Reply

Mitochondrial Functions, Cognition, and the Evolution of Intelligence: Reply to Commentaries and Moving Forward

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Received: 17 June 2020; Accepted: 3 December 2020; Published: 8 December 2020

Abstract: In response to commentaries, I address questions regarding the proposal that general intelligence (g) is a manifestation of the functioning of intramodular and intermodular brain networks undergirded by the efficiency of mitochondrial functioning (Geary 2018). The core issues include the relative contribution of mitochondrial functioning to individual differences in g; studies that can be used to test associated hypotheses; and, the adaptive function of intelligence from an evolutionary perspective. I attempt to address these and related issues, as well as note areas in which other issues remain to be addressed.

Keywords: mitochondria; cognition; intelligence; health; evolution

1. Introduction

I thank the editors and commenters for thoughtful critiques and questions regarding the proposal that mitochondrial functioning is the most fundamental biological process contributing to human cognition and links cognition to health and aging (Geary 2018, 2019a, 2019b). I cannot address all of the critiques and questions in detail and will focus on the most central of them: specifically, the relative contributions of mitochondria to individual differences in g (Debatin 2020; Matzel et al. 2020; Savi et al. 2020; Stankov 2020; Ujma and Kovacs 2020); empirical approaches to testing the hypothesis (Burgoyne and Engle 2020; Matzel et al. 2020; Sternberg 2020); and, the adaptive function of intelligence from an evolutionary perspective (De Boeck and Kovacs 2020; Sternberg 2020). Many of the more specific critiques and questions are addressed in the context of these broader issues and summary responses to them are provided in Table 1.
Table 1. Core critiques and summary of key replies.

| Authors                  | Core Critiques                                                                 | Key Reply Points                                                                                                                                 |
|--------------------------|-------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------|
| Burgoyne and Engle       | a. How can the hypothesis be falsified?                                        | a1. Strategies to falsify the hypothesis are in Table 1 of Geary (Geary 2018, p. 1030). See also Sections 2.2 and 3.                               |
|                          | b. What is the effect size directly related to variation in mitochondrial (mt) functioning? | a2. Unknown; some proportion of the variance associated with the g factor.                                                                             |
| Debatin                  | a. Systems approach to neuroenergetics is more comprehensive.                 | a1. Agreed, but mitochondrial functioning is central to this system. This is now noted in the introduction.                                      |
|                          |                                                                                 | a2. Mitochondrial functioning is central to this system. This is now noted in the introduction.                                                     |
| Matzel et al.            | a. Multiple lower-order systems influence brain and cognition.                 | a1. Correct, but their functioning is dependent on cellular energy production.                                                                     |
|                          | b. Variation in energy production is not sufficient to place constraints on brain and cognition. | b1. Probably true for many young people in wealthy and healthy populations.                                                                          |
|                          | c. Not sufficient variation in mtDNA genes to create a bottleneck in energy production. | b2. Probably not true with normal aging, various health conditions, nutritional deficits, parasite and toxin exposure, and myriad other stressors that are common outside of Western middle-class populations. |
|                          | d. Processes closer to (e.g., reaction time) mt should be more predictive of intelligence. | c1. Most mitochondrial functions are dependent on nuclear not mitochondrial genes.                                                                |
|                          | e. Should not smarter people should run faster?                                | d. The prediction is that the most complex processes will be the better predictor.                                                                    |
|                          | f. Should not intelligence be more strongly correlated with mothers’ than fathers’ intelligence? | e. No. Efficient energy production will only help if there is also sufficient muscle mass and mix of slow- and fast-twitch muscle fiber.            |
|                          |                                                                                 | f. No. Most mt genes are nuclear and inherited from both parents (Section 2.2).                                                                     |
| Savi et al.              | a. There are cyclical biological mechanisms other than mt.                     | a. True but their operation is dependent on energy availability.                                                                                     |
|                          | b. mt place a ceiling on brain functions but the ceiling might never be reached. | b. True in many cases. However, the ceiling appears to drop with normal aging and disorders that effect mt functions or substrates and may be raised with some interventions. |
|                          | c. This is verbal speculation.                                                  | c. Section 3 outlines ways to test the hypothesis.                                                                                                  |
|                          | d. A dynamic, network approach to intelligence is preferable.                  | d. This is not incompatible with mt; the more complex the network the more energy needed to build and maintain it.                                |
| Schubert and Hagemann    | a. Many cognitive systems rely on energy consuming long-distance brain networks and remain stable or gain with aging, such as vocabulary. | a. True, but retrieving a vocabulary definition, for example, has lower prefrontal engagement than using vocabulary knowledge to solve analogies. The differences in resource demands should be differentially compromised, with normal aging; the latter more than the former. |
|                          | b. Are intervention studies of healthy adults supportive of the hypothesis?    | b. See Section 3.                                                                                                                                   |
|                          | c. Should not intelligence be more strongly correlated with mothers’ than fathers’ intelligence? | c. No. Most mt genes are nuclear and inherited from both parents (Section 2.2).                                                                     |
Table 1. Cont.

| Authors            | Core Critiques                                                                 | Key Reply Points                                                                                                                                 |
|--------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------|
| Stankov            | a. Broad theories of intelligence are preferable to $g$, which is weaker than stated.  
                      | b. It is important to identify diseases that directly involve mt and directly link these to cognition.  
                      | c. Strong claims regarding mt and cognitive aging are premature.  | a. Agreed that the study of cognition must include multiple abilities, but a substantial $g$ factor remains important.  
                      |                                                                  | b. Agreed. See Section 3.  
                      |                                                                  | c. Agreed. This is a proposed mechanism, not a theoretical edict.  |
| Sternberg          | a. Correlational and not causal model.  
                      | b. Are individual differences in intelligence related to mt for any given age?  
                      | c. One’s living environments are important for cognition.  
                      | d. What is the evolutionary advantage of intelligence?  | a1. Agreed that much of the evidence is correlation.  
                      |                                                                  | a2. Methods to assess a causal relation are described in Section 3.  
                      |                                                                  | b. They could be, if people vary in factors that influence mt functioning; See Section 3.  
                      |                                                                  | c. True, as described in Section 2.1.3.  
                      |                                                                  | d. See Section 4.  |
| Ujma and Kovacs    | a. Genetic evidence does not strongly implicate mt functioning in intelligence.  
                      | b. There is no straightforward link between complexity of cognitive tasks and mt functioning.  | a1. These studies are focused on cognitive phenotypes and thus are biased toward identifying higher-level systems.  
                      |                                                                  | a2. The bottom-up focus on mt gene-product proteins reveals a relation between mt and cognition; see Section 2.2.1.  
                      |                                                                  | b. Correct. This is an hypothesis and some methods in Section 3 might be used to test it.  |

