarly, “treatment response” is a multifaceted concept that can incorporate many features in research settings, including changes in symptoms, assessment of functional level or metrics of health service use such as admission to hospital (Sajatovic et al., 2010; Salagre et al., 2018). In the absence of surrogate biomarkers to track drug response, the methods to measure these diverse outcomes become critical to advancing pharmacogenomic research.

In many areas of medicine the measurement of whether an individual benefits from a drug is conceptualised as a surrogate of survival or other “ultimate” treatment endpoints (Woodcock, 2010). However, these are generally less applicable in psychiatry as mortality is rarely a short term outcome and is not a primary part of pathophysiology (Leucht et al., 2012). The most popular methods to measure psychiatric drug response rely on defined changes in scores of pre- and post-treatment symptom rating scales (Leucht et al., 2009). While an in-depth review of their psychometric properties is outside the scope of this article, it is worth noting that most rating scales exchange sensitivity in favour of internal consistency and applicability in diverse settings (Fava et al., 2017). Additionally, when used on their own, symptom scales largely ignore the broad impacts psychiatric treatment can have on psychological and social function, as well as quality of life (Demyttenaere, 2019).

Valid assessment of treatment outcome requires longitudinal measurement of these multifaceted elements of symptom and functional level. This requirement has been recognised in several consensus outcome definitions in psychosis (Andreasen et al., 2005; Howes et al., 2017), with equivalent initiatives also developing in depression research (Gaynes et al., 2020). Nonetheless, whilst these measures have been adopted in some clinical trials and observational research, they have not been widely used in pharmacogenomic studies.

The majority of recommendations for the assessment of treatment response have relied on data from the prospective measurement of symptoms and functioning. Other initiatives have defined therapeutic outcomes by applying retrospective assessment of clinical records, notably for lithium treatment in bipolar disorder (Grof et al., 2002), and this approach has shown its utility for observational pharmacogenomic research (Hou et al., 2016). As an important note, ADRs are not always considered as part of drug response criteria (Jordan et al., 2018) despite the fact that they are often primary causes of treatment discontinuation (Lahijani and Harris, 2017; Legge et al., 2016). The standardisation of phenotypic definitions of ADRs has benefitted other areas of pharmacogenomic research by aiding replication and supporting accurate meta-analyses and psychiatry could benefit from similar initiatives (Carr and Pirmohamed, 2017). An alternative approach used in pragmatic trials has been reliance on all-cause treatment discontinuation as a primary outcome measure of medication response, given its ability to capture both the efficacy and tolerability of medications (Kahn et al., 2008; Lieberman et al., 2005).
A final crucial aspect in defining response is the assessment of treatment adherence. Whilst adherence can be assessed through patient self-reports or information on health records, for example prescription refills (Rossom et al., 2016; Sajatovic et al., 2010), the gold standard remains the measurement of serum or plasma levels of the drug. Many drug response analyses are not conducted in parallel with therapeutic drug monitoring (TDM) or serum level measurements, and thus lack a biochemical perspective that permits insights into drug metabolism, as well as confirmation of adherence (Hiemke, 2016). The integration of this information into consensus measures has already been recommended by several reviews on major psychiatric disorders (Fornaro et al., 2018; Gadad et al., 2018; Lally and Gaughran, 2019). Furthermore, the inclusion of ADRs and drug metabolism into treatment response measurements would result in more robust and generalisable evidence in psychiatric treatment research by following the triangulation principle that has been widely successful in epidemiological studies (Lawlor et al., 2017).

In summary, in order to advance pharmacogenomics research, operationalised definitions of treatment response are required that incorporate measures of symptomatic and functional change, ADRs and adherence, ideally through measurement of serum levels. This would allow the dissection of treatment response into its pharmacokinetic and pharmacodynamic elements which are currently conflated in much treatment response research. Such developments would help advance the investigation of the complex biological and clinical landscape underlying drug effectiveness (Nordon et al., 2016), with downstream benefits for genomic and epidemiological research seeking to elucidate causal mechanisms.

