Gender, aging and longevity in humans: an update of an intriguing/neglected scenario paving the way to a gender-specific medicine

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Abstract

Data showing a remarkable gender difference in life expectancy and mortality, including survival to extreme age, are reviewed starting from clinical and demographic data and stressing the importance of a comprehensive historical perspective and a gene–environment/lifestyle interaction. Gender difference regarding prevalence and incidence of the most important age-related diseases, such as cardiovascular and neurodegenerative diseases, cancer, Type 2 diabetes, disability, autoimmunity and infections, are reviewed and updated with particular attention to the role of the immune system and immunosenescence. On the whole, gender differences appear to be pervasive and still poorly considered and investigated despite their biomedical relevance. The basic biological mechanisms responsible for gender differences in aging and longevity are quite complex and still poorly understood. The present review focuses on centenarians and their offspring as a model of healthy aging and summarizes available knowledge on three basic biological phenomena, i.e. age-related X chromosome inactivation skewing, gut microbiome changes and maternally inherited mitochondrial DNA genetic variants. In conclusion, an appropriate gender-specific medicine approach is urgently needed and should be systematically pursued in studies on healthy aging, longevity and age-related diseases, in a globalized world characterized by great gender differences which have a high impact on health and diseases.

Key words: aging, centenarians, gender, gender-specific medicine, gut microbiome, longevity, mitochondrial DNA, X chromosome inactivation.

INTRODUCTION

Lifespan and longevity are complex and multifactorial traits resulting from an intriguing combination of ‘Nature’ and ‘nurture’, the unique reciprocal interaction between environmental, genetic, epigenetic and stochastic factors, each contributing to the overall phenotype [1,2].

Women live longer than men and this difference in life expectancy is a worldwide phenomenon indicating that human longevity seems strongly influenced by gender defined as the combination between biological sexual characteristics (anatomy, reproductive functions, sex hormones, expression of genes on the X or Y chromosome) and factors related to behaviour, social role, lifestyle and life experiences [3–6].

Following a historical perspective, in Europe in the 19th Century, life expectancy was less than 40 years and longevity of the two genders was generally very similar. The high female mortality due to pregnancy and childbirth corresponded to a higher male mortality from causes related to work, accidental injury or violence. Moreover, infectious and communicable diseases...
affected and killed men and women almost equally [7]. Throughout the 20th Century, mortality became concentrated in the older ages, non-communicable diseases became the prevailing causes of death, and a female survival advantage emerged and grew. This divergence in life expectancy can partly be explained by the declining rates in maternal mortality; however, a major contribution is due to differences in behaviour and biology between males and females [8].

Using historical data from 1763 birth cohorts from 1800 to 1935 in 13 developed countries, Beltrán-Sánchez et al. [9] showed that gender asymmetry emerged in cohorts born after 1880, that excess adult male mortality is rooted in a specific age group (50–70) and that heart disease is the main condition associated with increased excess male mortality in birth cohorts of 1900–1935. The authors have suggested that excess male mortality, found even after accounting for smoking-attributable deaths, may be explained by underlying traits of vulnerability to CVD (cardiovascular disease) that emerged with the reduction of infections and changes in diet and other lifestyle factors [9].

The maximum difference in life expectancy between males and females was found between the 1970s and the 1990s. The subsequent reduction of the gender gap can be attributed partly to the narrowing of differences in risk behaviours between males and females, along with the decline in mortality rates from CVD among men [10]. In the EU-28 countries, the difference in life expectancy between males and females was found between the 1970s and the 1990s. The maximum difference in life expectancy between males and females was 5.5 years in 2013 (http://ec.europa.eu/eurostat/statistics-explained/index.php/Mortality_and_life_expectancy_statistics); however, the gender gap varied largely across EU member states.

The survival advantage of women is counterbalanced by a worse quality of life in advanced age due to the increase in disability and degenerative diseases [12]. Therefore men and women have a diverse chance to attain longevity and, at the same time, the aging process is qualitatively different between genders.

The impact of gender difference in aging has been extensively assessed, but the study of the interaction between a series of fundamental aspects such as hormonal, immunological and metabolic pathways as well as genetic background remains largely unknown.

Accordingly, the present review aims to (i) give an accurate analysis of mortality causes and age-related diseases pattern in men and women; (ii) describe the most important mechanisms underpinning the gender difference in longevity and aging (sex hormones, immunity, genetic factors, nutrition and stress); (iii) attempt to explain the difference in longevity between males and females, in human models of extreme longevity such as centenarians and long-lived families, suggesting the importance of an integrated investigation of nuclear, mitochondrial DNA genetics and gut microbiome; and (iv) stress the urgent need for a gender-specific medicine, taking into account the profound differences in pathophysiological pathways, in clinical characteristics and in pharmacological response between men and women. In conclusion, the scientific world is obliged to revise all outcomes in all fields of medicine on the basis of gender differences.

### GENDER AND AGE-RELATED DISEASES

The epidemiology of age-related diseases is substantially different between genders and changes dramatically in women after menopause [13]. Table 1 reports mortality data by sex regarding the 14 leading causes of death in U.S.A. in 2013 and refers to all races and ages [14]. Women died at higher rates than men of chronic lower respiratory diseases, cerebrovascular diseases, AD (Alzheimer’s disease), influenza and pneumonia, septicaemia and hypertension-related diseases [14]. Even in the EU, a significant gender gap exists in mortality rates in all countries.

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Table 1: Number of deaths, percentage of total deaths by sex regarding the 14 leading causes of death in U.S.A. in 2013

| Rank | Cause of death                  | Male Number | Male Total deaths (%) | Female Number | Female Total deaths (%) | Higher mortality for |
|------|--------------------------------|-------------|-----------------------|---------------|-------------------------|----------------------|
| 1    | Heart disease                  | 321,347     | 24.6                  | 289,758       | 22.4                    | ♂                     |
| 2    | Malignant neoplasms            | 307,559     | 23.5                  | 277,322       | 21.5                    | ♂                     |
| 3    | Chronic lower respiratory diseases | 70,317     | 5.4                   | 78,888        | 6.1                     | ♀                     |
| 4    | Accidents (unintentional injuries) | 81,916     | 6.3                   | 48,641        | 3.8                     | ♂                     |
| 5    | Cerebrovascular diseases        | 63,691      | 4.1                   | 75,287        | 5.8                     | ♀                     |
| 6    | Alzheimer’s disease            | 25,836      | 2.0                   | 58,931        | 4.6                     | ♀                     |
| 7    | Diabetes mellitus              | 39,841      | 3.1                   | 35,737        | 2.8                     | ♂                     |
| 8    | Influenza and pneumonia        | 26,804      | 2.1                   | 30,175        | 2.3                     | ♂                     |
| 9    | Kidney diseases                | 23,493      | 1.8                   | 23,619        | 1.8                     | =                     |
| 10   | Suicide                        | 32,055      | 2.5                   | 9,094         | 0.7                     | ♂                     |
| 11   | Septicaemia                    | 17,994      | 1.4                   | 20,162        | 1.6                     | ♀                     |
| 12   | Chronic liver disease and cirrhosis | 23,709     | 1.8                   | 12,718        | 1.0                     | ♂                     |
| 13   | Essential hypertension-related diseases | 12,963 | 1.0                   | 17,807        | 1.4                     | ♀                     |
| 14   | Parkinson’s disease            | 15,088      | 1.2                   | 10,108        | 0.8                     | ♂                     |
| 15   | All other causes               | 253,421     | 19.4                  | 302,707       | 23.5                    | ♀                     |
In particular, death rates for IHD (ischaemic heart disease) and stroke are higher for men than for women [10]. There are important inequalities in healthy life years between men and women. In EU countries, life expectancy at age 50 reached 29.8 years for men and 34.6 years for women in 2010, but the average duration of life free from activity limitation remained practically the same in women (68.6 years) and men (67.9 years) [10,15,16], meaning that the almost 5 years of advantage in life expectancy of women are years of diseases and disability.

**Cardiovascular disease (CVD)**

Differences between women and men in the epidemiology, pathophysiology and symptoms of CVD are well-described. This gender gap should be taken into account because it strongly impinges on the effects of specific drugs and outcomes. Both factors linked to sex (gene expression from the sex chromosomes, sex hormones, metabolism of drugs by sex-specific cytochrome expression) and gender (sociocultural processes, behaviours, exposure to specific environment, nutrition, lifestyle and attitudes towards treatments and prevention) play a fundamental role in determining CVD risk [17].

Death rates for IHD are 70% higher for men than for women on average in all EU countries [10]. In addition, women showed a delayed onset of IHD (7–10 years on average) in several western EU states, even though, due to harmful lifestyle modifications, the prevalence of IHD is increasing in young women [17]. Moreover, IHD in women may show different symptoms and pain localization, and may need diverse diagnostic procedures and drugs [18].

HF (heart failure) is one of the major health threats of Western societies and affects up to 10% of the elderly, in absolute numbers more women than men [19]. However, women survive better than men and HF in women frequently occurs at older age and with less ischaemic aetiology than in men [20]. Recent data from the Framingham Heart Study showed that, in the latter half of the 20th Century, incidence of HF has declined by about one-third in all EU countries, even though, due to harmful lifestyle modifications, the prevalence of IHD is increasing in young women [17]. Moreover, IHD in women may show different symptoms and pain localization, and may need diverse diagnostic procedures and drugs [18].

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The difference in the epidemiology of hypertension between men and women deeply changes with age. In particular, hypertension has a low prevalence in young and adult age when it is more predominant among men. By contrast, hypertension is more common in women than in men in the elderly population [17]. Indeed, falling oestrogen production during and after menopause has been associated with hypertension in women [22].

During aging, there is a sex-specific ‘cardiac remodelling’. In particular, women develop more frequently concentric cardiac hypertrophy with smaller internal cavity and relatively larger wall thickness, preserving a better ejection fraction and myocardial contractility than men. On the other hand, men show more frequently eccentric hypertrophy leading to an increased stroke volume and dilatation [17].