Note: mt = mitochondria.
To begin, Debatin (2020) notes that mitochondria are not the only mechanism associated with energy production and there are multiple pathways involved in the generation (whether or not it is in the mitochondria) of adenosine triphosphate (ATP), including transport of substrates needed for this production. He rightly argues that the focus should be on the entire system involved in ATP production, of which mitochondria are only one part. I completely agree and noted this in the original proposal: “I have situated cellular energy production and functioning, largely supported by mitochondria, at the most basic level” (Geary 2018, p. 1032, italics added). I also noted that “[variation] in energy availability can result from differences in the substrates (e.g., pyruvate) available to fuel the process” (Geary 2018, p. 1033), but agree that my focus on mitochondria may have given the impression that the system is less complex than it actually is. Nonetheless, the mitochondria produce most but not all—glycolysis is an additional source of cellular energy (Pellerin and Pierre 1994)—of the energy needed for brain development and functioning and are thus a critical mechanism, although I fully agree that this is only part of a more complex system of energy dynamics.

Perhaps the most central question to emerge in the commentaries is the testability of the model and a correlated issue of whether mitochondrial functioning—broadly meaning ATP production, influences on the availability of associated substrates, control of oxidative stress and other processes—is influenced by environmental factors. As shown in Figure 1, mitochondrial functioning is very sensitive to a variety of social and environmental factors, including chronic psychosocial stress (Picard et al. 2014), glucose homeostasis as related to diet and activity levels (Picard and Turnbull 2013), and toxin exposure (Caito and Aschner 2015), among others. In Section 3, I illustrate this sensitivity with discussion of how glucose homeostasis (e.g., as related to diet) contributes to mitochondrial health and cognitive functioning, and in doing so illustrate one way in which the theory might be tested (Burgoyn and Engle 2020; Savi et al. 2020). Before turning to this literature, I consider issues regarding the relative importance of mitochondrial functions in the context of individual differences in g.

**Figure 1.** Mitochondrial health contributes to the functioning of brain systems at all levels from individual neurons to large-scale brain networks that support cognitive abilities. Mitochondrial health in turn is influenced by myriad factors, as illustrated by the surrounding ovals.

### 2. Relative Importance of Mitochondria

The key issues addressed in the first subsection concern the relative influence of mitochondrial functions on estimates of individual differences in g, broader cognitive abilities, and environment influences, including the Flynn effect (Burgoyn and Engle 2020; Matzel et al. 2020; Savi et al. 2020; Stankov 2020; Sternberg 2020). The key issue addressed in the second subsection concerns the
relation between mitochondrial genes and individual differences in \( g \) (Schubert and Hagermann 2020; Ujma and Kovacs 2020).

2.1. Cognition and Mitochondria

2.1.1. Mitochondrial Contributions as a Proportion of \( g \)

At this time, the relative contributions of individual differences in mitochondrial health to individual differences in cognitive functioning cannot be determined. The upper limit will be the proportion of variation in cognitive abilities that is captured by estimates of \( g \). Stankov (2020) argues that this upper limit could be lower, perhaps 35% or even lower still, than the 50% I suggested (Geary 2018). The variability in the estimates for \( g \) are related in part to differences in the specific measures included in the test batteries used in one study or another (Carroll 1993; Jensen 1998).

As described in the Aging and \( g \) section of Geary (2018), stronger correlations and thus a higher estimate of variance explained by \( g \) is anticipated for batteries that include more complex tasks. This is because mitochondrial energy production is hypothesized to provide a ceiling on optimal performance and the ceiling is more likely to be reached by more people (revealing greater individual differences) for tasks that require the sustained use of distributed (intermodular) brain systems, not simple processes (e.g., reaction time) as suggested by Matzel et al. (2020) and Schubert and Hagermann (2020); the former could reflect variation in the ability to maintain attentional control at the behavioral level, as suggested by Burgoyne and Engle (2020).

Moreover, the variance explained by a \( g \) factor should also depend on the diversity of the sample. Young, healthy and well-educated samples, as in the Pallier et al. (2000) study mentioned by Stankov (2020), will likely produce lower estimates than will more representative samples. By analogy, individual differences in cardiovascular fitness will be more apparent during a stress test than during a casual walk, and variation in fitness will be more apparent in the general population than in young, college-educated adults. In any event, a recent analysis revealed that the positive manifold is universal and explains approximately 46% of the covariance among cognitive measures (Warne and Burningham 2019), as I originally suggested (Geary 2018).

Even if we accept Stankov’s (2020) lower estimate for \( g \), there is still ample room for substantive contributions of mitochondrial health to cognitive functioning but estimating the extent of these contributions will require the types of interdisciplinary studies described in Section 3.

2.1.2. Mitochondria and Alternative Models of Human Cognition

Stankov (2020) argues that I chose to ignore the Cattell–Horn–Carroll (CHC) model and related theories in favor of \( g \), but this is not the case (see also Matzel et al. 2020). Broad factors other than \( g \) are clearly important for understanding human cognition. As was noted in Table 1 of the original article (Geary 2018, p. 1030), if the positive manifold is due to overlapping processes across different cognitive measures—an alternative model to \( g \)—and thus a statistical artifact, then this would falsify my model. Moreover, if my proposal regarding the structure of \( g \) is correct then models such as CHC become the central focus of the organization of human abilities (see Geary 2019a, Figure 1). This is because \( g \) will not manifest as a psychologically measurable ability, but rather as individual differences in abilities and intraindividual developmental (e.g., with normal aging) and disease-related (below) changes in abilities.

From this perspective, \( g \) might be considered a composite measure of the functioning of the brain networks that contribute to cognition (in agreement with Savi et al. 2020), systems whose functioning is ungirded by more basic mechanisms, including mitochondria and other factors that influence energy availability (Debatin 2020). Within-person longitudinal changes in estimates of \( g \), as with normal age-related changes in adulthood, might then provide a global estimate of the rate of degradation of these systems from their peak. As noted in the original proposal (Geary 2018), the most sensitive tests of this decline are predicted to be those that engage the largest distributed intermodular networks,
such as those associated with fluid ability (Jung and Haier 2007). This is because these systems are very energy intensive (Bullmore and Sporns 2012) and are therefore the most sensitive to declines in energy production. At the same time, deficits in these domains would be found in the context of broad declines across cognitive abilities, as appears to be the case (Tucker-Drob et al. 2019).