THE (A)TYPICAL STORY OF CLOZAPINE

The results of extensive research on the second-generation (or “atypical”) antipsychotic clozapine illustrate many of the challenges and opportunities to advance psychiatric pharmacogenomics research. Clozapine is the only evidence-based medication for treatment-resistant schizophrenia, a condition affecting up to a third of those diagnosed with this disorder (Flanagan et al., 2020; Siskind et al., 2021). It is efficacious in around 40% of treated individuals, with greater effectiveness if administered early in the course of the disorder, but it is widely underutilised due to the perceived complexity of its prescribing (Remington et al., 2016), including a gradual titration period which requires careful physical monitoring (Stanworth et al., 2012). Whilst there can be multiple dose-related ADRs, the most prominent adverse effect is a rare blood dyscrasia which can escalate to agranulocytosis, a frequently fatal loss of white blood cells (De Berardis et al., 2018). The unpredictability of this idiosyncratic reaction, which affects less than 1% of treated individuals, prompted the establishment of recurrent blood monitoring services, which have to be used by all those prescribed clozapine (Nielsen et al., 2016). Among several other ADRs, weight gain is common and frequently severe (Flanagan et al., 2020), and sedation constitutes the main reason for voluntary treatment discontinuation together with lack of monitoring compliance (Legge et al., 2016). The search for pharmacogenomic markers to facilitate the management of clozapine-treated individuals and lighten the burden of routine blood monitoring has been an explicit research goal for many years (Arranz et al., 2000; Legge and Walters, 2019; Lieberman et al., 1990), although the relationships between treatment efficacy, side effects and clozapine pharmacokinetics are still unclear (Mauri et al., 2018).
Clozapine undergoes first-pass metabolism in the liver in which up to 90% of the drug is converted to norclozapine (N-desmethylclozapine). This reaction is mediated primarily by the CYP1A2 enzyme, though CYP3A4 and CYP2C19 can also act as catalysts. The main design chosen by clozapine pharmacogenomic studies follows this pathway (Figure 2), tracking the plasma concentrations of clozapine and norclozapine in treated individuals, and genotyping or sequencing the exonic segments of the genes coding the aforementioned enzymes and perhaps a few others. Results of these “candidate” studies have accumulated during the past two decades but have led to inconsistent findings (Li et al., 2018), potentially contributing to erroneous pharmacogenomic guidance (Rahman et al., 2017) and have been unsupported by recent GWAS in larger samples (Pardiñas et al., 2019; Smith et al., 2020).

Hampering advances in this field is uncertainty around the biological mechanisms responsible for the efficacy and adverse effect profile of clozapine, and there is still debate about whether norclozapine is itself of any pharmacological relevance (Hellman et al., 2016; Nucifora et al., 2017). Additionally, the first-pass metabolism of clozapine leads to a secondary compound, clozapine N-oxide, that has rarely been studied in this line of research. This is likely because its plasma concentrations are not often assayed for monitoring treatment adherence, which is the main clinical use of clozapine and norclozapine TDM (Flanagan et al., 2020). Nevertheless, high clozapine doses can promote the formation of clozapine N-oxide, which is capable of interacting with a broad array of neurotransmitter receptors (Diaz et al., 2014). Furthermore, all clozapine metabolites can cross the blood-brain barrier through a yet-unknown transporter (Dickens et al., 2018; Hellman et al., 2016), participating in further biological pathways which are poorly understood. While these would be arguably the most relevant processes to study particularly in a genomic setting (McMillan and Tyndale, 2018), research is still scarce as directly measuring the amount of active clozapine in the in vivo brain is currently impractical outside of animal models, and predictive cheminformatics algorithms are still of limited application (Wong et al., 2019).

In summary, despite unparalleled effectiveness and after 30 years of intense research, clozapine remains a drug for which the active metabolite, site and mechanism of action remain unknown, demonstrating the challenges of moving from informative pharmacogenomic research studies to achieving actionable or clinically translatable evidence (Cremers et al., 2012). While by terminology it is an “atypical” antipsychotic, these knowledge gaps and translational challenges are typical of many other psychiatric drugs, limiting the advance of therapeutic developments (Alavijeh et al., 2005; Kose and Cetin, 2018). Nonetheless, in the last decade, there have been significant advances in our knowledge of clozapine’s metabolism and the origin of its adverse effect profile through the international application of unbiased genome-wide methods to routinely collected samples with clear phenotypic definitions (Goldstein et al., 2014; Legge et al., 2017; Pardiñas et al., 2019; Smith et al., 2020). For these insights and derived findings to advance research into the pharmacogenomics of treatment response, they will have to be explored further at scale. As obtaining high-quality data from large samples using traditional methods (i.e. personal follow-up) can be very costly and time-consuming, developing and consolidating population-scale phenotyping methods is an avenue with immense potential to advance psychiatric pharmacogenomics research (Smoller, 2018).

