Abstract

We have been testing the hypotheses that our unconscious behaviors are often more predominant than our conscious behaviors. We have tried to quantify the degree of influence of our genetic factors and unconscious signals from the environment in our behaviors. In this paper, the concept and the research framework of MyFinder proposed by the authors are discussed. MyFinder combines the notion of intelligent agent from the artificial intelligence (AI) community and personal genome research results from the biomedical community. One of the important objectives of MyFinder is to allow the person to be conscious of (i.e., be aware of) her/his unconscious behaviors. The on-going experiments related to the Quantified Self (QS) and Citizen Science, as well as author’s real experiences are introduced. We discuss the meaning of self-awareness and the social impact of our findings.

Introduction

Socrates once said, “know thyself.” A desire to know more about myself was the springboard for my concept. This desire can lead one to ponder the philosophical question, “who am I?” We take for granted the fact that we actively make life-orientating decisions using our own free will. However, what if these decisions are in fact controlled by our genetic disposition and unconscious signals from our societal surroundings?

The arrival of the Personal Genome Age has resulted in the current boom in the field of gene analysis. The concept of Personal Genome refers to the gene information about a single person. Individual gene differences are associated with such factors as susceptibilities to various disorder [PG 13]s, physical constitution [Ashley 10], and the effects of drugs [Butte 08]. Owing to revolutionary developments in gene analysis technology, we are approaching the age in which each one of us can obtain our personal gene information [Jun W 08].

From an evolutionary prospective, it is likely that rapid modern changes in lifestyle largely influence the human mind and body. Our fundamental motivation for this research is our desire to enrich the minds of modern people by clarifying the mechanisms of the brain, genes, and body and correcting them through the use of artificial intelligence [Kido 10].

What kind of influence does personal genome information have on an individual’s self-knowledge, our values, future lives, and research in artificial intelligence? This paper aims to outline the influence of personal genome information using the research framework of MyFinder.

MyFinder

MyFinder [Kido 11] has two objectives. The first objective is to find innate potential characteristics and personality and to bring out the maximum potential in one’s abilities (individual approach). The second objective is to create a research platform that facilitates scientific discoveries through community computing (collective intelligence approach).

In terms of the first objective of MyFinder, recent Personal Genome research has focused on the realization of custom-made medical care by finding disease risks and drug effects, but MyFinder is unique as emphasis is placed on aspects relating to wellness, mental sciences, and behavioral sciences. MyFinder is based on recent findings relating to gene expression control and the epigenome [Sasaki 05] and supports the hypothesis that our physical,
chemical, and psychological stress greatly influences the activity of our genes. For instance, recent research has reported that the act of laughing affects diabetes [Hayashi 06], and positive mental stress caused by laughing is associated with turning on and off the genetic switch that controls gene expression at the cellular level. It will be possible for an intelligent agent to monitor our daily physical, chemical, and mental stress by way of everyday observation and analysis of our daily habits including eating, sleeping, work style, time management, social interaction, skills, and preferences. The intelligent agent technology can be effective in learning individual behavioral characteristics and stress status. 

Regarding the other goal of MyFinder, to build a research platform for scientific discoveries through the use of community computing, MyFinder learns each user’s personality by monitoring daily behavior and aims to interactively inform the user of its findings by using psychology-based and behavioral sciences-based findings (e.g. including the Enneagram theory in psychology [Helen 96]). This function will aid an individual in rediscovering his/her innate potentials and personality. 

The Personal Genome information environment acts as an intermediary in the boundary region between artificial knowledge and the theory and practice of gene research. MyFinder is a new research framework which uses the Personal Genome information environment and integrates the concept of intelligent agent from artificial intelligence research and the concept of disease-related gene search in the community of genetics. This concept is shown in Fig. 1.

The ultimate objective of MyFinder is to identify the intelligent agent which watches over unconscious behaviors and state of a person and allows the person to increase his/her “awareness” by supporting the process of becoming conscious of his/her unconscious behavior. This means that the intelligent agent reads unconscious signals of the individual, finds a pattern from the signals,
constructs a model from the pattern, makes predictions based on the model, provides the person with predictions at the right timing, and continues to correct the model by receiving feedback in an interactive manner from the person. For instance, the revolutionary device such as the sociometer used in the research of “honest signals” by Pentland [Pentland 13] is an example of enabling the measurement of unconscious social signals of a person (nonverbal information such as a manner of speaking, tone of voice, and subtle actions).