The basic prediction is that the vulnerability of the cognitive system will be directly related to the amount of neural tissue needed to support the associated ability and the amount of time needed to complete any associated tasks. Schubert and Hagermann (2020) note that short-term memory (STM) is embedded in working memory (WM) tasks and that the latter involve the additional processes of information manipulation (Cowan 2017). Given that WM by definition engages a more complex system of cognitive processes, my prediction is that performance on these measures will decline more sharply than will performance on STM measures, if there are broad declines in mitochondrial health (e.g., as with normal aging).

2.1.3. Mitochondria, Environmental Conditions and the Flynn Effect

From an individual differences perspective, $g$ could be considered a composite estimate of the genetic and environmental influences on the development and current functioning of the brain systems that support cognition. Schubert and Hagermann (2020) and Sternberg (2020) broach the associated issue of whether this perspective is useful for understanding individual differences in cognition among healthy adults. The answer will have to await the development of more sensitive measures of individual variation in mitochondrial functions, and these are being currently developed (see Section 3.1).

Until that time, I note that the young and healthy samples described by Schubert and Hagermann (2020) are not representative of the condition of most people during our evolutionary history or even many people in developing nations today (for reviews see Geary 2015; Trumble and Finch 2019). The abundance of food, reduction in infections, improvements in living conditions and so forth are evolutionarily novel and likely mask the effects of individual differences in mitochondrial functions in the modern world, or at least do so for a larger segment of the population and the lifespan than in our ancestors.

On this view, broad improvements in public health, including better nutrition, reductions in disease and toxin exposure, and reductions in chronic social stressors will help to maintain mitochondrial health that in turn enable a fuller expression and maintenance of genetic potential (Sauce and Matzel 2018). The result is an improvement in cognitive abilities that essentially reflects a rescue of genetic potential by reduction in environmental risk factors, such as poor nutrition (Protzko 2017). This is in keeping with Schubert and Hagermann’s (2020) interpretation of Rae et al.’s (2003) finding that creatine supplementation—which increases the efficiency of mitochondrial energy production (among other things)—increased the cognitive performance of young healthy vegetarians who typically have below-average creatine levels, but similar treatments are not as helpful for healthy adults without creatine deficits (Avgerinos et al. 2018).

Improvements in nutrition and living conditions more broadly could have contributed to the well-known secular increase in performance on intelligence and cognitive tests in developed nations, that is, the Flynn effect (Flynn 1984; Dickens and Flynn 2001; Eppig et al. 2010; Pietschnig and Voracek 2015; Sauce and Matzel 2018). If so, we would expect larger Flynn-effect gains on measures of fluid than crystallized intelligence, and this seems to be the case. The approach could also explain at least some of the differences in average cognitive abilities across people living in developed and developing countries.

Before turning to the next section, Matzel et al. (2020) raised a related question, that is, if mitochondrial functioning is critical to $g$, then should not people high in $g$ also have advantages in other domains, such as running speed? The answer is: No. Evolutionary selection can independently act on traits above the level of mitochondria. There are different advantages to traits such as running speed and intelligence and thus we would expect them to be more or less elaborated in different contexts. Mitochondrial health will influence the extent to which these traits are fully expressed within
each individual, but this is not the same as high interindividual correlations between these traits. Nonetheless, a common link might be found with decrements in mitochondrial functioning, whereby intraindividual changes in performance across traits are correlated (e.g., with normal aging).

2.2. Genes and Mitochondria

2.2.1. Genome-Wide Association Studies, Mitochondrial Proteins, and g

Ujma and Kovacs (2020) argue that genome-wide association studies (GWAS) that have identified single-nucleotide polymorphisms (SNPs) associated with individual differences in cognition are incompatible with the mitochondria model (Geary 2018). One reason is that many of the identified SNPs are typically associated with various aspects of brain and neuronal functioning (e.g., Coleman et al. 2019; Davies et al. 2018). These types of studies use SNPs to predict cognitive or educational outcomes and thus might be biased to more readily detect SNPs associated with biological processes that are more directly related to these outcomes than are mitochondrial functions.

A complimentary and bottom-up approach would be to examine the relation between mitochondrial gene-product proteins and protein networks and individual differences in cognitive functions or intraindividual differences in the stability of cognitive functions with aging. Due to technical challenges (e.g., needing brain issue), these types of studies are not nearly as common as the standard GWAS studies noted by Ujma and Kovacs (2020). Nevertheless, they have the potential to test the mitochondrial model and to broaden the search for the biological foundations of cognitive ability more generally.

Wingo et al. (2019) conducted a protein-wide association study of longitudinal change (6 months to 20 years) in cognitive ability among older adults; proteins were analyzed from brain tissue after the participants had died. They found that 579 proteins and some interactions among them were associated with cognitive stability with aging. Stability was associated with low levels of expression for some proteins, such as those associated with chronic inflammation (see Section 3.2.2), and high levels of expression for other proteins. The most abundant of the latter included proteins associated with mitochondrial functions, neurons, and synaptic transmission; they note that “mitochondrial proteins were featured prominently among cognitive trajectory related proteins ... the 350 proteins with increased abundance in cognitive stability participate in mitochondrial activities and synaptic transmission in neurons” (p. 4). With control of pathological processes (β-amyloid plagues), protein networks associated with synaptic and mitochondrial functions remained significantly related to cognitive stability.

The results are consistent with the GWAS studies listed by Ujma and Kovacs (2020) and indicate that their dismissal of mitochondrial functions is premature. Indeed, Wingo et al. (2019) point out that core synaptic and neural functions are dependent on mitochondrial health, whereby “157 proteins of the 579 cognitive trajectory-associated proteins are mitochondrial proteins and of these, 122 are located in the mitochondria in either the pre- or postsynaptic density” (p. 9).

Ujma and Kovacs (2020) also noted that many of the SNPs identified in GWAS studies are from intergenic regions with no known function and have been identified due to linkage disequilibrium, that is, correlated transmission with functional genes. Gong et al. (2019) sought to address this issue by focusing on GWAS-identified SNPs correlated with intelligence and with known biological functions. Using Bayesian and other techniques, they identified the SNPs and associated genes most likely to be causally related to individual differences in intelligence. The most plausible SNP coded for an enzyme that is critical for the functioning of the mitochondrial electron transport chain associated with ATP production, that is, energy production expressed throughout the body. Other identified genes were associated with pyramidal cells and synapse formation; mitochondrial membrane potential; neural stem cells; and, cell growth and differentiation.
2.2.2. Parental Genetic Influences on Mitochondrial Functions

A more general question is whether the parent-child correlations for intelligence should be higher for mothers than for fathers, given that mtDNA are inherited solely from mothers (Matzel et al. 2020; Schubert and Hagermann 2020). There is no straightforward reason to believe there will be stronger mother-child than father-child correlations for intelligence, because the mtDNA include just 37 genes, only 13 of which are directly related to the electron transport chain that produces ATP. This is in comparison to more than 1000 biparental nuclear DNA (nDNA) genes that support mitochondrial energy production and other mitochondrial functions (Calvo et al. 2016): As noted in the original article (Geary 2018), the nDNA genes that support mitochondrial functions evolved as a compensatory mechanism for degradation of the mitochondrial genome (Havird and Sloan 2016).