Women are particularly susceptible to the deleterious impact of T2D (Type 2 diabetes) and hypertension on cardiovascular health. These conditions were associated with higher risk of HF in women with respect to men (T2D increases 3.4-fold and 2-fold the risk of HF in women and men respectively; hypertension increases 5-fold and 2-fold the risk of HF in women and men respectively) [21]. In addition, T2D worsens the coronary artery disease outcome more in women than in men [23]. Finally, some pregnancy-associated conditions, such as pre-eclampsia and other hypertensive disorders, further contribute to increased risk for future chronic hypertension, CVD, cerebrovascular diseases and death in women [24].

**Type 2 diabetes (T2D)**

Recent data show that the difference in the global estimates of T2D between men and women in terms of cases (male, 197.7 million; female, 184.1 million), prevalence (male, 8.7%; female, 8.1%) and age-specific prevalence [25] is small. Even if T2D prevalence is similar in men and women, it is slightly higher in men under 60 years of age and in women at older ages. Indeed, the longer survival of women is one of the factors leading to a higher prevalence of diabetes for women than for men at advanced age [26]. However, a stronger connection between diabetes and coronary heart disease has been demonstrated in women. In particular, the relative risk for mortality due to coronary heart disease in diabetic patients is 50% higher in women than in men, suggesting that T2D may induce a more unfavourable cardiovascular risk profile among women. Diabetic women have significantly higher levels of blood pressure and lipids than men with diabetes [27]. Moreover, findings from different studies conducted in the U.K. and the U.S.A. showed that the greater coronary risk associated with T2D observed in women may reflect a treatment bias that favours men. In particular, in these countries, diabetic men with cardiovascular problems are more frequently treated with hypoglycaemic drugs, aspirin, statins or anti-hypertensive drugs than women with similar pathological conditions [28–30].

Moreover, women over 65 years of age have a higher frequency of insulin resistance, dyslipidaemias, central adiposity and hypertension (named the metabolic syndrome) which in turn is a greater risk factor for CVD in women [31,32]. In particular, central adiposity tends to be more pronounced in postmenopausal women than in men playing a determinant role in the increase in CVD risk. It has been demonstrated that visceral adipose tissue contributes to insulin resistance secreting a variety of inflammatory mediators [such as IL (interleukin)-6, TNF-α (tumour necrosis factor α), leptin and resistin]. Moreover, lipid profile [HDL- (high-density lipoprotein) and LDL (low-density lipoprotein)-cholesterol as well as triacylglycerols] dramatically worsens after the menopause favouring atherosclerosis [32].

**Cancer**

Cancer mortality rates are higher for men than for women in industrialized countries. In some EU countries (i.e. Lithuania, Spain, Latvia, Estonia, the Slovak Republic, Portugal and Croatia), the mortality rates for neoplasms in men are dramatically increased. This gender difference can be explained partly by the greater prevalence of risk factors among men as well as by reduced availability/use of screening programmes for cancers affecting men, leading to lower survival rates after diagnosis [10]. For instance, lung cancer accounts for the greatest number of cancer deaths among men in almost all EU states. In 2011,
death rates from lung cancer among men were the highest in all EU countries where smoking habits among men remain very frequent (Hungary, Poland and Croatia). However, lung cancer mortality in American women has increased from 1950 to 1995 by 500% [18]. Evidence has suggested that the development of lung cancer is different in women in comparison with men. Non-smoking women have a 2.5-fold higher risk than men to develop lung cancer at a younger age, but they respond better to treatment. Women who smoke have a higher susceptibility to cigarette-smoking damage probably related to the polymorphism of Glutathione S-transferase (GST) Mu 1, which plays a role in detoxifying environmental carcinogens [33]. However, women with lung cancer survive longer than men, regardless of therapy and stage.

CRC (colorectal cancer) is the second leading cause of cancer death in both genders; in women it occurs 5 years later than in men. For this reason, the population screening for CRC should be extended beyond 70 years of age. Moreover, CRC in women is more often located in the right colon, the histology is mucinous, occult blood in stool may be negative until the last stages and it is frequently diagnosed in an urgent/emergency situation. Nevertheless, the survival is better in female patients with respect to male patients [34]. CRC in women more frequently expresses microsatellite instability showing a lower sensitivity to fluoropyrimidines, cornerstone drugs for the treatment of colorectal carcinoma [35].

Currently, prostate cancer has become the most common cancer among men after skin cancer in the majority of EU countries, particularly among men aged 65 years and over. However, death rates for prostate cancer are lower than for lung cancer. The primary risk factors are obesity, lack of exercise, age and family history [10].

Breast cancer is the second most common form of cancer in women after skin cancer in all EU countries. It can occur in both men and women, but it is very rare in men (10%). The incidence rates of breast cancer have increased in the last decade, but the death rates have diminished or remained stable, indicating an improving of survival rates due to earlier diagnosis and better treatment [36]. Numerous risk factors for breast cancer in women have been identified, including age, personal history of certain benign breast diseases or breast cancer, early menstrual or late menopause, never having been pregnant or having a first pregnancy after age 30, use of oral contraceptives, family history of breast cancer, presence of certain genetic mutations (BRCA1 and BRCA2), history of radiation therapy to the chest, long-term use of combined hormone therapy, use of DES (diethylstilbestrol), increased breast density, alcohol use and obesity after menopause. Risk factors for men for breast cancer include obesity, Klinefelter’s syndrome and an excess of breast tissue (http://www.cancer.gov/research/progress/snapshots/breast).

Neurodegenerative diseases

Women are more affected than men by dementia (definition comprising different conditions including AD and vascular dementia) showing a more frequent and rapid decline of cognitive function with aging. Prevalence rates among populations vary considerably because of methodological reasons (diagnostic criteria, sampling strategies and statistical analysis) [38]. Among people aged 90 years and over, the gender gap rises to 30% of prevalence for men and 47% for women [10].

The biological basis of gender impact on AD and neurodegeneration are still unclear. Indeed, the development and functioning of the central nervous system is strongly influenced by gender. The main risk factor for AD is age, and the fact that the majority of AD patients are females has been attributed to longer life expectancy. However, women are reported to have higher rates of AD than men, even after adjusting for survival [39,40]. The negative effect of the APOE (apolipoprotein E) ε4 allele, one of the most established genetic risk factors for AD, may explain, at least in part, this gender gap. Different studies have observed that female APOE ε4 carriers show a higher risk of AD compared with males [41,42]. A recent paper demonstrated that female APOE ε4 carriers presented widespread brain hypometabolism and cortical thinning compared with female non-carriers, whereas male APOE ε4 carriers showed only a small cluster of hypometabolism and regions of cortical thickening compared with male non-carriers, suggesting that the impact of APOE ε4 on brain metabolism and structure is strongly dependent on gender [43].

AD can be caused by defects in mitochondrial oxidative phosphorylation. Given that the mitochondrial genome (mtDNA) codes for polypeptides that are essential components of the respiratory chain, a number of studies have investigated the association between mtDNA-inherited variants and AD. In particular, research conducted on AD patients and controls from Italy has identified the sub-haplogroup H5 as a risk factor for AD for females in particular and independently of the APOE genotype [44].

It is also worth noting that sex hormones have a critical role in neurodegeneration processes. Oestrogen has been shown to be protective towards AD reducing amyloid β-peptide aggregation and improving neural functions [45–47]. During aging, the decrease in gonadal hormones production is gradual in men (testosterone), whereas in women, the fall of oestrogen is quick after menopause when the incidence of AD suddenly increases [39,48].

A neuroprotective effect of oestrogen on the risk of PD (Parkinson’s disease) onset and disease progression has also been reported. Both the prevalence and the incidence of the PD is higher in men than in women [49,50]. In women, the risk of PD is related to the fertile lifespan considering that a later age at menopause is associated with a later age at onset of PD [51,52], whereas a premature menopause increases the risk of PD [53]. These data suggest a relationship between the duration of endogenous oestrogen exposure and the susceptibility to develop PD in women.

Disability

It is important to underline that women pay for their survival advantage with a worse quality of life in their old age due to an increased prevalence of a variety of disabling non-lethal pathological conditions [15].

Diseases influencing the ADL (Activities of Daily Living) and IADL (Instrumental Activities of Daily Living) scales in women are the consequences of CVD, osteoarthritis, osteoporosis and cognitive decline. Women are more medicalized in terms of
frequency of medical visits, days of hospitalization and number of drugs routinely administered [13,54]. A recent Italian study on a cohort of hospitalized elderly patients (REPOSI) describes a gender dimorphism in the demographic and morbidity profiles as well as in the overall medication pattern of hospitalized elderly people [55,56]. In all EU countries, women reported a poorer self-perceived health, more long-standing illnesses and/or more health problems than men [10]. A possible explanation for this gender-associated health–survival paradox may be found in a higher female sensibility to physical discomfort that led the women to seek medical attention more frequently. Actually, the higher prevalence and severity of arthritis and musculoskeletal disease among older women widely contributes to their worse health and functional status. In particular, women are more frequently affected by severe forms of osteoarthritis affecting the hand, foot and knee, and the incidence of this condition highly increases at the time of menopause, suggesting a role for oestrogens in the pathogenesis of osteoarthritis. Moreover, gender disparities may also be caused by differences in bone strength, posture, ligament laxity, pregnancy and neuromuscular strength [57,58].

Stress and spousal bereavement
Owing to their longer life expectancy and the tendency to marry older men, women are more likely to become widows. Conjugal loss in advanced age is a stressful life experience able to drastically alter the social environment of the surviving spouse. Therefore widowhood is often associated with a feeling of loneliness, depression, loss of physical and cognitive functions, and poor nutritional status [59–62]. However, spousal bereavement may not have the same implications for women and men. For example, widows maintain higher levels of social contacts with family, friends and neighbours [63] than widowers and this behavioural difference may alleviate some of the undesirable effects of widowhood. Thus, even if widows are more numerous than widowers, it has been shown that widowhood has a more negative impact on health status and mortality in men than women [62,64–66].

