Review

Genetics and genomics of human ageing

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Ageing in humans is typified by the decline of physiological functions in various organs and tissues leading to an increased probability of death. Some individuals delay, escape or survive much of this age-related decline and live past age 100. Studies comparing centenarians to average-aged individuals have found polymorphisms in genes that are associated with long life, including APOE and FOXA3, which have been replicated many times. However, the associations found in humans account for small percentages of the variance in lifespan and many other gene associations have not been replicated in additional populations. Therefore, ageing is probably a highly polygenic trait. In humans, it is important to also consider differences in age-related decline that occur within and among tissues. Longitudinal data of age-related traits can be used in association studies to test for polymorphisms that predict how an individual will change over time. Transcriptional and genetic association studies of different tissues have revealed common and unique pathways involved in human ageing. Genomic convergence is a method that combines multiple types of functional genomic information such as transcriptional profiling, expression quantitative trait mapping and gene association. The genomic convergence approach has been used to implicate the gene MMP20 in human kidney ageing. New human genetics technologies are continually in development and may lead to additional breakthroughs in human ageing in the near future.

Keywords: human ageing; centenarians; tissue ageing; longitudinal studies; genomic convergence

1. INTRODUCTION

Ageing is characterized by a decline of multiple physiological functions leading to an increasing probability of death. Some ageing-related changes affect appearance, such as wrinkled skin and grey hair, whereas others affect organ function, such as decreased kidney filtration rate and decreased muscular strength. Ageing is a major risk factor in most human diseases, including heart disease and cancer. Average lifespan has increased at a steady pace of almost three months per year in both males and females since 1840 [1]. Japanese women have a life expectancy of 85 years, the highest in the developed world [1]. Initially, this increase in lifespan was probably attributed to decreases in juvenile mortality through treatment of infectious diseases [2]. In the latter half of the twentieth century, improvements in survival after age 65, due in part to improved treatments for ageing-related diseases, propelled the rise in lifespan [2]. For Japanese women, the chance of surviving from age 65 to 100 soared from less than 1 in 1000 in 1950 to 1 in 20 in 2002 [1]. Centenarians now constitute the fastest-growing segment of the US population, increasing in number from 3700 in 1940 to roughly 61 000 in 2006 [3]. Finding differences in genes between centenarians and average-aged individuals may point to molecular pathways important in the ageing process.

Ageing trajectories vary among individuals. Some individuals live to age 100, but most do not. Twin studies have shown that lifespan tends to be more correlated in monozygotic twin pairs than same-sex dizygotic twin pairs, with these differences yielding heritability estimates ranging from 0.23 to 0.33 [4,5]. Yet, little is known about specific genes that affect the rate of ageing or human lifespan.

2. CENTENARIAN STUDIES

One way to find potential ageing genes is to search for genetic differences between centenarians and average-aged individuals. The average lifespan in the US is 77 years and only one in 10 000 individuals survive to age 100 [6]. Siblings of centenarians have an eight to 17-fold greater relative risk of surviving to age 100 than those without centenarian siblings [7]. Alleles enriched in centenarians probably affect genes that are important for longevity. A summary of relevant gene association studies in long-lived individuals is shown in table 1. A genome-wide linkage scan of 400 short tandem repeats using 137 long-lived sibling groups of European descent from the Boston area identified a locus on chromosome 4 linked to exceptional longevity [8]. Genotyping 2000 single nucleotide polymorphisms (SNPs) within this 12 Mb locus revealed an association between microsomal transfer protein (MTP) and human lifespan [9]. The cases were 653 individuals aged 98 years and older.

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from the same Boston population and the controls were Caucasians less than 50 years old from several sources in the US [9]. This finding could not be replicated in French or German populations [10].

Most of the work comparing the genotypes of long-lived individuals to average-aged individuals has taken place in candidate genes rather than by genome-wide analysis. For example, the e4 allele of apolipoprotein E (APOE) is well known to increase risk of Alzheimer’s disease and cardiovascular disease [11,12]. The e4 allele (112Arg) of APOE differs from the most common e3 allele (112Cys) at a single amino-acid [13]. While the exact function of the e4 allele remains to be fully determined, evidence has been presented suggesting an interaction with the amyloid β protein. One characteristic of Alzheimer’s disease is the formation of amyloid beta plaques. In mice, the e4 form of APOE exhibits an impaired ability to promote amyloid β proteolysis compared to the e2 and e3 isoforms [14]. The e4 allele is found in significantly lower proportions of nonagenarians and centenarians, which suggests that individuals with this allele do not live as long as those without it [11,12,15]. By contrast, the e2 allele is enriched in long-lived individuals and may offer a protective effect for Alzheimer’s disease and cardiovascular disease [12,15].

