Information Needs in the Precision Medicine Era: How Genetics Home Reference Can Help

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Abstract

Precision medicine focuses on understanding individual variability in disease prevention, care, and treatment. The Precision Medicine Initiative, launched by President Obama in early 2015, aims to bring this approach to all areas of health care. However, few consumer-friendly resources exist for the public to learn about precision medicine and the conditions that could be affected by this approach to care. Genetics Home Reference, a website from the US National Library of Medicine, seeks to support precision medicine education by providing the public with summaries of genetic conditions and their associated genes, as well as information about issues related to precision medicine such as disease risk and pharmacogenomics. With the advance of precision medicine, consumer-focused resources like Genetics Home Reference can be foundational in providing context for public understanding of the increasing amount of data that will become available.

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KEYWORDS
individualized medicine; patient education as topic; databases, genetic; health resources

Introduction

In January 2015, President Obama announced the Precision Medicine Initiative [1], a research effort aimed at changing how diseases are diagnosed and treated in the United States. Its goal is to bring precision medicine—an approach to disease management that considers individual variability in genes, environment, and lifestyle—into the field of cancer and then ultimately into all areas of health and health care [2,3]. Integral to the initiative’s effectiveness is the recruitment of a longitudinal cohort of 1 million volunteers, who will be overseen by the National Institutes of Health (NIH). This cohort will provide genetic data, biological samples, and other health information to researchers [4]. Detailed analyses of this database of health information, including genome-sequencing data and lifestyle and environmental factors, will help determine the genetic contribution to disease development, identify disease risk factors, and generate effective treatments that incorporate the way genes affect a person’s response to drugs (a field of study called pharmacogenomics). This knowledge will enable clinicians to use genetic and other molecular information as part of routine medical care.

Reliable resources are necessary to support the public as they seek to educate themselves. Individuals searching online for information about the Precision Medicine Initiative may encounter unfamiliar concepts relating to health and genetics. Genetics Home Reference [5], an online resource from the US National Library of Medicine, provides consumer-focused information on various topics related to precision medicine, including how genetic variants relate to disease, pharmacogenomics, and genetic testing. In the era of precision medicine, online resources aimed at the general public, like Genetics Home Reference, are needed as more people become interested in the genetic aspects of health care.
Information Needs in the Precision Medicine Era

The need for new tools for researchers and clinicians to store, manage, and analyze large amounts of data has been discussed as a key factor in the implementation and success of precision medicine [2,6,7]. While storage and management of these data will be challenging, current tools may be helpful for data analysis. For example, to parse the meaning of newly identified genetic changes, researchers and clinicians can use tools such as Polymorphism Phenotyping (PolyPhen) [8] to determine whether a genetic change is likely involved in the development of disease. For pharmacogenomics associations, researchers and clinicians can use Pharmacogenomics Knowledgebase (PharmGKB) [9], a database that organizes information about genetic variants playing a role in drug response. Resources such as these will expand as more data are collected through precision medicine, making the databases more robust and increasingly useful for analysis.

While the utility of clinical resources for precision medicine has so far been paramount, the need for patient resources is equally important. Resources that are accessible for health care consumers can be used as a starting point for understanding precision medicine and its applications to health care. The influx of data generated by precision medicine means individuals will have access to more details about their health than ever before when making precision medicine-based health care decisions [10]. For people to make informed decisions in the era of precision medicine, it is imperative that they have an understanding of basic genetic principles; however, studies suggest that a substantial proportion of the general public lacks this understanding [11-13]. To support patient engagement in precision medicine and promote informed decision making, both clinicians and patients will require trusted online resources that provide easy-to-read information about genetic principles, genetic disorders, gene functions and their roles in disease, and pharmacogenomics. The use of Internet-based health tools increases patient engagement, which leads to better health outcomes [14].

The Internet is a major tool people use to research their health concerns: up to 80% of adults on the Internet report searching for health-related topics annually [15,16]. As precision medicine is adopted in clinical settings, it is inevitable that the public will turn to the Internet for information, as they have for other health inquiries. The Genetic and Rare Disease Information Center, an NIH resource that provides information targeted to patients and families with genetic disorders and to individuals interested in genetics who do not have a science background. Genetics Home Reference provides information targeted to patients and families with genetic disorders and to individuals interested in genetics who do not have a science background. Genetics Home Reference receives an average of 1.5 million visitors and 3.6 million page views each month. This website offers summaries of more than 1000 genetic conditions and more than 1300 genes. To construct these summaries, pertinent information is gleaned from scientific literature and written into summaries using language that can be understood by the lay public. Genetics Home Reference has information on dozens of topics aimed at educating the public about issues related to precision medicine, including genetic risk factors for disease and pharmacogenomics. For example, this resource provides information about the function of the BRCA1 and BRCA2 genes and explains how a mutation in either of these genes increases the risk of developing breast cancer and other types of cancer [22]. The presence of a mutation in either of these genes can help determine appropriate cancer screening and treatment approaches. Genetics Home Reference also offers information about genetic factors that alter a person’s response to a common blood-thinning drug called warfarin. These genetic variants predispose people who might need warfarin to develop either blood clots (warfarin resistance) [23] or abnormal bleeding (warfarin sensitivity) [24]. If a patient had one of these genetic variants, a doctor might target the initial warfarin dose for optimum effectiveness and reduce the risk of an adverse drug reaction.