RESEARCH WIDELY, IMPLEMENT LOCALLY
Advances in health informatics have made possible the routine collection, storage and analysis of electronic health records (eHRs) within healthcare systems, leading to this area of research becoming another pillar for the “precision medicine” approach (Denny and Collins, 2021). Datasets based on eHRs offer the potential for research at the level of the population, promising representativity and real-world relevance beyond the ascertainment and inclusion biases that have hindered clinical research to date. The integration of genomic data into eHRs offers many benefits but also challenges (Grebe et al., 2020), although pharmacogenomic alleles were some of the earliest data to be incorporated into routine records (Abul-Husn and Kenny, 2019). In addition to the potential utility of easily accessible pharmacogenomic variants to guide treatment decisions (Figure 1C), the longitudinal real-world phenotypes from eHRs could also become invaluable for pharmacogenomic research. As well as the availability of detailed prescribing information and therapeutic drug monitoring results, eHR data could help define treatment response, particularly in psychiatry where much of the relevant data is stored in free-text narrative notes (Aaslestad, 2016). While the analysis of clinical notes, and raw text in general, has traditionally been considered difficult, methodological advances in Natural Language Processing (NLP) and Artificial Intelligence give grounds for optimism and can feasibly unlock this data for clinical research purposes (Jackson et al., 2017). This would not, however, avoid the presence of confounders and biases, which could stem from patient, clinician or even healthcare system factors (Dueñas et al., 2020). Addressing these difficulties should take advantage of both the population-wide scale of clinical databases and the multidisciplinary nature of eHR-based research (Figure 3), as both open up opportunities for identifying these problems rapidly and early, as well as exploring solutions in which data curation and quality control algorithms are informed by clinical expertise (Rees et al., 2019).

Currently, several worldwide biobanks are working on incorporating information from eHRs into drug-specific pharmacogenomic studies or clinical trials (Hoffman et al., 2020), and recent results from Danish, British and Estonian Biobanks support previous observations that 90%-99% of the population has at least one clinically actionable pharmacogenomic allele (Lunenburg et al., 2021; McInnes et al., 2020; Reisberg et al., 2018). In psychiatry, the value of routinely collected phenotypes has been demonstrated by studies examining the antidepressant escitalopram (Jukić et al., 2018) and the antipsychotics aripiprazole and risperidone (Jukić et al., 2019), performed in thousands of records from a therapeutic drug monitoring service. This research used drug switching as one of its outcomes (referred to as “therapeutic failure”, equivalent to all-cause treatment discontinuation referred to above), and showed that individuals with certain CYP2C19 and CYP2D6 activity scores (ultrarapid and poor metabolisers) were more prone to a change of prescription early in the course of treatment. Importantly, this happened despite clinicians commonly altering doses during treatment in reaction to drug response, thus not being completely agnostic of metabolism and other underlying pharmacological processes. Similar studies conducted on child and adolescent cohorts replicated these results (Aldrich et al., 2019; Jallaq et al., 2020), suggesting that genetic variants determining enzyme activities could be beneficial in clinical decision-making even if non-genetic factors, such as the co-prescription of enzyme inhibitors, are already considered. While none of these experiments were explicitly designed for causal inference, complementary data of metabolite concentrations on those same individuals supported the hypothesis that drug switching was prompted by insufficient clinical response or the appearance of adverse effects. This triangulation and replication of evidence, based on a broad but consistently defined
treatment outcome, has now become part of pharmacogenomic guidelines for these medications (Bousman et al., 2021) and serve as exemplars for the field. While this is only the first step on the long road to eventual healthcare application, it demonstrates the willingness of the psychiatric pharmacogenomics community to embrace new findings based on robust research.