A study of 35 additional genes related to cardiovascular disease found that an allele in apolipoprotein C3 (APOC3) is enriched in centenarians and their offspring compared to average-aged controls in an Ashkenazi Jewish population [6]. APOC3 is a very low density lipoprotein (VLDL) involved in fat metabolism and may delay the catabolism of triglyceride-rich particles. The enriched allele in centenarians was associated with low levels of APOC3 and a favourable lipid profile [6].

Shortening of the telomeres at the ends of chromosomes has been associated with age-related disease and mortality [16–18]. A recent study identified a common haplotype of four SNPs in the human telomerase reverse transcriptase gene (hTERT) that is enriched in centenarians and associated with longer telomere length [19]. It was also shown that centenarians and their offspring maintain longer telomeres compared with controls and that longer telomeres are associated with protection from age-related diseases, better cognitive function and lipid profiles of healthy ageing [19].

In addition to testing genes known to be associated with age-related diseases and phenotypes for association with longevity, genes known to promote longevity in model organisms have been examined in human populations. Mutations in insulin or insulin-like signalling pathway genes have been shown to extend lifespan in Caenorhabditis elegans [20], Drosophila melanogaster [21,22] and mice [23,24]. The insulin-signalling pathway negatively regulates the forkhead (FOXO) transcription factor [25]. When insulin or insulin-like growth factor I receptor (IGFIR) mutations have been observed in centenarians [27]. These mutations are associated with reduced activity of IGFIR as measured in transformed lymphocytes [27].

The forkhead box O3A (FOXO3A) transcription factor contains alleles associated with longevity in multiple Asian and European populations [26,28–31]. In all cases, the alleles associated with longevity are introonic and unlinked to known coding SNPs, so the functional SNPs may affect gene expression rather than protein activity, but this has not yet been shown. In a male population of Japanese descent, the odds ratio for homoyzogous minor versus homoyzogous major alleles for SNP rs2802292 in FOXO3A between the long-lived individuals (greater than or equal to 95 years) and controls was 2.75 [26]. In two replication studies including both sexes, the odds ratios of the corresponding alleles were 1.26 for a German population and 1.36 for a Chinese population [29,30]. These enriched alleles in long-lived individuals may promote better health and contribute towards extended lifespan by increasing expression or activity of FOXO3A.

The only two genes associated with human longevity that have been replicated in multiple populations are FOXO3A and APOE [11,12,15,26,28–31]. The effect sizes of these two genes for longevity are small with odds ratios of 1.26 and 1.45 for survival to age 100 in replicate studies for FOXO3A and APOE, respectively [10,29]. These genes account for only a small portion of the genetic contribution to longevity measured through family heritability studies [4,5]. Therefore, much of the heritability of lifespan remains to be explained.

### 3. LONGITUDINAL STUDIES OF HUMAN AGEING

In addition to examining differences in lifespan in centenarian studies, important human ageing molecular pathways may be found by examining physiological ageing. Physiological age serves as an indicator of an individual’s general health status and may also serve
as an indicator of remaining healthy lifespan [32]. More important than simply reaching age 100 is the health of the individual. For example, some centenarians appear frail and look like they have lived 100 years whereas others appear more fit and have a physical appearance more like that of a 70 year old. Because the rate of ageing varies immensely among individuals, humans become increasingly different from each other with age. Thus, chronological age fails to provide an accurate indicator of the ageing process. Measuring the physiological age of the whole individual is difficult because tissues age at different rates. Therefore, it is easier to measure physiological age in individual organs or tissues in hopes of understanding the molecular basis of ageing.