GENDER AND THE IMMUNE SYSTEM
Much research has been carried out into the role of sex hormones in determining lifespan [67] and one hypothesis is that sex hormones appear to influence the immune system. This can determine a sexual dimorphism in the immune response in humans [68]. For instance, females produce more vigorous cellular and humoral immune reactions and are more resistant to certain infections. In contrast, men are more susceptible to many illness caused by viruses, bacteria, parasites and fungi. It is well known that oestrogens, androgens and progesterone affect cells of the innate and adaptive immune system differently during the reproductive phase of life [69]. Oestrogens inhibit NK (natural killer) cell cytotoxicity, reduce neutrophil chemotaxis and consequently inflammation [70,71]. Moreover, macrophages treated in vitro with oestradiol display a reduced production of pro-inflammatory cytokines, i.e. IL-1β, IL-6 and TNF-α [72]. Oestrogens and androgens are responsible for a reduced immature number of T-lymphocytes and thymus involution after puberty [73] and can also influence the adaptive immunity in an opposing way. Androgens polarize naïve CD4+ T-cells towards the Th1 subset and activate CD8+ T-cells; conversely, oestrogens stimulate Th2 responses and activate antibody production [74]. Testosterone increases IL-10 production, and men with androgen deficiencies have higher levels of IL-1β, IL-2 and TNF-α, higher antibody titres and higher CD4+/CD8+ T-cell ratios [75]. Oestradiol reduces the apoptosis of immature B-cells and also increases somatic hypermutation and isotype-switch recombination leading to high-affinity Ig-producing cells. These effects might contribute to an improved humoral response in women, but also favour the appearance of autoreactive clones and the susceptibility to autoimmune diseases [76]. Moreover, oestrogens down-regulate autoimmune regulator gene (AIRE) expression in mTECs (medullary thymic epithelial cells), that plays an important role in protection against autoimmunity, triggering the negative selection of self-reactive T-cells [77]. In addition AIRE induces Treg (regulatory T-cell) development; consequently oestrogens contribute to increased susceptibility to autoimmunity [77]. Several studies showed that females are 2–10-fold more susceptible than males to a series of disabling autoimmune diseases such as rheumatoid arthritis, multiple sclerosis, systemic rheumatoid arthritis, systemic lupus erythematosus, myasthenia gravis, Sjögren’s syndrome and Hashimoto’s thyroiditis [78–80]. The better immune response of females is also evident after vaccinations; women reveal higher levels of immunoglobulins and seroconversion and lower rates of disease [75]. In short, sex hormones have different effects on immune responses, with oestrogens exerting an immune-improving action, and progesterone and testosterone having an immune-suppressive effect.

The sudden loss of ovarian oestrogen and progesterone production that characterizes menopause, induces pathophysiological changes in different organs and systems [81]. Menopause reflects the inevitable final hallmark of a woman’s fertile lifespan and of the above-described beneficial effects of oestrogens on immune responses. Menopause affects various women’s health aspects, including bone density, breast cellular composition, cardiovascular health, mood/cognitive function and sexual wellbeing. Moreover, old women lose their immunological privilege towards infection [69] because the rapid reduction of oestrogen levels results in an increased susceptibility and mortality towards a series of infectious diseases (hepatitis, and meningococcal and pneumococcal infections) [69,79]. It is noteworthy that women will soon spend half of their life in post-menopause, if the current trend of increasing human life expectancy should persist. Various studies have reported an association between late-onset menopause and reduction in all causes of morbidity and mortality [82]. Both fecundity at an older age and a high age at menopause have been associated with longevity [83]. Several studies have suggested that ovarian sex steroid loss favours immunosenescence by contributing to the remodelling of the immune system. Immunosenescence is a multifaceted phenomenon that increases morbidity and mortality due to infections and age-related pathologies, and is characterized by changes in innate and adaptive immune responses to foreign antigens [84,85]. In Figure 1, the main aspects of immunosenescence [86–92] are shown and it is indicated that
Figure 1  The progression to immunosenescence characterized by age-related changes in immune cells and inflammatory mediators is faster in men than in women [86–93]

TLR, Toll-like receptor.

age-related changes in immune cells and inflammatory mediators, i.e. the progression to immunosenescence, are faster in men than in women [93].

Functional aspects of age/gender-specific differences of the immune system and its interplay with changing sex steroid hormone levels have not been investigated extensively. Post-menopausal women exhibit a reduced number of total lymphocytes, mainly B- and CD4+ T-lymphocytes [94] and an altered expression of inflammatory mediators such as an increased plasma level of IL-1β, IL-6, IL-10 and TNF-α [95–98]. After a transient rise in post-menopausal women, IFN-γ (interferon γ) levels gradually decrease with age. Yet the production of IL-10 increases during the post-menopausal period [95]. Moreover, in \textit{in vitro} stimulation studies, IFN-γ and IL-17 secretion is diminished in aged men in comparison with women [99]. In contrast, the anti-inflammatory cytokine IL-10 increases in aged women but not in men. Centenarians, mainly females, present markers of inflammation (e.g. increased plasma levels of IL-6 and CRP (C-reactive protein) and hypercoagulable state), but do not suffer most of the detrimental effects of inflammaging. Accordingly, centenarians seem to be equipped with gene variants that allow them to optimize the balance between pro- and anti-inflammatory molecules, thus minimizing the effects of the lifelong exposure to environmental insults and stressors [100].

GENDER AND NUTRIENT-SENSING PATHWAYS

DR (dietary restriction) without malnutrition, intended as a reduced intake of all dietary constituents except vitamins and minerals, is a well-known intervention to improve most aspects of health during aging and to extend lifespan in model organisms from invertebrates and rodents to primates, including humans [101]. However, in humans, this practice remains difficult, if not impossible, to sustain because it envisages unrealistic levels of self-deprivation, can impair reproductive function and libido, resistance to infection and wound healing, and can increase the risk of osteoporosis and fractures, anaemia and cardiovascular diseases [101]. Therefore interest in interventions able to recapture the beneficial effects of DR has grown. Among the mechanisms mediating the effects of DR, particular attention has to be paid to nutrient-sensing pathways, such as IIS (insulin and IGF-1 signalling) by their transcription factor FOXO (forkhead box O) or via mTOR (mammalian target of rapamycin), which are considered key modulators of lifespan and the aging process [102,103]. These highly conserved pathways are designated to couple nutritional status to energetically expensive processes, such as growth, reproduction and metabolism [104]. Several studies on experimental animal models have tried to disentangle the effect of IIS/TOR (target of rapamycin) signalling network on biological processes. Specifically, interventions aimed at the down-regulation of this pathway affect the expression of hundreds of genes involved in immunity and stress responses, activate anti-aging responses and are able to extend lifespan mimicking the action of DR.

On the whole, data on animal models have shown that genetic mutations inhibiting IIS and TOR nutrient sensing signalling have a stronger effects on lifespan extension in females [101]. For example, \textit{Drosophila} mutants with impaired insulin-like signalling have a significant life extension in females [105,106] and heterozygous IGF-1R (IGF-1 receptor)-knockout female mice are long-lived and show a higher oxidative stress resistance than wild-type mice, whereas the difference is not significant in males.
and more naive T-cells, lower plasma leptin levels and fat mass) in a number of age-sensitive biomarkers of aging (fewer memory leads to a significantly increased lifespan and to an improvement component of the nutrient-responsive mTOR signalling pathway, increases the lifespan of genetically heterogeneous UMHET3 mice more in females than in males at each dose evaluated [109].

In humans, IIS and mTOR signalling have been investigated for their role in the development of diseases, such as diabetes and cancer, and for their impact on longevity [110]. Large cohort studies have shown a significant interaction with gender. For example, genetic variation in IIS pathway components [GHRHR (growth hormone-releasing hormone receptor), GH (growth hormone), IGF-1, insulin, IRS1 (insulin receptor substrate 1)] have a higher influence on body size and are more beneficial for old age survival in women with respect to men [111]. Some human studies have investigated the role of sex hormones in regulating the somatotropic axis (GH and IGF-1) underlining gender differences in the impact of suppression of the nutrient-sensing pathways on aging and longevity. For instance, oestradiol reduces hepatic sensitivity to GH, whereas testosterone plays an opposite role enhancing the growth-promoting effects of the somatotropic axis [13,112,113] and increasing the risk of some age-related pathologies such as prostate cancer and cardiomyocyte hypertrophy [114,115].

To date, there is a lack of data on the effect of DR on human longevity and whether this practice has a different impact according to the gender is largely unknown. However, increasing interest has been paid to trials on the effects of IF (intermittent fasting) or adjusted rhythm of feeding on women’s health. A study in overweight or obese pre-menopausal women has demonstrated that IF (two non-consecutive days per week over a 6-month period) is an effective intervention to reduce weight, fat mass and waist circumference as well as to improve insulin sensitivity and other biomarkers such as total and LDL-cholesterol, triacylglycerols, CRP and arterial blood pressure [116]. Therefore IF may be considered as an alternative and more feasible practice than DR to reduce disease risk [116]. Interestingly, a ‘breakfast diet’ (980 kcal breakfast, 640 kcal lunch and 190 kcal dinner; 1 kcal = 4.184 kJ) on lean women with polycystic ovary syndrome improves glucose metabolism, decreases free testosterone and increases the ovulation rate with respect to an isocaloric ‘dinner diet’ (190 kcal breakfast, 640 kcal lunch and 980 kcal dinner) [117].

