The Disclosure of Direct-to-Consumer Genetic Testing: Sounding Out the Psychological Perspective of Consumers

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List of Abbreviations:

DTC: Direct-to-Consumer; FDA: Food and Drug Administration;
DTC-GC: Direct-to-Consumer Genetic Service

Commentary

Medicine is increasingly conceived as tailoring to the needs of each individual. Progress in this area comes, for instance, in the wake of human genetic testing, which intends to “profile” patients’ genetic predisposition towards specific diseases, promoting early and personalized interventions. This aspect is a fundamental component of the so-called pre-emptive approach [1].

In implementing genetic testing on a larger scale, there is a need to understand who the consumer of genetic testing really is. One must therefore consider the uniqueness of every individual’s psychological, social, cognitive and behavioral profiles. According to these premises, we assume that having information about the consumers’ psychosocial-cognitive state would help healthcare professionals and policy makers to find effective strategies that can be employed to successfully interact with consumers, and thereby strive to really enhance individual empowerment and shared decision making [2-4].

The transition from patient to consumer

Initially, genetic testing was provided only by specialized institutions, as a part of clinical and research studies and within a network of experienced professionals. The search by scientists for disease-linked genes began by studying DNA samples from disease families, i.e. families in which numerous relatives, over several generations, had developed the illness. Therefore, those who underwent this type of screening were people with a family history of disease.

Since the early 1980s, several private companies have been offering personalized medicine genetic testing to the public. Their perspective is not in the sale of the test (sometimes inexpensive at around $99) but rather in the creation of a DNA database of patients/citizens for the development of new treatments and new genetic tests. The 23andMe web page (the most popular genetic company in US) expressed it thus: “getting involved in a new way of doing research […] with enough data, we believe can produce revolutionary findings that will benefit all” (https://www.23andme.com/research/). Advertisements tend to highlight the benefits and minimize any possible limitations [5,6]. Companies offering DTC genetic tests employ highly aggressive marketing strategies, in which they explain that the data could be used: “to shed light on your ancestors, your family and, above all, on yourself”, to understand the most suitable diet or lifestyle for each person, to understand how one’s body reacts to nicotine for instance, or to put you in possession of “information to share with your family and friends” [7,8]. Genetic testing comes as a simple tool to acquire personal information, independent of the purpose for which it will be used.

In the outlined scenario, each individual, although not a professional researcher, can exercise the freedom to participate in scientific research, and at the same time can make available his/her genetic heritage, drawing an immediate advantage (e.g. gathering information about a condition from which he/she suffers). Consumers choose to be engaged in the biotechnology research field to use genetic analysis results for the protection of their own health [9].

The exploitation of the potentiality of genetic testing has led to the emergence of a category of consumers that medical science has termed “unpatients”. These are healthy people who show predispositions, at the genotypic level, concerning the onset of certain diseases whose genetic matrix is properly known [10].

Nevertheless, diseases caused by a single gene usually do not present any problems in interpreting risk probability, while common-complex disorders such as heart diseases, diabetes, arthritis, cancer and psychiatric disorders are usually the result of variation in many genes, each contributing to a small amount of genetic susceptibility, acting in concert with environmental or epigenetic factors [11,12]. In this case, tests present serious problems of interpretation because genes play a limited role (usually from 5% to 20% of the risk). Many genes can only increase susceptibility to disease development but they do not represent a certainty or a "sentence to disease".

Unsure about which is better to choose and how much (if any) trust to place in such services, consumers choose to consult several companies, aiming to gain greater certainty about the result. However, the analyses provided by DTC companies do not plumb the entire genome. The ‘variable’ points in DNA number millions throughout the genome, dozens in a single gene. It is sufficient that companies opt for different ones, although related to the same disease, to achieve radically different results. Moreover, consumers purchase tests without the obligatory involvement of the healthcare providers, leaving the consumers free to make their own interpretation and manage their genetic risk data. A free market of choices detached from a medical and therapeutic context takes shape [13].
The importance of being a consumer

It has been argued that tests may even cause psychological harm to consumers, since they provide complicated and potentially unreliable data, leaving people vulnerable to inappropriate health decisions [6,14]. A perceived high risk of disease may increase anxiety and a low risk may discourage healthy behavior [15,16]. Nevertheless, a recent review [17] concluded there was no evidence of harm by DTC genetic tests. The largest study on DTC genetic tests [18] found no significant difference in anxiety levels before testing, after testing and 3 months later, and no increase in the subsequent use of additional screening tests associated with the results of the DTC. Another study [19] found no significant changes in habits except for those regarding physician consultation. Concerning the profile of consumers, the literature shows that people who spontaneously decide to buy a genomic test are more likely to be “information seekers” [20], and we could infer that they might be “anxious” about their health and “anxious” to know. Indeed seeking information might serve both to reduce distress and to take concrete action against the risk [21,22]. Despite these inferences, the literature also shows that being an information seeker and being more informed about advances in genetics may be predictive of one’s decision on taking up genomic tests but not predictive of a concrete use of this information [20,23,24]. Consumers collect information which they then do not use, or at least they do not use the way we expect (change of habits, quality of life, screening etc.) [25]. The core issue is that if people do not believe that knowing more about a health condition, such as a genetic condition, will allow them to act something (to do something and face the condition), then they are not persuaded to effectively use this knowledge [26].

