Prioritising the application of genomic medicine

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The clinical translation of genomic sequencing is hampered by the limited information available to guide investment into those areas where genomics is well placed to deliver improved health and economic outcomes. To date, genomic medicine has achieved its greatest successes through applications to diseases that have a high genotype–phenotype correlation and high penetrance, with a near certainty that the individual will develop the condition in the presence of the genotype. It has been anticipated that genomics will play an important role in promoting population health by targeting at-risk individuals and reducing the incidence of highly prevalent, costly, complex diseases, with potential applications across screening, prevention, and treatment decisions. However, where primary or secondary prevention requires behavioural changes, there is currently very little evidence to support reduction in disease incidence. A better understanding of the relationship between genomic variation and complex diseases will be necessary before effective genomic risk identification and management of the risk of complex diseases in healthy individuals can be carried out in clinical practice. Our recommended approach is that priority for genomic testing should focus on diseases where there is strong genotype–phenotype correlation, high or certain penetrance, the effects of the disease are serious and near-term, there is the potential for prevention and/or treatment, and the net costs incurred are acceptable for the health gains achieved.

INTRODUCTION

Genomic sequencing is revolutionising research into human diversity and the causes of disease, resulting in improved diagnoses through the identification of molecular etiologies and the ability to manage many costly and disabling conditions. This is leading to increased investment by governments in genomic medicine, focussing initially on rare diseases and cancer, with the longer-term intent of incorporating genomic information into a new data ecology for biomedical discovery, more precise medical treatment, better stratification of patient populations and more efficient/cost-effective use of healthcare resources.

Currently genomic sequencing has achieved diagnostic rates for individually rare, but collectively common, monogenic diseases as high as in excess of 50%1 (with whole genome sequencing (WGS) typically producing a higher diagnostic rate than WES or earlier technologies),2 with the potential to reduce lengthy and expensive diagnostic pathways,3 as well as generate lifetime health benefits through application of appropriate treatments,4 when available. Clinical guidelines for pharmacogenomic testing are also available for a number of gene–drug pairs,5 while clinical genomic testing in the context of common complex diseases is more limited, with few robust polygenic risk scores being available.6

The clinical translation of genomic sequencing is further hampered by the limited information available to guide investment into those areas where genomics is well placed to deliver improved health and economic outcomes. There is a need for identification of the key elements of a framework for distinguishing applications of genomics that are likely to have demonstrable value and meet current effectiveness and cost-effectiveness evaluation guidelines, by prioritising the most pertinent genomic information to ensure genomic sequencing is most productively and efficiently applied in clinical practice.

FIRST STEPS IN IMPLEMENTING GENOMIC MEDICINE

Advancements in genomic sequencing enable more accurate diagnoses, rational disease prevention strategies, better treatment selection, and the development of novel therapies. This approach to medical care has been called many different names, such as genomic, personalised or precision medicine (see Table 1), but here we refer to genomic medicine as ‘the use of genomic information and technologies to determine disease risk and predisposition, diagnosis and prognosis, and the selection and prioritisation of therapeutic options’.7

When individuals with a condition with high penetrance (with near certainty the individual will develop the condition) are identified, there is a greater likelihood of preventative or mitigating interventions being successful. Accordingly, genomic medicine has achieved its greatest successes through applications to high penetrance monogenic diseases. Significant health benefits from the provision of genomic information using this criterion are already demonstrable, even when the behavioural modification required is arduous.

There are, as yet, few examples of the use of genomic sequencing with accompanying costs of the impact of change in disease trajectory, thus we will draw lessons from available studies, including in some cases conditions identified with older technologies such as biochemical or single gene tests.
Phenylketonuria (PKU) is an autosomal recessive disorder, caused by mutations in both alleles of the gene for phenylalanine hydroxylase. Although it is more cheaply diagnosed with a biochemical test, the management of this inborn error of metabolism is an important example of substantial therapeutic benefit being successfully achieved as a result of significant behavioural modification following the provision of genetic information. Delay in dietary treatment can have significant impacts on a child’s intellectual development. Maintaining a phenylalanine-restricted diet is quite onerous, yet despite this burden, the incidence of the cognitive impacts of PKU has been markedly reduced (by 92%) as a result of newborn screening and dietary advice. Genomic testing in the context of PKU is most likely to be most relevant as either a prenatal or preconception carrier screening test to restore reproductive confidence, however, this would also necessitate identifying the disease-causing mutation in the affected child first using genomic testing.