Self-Discovery by Personal Genome

The Personal Genome has brought about innovations in the community of prophylactic medical care. Among recent prominent examples of progress in Personal Genome analysis, a state-of-the-art technology is the integrative Personal Omics Profile (iPOP) project. Rui et al. [Rui 12] integrated multi-omics data (biomolecular information including epigenome, transcriptome, proteome, metabolome, etc.) including the Personal Genome of Dr. Michael Snyder of Stanford University and information collected over a period of 38 months (as of May, 2013) and reported the occurrence of genome edits caused by infection with HRV or RSV. In the 14th month, the results of changes in various omics data, including gene information caused by an allergy-induced fluctuation in cytokine, were reported [Rui 12]. Dr. Snyder has revealed gene information relating to various diseases primarily on diabetes and has tried to make use of the information for disease prediction, early diagnosis, monitoring, and therapy.

The first author is proceeding with a gene analysis of himself in cooperation with the bioinformatics team of Stanford University. The left section of Fig. 2 is one example of the mapping of gene variation and predominant physical constitution, drug efficacy, behavioral characteristics, character traits (disease risk, virus/bacterium resistance, drug efficacy, senses, thoughts, behavioral characteristics, physical constitution, aging, and physical ability) of the human body. The right section of Fig. 2 shows the positions of the genes on human chromosomes that increase disease risks due to gene variations. Common Variants are ordinary variations for which the gene variation is 1% or more, and Rare Variants are rare variations for which the gene variation is less than 1%. A universal scientific standard of measure has yet to be developed for Personal Genome analysis, although various results are obtained such as disease risk prediction as pointed out by Venter et al. [Venter 10].

Furthermore, it is known that disease risk can be dependent on racial differences [Kamatani 10], [Kido 10]. The first author and co-researchers compared disease risk evaluations of the Personal Genome services of three companies and reported on the results, such as degree and cause of difference, comparisons among risk prediction algorithms, difference in disease risk between a Japanese group and an American/European group. We are planning to conduct demonstrative research as well as research using mathematical models. As we pointed out in [Kido 06] and [Kido 11], there is ample room for adopting artificial intelligence technology for Personal Genome analysis.

The first author continues his attempts to annotate disease risk, drug efficacy and information concerning other genetic characterizations (e.g., reactions to alcohol, nature of hair, smoking habit, food preferences, life span, sensitivity to pain, and tendency to avoid failure) to better characterize his own genome information, as was partially introduced in [Kido 10]. For instance, he has reported research results based on themes such as heavy or light drinkers, smoker or non-smoker, long-lived or short-lived, gain weight easily or with difficulty, successful or unsuccessful with dieting, and tendency or not to avoid failure.

Examples of interpretation obtained from the Personal Genome of the first author and his father are introduced below.

- Difficulty in metabolizing alcohol (ALDH2)
- Special gene frame shift variation (TRPV1) reactive to capsaicin (pungent component of chili pepper).
- A gene variation (ACTN3) related to muscle performance that is the same as that of many world-class sprinters.
- Poor ability of avoid failure based on past experience (DRD2).
- High tendency to seek new experiences (DRD4).
- Plurality of disease candidates of higher risk than the average, although no serious disease risk can be found at present.
- The author’s father identified a risk for colorectal cancer from his Personal Genome, succeeded in early detection and excision of the cancer, and is currently very healthy.

Particularly evident with the last example of the author’s father is the fact that the Personal Genome can be applied in the future to prophylactic medical care.
The Personal Genome and related supporting sciences have the potential to discover a new approach in observing human beings, which will consequently lead to new perspectives on life. We would particularly like to note the possibility that mental stress exerts great influence on the activities of genes discovered through research on the epigenome in recent years [Sasaki 5]. We hope the study of individual differences (individuality) will lead to the enrichment of human nature, and not to discrimination. The aim of MyFinder is to help individuals to reconsider and positively understand themselves. In other words, it is intended to provide a device which enables us to experience the concept of “we are different and that is good” by identifying and accepting differences from others.