Measuring how a tissue changes with chronological age in a population can identify biomarkers that can be used as an index of physiological age [33]. The biomarker can then be used to determine if an individual is physiologically younger or older than his or her chronological age. Determining genes and pathways that associate with the measures of physiological age can reveal molecular processes important for the ageing of particular tissues. Studies in different tissues can be compared to find specific and common regulators of ageing.

Several longitudinal studies have been developed to collect data on many diverse phenotypes. The rationale for these studies is that the ageing process might be better understood by studying the same individuals throughout their lifespan, collecting serial assessments rather than by comparing individuals of different ages from different environments [34]. Along with multiple phenotypic measurements per individual, the cohorts have been genotyped for hundreds of thousands of SNPs across the genome.

The Baltimore Longitudinal Study of Ageing (BLSA) is the longest running scientific study of human ageing in the US [35]. Healthy volunteers aged 18 and older were enrolled in the study starting in 1958. These subjects visit the National Institute on Aging at regular intervals for two days of medical, physiological and psychological testing. Examples of measurements taken include electrocardiograms, memory tests, glomerular filtration rates and various serum metabolite levels [36–39]. Currently, the study population has 1450 active participants, aged 18–97 years (http://www.grc.nia.nih.gov/branches/blsa/blsa.htm).

Another longitudinal study of ageing is the InCHIANTI study, which consists of residents of two small towns in Tuscany, Italy [40]. The study includes 1320 participants (age range 20–102 years), who were randomly selected from the population registry of Greve in Chianti (population 11 709) and Bagno a Ripoli (population 4704) starting in 1998 [40]. Over 90 per cent of the population that were over the age of 65 participated in this study, and thus the cohort is a good representation of normal ageing. Since the baseline measurements were taken in 1998, three sets of follow-up measurements have been collected. The focus of the InCHIANTI study is the decline in mobility that occurs with age. Many physiological measurements are available for researchers including clinical histories, serum metabolite levels, glomerular filtration rates and muscle strength levels (http://www.inchiantistudy.net).

Within the past two years, both the BLSA and InCHIANTI cohorts have been genotyped for approximately 550 000 SNPs, making genome-wide association studies for various age-related traits possible [41,42]. Association studies using longitudinal data to investigate age-related traits possible 41,42]. Association studies using longitudinal data to investigate age-related traits possible 41,42]. Association studies using longitudinal data to investigate age-related traits possible. Table 2. Gene associations with age-related traits found using longitudinal study data.

<table>
<thead>
<tr>
<th>age-related trait</th>
<th>longitudinal dataset</th>
<th>N</th>
<th>gene</th>
<th>SNP</th>
<th>p-value</th>
<th>references</th>
</tr>
</thead>
<tbody>
<tr>
<td>serum iron</td>
<td>BLSA, InCHIANTI</td>
<td>1919</td>
<td>TMPRSS6</td>
<td>rs855791</td>
<td>3.9 × 10⁻⁷</td>
<td>[43]</td>
</tr>
<tr>
<td>serum B6</td>
<td>InCHIANTI</td>
<td>1178</td>
<td>ALPL</td>
<td>rs654748</td>
<td>1.2 × 10⁻⁴</td>
<td>[42]</td>
</tr>
<tr>
<td>serum B12</td>
<td>BLSA, SardiNIA, InCHIANTI</td>
<td>2927</td>
<td>FUT2</td>
<td>rs602662</td>
<td>2.4 × 10⁻¹²</td>
<td>[42]</td>
</tr>
<tr>
<td>plasma arachidonic acid</td>
<td>InCHIANTI</td>
<td>1075</td>
<td>FADS1</td>
<td>rs174537</td>
<td>6.0 × 10⁻⁴⁶</td>
<td>[48]</td>
</tr>
<tr>
<td>plasma eicosapentanoic acid weight</td>
<td>InCHIANTI</td>
<td>1075</td>
<td>FADS1</td>
<td>rs174537</td>
<td>1.1 × 10⁻¹⁴</td>
<td>[48]</td>
</tr>
<tr>
<td>BMI</td>
<td>Framingham heart study</td>
<td>2073</td>
<td>FTO</td>
<td>rs1121980</td>
<td>2.0 × 10⁻⁴</td>
<td>[52]</td>
</tr>
<tr>
<td>glomerular filtration rate</td>
<td>BLSA, InCHIANTI</td>
<td>2190</td>
<td>MMP20</td>
<td>rs1711437</td>
<td>3.6 × 10⁻⁵</td>
<td>[73]</td>
</tr>
</tbody>
</table>