However, to date, few studies have assessed the differences between men and women in response to nutritional interventions. Several papers have described the effects of a 4-week fully controlled isoenergetic Mediterranean diet on a group of 38 men and 32 pre-menopausal women (24–53 years). The results have shown an improvement in lipid profile, cardiovascular risk and inflammation markers which was significant in both genders [118,119]. Such a short-term consumption of Mediterranean diet significantly ameliorates insulin homoeostasis [120], leads to a favourable redistribution of LDL subclasses [121] and reduces adiponectin levels [122] only in men. The greater improvements in dietary intakes obtained in men with respect to women can explain, at least in part, these gender-related responses [123], but it is worth considering that gender differences in the remodelling, distribution and secretory activity of adipose tissue as well as the levels and ratio of androgenic and oestrogenic steroids may play a fundamental role in metabolism homoeostasis. These data underline the importance of considering gender in further studies evaluating the effects of dietary intervention on diseases, aging and longevity taking into account that men and women can show very different responses and require personalized treatments.

**HUMAN POPULATION MODELS TO STUDY GENDER EFFECT ON AGING AND LONGEVITY**

The particular combination of genetic, environmental, historical, anthropological, socio-economic and cultural factors as well as geographical origin could contribute to the longer female life expectancy worldwide. To increase our knowledge on these aspects, the model of centenarians could represent a useful approach. These extraordinary individuals (mostly women) are characterized by a peculiar and heterogeneous phenotype embodying the best example of longevity and successful aging. Most of them have survived, escaped or delayed the onset of major age-related diseases [124–126]. Centenarians are the outcome of a number of biological processes that exert their effects lifelong, from birth (and even before) until the extreme limits of human life. From a demographic point of view, the high number of centenarians in our societies is the integrated result of complex interactions between humans and their environment(s) which underwent consistent changes since the beginning of the 20th century and which are continuing today. Therefore the study of centenarians represents a sort of ‘historical probing’ that allows the tracing of the above-mentioned complex basis of the longevity today. A historical perspective of demographic data on gender and longevity in Italy is shown in Box 1.

The model of centenarians has some disadvantages due to their rarity, lack of an age-matched control group and phenotypical frailty related to their extreme age. Literature suggests that longevity ‘runs in families’ through different generations [136] and, indeed, centenarian offspring appear to be healthier [137,138] and to have a more favourable biological signature [139] with respect to age-matched controls, thus representing a suitable model to identify early biological factors/markers correlated to healthy aging and higher ‘risk’ of longevity. Thus female offspring of centenarian parents could represent a peculiar subgroup of women characterized by a survival advantage not accompanied by the worst quality of life typical of elderly women.

Within this scenario, where plenty of data have described the hormonal, immunological and metabolic gender differences, the study of long-lived families has allowed us to address peculiar aspects of the genetics of aging in women, following the ‘three genetics conceptualization’ we have proposed recently [140]. We have suggested that an integrated investigation of nuclear genetics, mitochondrial DNA genetics and gut microbiome is essential to grasp the genetic contribution to aging and longevity in humans considered as meta-organisms.
Demographic data on gender and longevity: a historical perspective in Italy

Low-mortality countries, as well as Italy, in the recent decades have seen a process of reduction in mortality at all ages of life that has allowed pronounced gains in life expectancy [127]. Currently, the average life expectancy at birth in Italy is among the highest in the world, having reached 80 years for men and 85 years for women. The improvement in living conditions, education, nutrition and lifestyles, and progress in the prevention, diagnosis and treatment of diseases, have been crucial in reducing the risk of death even in advanced ages of life. Indeed, mortality rates at older ages showed a linear downward trend between 1950 and 2012 [128].

The decline in old-age mortality is thought to be the main cause of the dramatic increase in centenarians [129], whose number doubles approximately every 7 years. According to data of the Italian National Institute of Statistics (http://demo.istat.it/), the number of centenarians has reached 19095 (i.e. over 31 per 100 thousand residents) on 1 January 2015. At the same date, according to the ISTAT register, 878 residents on Italian territory were semi-supercentenarians (persons aged 105 or over), whereas 17 were supercentenarians (persons aged 110 or over).

Increased levels of survival are linked to the long process of epidemiologic transition that saw a radical transformation of mortality in its gender, age and cause profiles [131,132]. As shown in Figure 2, life expectancy in Italy was very similar for males and females until the early 20th Century. Afterwards, the evolution of life expectancy, while being dramatically marked by a sharp drop during the two World Wars, showed a differentiation between males and females that reached a maximum of 6.75 years in 1979 and 1980.

The recent decrease in the gender gap is mainly due to the reduction of excess male mortality between the ages of 45 and 75 years. On the other hand, the disadvantage of men compared with women at older ages is emphasized. This phenomenon might be related to a generation effect: whereas in the younger generations with more healthy lifestyles the gap is reducing, the cohorts born in the early 20th Century and the mid-1930s are ‘carriers’ of an excess mortality [133]. The current centenarians emerge from these cohorts and show geographical differences in the female/male ratio (F/M), which is higher in the North-West and North-East areas (around 7:1 and 6:1 respectively), intermediate in the Centre (around 5:1) and lower in the South and Islands (around 4:1), according to the most recent ISTAT and census data. It is worth mentioning that in a mountainous zone of Sardinia (Nuoro province), an exceptionally high number of male centenarians was identified, together with an unusually low F/M ratio [134].

The pattern of distribution of extremely long-lived individuals is certainly affected by environmental factors which shaped the geography of life expectancy in Italy through a different impact on the main causes of death in the elderly [135]. However, a role of genetic factors is suggested by the finding of a correlation between centenarians’ gender ratio across the national territory and the first principal component obtained by studying the polymorphic variation at 95 different loci [5].

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**X chromosome inactivation (XCI) skewing in human aging and longevity**

In cells from females, one of the two X chromosomes is epigenetically and randomly inactivated in early embryonic life. Young women are a mosaic of two cell populations in which either the maternal or the paternal X chromosome is inactivated, and the ratio is close to 50% for each chromosome. A general concordance was seen in the XCI (X chromosome inactivation) pattern between haemopoietic tissue (blood and/or spleen) and several other tissues (e.g. brain, skin, heart, lung, muscle, kidney and gastrointestinal). According to the ‘Heterogametic Sex Hypothesis’, having two copies of the X chromosome may be advantageous for females because of possible selection with age or of the better X chromosome while inactivating the deleterious one [141]. In addition, previous data reveal that a small portion (~17%) of the genes on the inactivated X chromosome are partially active providing a further survival advantages for females [141]. During aging, a marked deviation from the equivalent ratio (50:50) between maternal and paternal X chromosome inactivation occurs (skewing of XCI) in blood cells and the concordance of XCI among tissues may weaken with age. In particular, comparing haemopoietic tissues and brain in the oldest women, the greatest difference between inactivation values of the two tissues were found [142]. The XCI patterns in brain are of particular clinical relevance, because the X chromosome is relatively enriched for genes involved in neuronal functioning [143]. Some authors suggested that age-associated XCI skewing could be involved in the pathogenesis of several diseases such as autoimmunity and cancer [144]. Our proposed experimental model of longevity-healthy aging consisting of female centenarians, their female offspring, female offspring born from non-long-lived parents (age-matched controls) and young women has allowed us to extend to centenarians the study of XCI skewing and to demonstrate that this process was significantly less severe and frequent in centenarian offspring compared with their age-matched controls [145]. These results highlight a possible detrimental link between the rate of XCI skewing and healthy aging/longevity, fitting the hypothesis that the balanced female mosaic is a winning strategy, sustaining a co-operative adaptive mechanism with possible biological advantages, whereas a skewed situation in favour of one of the two X chromosomes would represent an unfavourable condition to attain health and longevity. Conversely, the absence of a similar mosaic strategy in men might contribute to their shorter lifespan [1].

A recent paper has described the correlation between SEMs (stochastic epigenetic mutations) (i.e. rare or stochastic epimutations not shared among individuals) and XCI skewing during aging demonstrating that the number of SEMs was low in childhood and increased exponentially with age [146]. Moreover, a multivariate analysis has indicated a significant correlation between SEMs and degree of XCI skewing after adjustment for age, indicating for the first time that XCI skewing may not be a direct consequence of aging, but is mediated by the number of SEMs. The data from this study support the hypothesis that an increased number of SEMs might influence haemopoietic stem cells viability or might create conditions able to induce clonal stochastic loss of a specific type of haemopoietic cells [146].

**mtDNA and gender in human aging and longevity**

Mitochondria are considered to be important determinants of cell aging because they are involved in several fundamental processes such as cellular energy/ATP production, the urea cycle, heat production, apoptosis, inflammammasome activation and cell senescence. Mitochondria are also the main producers of ROS (reactive oxygen species), the most important by-products of OXPHOS (oxidative phosphorylation), which, besides their physiological role in cell signalling, have been suggested to play a role in the aging process as well as in age-related diseases. Data from primary culture of fibroblasts from long-living individuals, including female centenarians, indicate that longevity is characterized by a
preserved bioenergetics function probably attained by a successful mitochondria remodelling that can compensate for functional defects through an increase in mass, i.e. a sort of mitochondrial ‘hypertrophy’ [147].

Another aspect deserving particular attention in the study of female longevity is the complex and contradictory role of mtDNA variability. mtDNA is an active part of the genetic machinery of each cell and has an active cross-talk with the nuclear genome. Despite its limited length (16 569 bp), the mtDNA encodes few genes with a quantitatively relevant action because of the high copy number of mtDNA in each cell. mtDNA is inherited only through the mother and its germline variants (haplogroups), and D-loop mutations were found to be associated with longevity in several populations indicating a maternal component of longevity.