Molecular testing can also have an immediate and effective application to other common monogenic diseases that meet our recommended criteria of high penetrance and significant health effects (see below). Identifying patients with familial hypercholesterolaemia via genetic testing allows for the application of prophylactic therapy (statins), which has resulted in a reduction of death and major adverse cardiovascular events of 66% and 51%, respectively. Similarly, a reduction in the incidence of serious and costly monogenic conditions such as fragile X syndrome, cystic fibrosis, and β-thalassaemia of 47–90% has been reported as a result of informed family planning following preconception genetic testing, with the added benefit of restored reproductive confidence. This is a remarkable step forward from the time when a majority of families, even those with a family history (FH) of a serious genetic disorder, did not have a molecular diagnosis to inform reproductive management. With genomic testing, many more families could obtain an accurate assessment of their risk of having a child with a severe genetic disorder, and have their reproductive confidence restored through preconception care, screening, and preimplantation genetic diagnosis or invasive testing in pregnancy.

Additional examples demonstrate an emerging capacity to obtain a molecular diagnosis and offer treatment particularly in rare childhood diseases as new causal genes (e.g., there are more than 1000 monogenic causes of intellectual disability now known) are identified. New therapies are emerging such as a study reporting on 81 inborn errors of metabolism, with therapies including diet, co-factor/vitamin supplements, small molecule substrate inhibition, bone marrow and hematopoietic stem cell transplantation and gene therapy. With the exception of gene therapy and stem cell transplantation, these treatments may be relatively accessible and affordable. While there are no published studies yet of the long-term impact in terms of a molecular diagnosis on health outcomes and cost of change of management, there are a small number of published studies in clinical cohorts demonstrating that the diagnostic costs are lower when using genomic sequencing in childhood syndromes and neuromuscular disorders such as Stark et al., Tan et al. and Schofield et al. and while another by Tsiplova et al. reported similar findings for autism spectrum disorder, but using a hypothetical cohort. Sagoo et al. on the other hand reported a higher diagnostic rate, but at higher cost for a series of cohorts.

There is also potential for genomic sequencing to reduce the significant economic burden associated with adverse drug reactions, which can potentially be reduced by 4–30%. The annual cost of medical care associated with this group of preventable conditions is so high that the implementation of one genomic test early in life may be a cost-effective use of limited healthcare resources.

In cancer too we are beginning to see evidence of cost-effective interventions particularly in relation to screening, where the cost of prophylactic screening is much lower than treatment of the cancer itself. For example, Gallego et al. reported screening for Lynch syndrome using next generation sequencing panels was cost effective, while Li et al. reported screening women at risk of hereditary breast cancer with prophylactic intervention was similarly cost-effective.