Genetics Home Reference and Precision Medicine

Genetics Home Reference provides information targeted to patients and families with genetic disorders and to individuals interested in genetics who do not have a science background. Genetics Home Reference receives an average of 1.5 million visitors and 3.6 million page views each month. This website offers summaries of more than 1000 genetic conditions and more than 1300 genes. To construct these summaries, pertinent information is gleaned from scientific literature and written into summaries using language that can be understood by the lay public. Genetics Home Reference has information on dozens of topics aimed at educating the public about issues related to precision medicine, including genetic risk factors for disease and pharmacogenomics. For example, this resource provides information about the function of the BRCA1 and BRCA2 genes and explains how a mutation in either of these genes increases the risk of developing breast cancer and other types of cancer [22]. The presence of a mutation in either of these genes can help determine appropriate cancer screening and treatment approaches. Genetics Home Reference also offers information about genetic factors that alter a person’s response to a common blood-thinning drug called warfarin. These genetic variants predispose people who might need warfarin to develop either blood clots (warfarin resistance) [23] or abnormal bleeding (warfarin sensitivity) [24]. If a patient had one of these genetic variants, a doctor might target the initial warfarin dose for optimum effectiveness and reduce the risk of an adverse drug reaction.
Genetics Home Reference also covers other types of cancer, immune deficiencies and dysfunctions, enzyme deficiencies, and other drug sensitivities (see Table 1). A benefit to clinicians is the inclusion of numerous rare conditions that might never be covered during formal education, in addition to a variety of common disorders.

Table 1. A sample of conditions on Genetics Home Reference to which precision medicine could be applied.

| Condition                                           | Genetics Home Reference link                                                                 |
|-----------------------------------------------------|----------------------------------------------------------------------------------------------|
| **Cancers**                                         |                                                                                              |
| Breast cancer                                       | https://ghr.nlm.nih.gov/condition/breast-cancer                                               |
| Lynch syndrome                                      | https://ghr.nlm.nih.gov/condition/lynch-syndrome                                              |
| Prostate cancer                                     | https://ghr.nlm.nih.gov/condition/prostate-cancer                                             |
| Familial adenomatous polyposis                      | https://ghr.nlm.nih.gov/condition/familial-adenomatous-polyposis                              |
| Acute promyelocytic leukemia                        | https://ghr.nlm.nih.gov/condition/acute-promyelocytic-leukemia                               |
| Neuroblastoma                                        | https://ghr.nlm.nih.gov/condition/neuroblastoma                                              |
| Core binding factor acute myeloid leukemia          | https://ghr.nlm.nih.gov/condition/core-binding-factor-acute-myeloid-leukemia                 |
| **Immune system disorders**                         |                                                                                              |
| Celiac disease                                      | https://ghr.nlm.nih.gov/condition/celiac-disease                                              |
| Type 1 diabetes                                     | https://ghr.nlm.nih.gov/condition/type-1-diabetes                                             |
| Autoimmune Addison disease                         | https://ghr.nlm.nih.gov/condition/autoimmune-addison-disease                                 |
| Rheumatoid arthritis                                | https://ghr.nlm.nih.gov/condition/autoimmune-addison-disease                                 |
| Graves disease                                      | https://ghr.nlm.nih.gov/condition/graves-disease                                              |
| Autoimmune lymphoproliferative syndrome             | https://ghr.nlm.nih.gov/condition/amyotrophic-lateral-sclerosis                              |
| Systemic lupus erythematosus                        | https://ghr.nlm.nih.gov/condition/systemic-lupus-erythematosus                               |
| **Enzyme deficiencies**                             |                                                                                              |
| Lactose intolerance                                 | https://ghr.nlm.nih.gov/condition/lactose-intolerance                                         |
| Glucose-6-phosphate dehydrogenase deficiency        | https://ghr.nlm.nih.gov/condition/glucose-6-phosphate-dehydrogenase-deficiency                |
| Hereditary antithrombin deficiency                  | https://ghr.nlm.nih.gov/condition/hereditary-antithrombin-deficiency                         |
| Familial hypercholesterolemia                       | https://ghr.nlm.nih.gov/condition/hypercholesterolemia                                        |
| Protein C deficiency                                | https://ghr.nlm.nih.gov/condition/protein-c-deficiency                                       |
| Autosomal recessive congenital methemoglobinemia    | https://ghr.nlm.nih.gov/condition/autosomal-recessive-congenital-methemoglobinemia            |
| Gaucher disease                                     | https://ghr.nlm.nih.gov/condition/gaucher-disease                                             |
| **Adverse drug reactions**                          |                                                                                              |
| Warfarin sensitivity                                | https://ghr.nlm.nih.gov/condition/warfarin-sensitivity                                        |
| Warfarin resistance                                 | https://ghr.nlm.nih.gov/condition/warfarin-resistance                                         |
| Malignant hyperthermia                              | https://ghr.nlm.nih.gov/condition/malignant-hyperthermia                                      |
| Pseudocholinesterase deficiency                     | https://ghr.nlm.nih.gov/condition/pseudocholinesterase-deficiency                             |
| Dihydropyrimidinase deficiency                      | https://ghr.nlm.nih.gov/condition/dihydropyrimidinase-deficiency                              |
| Thiopurine S-methyltransferase deficiency            | https://ghr.nlm.nih.gov/condition/thiopurine-s-methyltransferase-deficiency                   |
| Dihydropyrimidine dehydrogenase deficiency          | https://ghr.nlm.nih.gov/condition/dihydropyrimidine-dehydrogenase-deficiency                 |
| Stevens-Johnson syndrome/toxic epidermal necrosis   | https://ghr.nlm.nih.gov/condition/stevens-johnson-syndrome-toxic-epidermal-necrosis          |