Scientific Discovery through the use of Community Computing: Citizen Science

Using the crowdsourced health study platform, Genomera, the Genetic Predisposition for Healthy Sleep (available at: http://genomera.com/studies/sleep-genetic-predisposition-for-healthy-sleep), a community computing peer study was launched in September 2012. As of January 2014, 26 participants had shared their 23andMe genomic data relevant to the study, and 12 had fully completed the supporting phenotypic questionnaire. The phenotypic questionnaire is the Pittsburgh Sleep Quality Index, a standard tool for measuring sleep quality in academic research studies. The aim of the study is to investigate possible linkage between personalized genomic profiles and sleep quality. The study was inspired by similar recent studies that linked genetic SNPs to sleeping profiles in healthy adults [Bodenmann 12, Nova 12, Landolt 11, Ciarleglio 8].

Demographically, the study had 12 participants (six males, four females, two undisclosed), nine of whom disclosed their age. The age range was 24-61 years, with three participants in their 20s, three in their 30s, two in their 50s, and one over 60. Participants primarily lived in a variety of cities in the United States, and a few lived internationally. Most participants were of European descent, and a few were of African and Asian descent.

The findings of the study appear in Figure 3. The phenotypic results are in the top line (“Instrument Score”), consisting of 15 questions which participants answered on a four-tier scale of bad-to-good. Given a maximum score of 45, these study participants indicated that they are already enjoying a fairly good quality of sleep (scores ranging from 21-42). The x-axis of Figure 3 represents the 12 participants with their phenotypic score and different genomic scores. The second line from the top indicates the total genomic score. This was calculated based on the total number of favorable alleles each participant had, based on the 11 SNPs (single nucleotide polymorphisms) examined in the study that are listed in Table 2. For each SNP, a value of 0, 1, or 2 was assessed per the number of favorable alleles. A maximum value of 22 was possible. The study participants had an average score of 14 and a range of 9-18, indicating a higher than random or average presence of the favorable alleles. However, it can be seen readily from Figure 3 that no correlation was found between the phenotypic survey result and the genomic profile. Higher sleep survey scores were not indicative of a higher presence of the favorable alleles in the underlying genomic profile.

Figure 3. Genomics and Healthy Sleep Study Results

Three studies had found a correlation between genomic profile and sleep quality, and the lower three lines in Figure 3 enumerate them. First, the bottom line, looking at the two ADORA2 SNPs investigated by Bodenmann does show a slight incline in the right half of the line, but it is not striking enough to suggest a correlation. The second line from the bottom represents the BDNF, COM VALMET, and PRNP SNPs evaluated by Landolt, and shows no correlation with reported results from the phenotypic study. Likewise, the third line from the bottom represents the circadian rhythm SNPs delineated by Ciarleglio and shows no correlation with participant sleep quality.

Table 2. Genomic SNPs Examined

<table>
<thead>
<tr>
<th>Gene</th>
<th>SNP</th>
<th>Favorable Allele</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADORA2A</td>
<td>rs5751876</td>
<td>T</td>
<td>Bodenmann 12</td>
</tr>
<tr>
<td>ADORA2A</td>
<td>rs2236624</td>
<td>C</td>
<td>Bodenmann 12</td>
</tr>
<tr>
<td>BDNF</td>
<td>rs6265</td>
<td>C</td>
<td>Landolt 11</td>
</tr>
<tr>
<td>COMT</td>
<td>rs4680</td>
<td>A</td>
<td>Landolt 11</td>
</tr>
<tr>
<td>VALMET</td>
<td>rs799990</td>
<td>A</td>
<td>Landolt 11</td>
</tr>
<tr>
<td>PRNP</td>
<td>rs4964059</td>
<td>A</td>
<td>Ciarleglio 8</td>
</tr>
<tr>
<td>ARNTL2</td>
<td>rs1801260</td>
<td>A</td>
<td>Ciarleglio 8</td>
</tr>
<tr>
<td>CLOCK</td>
<td>rs2304669</td>
<td>A</td>
<td>Ciarleglio 8</td>
</tr>
<tr>
<td>PER2</td>
<td>rs228697</td>
<td>C</td>
<td>Ciarleglio 8</td>
</tr>
<tr>
<td>PER3</td>
<td>rs10462021</td>
<td>A</td>
<td>Ciarleglio 8</td>
</tr>
<tr>
<td>ADA</td>
<td>rs73598374</td>
<td>T</td>
<td>Nova 12</td>
</tr>
</tbody>
</table>