Thus, we argue, the priority for genomic testing should be to identify individuals at high risk of imminent, serious, preventable (or reversible) disorders that are cost effective to treat. These patients and their families are well placed to benefit from genomic medicine through sequencing of affected children and provision of targeted therapies where available. There are already population-specific screening programmes demonstrated to be effective in identifying parents at high risk of having children with such disorders, but there is a need to assess the feasibility and cost-effectiveness of expanded genomic carrier screening. Existing genomic medicine and carrier screening programmes are not universally available. They are often still within a research context, which in time, will provide evidence on incremental health gains and cost-effectiveness, which is valuable for making the case for universal access and public funding.
| Disease                          | Incidence                | Intervention                                      | Reduction in incidence | Lifetime per patient direct medical costs of disease (USD 2016) | Annual direct medical costs of disease incurred by health system (USD 2016) |
|---------------------------------|--------------------------|---------------------------------------------------|------------------------|---------------------------------------------------------------|---------------------------------------------------------------------|
| Phenylketonuria (PKU)           | 1/10,000 \(^{38}\)      | Dietary treatment with a phenylalanine restricted diet | 92%\(^{6}\)           | $32,930/patient (up to 36 years of life) \(^{39}\)             | —                                                                   |
| \(\beta\)-thalassemia major     | 1/100,000 \(^{38}\)     | Reproductive planning                              | 90%\(^{10}\)          | $1,370,182/patient (60 year lifespan) \(^{50}\)                | —                                                                   |
| Cystic fibrosis (CF)            | 1/8,000 to 1/10,000 \(^{38}\) | Reproductive planning                              | 65%\(^{9}\)           | $511,961/patient (28 year lifespan) \(^{51}\)                 | —                                                                   |
| Fragile X syndrome (FXS)        | 1/4,000 to 1/5,000 \(^{38}\) | Reproductive planning                              | 47%\(^{8}\)           | $679,469/patient (72 year lifespan) \(^{52}\)                 | —                                                                   |
| Familial hypercholesterolaemia  | 1/500 for heterozygous dominantly inherited; 1/1,000,000 for homozygous autosomal recessive inherited \(^{38}\) | Prophylactic statin therapy                         | 51%\(^{7}\)           | —                                                             | $2,931,345,622 \(^{43}\) (per annum)                              |
| Adverse drug reactions (ADRs)   | 2,216,000 hospitalised patients in the US had serious ADRs; 106,000 had fatal ADRs \(^{44}\) 1/16 hospitalisations in the UK are a result of ADRs \(^{45}\) | Informed prescribing                                | 17% (4–30%)\(^{17}\) | —                                                             | $242,392,437,330 \(^{46}\) (per annum)                              |

\(^{a}\) For PKU percentage refers to the reduction in PKU patients with low IQ (<90)

\(^{b}\) For FH percentage refers to the reduction in major adverse cardiovascular events in patients with homozygous FH after receiving lipid-lowering therapy

\(^{c}\) All costs were inflated to 2016 prices using country-specific (United Kingdom or United States) OECD all items non-food, non-energy Consumer Price Indexes (http://stats.oecd.org/Index.aspx?DatasetCode=MB_PRICES) and converted to United States dollars (1 GBP = 1.29 USD) as to report all costs in a common currency and year

\(^{d}\) For PKU the cost refers to the per-patient lifetime direct medical costs of a PKU patient that does not receive a phenylalanine restricted diet, thereby representing the direct medical costs of not identifying an individual with PKU

\(^{e}\) For FH the cost refers to the annual medical costs of coronary heart disease across all of the United Kingdom, which highlights the magnitude of the direct medical costs that are associated with the condition. The argument being, if prophylactic statin therapy were used in all individuals identified to have either heterozygous or homozygous FH through genomic sequencing, a 51% reduction in the incidence of major cardiovascular events would therefore result in large savings to health care systems in terms of direct medical costs

\(^{f}\) For ADRs the cost refers to the annual medical costs of drug-related morbidity and mortality in the United States, which highlights the magnitude of the direct medical costs that are associated with ADRs. Even a modest reduction of 17% in the incidence of these events through more informed prescribing after genomic testing would therefore result in large savings to health care systems in terms of direct medical costs
EXPANDING GENOMIC MEDICINE TO COMPLEX DISEASES

There has been considerable hope for the successful application of genomic medicine to common complex diseases, such as heart disease, cancer, obesity, diabetes and lung disease. It has been suggested that adding personal genetic risk (usually based on a combination of low penetrance alleles) to general risk information (i.e., lifestyle factors, medical and FH) could impact on individual behaviour and potentially prevent such diseases from developing, however evidence for the utility of genomic testing in this context is not yet available.