Clinical trials, particularly randomised controlled trials (RCTs), are the gold standard experimental designs to evaluate the robustness of research findings and, ultimately, the utility of interventions derived from them (Lawlor et al., 2017). The design of pharmacogenomic RCTs is not straightforward and financial incentives to support them are not always apparent (Russell et al., 2021), which has likely limited the scale of these studies in psychiatry. In fact, most of those carried out to date have been industry-sponsored, focusing on antidepressant use and following the more numerous PharmGKB guidelines for these medications (Table 1). A recent synthesis of this RCT evidence, including the GUIDED trial discussed before, showed that individuals who receive genotype-guided drug dosing are 1.71 times more likely to achieve remission than those on standard care (Bousman et al., 2019), although the context of these findings needs to be considered in light of frequent multiple secondary outcomes and post-hoc analyses undertaken as part of the primary research studies. Furthermore it is worth noting that several of these trials recruited only participants with prior inadequate treatment response, although there is little reason to think that the benefits of pharmacogenomics will be restricted to this patient group. Basic research to demonstrate this wider utility could explore, for example, the stratification of patient cohorts by treatment response or outcome (Figure 3), which could result in further insights regarding the genetic basis of these traits (Fabbri et al., 2021).

The declaration of a drug-gene association as therapeutically useful, even within regulator-approved product labelling, does not ensure its adoption into routine clinical work (van Schaik et al., 2020). Importantly, pharmacogenomic guidelines do not advise on which therapeutic situations, if any, should prompt a clinician to order a genetic test, instead working as “decision-support tools” when testing is already available in a particular healthcare setting (McMahon et al., 2019). This is still uncommon and can result in substantial public and private costs and implementation challenges, which at the moment are being explored in health systems across the world (Krebs and Milani, 2019). The application of these tests and the need for patient support in the form of counselling, as well as other resource implications, are yet to be fully assessed within psychiatry (Moldovan et al., 2019). Recent surveys show that psychiatrists are generally accepting of incorporating pharmacogenomic information in their routine clinical work, though they also recognise that their training is likely inadequate to take on the added responsibilities that this would entail (Ward et al., 2019). Public perceptions, on the other hand, seem to be more cautious and mostly interested in the effects test results could have on clinical decisions (Liko et al., 2020). Responding to these views requires psychiatric pharmacogenomics to be implemented in a multidisciplinary way, involving psychiatrists, genetic counsellors, pharmacists, and patients, with a strong emphasis on outreach and educational practices (Hicks et al., 2019). The experience of some clinics applying these principles shows that local pharmacogenomic support services can sustainably contribute to psychiatric community healthcare (Arwood et al., 2020; Dunnenberger et al., 2016), effectively reducing the amount of trial-and-error needed to find efficacious medications in a sizeable
portion of individuals. Similar outcomes have been obtained, though not specifically in psychiatry, by integrating pre-emptive genetic testing services into existing clinical pharmacy frameworks (Thornley et al., 2021). Thus, it becomes plausible that the development and adoption of these specialist services could act as the gateway through which pharmacogenomics knowledge becomes integrated into the psychiatric standard of care (Figure 1A, Figure 1B), revealing a road from “precision medicine” into “precision psychiatry”.

PHARMACOGENOMIC PHENOTYPES: SIMPLE TRAIT GENOMICS?

In basic science settings, the next stage of pharmacogenomics research is likely to be focussed on ascertaining rare and structural genomic variation primarily within key ADME proteins (Figure 2), and ideally in diverse worldwide populations to fully characterise pharmacogenomic variation across ancestries (Lauschke and Ingelman-Sundberg, 2019; Schwarz et al., 2019). However, rare variant associations can require novel effect annotations, complex sequencing protocols and larger sample sizes than GWAS to achieve strong statistical support, which also complicates the evaluation of their impact in RCT settings and for clinical implementation (van Schaik et al., 2020). Nonetheless, as we have outlined, we believe that confronting these challenges will help demonstrate biologically-valid strata within many of the complex, heterogeneous psychiatric disorders, and thus invigorate mechanistic research to identify the causal basis of treatment response. Whilst this may appear more tractable for pharmacogenomic phenotypes, given their relatively simpler genetic architectures, the key hurdles of human genomics research remain particularly in moving from genetic association to biological understanding, supporting the need for functional genomic and biological experimental follow up (Figure 3). A recent cautionary example from the analysis of a diabetes biomarker shows how a putatively causal variant in a membrane transporter ended up being a confounded signal from an unrelated population-specific blood trait (Chai et al., 2020). Without similar and careful evaluations, genomic studies are vulnerable to attempting to derive causal mechanisms from incomplete information on gene function, tempted by the “narrative potential” of the human genome (Goldstein et al., 2013). Lack of mechanistic follow-up on pharmacogenomic studies could be partly behind the current heterogeneity seen on genetic testing panels, which use a variety of SNPs and sequence variants to call the same alleles or enzyme activity scores (Tillemann et al., 2019). This is an issue in both clinical and research settings, as it further complicates the accurate determination of metaboliser status in some particularly complex genes, such as CYP2D6, risking inaccurate results if panels lacking the appropriate coverage of structural variants are used (Jarvis et al., 2019). Finally, resolving genomic associations into single or sets of putatively causal markers would allow for using statistical approaches such as Mendelian Randomisation (MR) to facilitate the design of downstream biological experiments and clarify the relationships between different metabolic traits (Tin and Köttgen, 2020). If enough potentially causal variants are available, this latter use could help in the identification of potential drug-drug interactions at the pharmacokinetic level (Pardiñas et al., 2019), which are an important cause of ADRs in psychiatric pharmacotherapy (Hefner et al., 2020).