In particular, the EU large project GEHA (Genetics of Healthy Ageing) studied 2200 ultra-nonagenarians (90+) from different EU countries belonging to 90+ sibpairs together with the same number of sex- and geographically-matched younger controls, and was able to identify different haplogroups related to longevity in males and females. The J2 haplogroup was associated with male longevity, whereas the H2 and T2 haplogroups were associated with female longevity [148]. Taking advantage of the complete sequencing of a high number of mtDNA molecules, it was also possible to evaluate for the first time the cumulative effect of specific and comonont mtDNA mutations, including those that have a low or very low impact. The analysis of the mutations occurring in different OXPHOS complexes showed a complex scenario with a different mutation burden in nonagenarian persons compared with controls. In particular, mutations in subunits of OXPHOS complex I had a beneficial effect on longevity, whereas the simultaneous presence of mutations in complex I and III and in complex I and V seemed to be detrimental [148]. The final conclusion was that “particular rare mtDNA mutations present only in specific populations might be beneficial (or detrimental) for longevity and may explain part of the genetic component of longevity in that population, similarly to what has been suggested for private nuclear DNA polymorphisms” [148].

mtDNA mutations are transmitted from centenarian mother to the progeny. One of the factors that can contribute to aging and longevity is the accumulation with age of mtDNA mutations. mtDNA heteroplasmy, i.e. the presence in the same cell of wild-type and mutated mtDNA molecules, has been supposed to have a double role, fuelling mitochondrial dysfunction and, at the same time, functioning as a reservoir of genetic variability helping the cells to cope with environmental and physiological stressors during life [149,150]. To test the hypothesis that mtDNA heteroplasmy could play a role in human aging and longevity, Giuliani et al. [151] exploited two approaches: (i) the previously described informative model, i.e. 31 centenarian families constituted by the centenarian mother plus the female offspring, in comparison with 28 female offspring of not long-lived parents; (ii) the most recent technology of ultra-deep mtDNA sequencing (average coverage of 49334-fold for each 853bp mtDNA fragment examined). This method allowed the detection of 119 heteroplasmic positions with a minor allele frequency $\geq 0.2\%$. The results indicate that a low
level of heteroplasmies are transmitted and maintained within families until extreme age. However, a non-heteroplasmic variant associated with longevity and healthy aging was identified but a particular and unique heteroplasmia profile for each family was drawn. Therefore mtDNA heteroplasmia appears to be a familial trait transmitted by the mothers which can contribute to healthy aging and longevity [151].

On the other hand, a number of studies have investigated the association between mtDNA inherited variants and multifactorial diseases, such as diabetes [152], ischaemic disease [153] and neurodegenerative diseases such as PD [154] and AD [44]. As described previously, a high-resolution analysis (sequencing of displacement loop and restriction analysis of specific markers in the coding region of mtDNA) found that sub-haplogroup H5 is a risk factor for AD in particular for females and independently of the APOE genotype partially explaining the higher prevalence of AD in women [44].

**Gut microbiota and gender in human aging and longevity**

Humans have to be considered as metaorganisms due to symbiotic relationship with the numerous microbial communities (‘microbiota’) present in various anatomical locations of the human body. Several hundreds of individual bacterial species colonize mouth, upper airways, skin, vagina and intestinal tract constituting a complex and dynamic ecosystem which cross-talk with the environment as well as the rest of the body, including liver and brain among others. At present, the microbiota associated with the intestinal tract (GM (gut microbiota)) is the most studied. The GM are essential for the synthesis of some fundamental nutrients and energy production from food and are able to strongly modulate innate and specific immunity. The gastrointesinal tract of newborns becomes colonized immediately after birth with microorganisms, mainly from the mother. The composition of vaginal tract microbiota of the mother, the mode of delivery (natural or Caesarean) and breast or formula feeding have a deep impact on the GM of human offspring since the very beginning of life. Strong evidence has suggested that the early composition of the microbiota of newborns plays an important role for the postnatal development and functionality of the immune system [155].

Data regarding the association between genders and specific GM communities are still unreliable even if some reports found that some specific taxa (Bacteroides, Ruminococcus, Eubacterium and Blautia) are more abundant in men, whereas Treponema is prevalent in women [78,156]. Probably, these differences in GM composition are due to lifestyle and dietary factors as well as cultural gender-related habits rather than sex hormone effects [78]. Alterations of the GM have been observed in numerous diseases such as obesity, T2D, inflammatory bowel disease and CRC. In particular, specific signatures of GM patterns are associated with autoimmunity affecting prevalently women and contributing to the increase in their morbidity [78]. Thus there is an urgent need to consider the role of gender background in the GM ecology and its relationship with autoimmunity disease onset and therapy effects. This consideration is reinforced by the fact that the importance of GM in human aging is dramatically emerging. This endogenous ecosystem, together with the external antigenic load, is coming out as a crucial driving force of the homeoeostasis of the immune system, and lifelong GM changes, from newborns to centenarians, can represent an important source of inflammatory stimuli. Our group has shown that female centenarians have a different composition of the GM in comparison with sex-matched younger persons, which is associated with an increase in inflammaging (high plasma levels of pro-inflammatory cytokines such as IL-6 and IL-8). In general, with aging, a decrease in the biodiversity of the composition of the GM is observed, with a trend towards an increase in potentially pathogenic bacteria (pathobionts) with respect to the beneficial ones (symbionts producing butyrate and other short-chain fatty acids) [157]. However, data on the remodelling of the GM and its association with inflammaging are still lacking in men, underlining again the importance of conducting gender-specific studies to fill this gap.

**AN AGENDA FOR THE FUTURE: A MANDATORY NEED FOR A GENDER-SPECIFIC MEDICINE**

The aging process starts ‘in utero’ and early events exert potent effects later in life both in adult age and in old age. This lifelong perspective of aging and age-related diseases let emerge the importance of going beyond sex and to consider ‘gender’. Indeed, men and women differ not only biologically (biology, physiology and genetics), but also regarding lifestyle and habits (smoking, nutrition, physical activity, type of work and education, among others) as well as regarding the capability of coping with stress (spousal bereavement, serving as care-givers to family members). These biological and non-biological factors interact continuously lifelong, playing an overwhelming role in modulating health and/or the propensity to diseases and disabilities later in life.

From basic to clinical sciences, there is a mandatory need for studies where gender is appropriately and fully considered. The enormous progress of medicine in the last 50 years has been reached by scientific investigations and publications where gender has been rather neglected: ‘put gender on the agenda’ has been repeatedly stated by top journals such as Nature since 2010 [158,159].

Gender medicine can be considered quite a new but mandatory dimension of medicine that has to go much deeper in understanding the differences between men and women regarding all pathophysiological pathways, clinical characteristics and pharmacological responsiveness, as well as the importance of lifestyle and cultural aspects [18].

Within this scenario, it is even better to speak about a ‘gender-specific medicine’ and not only an indefinite and/or separated ‘gender medicine’ since the gender perspective is broader, should be more pervasive and penetrate all specialties of medicine. Gender medicine is not a separate exercise, or a separate branch of medicine. Therefore gender should be the focus for the clinical approach and this task requires a deep cultural change of mind as well as a reorganization of clinical services in all countries and health systems. Gender medicine is even more necessary in neglected countries such as in Africa where the conditions of
women are worse and gender differences are stronger and have a higher impact on the health status. At the same time, it is currently no longer possible to conduct medical as well as biological sciences and education programmes without taking into considerations gender differences in the medical schools as well as in all educational programmes.

The knowledge of the biology of gender differences in humans are still in their infancy and there is an urgent need for specifically targeted large studies across countries, to take into account the above-mentioned cultural and anthropological differences in a globalized world where migration of persons from countries characterized by different genetic, cultural and anthropological traits and habits is a hot topic.

In conclusion, the development of a gender-specific medicine is of the utmost importance in order to complete our understanding of the main mechanisms of aging as well as the differences in prevention, care, treatment, evolution and outcomes of non-communicable diseases in both genders.

ACKNOWLEDGEMENTS

Thanks are due to Francesco Scalone for his suggestions about Italian demographic data.

FUNDING

This study was supported by the European Union’s Seventh Framework Programme [grant numbers 266486 (“NU-AGE: New dietary strategies addressing the specific needs of the elderly population for healthy aging in Europe”), 602757 (“HUMAN: Health and the understanding of Metabolism, Aging and Nutrition”) and 259679 (“IDEAL: Integrated research on DEvelopmental determinants of Aging and Longevity”) (to C.F.)], by the Italian Ministry of Health [grant number 259679 (“IDEAL: Integrated research on DEvelopmental determinants of Aging and Longevity”) (to C.F.)], by the Italian Ministry of Health [grant number 602757 (“HUMAN: Health and the understanding of Metabolism, Aging and Nutrition”) and 259679 (“IDEAL: Integrated research on DEvelopmental determinants of Aging and Longevity”) (to C.F.)], and by the Italian Ministry of University and Research [grant number CTNO1_00230_413096 (to C.F.)] and by the University of Florence (to D.M.), and by ‘Progetto di Ricerca Sanitaria Finalizzata’ Veneto Region, Italy.