My point in the original article was that the uniparental inheritance of mtDNA could contribute to the greater variation in intelligence among males than females (Johnson et al. 2008), and especially contribute to the over-representation of males at the lower end of the distribution. This is because uniparental inheritance means that mutations that disadvantage males but are neutral or beneficial to females cannot not be eliminated by natural selection (Beekman et al. 2014). These mutations would presumably result in more deficits in brain and cognition for males than females, but the evolution of compensatory nDNA and the potential importance of mtDNA–nDNA interactions as related to mitochondrial functions do not result in a straightforward relation between maternal intelligence and offspring intelligence.

Moreover, mt mutations that differentially compromise male cognition will not necessarily be expressed in their mothers’ intelligence, because mutations that compromise the latter will eventually be eliminated by evolutionary selection. The associated process of purifying selection (for females) would disrupt any mtDNA-related correlations between mothers’ and sons’ intelligence. Rather, these correlations would be driven by nDNA and as such should be similar to father-child correlations, as appears to be the case (Whitley et al. 2011). Note that this does not mean that maternally inherited mtDNA mutations cannot affect boys’ cognition more strongly than that of girls. Rather, any such mutations will not necessarily be detected with simple mother-son phenotypic correlations.

3. Empirical Studies and Testing the Hypothesis

As noted in several commentaries (Burgoyne and Engle 2020; Matzel et al. 2020; Savi et al. 2020; Sternberg 2020; Stankov 2020), the critical next steps include empirical tests of predictions derived from the model. The key hypothesis is that mitochondrial health is linked to cognitive performance. One specific prediction is that biomarkers that are indicative of good mitochondrial functioning will predict better cognitive performance, especially on complex tasks. A second specific prediction is that the progression of diseases that compromise mitochondrial functioning will be associated with parallel declines in cognitive functioning. Finally, interventions that promote mitochondrial health should result in improvement in cognitive performance. Overviews of research related to each of these specific predictions is provided below.

3.1. Mitochondrial Biomarkers

Although much remains to be determined, efforts are underway to develop biomarkers of mitochondrial health and dysfunction. In the near future, these biomarkers might prove to be useful in the study of individual differences in cognitive functioning and longitudinal change in this functioning (Picard et al. 2019). One such marker is mtDNA copy number. The latter is an estimate of the number of functioning mitochondria per cell, which generally declines with aging and the progression of various diseases (Montier et al. 2009). There are currently several methods available for the assessment of copy number and others in development (Longchamps et al. 2020); other potential markers include indicators of systemic inflammation (below) and the proportion of wild-type (original) and mutated mtDNA.
There are several preliminary studies that suggest this might be a useful approach (Mengel-From et al. 2014; Silzer et al. 2019), with the caution that the relation between mtDNA copy number measured in peripheral cells (e.g., white blood cells) in many of these studies and mtDNA copy number in brain cells is not well understood. In the larger of these studies ($N = 1067$), Mengel-From et al. found steady mtDNA copy numbers through age 48 years in healthy individuals and age-related declines thereafter. For older individuals, higher mtDNA copy numbers were associated with better physical health and higher cognitive performance; “higher mtDNA copy number was consistently associated with higher cognitive composite score and MMSE [Mini Mental State Examination]” (Mengel-From et al. 2014, p. 1152). In a sample of older women, J. W. Lee et al. (2010) found that higher mtDNA copy number was associated with higher MMSE scores ($r = 0.33$), controlling age and years of education, and Silzer et al. (2019) found that higher copy number was associated with better list learning and memory scores in older adults ($r = 0.46$).

In a unique study noted by Matzel et al. (2020), Bijnens et al. (2019) examined the relation between placental mtDNA copy number taken at the time of birth and intelligence 8 to 15 years later. A doubling of copy number was associated with a 2-point gain in IQ. Among monozygotic twins who shared the same placenta, the twin with higher mtDNA levels (in cord blood) had higher IQ scores than the cotwin ($r = 0.41$). The potential mechanism is prenatal competition between the twins for maternal resources; the twin receiving more resources potentially having higher mtDNA copy numbers in cord blood that in turn may have influenced prenatal brain development. Matzel et al. suggested that this was a prenatal environmental effect and not related to specific advantageous maternal mtDNA. I agree. The point is that, whatever the source, additional mitochondria might provide benefits to prenatal brain development. [The related issues regarding the contributions of mtDNA and nDNA to mitochondrial functioning are addressed in the section Uniparental Inheritance of Mitochondria and Variation in $g$ (see Geary 2018).]

At the same time, there are tissue-specific variations in mtDNA copy number and thus the best noninvasive method (e.g., in blood cells) for their measurement in the central nervous system is not yet known (Wachsmuth et al. 2016). Moreover, there are some disease conditions that can result in a compensatory increase in mtDNA copy number, possibly with a corresponding increase in the number of mtDNA mutations, and thus a linear relation between copy number and cognitive functions might not always emerge (Hulgan et al. 2019). Despite these caveats, the development of biomarkers of mitochondrial health will provide unique opportunities for future interdisciplinary work involving cognitive and biomedical scientists.

3.2. Mitochondrial-Related Disorders and Cognition

There are progressive diseases, including mitochondrial disorders, that compromise mitochondrial functions and are associated with cognitive declines. Prospective studies of cognitive changes associated with disease progression provide an indirect test of the hypothesized relation between mitochondrial health and cognitive function. Moreover, the pattern of cognitive decline should be similar to that found for normal age-related cognitive changes and perhaps the Flynn effect, including larger changes for more attention-demanding tasks (e.g., fluid intelligence) relative to less difficult tasks, such as short-term memory (e.g., forward digit span), memory retrieval (e.g., vocabulary), or simple reaction time.