Though these considerations pertain to pharmacokinetic phenotypes, they do not necessarily apply to the analysis of pharmacodynamic outcomes, including drug response, which may well have more complex, polygenic genetic architectures given their multifactorial aetiology.
(Roden et al., 2019; Rohde and Kristensen, 2020). Robust pharmacogenomic analyses of response to psychiatric drugs are pressingly needed, as no replicated pharmacodynamic associations have yet been found (Bousman et al., 2021). Indeed it has been recently argued that the focus of new genomic studies should be disease outcomes, rather than the presence of disease as traditionally assayed with case-control experimental designs, if such studies are to deliver the goal of target identification and improved therapeutics (Paternoster et al., 2017).

The field of depression research has progressed this idea by extensively exploring mechanisms behind associations between genes in the hypothalamus-pituitary-adrenocortical system and treatment response (Ising et al., 2019; O’Connell et al., 2018). More widely, a retrospective study has shown that pharmacodynamic variants on genes encoding drug targets are predictive of ADRs affecting specific organ systems (Nguyen et al., 2019b). There is also preliminary evidence that rare genetic variation could be associated with differential drug response, supporting the use of mutation burden analyses to highlight pharmacodynamic genes (Wolking et al., 2020). This approach is increasingly tractable with the advent of whole-exome and whole-genome sequencing but again requires well-characterised samples stratified for response and non-response. Another interesting avenue is that drug metabolism, rightly considered a strong mediator of drug response, could also act as a moderator in certain cases. This would manifest as the pharmacogenomic effect sizes of variants in the targets of a given drug depending on the activity of relevant metabolic enzymes (Baranova et al., 2017). Needless to say, very large samples would be needed to detect such dependencies with reliability, of at least four times the size needed to detect the main pharmacogenomic effects (Brookes et al., 2004).

A way of partially addressing this scenario in more limited conditions could be to include predictors of drug metabolism, such as phenoconversions, as part of drug response analyses, because the ignorance of these is known to hinder the discoverability of pharmacogenomic markers (Matthaei et al., 2016). Other predictors could also be genomic variants themselves or composite markers, inspired by the growing body of literature on genetic predisposition in psychiatry and the potential benefits for research of sample stratification in polygenic extremes (Murray et al., 2021). While these holistic experimental designs are less common than other approaches, they could be facilitated by the ongoing development of health informatics algorithms to exploit routinely-collected eHRs (Hoffman et al., 2020), and would follow recent recommendations for the inclusion of more individual- and drug-related characteristics in pharmacogenomic analyses (Thorn et al., 2019).