REFERENCES

1 Ostan, R., Monti, D. and Franceschi, C. (2015) Gender and longevity. Ital. J. Gender-Specific Med. 1, 10–14
2 Cevenini, E., Bellavista, E., Tieri, P., Castellani, G., Lescai, F., Francesconi, M., Mishto, M., Santoro, A., Valensin, S., Salvioi, S. et al. (2010) Systems biology and longevity: an emerging approach to identify innovative anti-aging targets and strategies. Curr. Pharm. Des. 16, 802–813 CrossRef PubMed
3 Regitz-Zagrosek, V. (2012) Sex and gender differences in heart. EMBO Rep. 13, 596–603 CrossRef PubMed
4 Franceschi, C., Motta, L., Valensin, S., Rapisarda, R., Franzone, A., Berardelli, M., Motta, M., Monti, D., Bonafé, M., Ferrucci, L. et al. (2000) Do men and women follow different trajectories to reach extreme longevity? Italian Multicenter Study on Centenarians (IMUSCE). Aging (Milano) 12, 77–84 PubMed
5 Passarino, G., Calignano, G., Vallone, A., Franceschi, C., Jeune, B., Robine, J.M., Yashin, A.I., Cavalli Sforza, L.L. and De Benedictis, G. (2002) Male/female ratio in centenarians: a possible role played by population genetic structure. Exp. Gerontol. 37, 1283–1289 CrossRef PubMed
6 Barford, A. and Dorling, D. (2006) Life expectancy: women now on top everywhere. BMJ 332, 808 CrossRef PubMed
7 Bacci, M., (2015) La differenza di genere nella longevitá: si attua il vantaggio delle donne. In Longevitá, Vecchiaia, Salute. (Salvini, S., ed.), pp. 34–38, Neodemos, Firenze
8 Wisser, O. and Vaupel, J.W. (2014) The sex differential in mortality: a historical comparison of the adult-age pattern of the ratio and the difference, MPIDR Working Paper WP 2014-005., Max Planck Institute for Demographic Research, Rostock
9 Beltrán-Sánchez, H., Finch, C.E. and Crimmins, E.M. (2015) Twentieth century surge of excess adult male mortality. Proc. Natl. Acad. Sci. U.S.A 112, 8993–8998 CrossRef PubMed
10 OECD (2014) Life expectancy and healthy life expectancy at birth., OECD Publishing, Paris 16–17, Health at a Glance: Europe 2014
11 Reference deleted
12 Van Oyen, H., Nusselder, W., Jagger, C., Kolip, P., Cambois, E., Robine, J.M. (2013) Gender differences in healthy life years within the EU: an exploration of the “health-survival” paradox. Int. J. Public Health 58, 143–155 CrossRef PubMed
13 Austad, S.N. and Bartke, A. (2015) Sex differences in longevity and in responses to anti-aging interventions: a mini-review. Gerontology 62, 40–46 CrossRef PubMed
14 Heron, M. (2016) Deaths: leading causes for 2013. Nati. Vital. Stat. Rep. 65, 1–95
15 Jagger, C., Gillies, C., Moscone, F., Cambois, E., Van Oyen, H., Nusselder, W. and Robine, J.M. (2008) Inequalities in healthy life years in the 25 countries of the European Union in 2005: a cross-national meta-regression analysis. Lancet 372, 2124–2131 CrossRef PubMed
16 Fouweather, T., Gillies, C., Wohland, P., Van Oyen, H., Nusselder, W., Robine, J.M., Cambois, E. and Jagger, C. (2015) Comparison of socio-economic indicators explaining inequalities in Healthy Life Years at age 50 in Europe: 2005 and 2010. Eur. J. Public Health 25, 976–983 CrossRef PubMed
17 Regitz-Zagrosek, V., Oertelt-Prigione, S., Prescott, E., Francioni, F., Gerdtz, E., Foryst-Ludwig, A., Maas, A.H., Kautzky-Willer, A., Knappe-Wegner, D., Kintscher, U. et al. (2016) Gender in cardiovascular diseases: impact on clinical manifestations, management, and outcomes. Eur. Heart. J. 37, 24–34 CrossRef PubMed
18 Baggio, G., Corsini, A., Florenani, A., Gianni, S. and Zagorov, V. (2013) Gender medicine: a task for the third millennium. Clin. Chem. Lab. Med. 51, 713–727 CrossRef PubMed
19 Ambrosy, A.P., Fonarow, G.C., Butler, J., Chioncel, O., Greene, S.J., Vaduganathan, M., Nodari, S., Lam, C.S., Sato, N., Shah, A.N. and Gheorghiade, M. (2014) The global health and economic burden of hospitalizations for heart failure: lessons learned from hospitalized heart failure registries. J. Am. Coll. Cardiol. 63, 1123–1133 CrossRef PubMed
20 Levy, D., Kchaiaia, S., Larson, M.G., Benjamin, J.E., Kupka, M.J., Ho, K.K., Murabito, J.M. and Vasan, R.S. (2002) Long-term trends in the incidence of and survival with heart failure, N. Engl. J. Med. 347, 1397–1402 CrossRef PubMed
21 KCheiaia, S. and Vasan, R.S. (2015) Heart failure in women: insights from the Framingham Heart Study. Cardiovasc. Drugs Ther. 29, 377–390 CrossRef PubMed
22 Barton, M. and Meyer, M.R. (2009) Postmenopausal hypertension: mechanisms and therapy. Hypertension 54, 11–18 CrossRef PubMed
23 Barrett-Connor, E.L., Cohn, B.A., Wingard, D.L. and Edelstein, S.L. (1991) Why is diabetes mellitus a stronger risk factor for fatal ischemic heart disease in women than in men? The Rancho Bernardo Study. JAMA 265, 627–631 CrossRef PubMed
24 Leslie, M.S. and Briggs, L.A. (2016) Preeclampsia and the risk of future vascular disease and mortality: a review. J. Midwifery Womens Health 61, 315–324 CrossRef PubMed
25 Guariguata, L., Shaw, J.E., Whiting, D.W. and Linnenkamp, U. (2014) Determinants of gender differences in the prevalence of diabetes. Diabetes Res. Clin. Pract. 106, e14–e16 CrossRef PubMed
26 Wild, S., Roglic, G., Green, A., Sicree, R. and King, H. (2004) Global prevalence of diabetes: estimates for the year 2000 and projections for 2030. Diabetes Care 27, 1047–1053 CrossRef

27 Huxley, R., Barzi, F. and Woodward, M. (2006) Excess risk of fatal coronary heart disease associated with diabetes in men and women: meta-analysis of 37 prospective cohort studies. BMJ 332, 73–78 CrossRef PubMed

28 Cull, C.A., Neil, H.A. and Holman, R.R. (2004) Changing aspirin use in patients with Type 2 diabetes in the UKPDS. Diabet Med. 21, 1368–1371 CrossRef PubMed

29 Wexler, D.J., Grant, R.W., Meigs, J.B., Nathan, D.M. and Cagliero, E. (2005) Sex disparities in treatment of cardiac risk factors in patients with type 2 diabetes. Diabetes Care 28, 514–520 CrossRef PubMed

30 Persell, S.D. and Baker, D.W. (2004) Aspirin use among adults with diabetes: recent trends and emerging sex disparities. Arch. Intern. Med. 164, 2492–2499 CrossRef PubMed

31 Maggi, S., Noale, M., Gallina, P., Bianchi, D., Marzari, C., Limongi, F. and Crepaldi, G. (2006) Metabolic syndrome, diabetes, and cardiovascular disease in an elderly Caucasian cohort: the Italian Longitudinal Study on Aging. J. Gerontol. A Biol. Sci. Med. Sci. 61, 505–510 CrossRef PubMed

32 Mottillo, S., Filon, K.B., Genest, J., Joseph, L., Pilote, L., Poirier, P., Rinfret, S., Schiffin, E.L. and Eisenberg, M.J. (2010) The metabolic syndrome and cardiovascular risk: a systematic review and meta-analysis. J. Am. Coll. Cardiol. 56, 1113–1132 CrossRef PubMed

33 Donington, J.S. and Colson, Y.L. (2011) Sex and gender differences in non-small cell lung cancer. Semin. Thorac. Cardiovasc. Surg. 23, 137–145 CrossRef PubMed

34 Nelson, R.L., Dollear, T., Freels, S. and Persky, V. (2007) The relation of age, race, and gender to the subside location of colorectal carcinoma. Cancer 105, 103–107

35 Strimpakos, A.S., Syrigos, K.N. and Saif, M.V. (2009) Pharmacogenetics and biomarkers in colorectal cancer. Pharmacogenomics J. 9, 147–160 CrossRef PubMed

36 Ferlay, J., Steliarova-Foucher, E., Lortet-Tieulent, J., Rosso, S., Coebergh, J.W., Comber, H., Forman, D. and Bray, F. (2013) Cancer incidence and mortality patterns in Europe: estimates for 40 countries in 2012. Eur. J. Cancer 49, 1374–1403 CrossRef PubMed

37 Reference deleted PubMed

38 Alzheimer Europe (2013) Dementia in Europe Yearbook 2013., 115–159 CrossRef PubMed

39 Sampedro, F., Vilaplana, E., de Leon, M.J., Alcolea, D., Pegueroles, J., Montal, V., Carmona-Iragui, M., Sala, I., Sanchez-Saudinos, M.B., Antón-Agurrie, S. et al. (2015) Alzheimer's Disease Neuroimaging Initiative: APOE-by-sex interactions on brain structure and metabolism in healthy elderly controls. Oncotarget 6, 26663–26674 CrossRef PubMed

39 Santoro, A., Balbi, V., Balducci, E., Pirzirini, C., Rosini, F., Tavano, F., Achilli, A., Siviero, P., Minicucci, N., Bellavista, E. et al. (2010) Evidence for sub-haplgroup H5 of mitochondrial DNA as a risk factor for late onset Alzheimer's disease. PLoS One 5, e12037 CrossRef PubMed

40 Jaffe, A.B., Toran-Allerand, C.D., Greengard, P. and Gandy, S.E. (1994) Estrogen regulates metabolism of Alzheimer amyloid β precursor protein. J. Biol. Chem. 269, 13065–13068 PubMed

41 Weng, Q., Santizo, R., Baughman, V.L., Peligrino, D.A. and Iadecola, C. (1999) Estrogen provides neuroprotection in transient forebrain ischemia through perfusion-independent mechanisms in rats. Stroke 30, 630–637 CrossRef PubMed

42 Aenlle, K.K., Kumar, A., Cui, L., Jackson, T.C. and Foster, T.C. (2009) Estrogen effects on cognition and hippocampal transcription in middle-aged mice. Neurobiol. Aging 30, 932–945 CrossRef PubMed

43 Rocca, W.A., Grossardt, B.R. and Shuster, L.T. (2010) Oophorectomy, menopause, estrogen, and cognitive aging: the timing hypothesis. Neurodegener. Dis. 7, 163–166 CrossRef PubMed

44 Nitkowska, M., Czyzyk, M. and Friedman, A. (2014) Reproductive life characteristics in females affected with Parkinson's disease and in healthy control subjects: a comparative study on Polish population. Neurol. Neurochir. Pol. 48, 322–327 PubMed

45 Ragonese, P., D'Amelio, M., Saleni, G., Aridon, P., Gammino, M., Epifanio, A., Morgante, L. and Savettieri, G. (2004) Risk of Parkinson disease in women: effect of reproductive characteristics. Neurology 62, 2010–2014 CrossRef PubMed