The parallel between cognitive declines with disease progression and normal age-related changes follows from the hypothesis that compromises to mitochondrial functioning are a common underlying and causal mechanism, and inconsistent with Matzel et al.’s (2020) argument that the relation between cognition and health is simply due to health-related decisions and behaviors (see also Deary et al. 2010). At this time, only indirect evidence can be marshalled to evaluate the hypothesis. The study of mitochondrial disorders is one approach, although it is not as straightforward as it might seem. The study of cognitive changes associated with obesity-related compromises in mitochondrial health is another such disease (Picard and Turnbull 2013).
3.2.1. Mitochondrial Disorders

On the basis of my proposal (Geary 2018, 2019a), mitochondrial disorders should be associated with lower performance across cognitive domains, as noted by Matzel et al. (2020) and Stankov (2020). As an example, Stankov cites one study in which some but not all individuals with a mitochondrial disorder showed cognitive impairments. These types of studies are not necessarily evidence against my proposal, because there are different forms of mitochondrial disorder with different degrees of severity and different rates of disease progression (Chinnery and Turnbull 2001; Set et al. 2019).

At this point, a well-designed study of the cognitive competencies of individuals with mitochondrial disorders and changes in these competencies as the disease progresses remains to be conducted. A recent review of extant studies shows that cognitive deficits are more common on complex attention-demanding measures (e.g., executive functions) than on measures of crystallized intelligence; that asymptomatic individuals often show only subtle deficits but deficits are common in symptomatic individuals (e.g., showing vision difficulties, seizures); and, cognitive abilities decline as the disease progresses (Moore et al. 2020; Moore et al. 2019; see also Finsterer 2012).

3.2.2. Obesity, Diabetes, and Inflammation

The availability of energy substrates (lipids, glucose), including chronic over availability resulting in obesity and physical exercise that consumes these substrates (below), can influence long-term mitochondrial health and functioning (Picard and Turnbull 2013). There are many details that cannot be adumbrated here but the key is that mitochondria are highly responsive to metabolic state, and chronic obesity-related disruption of glucose homeostasis is associated with cognitive declines and accelerated aging that are at least in part related to compromised mitochondrial functions.

An overabundance of calories is associated with increased risk of insulin resistance, Type II diabetes, atherosclerosis, and other health issues. Among other things, swollen adipose tissue leaks lipids (fatty acids) that can trigger an immune response and chronic inflammation (Spyridaki et al. 2016). The oversupply of energy also overwhelms mitochondrial energy production systems that in turn exacerbate inflammation and increase cell and DNA damaging oxidative stress (de Mello et al. 2018; Sripetchwannsee et al. 2018). The mechanisms that would typically keep this damage in check are simultaneously compromised, resulting in further damage (Picard and Turnbull 2013).

The associated inflammatory molecules can disrupt the blood brain barrier and their movement into the central nervous system can result in neuroinflammation (Pugazhenthi et al. 2017). One consequence is a reduction in the energy producing capacity of mitochondria, among other things (e.g., disruption of intracellular signalling; Bhatti et al. 2017). There are defensive mechanisms to maintain energy supplies—for instance increased generation of new mitochondria but often with the cost of increased levels of mtDNA mutations—but declines continue (without intervention) and eventually impact system functioning. Picard and Turnbull (2013) estimated that these changes, if chronic, are equivalent to 5 to 20 years of normal aging moving from middle to old age (see also Stevenson et al. 2019).

As with mitochondrial disorders, the definitive study of obesity-related disruptions in mitochondrial functioning as related to cognition remains to be conducted. There is nevertheless good reason to suspect a substantive relation. There are moderate but consistent (Effect Size, $ES \approx 0.3$ to 0.45) differences in various components of executive functions comparing obese to normal weight individuals and even moderate deficits in inhibition and working memory comparing overweight to normal weight individuals ($ES \approx 0.1$ to 0.2; Yang et al. 2018). These patterns are likely bidirectional, with poor executive functions associated with difficulties in regulating food intake, as well as compromises in executive functions resulting from the just described biological mechanisms (Shields et al. 2017; Spyridaki et al. 2016).

In a sample of healthy adults ($N = 180$), Spyridaki et al. (2014) found lower fluid intelligence in obese as compared to normal weight individuals. A series of path models suggested that the differences were mediated by inflammation, one indicator of compromised mitochondrial functioning;
an alternative model in which fluid intelligence caused weight differences did not fit the data as well as the inflammation model. Stronger evidence is found with prospective studies of the relation between baseline risk factors, such as chronic inflammation, and rate of cognitive decline and impairment with aging (Cheng et al. 2012; Levine et al. 2018; R. Schmidt et al. 2002; Yaffe et al. 2004). As an example, a 10 year longitudinal study of 45- to 69 year olds revealed accelerated declines in reasoning and memory abilities for individuals with indications of chronic inflammation at baseline (Singh-Manoux et al. 2014). The declines were equivalent to an additional 1.7 years of normal age-related cognitive changes over the 10 years of the study.

Studies using Positron Emission Tomography (PET) indicate declines in glucose utilization with the progression of metabolic diseases (e.g., diabetes). Volkow et al. (2009) found that obesity was associated with lower baseline levels of glucose utilization among young, healthy adults ($M = 34$ years) that in turn was associated with lower performance on a battery of neuropsychological tests. They suggested that the lower baseline glucose utilization was due to reduced grey matter volume, but this remains to be determined.

In a large study of older adults ($M = 61$ years), Willette et al. (2015) found that insulin resistance was associated with lower overall ($\beta = -0.29$) and regional differences in brain glucose metabolism. The strongest changes were in the hippocampus and medial temporal lobe, which has a high density of insulin receptors and high energy demands because of continual neurogenesis. Higher glucose metabolism (an indication of mitochondrial functioning) in these regions was associated with better immediate memory ($\beta = 0.32$) and verbal learning ($\beta = 0.31$) but had weaker relations to working memory ($\beta = 0.12$) and processing speed ($\beta = 0.20$).

With disease progression these deficits become widespread and are found at multiple levels, including reductions in grey and white matter, reduced hippocampal neurogenesis, and reduced mitochondrial trafficking within axons that in turn disrupts synaptic functions, among other things (O’Brien et al. 2017; Dye et al. 2017). Studies of older at-risk individuals (e.g., diabetes) indicate declines in glucose metabolism throughout the prefrontal cortex and parietal regions (Baker et al. 2011; Kuczynski et al. 2009), that is, regions associated with working memory and fluid intelligence (Jung and Haier 2007).