**Table 2.** Gene associations with age-related traits found using longitudinal study data.
SardiNIA cohort, 98 quantitative traits have been measured, including blood chemistry components, cardiovascular traits and personality facets (http://sardinia.nia.nih.gov). This particular genome-wide association study sought to find genetic variants that affect circulating levels of B vitamins. B vitamins have a protective effect against mortality and age-related diseases, like coronary artery disease and colon cancer [46,47]. In this study, serum vitamin B6 levels were significantly associated with SNPs in tissue-nonspecific alkaline phosphatase (ALPL) and serum vitamin B12 levels were significantly associated with SNPs in fucosyltransferase 2 (FUT2) [42]. ALPL, probably influences the catabolism of vitamin B6 while FUT2 interferes with absorption of vitamin B12 [42]. The alleles associated with lower vitamin levels may indicate a risk for vitamin deficiencies and age-related diseases.

In a third genome-wide association study, plasma polyunsaturated fatty acid (PUFA) levels were found to associate with SNPs in the genes FADS1 in the InCHIANTI cohort and the GOLDN replication cohort [48]. GOLDN is the Genetics of Lipid Lowering Drugs and Diet Network study and this population consists of 1120 white men and women from the US [49]. GOLDN is not a longitudinal study, but has measurements of PUFA levels from one time point. FADS1 is a fatty acid desaturase involved in the metabolism of PUFAs. The alleles associated with higher PUFA levels may help protect against age-related diseases. High plasma PUFA concentrations have been shown to reduce the risk of mortality and cardiovascular disease [50].

Genome-wide association models that incorporate information about how an individual's phenotype changes over time have been developed [51]. Instead of simply finding genotype–phenotype associations, longitudinal methods have the potential to reveal alleles that are associated with the rate of ageing. Both the age at which physiological function begins to decline and the rate of such decline varies among individuals. In the BLSA, one third of individuals showed no decrease in glomerular filtration rate measured over a 20-year period, whereas the remainder of the population showed a distinct decline [37]. For those individuals who showed a significant decline in filtration rate, the slope of the decrease varied widely [37].

One association study examined changes in weight and body mass index (BMI), two traits that increase with age, in the Framingham Heart Study cohort [52]. The cohort included 7130 Caucasian individuals each with weight and BMI data at up to four time points. An allele of FTO (fat mass and obesity associated), a gene of unknown function, was previously found to associate with weight and BMI in two cross-sectional genome-wide association studies [53,54]. Luan et al. found that homozygotes for the minor allele of rs1121980 in FTO had the most pronounced increase in weight and BMI [52].

4. TRANSCRIPTIONAL STUDIES OF HUMAN AGEING

To elucidate molecular differences associated with ageing, an attractive approach is to use DNA microarrays to scan the entire genome for genes that change expression with age (age-regulated genes). Changes in gene expression levels may lead to important functional consequences in the process of ageing. Transcriptional profiles of many human tissues, including brain [55,56], blood [57], eye [58], kidney [59,60], muscle [61–63] and skin [64] have been generated. A major aim of these studies is the identification of novel biomarkers that can be used as indices of physiological age. Determining genes and pathways that associate with the measures of physiological age have revealed molecular processes important for the ageing of particular tissues. Studies in different tissues have been compared to find specific and common regulators of ageing. Ageing DNA microarray studies have been reviewed extensively [65,66], but two studies will be highlighted here because they have led to a new type of analysis known as genomic convergence.

In their analysis of kidney ageing, Rodwell et al. showed that age-regulated markers could predict physiological age [60]. They measured levels of gene expression in kidney samples from 72 patients between 27 and 92 years of age, and they identified a set of 447 age-regulated genes. Next, they tested whether the overall level of expression of the 447 genes was a signature of physiological age for different individuals. The relative amount of physiological ageing for each of the patients used in the study was measured by examining the morphological and histological appearance of the kidney. A remarkable similarity was found between the transcriptional ageing profiles determined using chronological age and the physiological states of the kidney in different individuals. Some elderly persons displayed an ageing signature similar to that of people who were 30–40 years younger, and these individuals also had relatively healthy kidneys. Others displayed an ageing signature similar to people who were 10–20 years older, and these patients showed poor kidney physiology for their chronological age.