Hence, we need a comprehensive examination of “who are these consumers?” [27].

The last few years of scientific and policy debate on direct-to-consumer genetic testing culminated in a series of studies indicating that, contrary to initial intuitive expectations, people are substantially resilient to the long-term consequences of positive results to genetic testing [28]. A little anxiety, when a health-related outcome is identified, is an almost predictable index to something unknown because people have to face the uncertainty of the onset of a disease [29], the possibility of passing the faulty gene on to their children, and the potential for genetic discrimination. However, we suggest that non-clinical anxiety can be considered as a precursor of patients’ health engagement and behavioral change, but only in the light of other complex cognitive and emotional mechanisms whose investigation in genetic risk is of paramount importance [29].

Both emotional and cognitive factors affect people’s resilience to gene test results [30-32]. Ho et al. [33] demonstrated that people who undergo HCRC (hereditary colorectal cancer) genetic testing and have a high level of hopefulness would have a higher tendency to show a resilience outcome trajectory pattern than their low-hope counterparts. A mindful and accepting orientation toward experience may facilitate psychological resilience in response to negative outcomes and illness [34,35]. Predictors of resilience include personality characteristics, positive or negative personal and family experiences [29] and being flexible in the coping strategies used.

Moreover, we need to take into account the fact that consumers’ attitudes towards genetic testing differ by country, based on cultural differences, and are related to individual characteristics such as age, gender and educational level [36,37]. People are more worried about privacy and discrimination problems in the USA and northern Europe [38-40], whereas they are more concerned about the influence such tests may have on future plans and on their reliability in southern Europe [37,41,42]. Younger people have a higher level of interest in genetic testing and DTCGT, along with a higher attitude towards new health technologies in general [43].

According to all these evidences, we argue that future studies on direct-to-consumer genetic testing should provide a rich conceptual framework about the worries and concerns that consumers may feel about genetic testing, and identify their psycho-emotional profiles and health-related behaviors. In this way we would hope to obtain a clearer picture of their different interests perceptions of genetic risk. We furthermore suggest the implementation of a longitudinal perspective, with regular follow-up assessments to empirically measure how consumers’ experiences, perceptions, attitudes and preferences regarding risk communication change over time [39].

Perception of risk, uncertainty, and responsibility

Risk is linked to the knowledge of the probability of possible outcomes, whereas in a state of uncertainty the probability of an event is unknown. However, in medical contexts the term “risk” is often used even when probabilities are unknown and the distinction between risk and uncertainty becomes unclear. DTC is such a case.

The introduction of DTC genetic information raises questions on risk communication, health responsibility, and management of the gap between risks and lack of therapeutic options. Typically, decisions about appropriate risk levels for genetic information are decided by expert committees [13]. There is an almost unanimous belief that the spread of this kind of information is justified only if there is a useful way to pre-emptively react (e.g. when therapeutic effectiveness is mainly linked to an early diagnosis and genetic results are able to ensure a susceptibility diagnosis for curable disease).

Nevertheless, although for many genetic conditions (e.g. Alzheimer or Huntington disease) there are no effective therapies or unproven therapies (whose effectiveness and safety have yet to be demonstrated), the results of a genetic test, even if unfavorable, may have salient personal implications, since such results could influence reproductive decisions, decisions on family planning and on life planning in general. The utility of a genetic test cannot therefore be assessed by the sole criterion of its medical implications, but one should also consider the broader implications involving aspects of the life of the consumer, who should receive full information on the meaning of the genetic investigation and who should have the space for an independent evaluation.

We should strive to truly enhance individual empowerment and shared decision making and promote a model that is not opposed to the implementation of DTC genetic testing but imposes informational commitments to the companies which offer them (Figure 1).
Figure 1: Philosophical, social and ethical challenges of Direct-to-Consumer Genetic Testing. Web availability of genetic information and DTC services, without the involvement of health care professionals, associated with the informational richness, complexity and frequently uncertain meaning of these data, have generated a range of concerns about the ethical management of genomic information. Important issues about the concept of risk, uncertainty and responsibility distribution between the healthcare system and consumer arise. In this framework the psycho-behavioural identikit of the consumer plays a crucial role in policy regulation.

Competing Interests:

The authors declare no competing interests.

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