A recent meta-analysis suggested that for otherwise healthy people, many of these declines may not be detectable until years after the onset of inflammation or other aspects of disrupted glucose homeostasis (Attie et al. 2019; Cheng et al. 2012), although high fat diets might result in changes fairly quickly (Edwards et al. 2011). As cognitive and functional abilities begin to decline, there is often an increase in brain glucose metabolism that likely reflects use of compensatory cognitive strategies and biological mechanisms (e.g., increased mitochondrial biogenesis; Iozzo and Guzzardi 2019). The compensatory mechanisms temporarily maintain functions but could accelerate mitochondrial and neural damage and eventually result in precipitous declines in cognition and day-to-day functioning.

From the perspective of my model, deficits associated with disrupted glucose homeostasis (e.g., as indexed by measures of systemic inflammation) should result in a pattern of cognitive decline that varies with the energy demands of the cognitive task—these could be indexed by PET studies or the size and complexity of the supporting brain systems as identified in fMRI studies (see critique by Ujma and Kovacs 2020)—and track the pattern of age-related cognitive declines. This would include steeper declines in fluid intelligence, measures of active learning, and attentional control and later and more shallow declines in crystallized abilities, such as vocabulary (Geary 2018). To be sure, there are many pathological changes associated with chronic disruptions in glucose homeostasis, above and beyond declines in mitochondrial functioning, but declining mitochondrial health is an important component of the disease process (Picard and Turnbull 2013), and the study of the pattern of cognitive declines associated with this progression provides a means to assess the plausibility of my model (Geary 2018, 2019a).

An important corollary noted by Matzel et al. (2020) is that the relation between mitochondrial declines and cognitive declines is not linear, because of compensatory mechanisms that maintain
energy production. As noted, the maintenance of energy production may be associated with increased mitochondrial numbers but with increases in the proportion of mutated mtDNA (vs. wild type). If so, then the proportion of mutated to wild-type mtDNA copy numbers might eventually prove to be a useful marker of impending cognitive declines.

3.3. Mitochondrial Health and Cognition

If disruption of glucose homeostasis and associated issues with inflammation and mitochondrial dysfunction is causally related to the just-described cognitive declines, then interventions that address the disruption should improve cognitive functioning or reduce the rate age-related decline. These types of interventions exist and range from substantive weight loss (e.g., involving bariatric surgery), insulin administration, use of drugs to reduce oxidative stress, and exercise, among others (Bhatti et al. 2017; Shemesh et al. 2012; Spitznagel et al. 2015). There is evidence that many of these interventions can improve cognitive functions, although individuals differ in their responsiveness to such interventions for reasons that are not yet fully understood.

I focus here on weight loss, exercise, and their combination, because studies of nonhuman animals and humans have provided key insights into the biological changes associated with such interventions and correlated changes in cognition. Overall, a combination of calorie restriction, weight loss, and exercise can reverse many of the above-described effects associated with too much energy and disrupted glucose homeostasis. These include the upregulation of a variety of protective mechanisms, including upregulation of mechanisms that support mitochondrial energy production, DNA repair and antioxidant production, as well as removal of damaged cells and damaged mitochondria within cells. The result is improvements in functioning across intracellular to intermodular systems (Mattson 2012).

In a large-scale randomized controlled trial (RCT), Ngandu et al. (2015) followed nearly 1200 older individuals (66 to 77 years old) at-risk for dementia for 24 months; these individuals were average or lower in overall cognitive ability. The intervention included diet, exercise, and cognitive stimulation. The combination resulted in modest (ES = 0.13) cognitive benefits overall but more substantive benefits for executive functions, processing speed, and delayed memory recall. The results are similar to those found in other RCTs with healthier adults, where ESs are approximately 0.15 (Smith et al. 2010). A more recent meta-analysis included estimates of longitudinal change in cognition following weight loss and RCTs for very obese individuals, and found more substantive benefits (Veronese et al. 2017). For the longitudinal studies, the ESs for various attention, executive functions, and memory measures ranged from 0.30 to 0.66, with no improvement on language measures (language was not frequently assessed in these studies and thus the ES is unreliable). The repeated assessment in such studies could, however, inflate the ES estimates, but the RCTs also showed improved cognitive performance, especially with the combination of diet and exercise; attention (ES = 0.44), memory (ES = 0.35), language (ES = 0.21), with no effect for executive functions but this was assessed in only two studies. The authors concluded that the gains might be related to reductions in insulin resistance and improvements in glucose metabolism, as well as reductions in inflammation and oxidative stress, among other things.

In one of these studies, Napoli et al. (2014) examined the relation between specific health parameters and cognitive improvements for obese, high-risk older adults. The combination of diet and exercise resulted in improvements in overall cognitive ability (composite), executive functions, and word fluency. Within the exercise groups (with and without diet) increases in VO\textsubscript{2}max—which is related to the flow of oxygen to mitochondria—and increased strength—physical exercise is helpful for insulin regulation—explained 19% to 24% of the improvement in global cognitive functions. The interventions did not, however, improve inflammation, possibly because many participants remained obese, although they had lost significant amounts of weight.

Although much remains to be learned, studies that restore glucose homeostasis are associated with improvement in cognitive functions or slower age-related declines in cognition. Various aspects of mitochondrial functioning, including energy production, are integral to these improvements but are not the only source of them. At this time, it may be difficult to disentangle the relative contributions
of one mechanism (e.g., mitochondrial energy production) or another (e.g., improved sensitivity to insulin) to cognitive improvements, given their interdependency. Nonetheless research in this area provides a means to test the predictions laid out in my original proposal (Geary 2018).

4. Adaptive Function of Intelligence

Matzel et al. (2020) correctly note that there are evolved mechanisms to ensure an adequate energy supply, especially to key organs such as the heart and brain. They further suggest that most individuals, during our evolutionary history, were healthy and not subject to conditions that would compromise mitochondrial energy production or other functions. This is not correct. Common threats to populations living in natural conditions, human and nonhuman, include chronic parasitic infections, nutritional and caloric deficits, and chronic social stressors (e.g., competition for mates, parenting; Geary 2015; Trumble and Finch 2019), all of which can compromise one or several aspects of mitochondrial functioning (Geary 2017; Hill 2014; Koch et al. 2017). As noted, reductions in the severity of these threats over the 19th and 20th centuries might have contributed, at least in part, to the Flynn effect.

Sternberg (2020) suggested that there was no adaptive use for intelligence, at least from an evolutionary perspective. Most generally, the large caloric costs associated with the development and maintenance of the human brain must be balanced against substantial benefits, at least in some contexts (Leonard and Robertson 1994); such a costly organ could not have evolved without adaptive benefits.