46 Rocca, W.A., Bower, J.H., Maragalone, D.M., Ahlskog, J.E., Grossardt, B.R., de Andrade, M. and Melton, L.J. (2008) Increased risk of parkinsonism in women who underwent oophorectomy before menopause. Neurology 70, 200–209 CrossRef PubMed

47 Christensen, K., Dobblhammer, G., Rau, R. and Vaupel, J.W. (2009) Ageing populations: the challenges ahead. Lancet 374, 1196–1208 CrossRef PubMed

48 Corrao, S., Santalucia, P., Argano, C., Dade, C.D., Barone, E., Tettamanti, M., Petrucci, L., Franchi, C., Kamal Eldin, T., Marengoni, A. et al. (2014) Gender-differences in disease distribution and outcome in hospitalized elderly: data from the REPOSI study. Eur. J. Intern. Med. 25, 617–623 CrossRef PubMed

49 Santalucia, P., Franchi, C., Dade, C.D., Tettamanti, M., Petrucci, L., Corrao, S., Marengoni, A., Marucci, M., Nobili, A. and Mannucci, P.M. (2015) Gender difference in drug use in hospitalized elderly patients. Eur. J. Intern. Med. 26, 483–490 CrossRef PubMed

50 Johnson, V.L. and Hunter, D.J. (2014) The epidemiology of osteoarthritis. Best Pract. Res. Clin. Rheumatol. 28, 5–15 CrossRef PubMed

51 Srikanth, V.K., Fryer, J.L., Zhai, G., Winzenberg, T.M., Ahsan, H., Jones, G. et al. (2009) A meta-analysis of sex differences in non-small cell lung cancer. Semin. Thorac. Cardiovasc. Surg. 22, 200–209 CrossRef PubMed

52 Sundström, A., Westerlund, O., Mousavi-Nasab, H., Adolfsson, R. and Nilsson, L.G. (2014) The relationship between marital and parental status and the risk of dementia. Int. Psychogeriatr. 26, 749–757 CrossRef PubMed

53 Heuberger, R. and Wong, H. (2014) The association between depression and widowhood and nutritional status in older adults. Geriatr. Nurs. 35, 428–433 CrossRef PubMed
Gender, aging and longevity in humans

61 Cole, M.G. and Dendukuri, N. (2003) Risk factors for depression among elderly community subjects: a systematic review and meta-analysis. Am. J. Psychiatry 160, 1147–1156 CrossRef PubMed

62 Dahlberg, L., Andersson, L., McKee, K.J. and Lennartsson, C. (2015) Predictors of loneliness among older women and men in Sweden: a national longitudinal study. Aging Ment. Health 19, 409–417 CrossRef PubMed

63 Dykstra, P.A. and Fokkema, T. (2007) Social and emotional loneliness among divorced and married men and women: comparing the deficit and cognitive perspectives. Basic Appl. Social Psychol. 29, 112 CrossRef

64 Lennartsson, C. and Lundberg, O. (2007) ‘What’s marital status got to do with it?’ Gender inequalities in economic resources, health and functional abilities among older adults. In Health Inequalities and Welfare Resources: Continuity and Change in Sweden. (Fritzell, J. and Lundberg, O., eds), pp. 179–198, The Policy Press, Bristol

65 Moon, J.R., Kondo, N., Glymour, M.M. and Subramanian, S.V. (2011) Widowhood and mortality: a meta-analysis. PLoS One 6, e23465 CrossRef PubMed

66 Clouston, S.A., Lawlor, A. and Verdery, A.M. (2014) The role of partnership status on late-life physical function. Can. J. Aging 33, 413–425 CrossRef PubMed

67 Vina, J., Borras, C., Gambini, J., Sastre, J. and Pallardo, F.V. (2005) Why females live longer than males: control of longevity by sex hormones. Sci. Aging Knowledge Environ. 23, pe17

68 Bouman, A., Heineman, M.J. and Faas, M.M. (2005) Sex hormones and the immune response in humans. Hum. Reprod. Update 11, 411–423 CrossRef PubMed

69 Giefing-Kröl, C., Berger, P., Lepperdinger, G. and Grubeck-Loebenstein, B. (2015) How sex and age affect immune responses, susceptibility to infections, and response to vaccination. Aging Cell 14, 309–321 CrossRef PubMed

70 Hao, S., Zhao, J., Zhou, J., Zhao, S., Hu, Y. and Hou, Y. (2007) Modulation of 17β-estradiol on the number and cytotoxicity of NK cells in vivo. Int. Immunopharmacol. 7, 1765–1775 CrossRef PubMed

71 Ashcroft, G.S., Greenwell-Wild, T., Horan, M.A., Wahl, S.M. and Ferguson, M.W. (1999) Topical estrogen accelerates cutaneous wound healing in aged humans associated with an altered inflammatory response. Am. J. Pathol. 155, 1137–1146 CrossRef PubMed

72 Kramer, P.R., Kroner, S.F. and Guan, G. (2004) 17β-Estradiol regulates cytokine release through modulation of CD16 expression in monocytes and monocyte-derived macrophages. Arthritis Rheum. 50, 1967–1975 CrossRef PubMed

73 Olsen, N.J. and Kovacs, W.J. (2011) Evidence that androgens modulate human thymic T cell output. J. Investig. Med. 59, 32–35 CrossRef PubMed

74 Gonzalez, D.A., Diaz, B.D., Rodriguez Perez Mdel, C., Hernandez, A.G., Chico, B.N. and de Leon, A.C. (2010) Sex hormones and autoimmunity. Immunol. Lett. 133, 6–13 CrossRef PubMed

75 Klein, S.L., Jedlicka, A. and Pekosz, A. (2010) The Xs and Y of immune responses to viral vaccines. Lancet Infect. Dis. 10, 338–349 CrossRef PubMed

76 Sakiani, S., Olsen, N.J. and Kovacs, W.J. (2013) Gonadal steroids and humoral immunity. Nat. Rev. Endocrinol. 9, 56–62 CrossRef PubMed

77 Bakhru, P. and Su, M.A. (2016) Estrogen turns down “the AIRE”. J. Clin. Invest. 126, 1239–1241 CrossRef PubMed

78 Gomez, A., Luckey, D. and Taneja, V. (2015) The gut microbiome in autoimmunity: sex matters. Clin. Immunol. 159, 154–162 CrossRef PubMed

79 Fish, E.N. (2008) The X-files in immunity: sex-based differences predispose immune responses. Nat. Rev. Immunol. 8, 737–744 CrossRef PubMed

80 Zandman-Goddard, G., Peeva, E. and Shoenfeld, Y. (2007) Gender and autoimmunity. Autoimmun. Rev. 6, 366–372 CrossRef PubMed

81 Vrachnis, N., Zygozuris, D., Iliodromiti, Z., Danilidis, A., Valsamakis, G. and Kalantarioud, S. (2014) Probing the impact of sex steroids and menopause-related sex steroid deprivation on modulation of immune senescence. Maturitas 78, 174–178 CrossRef PubMed

82 Ossewaarde, M.E., Bots, M.L., Verbeek, A.L., Peeters, E.H., van der Graaf, Y., Grobbee, D.E. and van der Schouw, Y.T. (2005) Age at menopause, cause-specific mortality and total life expectancy. Epidemiology 14, 556–562 CrossRef

83 Gagnon, A., Smith, K.R., Tremblay, M., Vézina, H., Paré, R.P. and Desjardins, B. (2009) Is there a trade-off between fertility and longevity? A comparative study of women from three large historical databases accounting for mortality selection. Am. J. Hum. Biol. 4, 533–540 CrossRef

84 Weinberger, B. and Grubeck-Loebenstein, B. (2012) Vaccines for the elderly. Clin. Microbiol. Infect. 18 Suppl. 5, 100–108 CrossRef PubMed

85 Scholz, J.L., Diaz, A., Riley, R.L., Cancro, M.P. and Frasca, D. (2013) A comparative review of aging and B cell function in mice and humans. Curr. Opin. Immunol. 25, 504–510 CrossRef PubMed

86 Solana, R., Tarazona, R., Gasioso, I., Lesur, O., Dupuis, G. and Fulop, T. (2012) Innate immunosenescence: effect of aging on cells and receptors of the innate immune system in humans. Semin. Immunol. 24, 331–341 CrossRef PubMed

87 Shaw, A.C., Panda, A., Joshi, S.R., Qian, F., Allore, H.G. and Montgomery, R.R. (2011) Dysregulation of human Toll-like receptor function in aging. Ageing Res. Rev. 10, 346–353 CrossRef PubMed

88 Selmin, A., Ojala, J., Kaarniranta, K. and Kauppinen, A. (2012) Mitochondrial dysfunction and oxidative stress activate inflammasomes: impact on the aging process and age-related diseases. Cell. Mol. Life Sci. 69, 2999–3013 CrossRef PubMed

89 Sansoni, F., Vescovini, R., Fagnoni, F., Biasini, C., Zannì, F., Zanì, L., Teleria, A., Lucchinì, G., Passeri, G., Montì, D., Franceschi, C. and Passeri, M. (2008) The immune system in extreme longevity. Exp. Gerontol. 43, 61–65 CrossRef PubMed

90 Franceschi, C., Bonafé, M., Valensin, S., Ojala, J., Kaarniranta, K. and Kauppinen, A. (2008) Inflamm-aging: an evolutionary perspective on immunosenescence. Ann. N.Y. Acad. Sci. 1108, 244–254 CrossRef PubMed

91 Cevenini, E., Monti, D. and Franceschi, C. (2013) Inflamm-aging. Curr. Opin. Clin. Nutr. Metab. Care 16, 14–20 CrossRef PubMed

92 Franceschi, C. and Campisi, J. (2014) Chronic inflammation (inflammaging) and its potential contribution to age-associated diseases. J. Gerontol. A Biol. Sci. Med. Sci. 69 (Suppl. 5), S4–S9 CrossRef PubMed