However, the evidence from an updated Cochrane review and meta-analysis provides little support for behavioural change when healthy individuals are presented with genetic information compared to general risk information concerning their risk of various complex diseases. Since the publication of the Cochrane review in 2016 evidence of behavioural change has been noted in individual studies, but generally the impact on disease incidence is small. For example, one study indicated that the use of statins with the provision of both genetic and general risk information for coronary heart disease (CHD), compared to only general risk information alone, would result in a reduction in CHD incidence over 10-years in a high genetic risk population of only ~5%,23 assuming a ~45% relative-risk reduction from high-intensity statin therapy.

In another recent study, individuals receiving genetic risk information for Alzheimer’s disease and coronary artery disease together (compared to only receiving genetic risk information for Alzheimer’s disease alone) reported more health behaviour changes related to diet, exercise, medications, dietary supplements and stress reduction. This one study alone indicates a greater likelihood of behavioural change when multiple genetic risks are presented together and at least one of the disease risks are modifiable through medical intervention. Although this study did not report long enough follow-up to determine whether the intervention reduced Alzheimer’s disease or coronary artery disease incidence and does not supersede the negative effect noted in the majority of the evidence synthesised in the updated Cochrane review. Similarly, Vassy et al. report on the use of WGS plus FH over FH alone in primary care for health adult patients with new clinical actions in 34% of the WGS plus FH patients compared 16% for FH alone. Only 2 out of the 11 patients with an identified Mendelian allele manifested a phenotype consistent with at least 80% non-penetrance. The study was also limited by a small sample size (100 patients) with no follow-up data reporting on whether patients obtained a significant health benefit.

Currently there is limited clinical and economic evidence of the utility of genomic sequencing in common complex diseases, and of those studies reported some have significant limitations. For example, Dauzé et al. estimated that reporting genomic variants in these diseases would lead to modulation of health-related behaviours with a 10–50% reduction in disease incidence, valued at $US33–607 (£22–405) billion per condition. However, this estimate was predicated on the assumption that the provision of genomic information will overcome the limitations of current lifestyle interventions where adherence to such programmes has traditionally been poor on a population level. As outlined above, the evidence does not support this assumption, and thus the projected benefits are in all likelihood vastly overestimated.

This is not to say that genomics is not relevant to the reduction of complex diseases, but rather it is necessary to better understand the genotype–phenotype relationship in complex diseases before effective genomic risk identification and management of complex diseases can be implemented in clinical practice.

Furthermore, some conditions traditionally classified as complex diseases are in fact disease clusters with many causes. For example, there are nearly 200 genetic variants known to have an effect on blood cholesterol, which may, in part, impact response to lipid lowering medication. As an example, in some populations receiving statins, only a small proportion of patients achieve target reductions in plasma cholesterol levels. Thus, targeting group health risk modification at a large cluster of conditions categorised under a single heading may obscure the potential of genomic medicine in complex diseases. Applying genomic medicine to stratify and target specific interventions to individual genetic variation, and along with better knowledge of the genotype–phenotype correlations will enable better outcomes, both in therapeutic and behavioural responses.

A FRAMEWORK FOR DISTINGUISHING APPLICATIONS OF GENOMICS LIKELY TO HAVE DEMONSTRABLE VALUE IN REDUCING DISEASE BURDEN AND COSTS OF DISEASE

While there are well established guidelines for the conduct of an evaluation of clinical and cost-effectiveness within medicine such as those from the National Institute for Health and Clinical Excellence, as well as more recent frameworks for the evaluation of genomics, there is little to guide researchers on which areas of medicine are likely to yield outcomes that might meet the guidelines. The evidence to date suggests that based on our current knowledge, there is greater potential for deriving value from genomic sequencing (with its capacity to test many conditions simultaneously) when applied to monogenic disorders, with enormous potential to improve health outcomes through screening, disease prevention and change of management. Our synthesis of the examples included in this paper suggest that the distinguishing characteristics of applications of genomics with significant capacity to reduce the incidence of costly illness (Fig. 1) are the:

- strength of the genotype–phenotype correlation,
- high penetrance,
- imminence of severe illness,
- severity of the disease impact,
- relatively high diagnostic yield,
- availability of prevention or targeted treatment, and
- the net costs incurred are acceptable for the health gains achieved.