There are several models that attempt to understand the adaptive importance of intelligence, most of which focus on foresight, planning, and the ability to cope with social and ecological change and novelty (Alexander 1989; Ash and Gallup 2007; Bailey and Geary 2009; Flinn et al. 2005; Geary 2005; Kanazawa 2004; Kaplan et al. 2000; Potts 1998; Richerson et al. 2001). The models thus primarily focus on fluid intelligence rather than g per se, but discussion of these models is nevertheless useful as related to Sternberg’s question.

The foci of different models of the evolution of intelligence range from the ability to anticipate and prepare for variation in seasonal changes (e.g., winter, Kanazawa 2008; Potts 1998) to the protracted learning needed to become proficient in obtaining the staples of life (e.g., hunting; Kaplan et al. 2000) to competition with other people or groups of people (Alexander 1989; Flinn et al. 2005; Geary 2005). It is likely that some combination of these factors contributed to the evolution of fluid intelligence, although their relative importance may have changed across evolutionary time.

My attempt to integrate these different models is summarized in Figure 2. In keeping with earlier proposals (Geary 2005; Kanazawa 2004) and Cattell’s (1963) original insight, the key idea is that the evolutionary function of fluid intelligence is to cope with social and ecological variability or situations that cannot be addressed by evolved or learned responses. These are situations in which automatically invoked solutions need to be inhibited and new solutions constructed. This rudimentary ability to solve novel problems is found in many species of primate and thus was almost certainly present in our hominin ancestors (Burkart et al. 2017).

One key issue that needs to be addressed with these models is the three-fold increase in brain size over the past 4 million years of hominin evolution and the accelerated pace of enlargement over the past 2 million years, with the emergence of key Homo species (e.g., Antón et al. 2014; McHenry 1994). Although brain size and fluid intelligence are only moderately correlated (Gignac and Bates 2017; J. J. Lee et al. 2019) and other factors (e.g., neural density) are important (Dicke and Roth 2016), there were almost certainly evolutionary gains in fluid intelligence during hominin evolution.

One argument is that the evolutionary enlargement of hominin brain size was driven by ecological change—reductions in forested area and increases in savannah in sub-Saharan Africa—associated with climatic fluctuations before the emergence of Homo (e.g., Vrba 1995). One difficulty with the climate argument is that sympatric (living in same ecology) species of primate did not experience the same change in brain size (Elton et al. 2001). In other words, why did our australopithecine and
Homo ancestors show marked increases in brain volume when other primate species living in the same ecologies did not?

![Figure 2](image-url)

**Figure 2.** A cyclical within-species arms race is the most plausible account of the evolutionary enlargement of the human brain and enhancement in fluid intelligence.

The most straightforward answer is a within-species arms race that accelerated after the emergence of Homo. Species within this genus are important because they developed relatively sophisticated tools that likely reduced mortality rates and supported increased population sizes (Alexander 1989; Flinn et al. 2005)—improvement in the ability to extract resources from the ecology and reduce predatory threats is what Alexander called ‘ecological dominance’.

There is in fact consistent archeological evidence for changes in our ancestors’ ability to obtain resources from the ecology (see Foley 1987; Foley and Lahr 1997). One result of these and related innovations, especially cooking, is an increase in the availability of calories and the energy produced by mitochondria, which would have removed an evolutionary constraint on the expansion of the energy-demanding brain (Aiello and Wheeler 1995). Another result is an increase in the carrying capacity of these ecologies that in turn is typically associated with population increases and expansions. As Homo erectus and later Homo sapiens expanded into new ecologies and became increasingly skilled at exploiting them, an evolutionary Rubicon was crossed:

the ecological dominance of evolving humans diminished the effects of ‘extrinsic’ forces of natural selection such that within-species competition became the principle ‘hostile force of nature’ guiding the long-term evolution of behavioral capacities, traits, and tendencies. (Alexander 1989, p. 458)

Evidence in support of this argument is found in the pattern of human migration and the subsequent extinction of megafauna (Martin 1967, 1973), and parallels Mac Arthur and Wilson’s (1967) analysis of island biogeography. When a species migrates into an unexploited region (e.g., an island) there are few constraints on population expansion, low levels of social competition, and a rapid increase in population size, as shown by the top oval in Figure 2. As the population expands, the quantity of per capita resources necessarily declines and competition for access to these diminishing resources necessarily intensifies.

The increase in social competition is represented by the rightmost oval of Figure 2. In this circumstance, Darwin and Wallace’s (1858, p. 54) conceptualization of natural selection as a “struggle for existence” becomes in addition a struggle with other human beings for control of the resources that support life and allow one to reproduce (Geary 2005). The eventual result of this struggle is a population crash, as was described by Malthus (1798) for many human populations in developing nations before the demographic shift and Hamilton and Walker (2018) for hunter-gatherer societies. Following the crash, the per capita availability of resources is higher than it was before the crash and thus another
cycle of population expansion to the carrying capacity of the ecology and later contraction begins (Fanta et al. 2018).

The repeating cycle accelerates evolutionary selection and will favor those individuals who gain competitive advantage over others; the latter specifically refers to greater social influence and enhanced control over culturally important resources (Geary 2021). There are many ways to achieve advantage and at least some of these are likely facilitated by fluid intelligence. As shown in Figure 2, my proposal is that fluid intelligence facilitated the creation of novel technologies, social strategies, and patterns of social organization that not only resulted in competitive advantage but further gains in these areas. We cannot of course directly study the associated social dynamics among early species of Homo or early modern humans, but we can look for such dynamics in the historical record.

The development of large-scale agriculture provides an apt example. Although the quality of the associated diet was often lower than could be achieved through hunting and gathering, it enabled a change in human social organization—larger groups have a competitive advantage over smaller ones—and increased the overall availability of calories that in turn supported population expansions (Clark 2008). Critically, these additional calories could be stored as grain reserves or in livestock and these created a tempting source of wealth for the taking (Hirschfeld 2015; Turchin 2009). In areas in which steppes occupied by nomadic herders abutted fertile agricultural lands, nomadic groups and agricultural communities came into contact and oftentimes eventual conflict (Turchin 2009). The historical record shows that many nomadic groups began to raid agricultural settlements and the theft of these communities’ resources created benefits for the formation of larger agricultural communities. To counter the defensive advantage of larger communities, smaller nomadic groups had to unite to continue their raiding.

This type of cycle appears to have independently emerged in many parts of the world and was associated with advances in social organization, military strategy and technology (e.g., chariot), and eventually led to the formation of empires (see Currie et al. 2020; Hermann et al. 2020; Turchin 2009; Turchin et al. 2013). The historical record and the implied intensity of social competition (especially for men; see Geary 2021) is supported by numerous population genetic studies showing the replacement of one human population by another in all parts of the world in which it has been studied (Zeng et al. 2018; Zerjal et al. 2003). In other words, studies in these areas are consistent with a long-term within-species arms race. The cultural advances that emerged during the arms race must have been aided, at least in part, by individuals with a combination of strong fluid abilities and other traits (e.g., conscientiousness and planning) that support innovations in social organization and technology.