**PHARMACOCHEMGENOMICS NEEDS TO BE GLOBALLY FOCUSSED**

One issue that remains a pressing concern (Denny and Collins, 2021). is the realisation that since the allele frequencies of actionable pharmacogenomic variants often differ by ancestry, their clinical use needs to be implemented in such a way as to avoid reinforcing the structural inequalities that lead to treatment gaps and health disparities among ethnic groups in many countries (Martin et al., 2017; Olivier and Williams-Jones, 2014). Increasing awareness of this issue among drug industry regulators is encouraging (Tan-Koi et al., 2019), but the research community must contribute to this discussion and to rectifying the under-representation of those of non-European ancestries in pharmacogenomic studies, a disappointing constant across medical genetics research (Mills and Rahal, 2020). It is an indictment that only 5% of pharmacogenomics research publications explicitly state the ethnicity or ancestry of their participants (Popejoy, 2019). Without these data, the reliable assessment of population-specific
pharmacogenomic effects, known in principle for at least the last four decades, is impossible. Moreover, genetic markers with strong ancestral stratification, of uncertain utility in global populations when applied to predictive tasks, can mistakenly become incorporated in genetic testing panels marketed to large amounts of individuals (Manrai et al., 2016). Thus, improved consistency in the reporting of genetic ancestry and ethnic variability is required, ideally following standardised criteria (Huddart et al., 2019), a necessity if the full potential of pharmacogenomic interventions is to be realised across the world.

CONCLUSIONS

Genomics offers two main prospects for psychiatry. First, it provides a potentially unbiased route to understanding pathogenesis and pathophysiology. Second, it can potentially provide novel predictive tests for research and clinical practice (Owen and O’Donovan, 2020). These two elements come together in pharmacogenomics, which will be central to delivering precision medicine and within it, precision psychiatry. Pharmacogenomics offers a way to improve our understanding of the metabolism, safety and efficacy of psychiatric drugs, whilst potentially contributing to their development or repurposing (Schuck and Grillo, 2016; Yin and Vandell, 2019). It will also allow us to delineate groups of individuals who could benefit from particular therapeutic strategies (Rees and Owen, 2020). Psychiatric pharmacogenomic insights are likely closer to clinical implementation than those derived from wider psychiatric genomics, given the relative simplicity of the genomic architectures of many pharmacogenomic phenotypes (particularly those of ADME traits) and their manifest clinical applicability. If psychiatry can overcome the obstacles and roadblocks described in this review and seize the opportunities offered by new technologies (Figure 3), we can be optimistic that the rate at which these insights are generated will increase.

ACKNOWLEDGEMENTS

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DECLARATION OF INTERESTS

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Table 1: Summary of psychiatric pharmacogenomic drug-gene associations reflected in current PharmGKB guidelines, all with CPIC evidence codes “A” or “B” indicating a recommendation for genotype-guided prescribing. Original data from Bousman et al. (2021). Advice based on regulator-approved product labels is not reflected here as inconsistencies have been noted in some psychiatric drugs (van Schaik et al., 2020). Further details about PharmGKB recommendations and their adherence to evidence-based medical guideline standards are available in Caudle et al. (2014).

Note: “PM” - “Poor metabolisers”; “UM” - “Ultrarrapid metabolisers”

<table>
<thead>
<tr>
<th>Medication class</th>
<th>Drug(s)</th>
<th>Biological mechanisms</th>
<th>Gene</th>
<th>Main supporting evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antidepressants</strong></td>
<td>Amytriptyline, citalopram, clomipramine, doxepin, escitalopram, imipramine, sertraline, trimipramine,</td>
<td>Pharmacokinetics</td>
<td>CYP2C19</td>
<td>Treatment discontinuation more likely in PMs/UMs</td>
</tr>
<tr>
<td></td>
<td>Amytriptyline, clomipramine, desipramine, doxepin, fluvoxamine, imipramine, nortriptyline, paroxetine, trimipramine, venlafaxine</td>
<td>Pharmacokinetics</td>
<td>CYP2D6</td>
<td>Treatment discontinuation more likely in PMs/UMs</td>
</tr>
<tr>
<td><strong>Antipsychotics</strong></td>
<td>Aripiprazole, brexiprazole, haloperidol, pimozide, risperidone, zuclopenthixol</td>
<td>Pharmacokinetics</td>
<td>CYP2D6</td>
<td>Treatment discontinuation more likely in PMs</td>
</tr>
<tr>
<td><strong>Mood stabilisers</strong></td>
<td>Phenytoin</td>
<td>Pharmacokinetics</td>
<td>CYP2C9</td>
<td>ADRs more likely in PMs</td>
</tr>
<tr>
<td></td>
<td>Carbamazepine, oxcarbazepine, phenytoin</td>
<td>Inmunological</td>
<td>HLA-A, HLA-B</td>
<td>HLA-A<em>31:01 and HLA-B</em>15:02 likely causal for severe ADRs</td>
</tr>
<tr>
<td><strong>ADHD medications</strong></td>
<td>Atomoxetine</td>
<td>Pharmacokinetics</td>
<td>CYP2D6</td>
<td>ADRs more likely in PMs</td>
</tr>
</tbody>
</table>
**Figure 1:** Journey of 100 people through an initial diagnosis of Major Depressive Disorder under several current and potential healthcare models. (A) Standard genetics-agnostic care in high-income countries with extensive public health systems. (B) Near-future scenario in which the clinical implementation of psychiatric pharmacogenomics (PGx) testing has cleared all current barriers and became introduced before an eventual antidepressant prescription. (C) Future (post-2030, following Denny and Collins, 2021) scenario following the above development and in which information from electronic health records (eHR) systems is easily accessible before drug prescription and enriched with genetic data.