Genetics Home Reference also provides a primer called Help Me Understand Genetics for individuals who need foundational information. This primer has multiple chapters, covering topics from basic biology to the application of genetics in medicine. The precision medicine section of the primer explains this new approach to health care as well as the goals, benefits, and limitations of the Precision Medicine Initiative (see Table 2). Also of interest are health care–based issues, such as chapters on mutations and health, pharmacogenomics, and genetic testing. Help Me Understand Genetics provides information on the
many facets of genetic testing that individuals will need to become familiar with as genetic testing becomes more routine for disease diagnosis and defining treatment options, such as indications for testing, interpretation of test results, and the difference between research and clinical testing.

Table 2. Background information about precision medicine from Genetics Home Reference.

| Topic                                                                 | Genetics Home Reference link                                                                 |
|----------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|
| What is precision medicine?                                          | https://ghr.nlm.nih.gov/handbook/precisionmedicine/definition                                  |
| What is the difference between precision medicine and personalized medicine? | https://ghr.nlm.nih.gov/handbook/precisionmedicine/precisionvspersonalized                      |
| What is the Precision Medicine Initiative?                           | https://ghr.nlm.nih.gov/handbook/precisionmedicine/initiative                                  |
| What are some potential benefits of precision medicine and the Precision Medicine Initiative? | https://ghr.nlm.nih.gov/handbook/precisionmedicine/potentialbenefits                           |
| What are some of the challenges facing precision medicine and the Precision Medicine Initiative? | https://ghr.nlm.nih.gov/handbook/precisionmedicine/challenges                                  |
| What is pharmacogenomics?                                            | https://ghr.nlm.nih.gov/handbook/genomicresearch/pharmacogenomics                              |

Conclusion
Consumers are required to take an increasingly active role in their health care decisions, and they turn to the Internet to gather information regarding health issues. In the era of precision medicine, individuals will search for information to understand their genetic profiles and other health concerns. Various aspects of precision medicine are covered in online resources, including ClinVar [25], Genetic Testing Registry (GTR) [26], GeneReviews [27], Online Mendelian Inheritance in Man (OMIM) [28], and Orphanet [29]. However, these resources are designed primarily for researchers and clinicians, using technical information and language that can be overwhelming for most lay individuals.

Few consumer-focused resources about precision medicine exist, and the need for such resources will only increase. Resources are needed to put into context the growing amount of genetic and other health data that are becoming available [5]. Genetics Home Reference provides consumer-friendly information on topics relevant to precision medicine, including genetic conditions, gene function, and the effects of genetic variation on health, genetic testing, and pharmacogenomics. This information is useful to a variety of people, patients and clinicians alike, as the public increasingly turns to the Internet as a health resource. During this era of precision medicine, Genetics Home Reference seeks to facilitate health consumers in becoming well informed.

Research into consumers’ specific information needs related to precision medicine could help guide the evolution of existing educational resources and the development of new resources [30]. These studies should first assess existing resources to identify areas that are not covered. For example, we are unaware of any comprehensive consumer databases that connect specific genetic variants to the development of disease or that outline treatment options based on particular genetic profiles. Such resources would help patients gather targeted information that is specific to their health situation and would be valuable additions in the age of precision medicine.

Genetics Home Reference is committed to supporting patient engagement through the ongoing addition of new information that is relevant to precision medicine. To determine which topics are relevant for inclusion on the website, Genetics Home Reference staff frequently collaborates with outside groups that include other NIH institutes, advocacy and support groups, and unsolicited user feedback. These collaborations ensure that the content on Genetics Home Reference appeals to a wide range of audiences. Genetics Home Reference will also continue to provide links to other reputable online resources that offer information beyond our scope. Developers of consumer-focused health resources, including Genetics Home Reference, have the opportunity to be proactive in providing education about precision medicine to the public concurrently with the implementation of this new approach to care.

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Conflicts of Interest
None declared.

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Abbreviations

GTR: Genetic Testing Registry
NIH: National Institutes of Health
OMIM: Online Mendelian Inheritance in Man
PharmGKB: Pharmacogenomics Knowledgebase
PolyPhen: Polymorphism Phenotyping