93 Hirokawa, K., Utsuyama, M., Hayashi, Y., Kitagawa, M., Makinodan, T. and Fulop, T. (2013) Slower immune system aging in women versus men in the Japanese population. Immunity 39, 10–19 CrossRef PubMed

94 Gligio, T., Imro, M.A., Filaci, G., Scudeletti, M., Puppo, F., De Cecco, L., Indiveri, F. and Costantini, S. (1994) Immune cell circulating subsets are affected by gonadal function. Life Sci 54, 1305–1312 CrossRef PubMed

95 Deguchi, K., Kamiada, M., Iraraha, M., Maegawa, M., Yamamoto, S., Ohmoto, Y., Murata, K., Yasui, T., Yamano, S. and Aono, T. (2001) Postmenopausal changes in production of type 1 and type 2 cytokines and the effects of hormone replacement therapy. Menopause 8, 266–273 CrossRef PubMed

96 Kamada, M., Iraraha, M., Maegawa, M., Ohmoto, Y., Takeji, Y., Yasui, T. and Aono, T. (2001) Postmenopausal changes in serum cytokine levels and hormone replacement therapy. Am. J. Obstet. Gynecol. 184, 309–314 CrossRef PubMed

97 Vural, P., Alegul, C. and Canbaz, M. (2006) Effects of hormone replacement therapy on plasma pro-inflammatory and anti-inflammatory cytokines and some bone turnover markers in postmenopausal women. Pharmacol. Res. 54, 298–302 CrossRef PubMed

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Gender, aging and longevity in humans

133 di Fraia, G. (2011) Il recupero dello svantaggio maschile nella sopravvivenza. In Rapporto Sulla Popolazione, L’Italia a 150 Anni dall’Unità. (Salvini, S. and De Rose, A., eds), pp. 88–90, Il Mulino, Bologna

134 Poulain, M., Pes, G.M., Grasland, C., Carru, C., Ferrucci, L., Baggio, G., Franceschi, C. and Deiana, L. (2004) Identification of a geographic area characterized by extreme longevity in the Sardinia island: AKEA study. Exp. Gerontol. 39, 1423–1429 CrossRef PubMed

135 Lippi, R.M. and Caselli, G. (2002), Evoluzione della Geografia della Mortalità in Italia: Tavole Provinciali e Probabilità di Morte per Causa. Anni: 1971–1973, 1981–1983, 1991–1993, Dipartimento di Scienze Demografiche, Università degli Studi La Sapienza, Roma

136 Caselli, G., Poizzi, L., Vaupel, J.W., Deiana, L., Pes, G., Carru, C., Franceschi, C. and Baggio, G. (2006) Family clustering in Sardinian longevity: a genealogical approach. Exp. Gerontol. 41, 727–736 CrossRef PubMed

137 Guerini, P., Miglio, R., Monti, D., Mari, D., Sansoni, P., Caruso, C., Bonafede, E., Bucco, L., Cevenini, E., Ostani, R. et al. (2013) Does the longevity of one or both parents influence the health status of their offspring? Exp. Gerontol 48, 395–400 CrossRef PubMed

138 Bucco, L., Ostani, R., Cevenini, E., Pini, E., Scurti, M., Vitale, G., Mari, D., Caruso, C., Sansoni, P., Fanelli, F. et al. (2016) Centenarians’ offspring as a model of healthy aging: a reappraisal of the data on Italian subjects and a comprehensive overview. Aging (Albany NY) 8, 510–520 CrossRef PubMed

139 Horvath, S., Pirazzini, C., Baccalini, M.G., Gentilini, D., Di Blasio, A.M., Delle Donne, M., Mari, D., Arosio, B., Monti, D., Passarino, G. et al. (2015) Mitochondrial DNA backgrounds might modulate the status of their offspring. Aging (Albany NY) 7, 1159–1170 CrossRef PubMed

140 Garagnani, P., Pirazzini, C., Baccalini, M.G., Candela, M., Brignoli, P., Sevini, F., Luiselli, D., Baccalini, M.G., Salvioni, S., Capri, M. et al. (2014) The three genetics (nuclear DNA, mitochondrial DNA, and gut microbiome) of longevity in humans considered as metagenomics. Biomed. Res. Int. 2014, 560340 CrossRef PubMed

141 Austad, S.N. (2006) Why women live longer than men: sex differences in longevity. Gend. Med. 3, 79–92 CrossRef PubMed

142 Bittel, D.C., Theodore, M.F., Kibiryeva, N., Fischer, W., Talebzadeh, Z. and Butler, M.G. (2008) Comparison of X-chromosome inactivation patterns in multiple tissues from human females. J. Med. Genet. 45, 309–313 CrossRef PubMed

143 Deng, X., Berletch, J.B., Nguyen, D.K. and Disteche, C.M. (2014) X chromosome regulation: diverse patterns in development, tissues and disease. Nat. Rev. Genet. 15, 387–378 CrossRef PubMed

144 Ozcelik, T. (2008) X chromosome inactivation and female predisposition to autoimmunity. Clin. Rev. Allergy Immunol. 34, 348–351 CrossRef PubMed

145 Gentilini, D., Castaldi, D., Mari, D., Monti, D., Franceschi, C., Di Blasio, A.M. and Vitale, G. (2012) Age-dependent skewing of X chromosome inactivation appears delayed in centenarians’ offspring: is there a role for allele imbalance in healthy aging and longevity? Aging Cell 11, 277–283 CrossRef PubMed

146 Gentilini, D., Garagnani, P., Pisini, S., Baccalini, M.G., Calzari, L., Mari, D., Vitale, G., Franceschi, C. and Di Blasio, A.M. (2015) Stochastic epigenetic mutations (DNA methylation) increase exponentially in human aging and correlate with X chromosome inactivation skewing in females. Aging (Albany NY) 7, 568–578 CrossRef PubMed

147 Sgarbi, G., Materrese, P. Rinti, M., Lanzarini, C., Ascione, B., Gibellini, L., Dika, E., Patrizi, A., Tommasino, C., Capri, M. et al. (2014) Mitochondria hyperfusion and elevated autophagic key activity are major mechanisms for cellular bioenergetic preservation in centenarians. Aging (Albany NY) 6, 296–310 CrossRef PubMed

148 Raule, N., Sevini, F., Li, S., Barbieri, A., Tallaro, F., Lomartire, L., Vianello, D., Montesanto, A., Molianen, J.S., Bezrukov, V. et al. (2014) The co-occurrence of mtDNA mutations on different oxidative phosphorylation subunits, not detected by haplogroup analysis, affects human longevity and is population specific. Aging Cell 13, 401–407 CrossRef PubMed

149 Rose, G., Passarino, G., Scorna Ivenchi, V., Romeo, G., Dato, S., Bellizzi, D., Mari, M., Feraco, E., Maletta, R., Bruni, A. et al. (2007) The mitochondrial DNA control region shows genetically correlated levels of heteroplasmy in leukocytes of centenarians and their offspring. BMC Genomics 8, 293 CrossRef PubMed

150 Rose, G., Romeo, G., Dato, S., Crocco, P., Bruni, A.C., Hervonen, A., Majamaa, K., Sevini, F., Franceschi, C. and Passarino, G. (2010) Somatic point mutations in mtDNA control region are influenced by genetic background and associated with healthy aging: a GEHA study. PLoS One 5, e13395 CrossRef PubMed

151 Giuliani, C., Barbieri, C., Li, M., Bucco, L., Monti, D., Passarino, G., Luiselli, D., Franceschi, C., Stoneking, M. and Garagnani, P. (2014) Transmission from centenarians to their offspring of mtDNA heteroplasmy revealed by ultra-deep sequencing. Aging (Albany NY) 6, 454–467 CrossRef PubMed

152 Achilli, A., Olivieri, A., Pala, M., Hooshier Kashani, B., Carossa, V., Pereg, U.A., Gandini, F., Santoro, A., Battaglia, V., Grugni, V. et al. (2011) Mitochondrial DNA backgrounds might modulate diabetes complications rather than T2DM as a whole. PLoS One 6, e21029 CrossRef PubMed

153 Chinnery, P.F., Elliot, H.R., Syed, A. and Rothwell, P.M. (2010) Mitochondrial DNA haplogroups and risk of transient ischaemic attack and ischaemic stroke: a genetic association study. Lancet Neurol. 9, 498–503 CrossRef PubMed

154 Gaweda-Walerych, K., Maruszak, A., Safranow, K., Bialecka, M., Klodowska-Duda, G., Czyzewski, K., Slawek, J., Rudzinska, M., Styczynska, M., Opala, G. et al. (2008) Mitochondrial DNA haplogroups and subhaplogroups are associated with Parkinson’s disease risk in a Polish PD cohort. J. Neural. Transm. 115, 1521–1526 CrossRef PubMed

155 Lee, Y.K. and Mazmanian, S.K. (2010) Has the microbiota entered the brain? Nature 468, 931–937 CrossRef PubMed

156 Mueller, S., Saumer, K., Hanisch, C., Norin, E., Alm, L., Midttvedt, T., Cresci, A., Silvi, S., Orpiansci, C., Verdenelli, M.C. et al. (2006) Differences in fecal microbiota in different European study populations in relation to age, gender, and country: a cross-sectional study. Appl. Environ. Microbiol. 72, 1027–1033 CrossRef PubMed

157 Biagi, E., Nylund, L., Candela, M., Ostani, R., Bucco, L., Pini, E., Nikkila, J., Monti, D., Satokari, R., Franceschi, C. et al. (2010) Through ageing, and beyond: gut microbiota and inflammatory status in seniors and centenarians. PLoS One 5, e10667 CrossRef PubMed

158 Anon (2010) Putting gender on the agenda. Nature 465, 665

159 Schiebinger, L. (2014) Scientific research must take gender into account. Nature 507, 9 CrossRef PubMed