For reference, panel A contains the following assumptions: (i) 79% of those with a diagnosis of depression are initially prescribed pharmacotherapy (Treviño et al., 2017); (ii) at least 60% of people discontinue their first antidepressant within three months (Jung et al., 2016; Rossom et al., 2016); (iii) while the symptomatology of some individuals can improve or remit without treatment (Kato et al., 2021), 40% of people who adhere to antidepressant treatment do not obtain any therapeutic benefit (Rottenberg et al., 2018), and early discontinuation is associated with an 8-fold increase in this rate (Ho et al., 2016). Panel B follows the same proportions, additionally assuming that: (i) 62.4% of individuals in the population carry a PGx actionable variant for commonly prescribed antidepressants, as shown in blue (rate in European UK Biobank participants; Dr Gregory McInnes, personal communication); (ii) PGx testing is freely available to everyone, detects genotypes without error and clinicians always follow genotype-dosing guidelines when they are available. (iii) PGx-informed antidepressant treatment leads to a 1.7-fold improvement in therapeutic benefit (Bousman et al., 2019) and has a similar effect on treatment adherence (Jessel et al., 2020). Finally, for panel C we have optimistically considered that: (i) in eHR-driven healthcare, the therapeutic improvement of PGx-guided care increases to 2.5-fold and (ii) even standard prescriptions would be more effective (1.5-fold) than in current practice due to the use of routinely collected personal and exposure data in clinical decision-making (Denny and Collins, 2021).
**Figure 2:** Schematic depiction of a hepatocyte illustrating current knowledge about the clozapine metabolic pathway, which highlights proteins identified in the latest PharmGKB review (Thorn et al., 2018) and those encoded by ADME genes found in GWAS approaches (Legge et al., 2017; Pardiñas et al., 2019; Smith et al., 2020). Following the analysis of the ExAC database by Ingelman-Sundberg et al. (2018), boxes within each protein indicate the percentage of functional genic variation expected to be common (MAF≥1%, left box, blue) or rare (MAF<1%, right box, red).
Sample collection
Manual phenotyping of small samples
Strong ancestry/ethnicity bias in recruitment
Lack of funding for longitudinal studies

Psychiatric disorders
Multifactorial aetiology
Socioeconomic risk factors
Complex genetic architecture
Large variability in drug response

Study design
Inconsistent definitions of treatment response
Limited knowledge of drug effects in brain
Focus on “candidate” genes and variants

Results
Harder replication of studies
Uncertain generalisability
Slow progress and lack of impact

Sample collection
Semi-automatic phenotyping enabled by NLP
Population-scale routine electronic health records
Resources and opportunities to involve underrepresented populations and disorders

Study design
Operationalised drug response definitions
Pharmacological data for assessing outcomes, adverse reactions and treatment adherence
Genome/exome-wide analyses, inference of credible causal variants and biological follow-up

Results
Potentially straightforward replication
Real-world parallels favour designing RCTs
Faster progress from basic to translational research
Figure 3: Inherent challenges of psychiatric pharmacogenomics research (center, red), its usual roadblocks and limitations (left, silver) and needed solutions to advance into the precision psychiatry framework (right, gold). Coloured spheres illustrate the demographic, genetic, and socioeconomic diversity of the population of those affected by mental illness. Note that the study designs that we advocate for future psychiatric pharmacogenomics are not limited to assessing larger samples, but contain methodological innovations which would implicitly make these samples more diverse and representative.
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