Genes that are age-regulated in all tissues would reveal genes involved in core mechanisms that underlie cellular ageing. Zahn et al. [63] discovered genetic pathways that show common age regulation in human kidney, brain and muscle. They used microarrays to analyse expression in 81 skeletal muscle samples from patients aged 16–86 years and found 250 age-regulated muscle genes [63]. Similar to the ageing expression profile for the kidney, the overall expression behaviour of this set of age-regulated muscle genes correlated with the physiological as well as chronological age of the muscle sample. Next, they compared their muscle-ageing results to previously published data on kidney and brain ageing of similarly large sample size [56,60]. Although most of the age-related changes were tissue specific, they found evidence for common age regulation of six genetic pathways in all three tissues. Specifically, there is an overall increase in expression of the extracellular matrix genes, the ribosomal genes, the cell growth genes and the complement activation genes in all three tissues. Increased overall expression of the extracellular matrix and complement activation gene sets with advancing age may contribute to widespread

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fibrosis and inflammation in the elderly. There is an overall decrease in expression of the chloride transport genes and the electron transport genes in all three tissues. Decreased overall expression of electron transport chain genes with age might support the mitochondrial free-radical theory of ageing [67], as free-radical generation by mitochondria would preferentially damage the electron transport chain protein complexes. Decreased expression of the electron transport genes (encoded in the nucleus) might be caused by feedback regulation from damage to the electron transport chain protein complexes [63]. However, it is also possible that increased oxidative damage occurs as a consequence of the decreased expression of the electron transport chain genes. In addition, an increasing number of studies in model organisms have critically challenged the mitochondrial free-radical theory of ageing [68].

Importantly, these comparisons of transcriptional networks among different tissues were not done at the single gene level, but rather at the genetic pathway level [63]. With this approach, known as gene set enrichment analysis (GSEA), age regulation for every gene in a pathway is combined to determine whether there is an overall regulation of the entire pathway [69]. Screening for coordinated age regulation of genetic pathways increases the sensitivity of the analysis, as the combined effects of small regulation of many genes in a pathway can be significant. This systems biology method is especially powerful in studies of ageing because of the polygenic nature of the phenotype.

5. GENOMIC CONVERGENCE IDENTIFIES A GENE INVOLVED IN KIDNEY AGEING

In genome-wide association studies, hundreds of thousands of SNPs are tested. Thus, the penalty for multiple hypothesis testing is a large obstacle to overcome. A powerful alternative to genome-wide association studies is genomic convergence, which selects candidate genes for a specific phenotype based on genome-wide expression studies [70–72]. Gene expression microarrays can be used to identify genes whose expression levels increase or decrease in affected individuals compared to controls. Differential expression between cases and controls may indicate that the gene is functionally involved in disease pathogenesis. For example, a set of genes that is overexpressed in a type of neural disorder may include the genes that cause that disorder. Therefore, the genes that change expression between cases and controls can be used as candidates in genetic association studies. The power of the genetic association study is increased because a subset of genes rather than the whole genome is tested. Genomic convergence was first used to identify genes associated with Parkinson’s disease, schizophrenia and Alzheimer’s disease [70–72].