Overall, the citizen science study was unable to corroborate a correlation between genomic profiles and sleep quality in healthy persons. There could be many reasons, especially that the small sample size of 12 may be insufficient to replicate the results of the other studies. This
could be due to the fact that despite age, gender, and geographical differences, the study cohort may be a homogeneous group of early-adopter highly-interested persons who are already experiencing high sleep quality and better than average favorable genomic profiles for sleep. There may not be a correlation between this particular phenotypic instrument (Pittsburgh Sleep Quality Index) and the SNPs implicated for sleep quality on a general basis as explored in this study. Finally, a great deal of research has focused on genomic linkage to sleep pathologies, and sleep quality in healthy individuals is a new field where additional efforts may lead to replication. However, it is important to notice the potential value of crowdsourced citizen science studies as a resource for attempting to replicate the results of academic studies in additional cohorts, with greater permutation, and also as a potential mechanism for new discovery.

**Discussion and Future Perspectives**

**What is “Awareness”?**

MyFinder has the theme of “information environment that gives awareness to a person”. In order to realize the theme, it is necessary to understand what the “awareness” actually is. In this paper, awareness is defined as “to let an individual be conscious of her/his unconscious behavior”. The QS experience and the personal genome experience of the author were described in the third section as awareness was obtained from the viewpoint of the “individual”. Next, the “scientific discoveries by community computing” were described in fourth section as awareness obtained from the viewpoint of the “collective intelligence”.

The awareness can be systematized by the two axes of consciousness/unconsciousness and myself/others. In view of the “Johari window” (a graph model of awareness in interpersonal relationship), which is often referred to in communication psychology and the like, the self can be divided into (1) Open Self (self recognized by the person herself/himself and by others), (2) Hidden Self (recognized by the person herself/himself but not by others), (3) Blind Self (not recognized by the person herself/himself but is recognized by others), and (4) Unknown Self (not recognized by the person herself/himself and others), and the efforts for expanding the region of Open Self and Hidden Self can be considered as the process of bringing about awareness.

MyFinder supports the expansion of the range of self recognized by the person and the self recognized by others by letting an individual know the differences between her/his behaviors, senses, and thinking characteristics (i.e., supports understanding Open Self). Also, in cases where MyFinder is used to become aware of the differences in behaviors, senses, or thinking characteristics, MyFinder gives opportunities for developing mutual understanding by becoming aware of prejudice and sense of discrimination which have unconsciously been generated.

An important aim of MyFinder is to support the process of gaining awareness that allows a deeper mutual understanding.

**Social Significance of MyFinder: Design of Future Society**

In the Introduction, we raised the question “what if our decisions are in fact controlled by characteristics caused by personal genes and unconscious signals from the society surrounding us to considerable degrees?” In essence, we suggested the following two points:

- We do not know our own minds as well as we think we do. Supporting technology that enables a person to be conscious of her/his unconscious behaviors will become important.
- From now on, research on brain science and genes will make progress to scientifically clarify the differences among individuals. Society should be directed into respecting diversity (the concept of “gifted”: “we are different and that is good”, not toward prejudice and discrimination).

In relation to the question above, the following challenging questions would be raised.

- If decision-making is indeed influenced by genes and unconscious social signals, are “freedom” and “responsibility” merely based on nothing but the illusion of freedom?
- Is it really beneficial for a person to know her/his abilities and characteristics?
- How can a person be conscious of his/her self if he/she is different in comparison to others?
- How can the act of knowing herself/himself be converted into her/his happiness?
- What is essential in order to direct our society towards respecting personal differences as personality, rather than towards discrimination?
- How should we share the personal data in view of the above-identified questions?

These challenges will promote artificial intelligence research and contribute to improvements in the quality of life of human kind.

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The paper was primarily conceived and written by Kido. Swan contributed to the citizen science study section.

References