Promising areas that with further evidence in relation to health outcomes and/or cost impacts meet all or most of the criteria set out in our framework include: inborn errors of metabolism, neuromuscular disorders, neurodevelopment disorders, inherited cancer risk, preconception carrier screening or prenatal screening, and in the paediatric and neonatal intensive care unit. The importance of critical criterion in our framework such as diagnostic rate, health outcomes due to prevention or
therapy and health economic outcomes are highlighted in a paper by Gaff et al.\textsuperscript{34} The urgent need for evidence related to cost-effectiveness that takes account of both diagnostic rate and health outcomes as described in our framework is highlighted by Sagoo et al: the main limitation of the studies to date is that while it is possible to derive an incremental cost-effectiveness ratio from a relative diagnostic rate and cost for genomic sequencing, there is “no universally acknowledged willingness-to-pay threshold for a diagnosis”, and this makes an incremental cost per additional diagnosis difficult to interpret.\textsuperscript{18}

**IMPROVING ACCESS AND USABILITY OF GENOMIC DATABASES**

While existing databases such as OMIM\textsuperscript{35} provide a wealth of descriptive information concerning genomic variants, they do not allow the effect of a specific genomic variant to be predicted. In contrast, the ClinVar database provides specific variant information that predicts the relationship between human genomic variation and expressed phenotypes using supporting evidence. These are, however, limited in their scope, in that, ClinVar is currently largely incomplete and the ACMG list is limited to only genes that should be reported as additional findings rather than variants that might be of interest in the primary use of testing.\textsuperscript{35}

To improve accessibility of genomic information, there is a need for comprehensive, evidence-based, continuously curated, user-friendly genotype–phenotype databases to aid diagnosis and prognosis, and thereby appropriate treatment and prevention to capitalise on the promise of genomic medicine. Applying genomics to conditions that meet one of our main criterion for prioritisation of receiving genomic testing (i.e., a strong genotype–phenotype relationship) reduces the potential for false positives, while continuous curation is important in reducing the potential for both false positives and false negatives. With this information readily available from such databases in the future, numerous rare but collectively common and very costly diseases could be prevented (such as those listed in Table 2 and others fitting our recommended criteria), and treating patients with expensive but ineffective treatments could be avoided, offering large cost savings that create headroom to treat other patients with effective targeted interventions.

**FULFILLING THE PROMISE OF GENOMIC MEDICINE**

We are in a period of tremendous innovation in genomic medicine and large population studies such as Genomics England’s 100,000 Genomes Project and the All of Us programme in the United States hold great promise in identifying further conditions where highly penetrant variants causing serious disease might be much more effectively treated. To facilitate rapid translation of evidence based genomic medicine, there will also need to be increased capacity within the health system, particularly in laboratory and genetic services. The evidence suggests that deriving value from genomic medicine, in the short term at least, will be a function of the strength of the genotype–phenotype correlation (high penetrance), the severity of the disease impact, the availability of prevention or targeted treatment, and the net costs incurred for the health gains achieved. Further, widely available comprehensive, evidence-based, continuously curated, user-friendly genotype–phenotype databases of genomic approaches to treatment and prevention, which will be supported by data collected in the large population studies currently ongoing, will maximise the benefits from genomic medicine.

**ACKNOWLEDGEMENTS**

The authors would like to thank Owen Tan for his advice on the cost-effective use of next generation sequencing panels in cancer.

**AUTHOR CONTRIBUTIONS**

B.D. designed the study protocol, extracted, analysed and interpreted the data from a review of the literature; drafted and revised this article; and gave final approval of this version to be published. D.S., T.R. and J.S.M. conceptualised the study, assisted with its design and the interpretation of data, critically reviewed and edited the draft article and gave final approval of this version to be published. B.D., D.S., T.R. and J.S.M. are accountable for all aspects of the work and will ensure that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

**ADDITIONAL INFORMATION**

**Competing interests:** The authors declare that they have no competing financial interests.

**Publisher's note:** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

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