In a recent study, the genomic convergence approach was used to find genes associated with kidney ageing [73]. This genomic convergence approach consisted of three steps (figure 1). First, a genome-wide transcriptional profile of ageing in the human kidney gave an unbiased view of gene expression changes that occur with age [60]. These age-regulated genes also associated with kidney physiological age [60]. Additional genes from age-regulated pathways were included in the subsequent analysis [63]. The 630 age-regulated genes were probably enriched for genes that contribute to kidney ageing. The second step of the genomic convergence approach was mapping of expression quantitative trait loci (eQTLs). If a gene is functionally involved in kidney ageing and if DNA differences in the gene cause variation in expression among individuals, then there may be an association between the specific allele carried by an individual and that individual’s physiological ageing trajectory. Wheeler et al. found 101 genes that have SNPs that associate with different expression levels [73]. In the third and final step, the 101 genes with eQTLs were tested for association with glomerular filtration rate (GFR), a phenotype of kidney ageing. Data from the BLSA and InCHIANTI cohorts were used to test SNPs in the 101 eQTLs for association with GFR. SNPs in one of these genes, matrix metalloproteinase 20 (MMP20), showed a statistically significant association with normal kidney ageing. Matrix metalloproteinases degrade extracellular matrix proteins and changes in the extracellular matrix play a key role in ageing of the kidney [74]. Interstitial fibrosis occurs during ageing in the kidney because of an increase in matrix [75] and different activities of MMP20 may play a role in this process. However, this genetic association has not yet been replicated in additional populations. Similar genomic convergence approaches that combine multiple types of genomic information (i.e. gene expression, eQTLs, protein levels) with genetic association studies can be used for all complex phenotypes, including other ageing-related phenotypes to find genes and pathways that affect ageing.

Figure 1. The genomic convergence approach. Genes are filtered in each step for those the most likely to be functional. In the first step, a genome-wide transcriptional profile defines genes that show age-related changes in expression. In the second step, SNPs are tested for association with the expression of age-regulated genes. In the final step, the expression quantitative trait loci (eQTLs) are tested for association with a phenotype of ageing (e.g. age-dependent decline in kidney glomerular filtration rate in Wheeler et al. [73]).

Genome-wide transcriptional profile of ageing in relevant tissue
(kidney example)

Expression QTL mapping of age-regulated genes
(630 genes in kidney)

test eQTLs for association with age-related trait
(2038 eSNPs)

Ageing genes
(MMP20)
6. FUTURE DIRECTIONS IN THE GENETICS OF HUMAN AGEING

The application of powerful new human genetics technologies to the study of ageing has just begun. Unlike most other biological processes, the genetic factors that influence ageing may not be evolutionarily conserved because wild animals usually die from predation and infection, not ageing [76]. Thus, animal models for ageing may have limited relevance to human ageing. Therefore, genetic studies of ageing in humans are necessary. Luckily, the field of human genetics has experienced rapid advancement. It is now possible to interrogate thousands of samples across the genome in order to find genes associated with a particular phenotype.

Finding new human ageing genes contributes to our understanding of molecular mechanisms underlying the human ageing process. Like other complex human traits, finding common variants that account for the entire genetic component of human lifespan variability has proved difficult. If rare variants rather than common variants explain most of the genetic variation in ageing among humans, new genotyping techniques and new analysis methods must be developed to find genes and pathways involved in ageing. Next-generation sequencing technologies are faster and cheaper than traditional capillary sequencers. Several complete genomes have been sequenced using these next-generation technologies and many rare variants have been identified [77, 78].

Methods are needed that can combine rare variants from multiple individuals that are in the same gene into a score that can be tested for association with different phenotypes of ageing and longevity. Information about how that rare variant might affect function could be included in the score. This information could include whether the rare variant causes a stop codon, a nonconservative amino-acid change or affects expression. Several association test methods for rare variants using this logic have been proposed that could be used for any complex trait including ageing [79–81]. In addition to scoring individual genes, combining information from molecular pathways may elucidate additional associations. If differences in a particular gene affect a phenotype, it is probable that other genes in the same pathway also affect the phenotype. Therefore, rare variants in different genes of the same pathway could be combined to test for association of the pathway with an ageing trait of interest. New technologies and many rare variants have been identified [77, 78].

Once key ageing genes and pathways are found, an individual’s genotype could predict the rate of overall health decline. Genotypes of tissue-specific ageing genes could predict which tissues are likely to decline rapidly and which are likely to decline more gradually. Among young individuals, an unfavourable SNP genotype may indicate risk for rapid decline in a particular organ or tissue and this information could be extremely useful to identify patients who may require early intervention. Among older individuals, a favourable SNP genotype may indicate that they may still be eligible as organ donors even though they are currently over the upper age limit. For some ageing genes, one allele may adversely affect tissue function in old age because it increases the activity of the gene above a healthy level. In these cases, one could develop a drug to target the gene or pathway in individuals carrying the overactive allele, and thereby preserve function in old age. The goal of human ageing research is not necessarily to extend lifespan, but instead to extend the healthy years of life.

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