5. Discussion

The positive manifold and the associated covariation among cognitive and academic measures is one of the most replicated findings in the behavioral sciences (Warne and Burningham 2019). Considerable advances have been made in identifying the brain and cognitive systems that contribute to the manifold (e.g., Jung and Haier 2007) but, as noted in all of the commentaries, much remains to be determined. The bulk of the associated theory and research has been focused on proximate cognitive processes, the broad neural networks that support these processes, and on real-world correlates, such as performance in occupational settings. These are all critical aspects of theoretical and empirical work in the field of intelligence, but there is much to be gained by expanding this work to different levels of analysis, from the dynamics of cell biology to the evolutionary function of intelligence and cognition more broadly.

My proposals here and elsewhere (Geary 2005, 2018) are first approximations of how we might more fully understand intelligence by expanding the nomological net to capture a wider range of literatures and phenomena. The first point for now is that the study of cognitive changes associated with the progression of diseases that undermine mitochondrial health provide a means to assess the plausibility of my model that in turn might provide links between these diseases and normal age-related cognitive declines and perhaps to the pattern of secular changes associated with the Flynn
Cognitive scientists in turn can contribute to our understanding of disease progression by designing psychometric assessments that cover the broad range of cognitive abilities, such as those identified in the CHC model, and experimental measures that might be more sensitive to individual and developmental differences in core abilities (e.g., attentional control) than are commonly used neuropsychological assessments (Engle 2002).

The second point is that our understanding of intelligence will be enhanced by considering the challenges that different aspects of cognition helped to address during our evolutionary history. The proposal that the evolution of fluid abilities was driven by the benefits of enhanced cognitive and behavioral flexibility as related to coping with variable or novel social and ecological conditions (Geary 2005; Kanazawa 2004) is consistent with recent comparative studies (Burkart et al. 2017). The consideration of comparative work and potential evolutionary processes that drove changes in brain and cognition may be useful for placing psychometric and cognitive studies of human abilities in broader perspective.

**Funding:** The author did not receive funding for the preparation of this article.

**Conflicts of Interest:** The author declares no conflict of interest.

**References**

Aiello, Leslie C., and Peter Wheeler. 1995. The expensive-tissue hypothesis: The brain and the digestive system in human and primate evolution. *Current Anthropology* 36: 199–221. [CrossRef]

Alexander, Richard. D. 1989. Evolution of the human psyche. In *The Human Revolution: Behavioural and Biological Perspectives on the Origins of Modern Humans*. Edited by Paul Mellars and Chris Stringer. Princeton: Princeton University Press.

Antón, Susan C., Richard Potts, and Leslie C. Aiello. 2014. Evolution of early Homo: An integrated biological perspective. *Science* 345: 1236828. [CrossRef] [PubMed]

Ash, Jessica, and Gordan G. Gallup. 2007. Paleoclimatic variation and brain expansion during human evolution. *Human Nature* 18: 109–24. [CrossRef] [PubMed]

Attì, Anna R., Stefano Valente, Antonia Iodice, Llaria Caramella, Barbara Ferrari, Umberto Albert, Laura Mandelli, and Diana De Ronchi. 2019. Metabolic syndrome (MetS), mild cognitive impairment (MCI) and dementia: A meta-analysis of longitudinal studies. *The American Journal of Geriatric Psychiatry* 27: 625–37. [CrossRef] [PubMed]

Avgerinos, Konstantinos I., Nikolaos Spyrou, Konstantinos I. Bougioukas, and Dimitrios Kapogiannis. 2018. Effects of creatine supplementation on cognitive function of healthy individuals: A systematic review of randomized controlled trials. *Experimental Gerontology* 108: 166–73. [CrossRef] [PubMed]

Bailey, Drew H., and David C. Geary. 2009. Hominid brain evolution: Testing climatic, ecological, and social competition models. *Human Nature* 20: 67–79. [CrossRef]

Baker, Laura D., Donna J. Cross, Satoshi Minoshima, Dana Belongia, G. Stennis Watson, and Suzanne Craft. 2011. Insulin resistance and Alzheimer-like reductions in regional cerebral glucose metabolism for cognitively normal adults with prediabetes or early type 2 diabetes. *Archives of Neurology* 68: 51–57. [CrossRef] [PubMed]

Beekman, Madeleine, Damian K. Dowling, and Duur K. Aanen. 2014. The costs of being male: Are there sex-specific effects of uniparental mitochondrial inheritance? *Philosophical Transactions of the Royal Society B: Biological Sciences* 369: 20130440. [CrossRef] [PubMed]

Bhatti, Jasvinder S., Gurjit K. Bhatti, and P. Hemachandra Reddy. 2017. Mitochondrial dysfunction and oxidative stress in metabolic disorders—A step towards mitochondria based therapeutic strategies. *Biochimica et Biophysica Acta (BBA)-Molecular Basis of Disease* 1863: 1066–77. [CrossRef] [PubMed]

Bijnens, Esmeer M., Catherine Derom, Steven Weyers, Bram G. Janssen, Evert Thiery, and Tim S. Nawrot. 2019. Placental mitochondrial DNA content is associated with childhood intelligence. *Journal of Translational Medicine* 17: 361. [CrossRef] [PubMed]

Bullmore, Ed, and Olaf Sporns. 2012. The economy of brain network organization. *Nature Reviews Neuroscience* 13: 336–49. [CrossRef] [PubMed]

Burgoyne, Alexander P., and Randall W. Engle. 2020. Mitochondrial functioning and its relation to higher-order cognitive processes: Commentary on Geary (2018; 2019). *Journal of Intelligence* 8: 14. [CrossRef] [PubMed]
Burkart, Judith M., Michele N. Schubiger, and Carel P. van Schaik. 2017. The evolution of general intelligence. *Behavioral and Brain Sciences* 40: e195. [CrossRef] [PubMed]

Caito, Samuel W., and Michael Ashchner. 2015. Mitochondrial redox dysfunction and environmental exposures. *Antioxidants & Redox Signaling* 23: 578–95. [CrossRef]

Calvo, Sarah E., Karl R. Clauser, and Vamsi K. 2016. Mootha. MitoCarta2. 0: An updated inventory of mammalian mitochondrial proteins. *Nucleic Acids Research* 44: D1251–57. [CrossRef]

Carroll, John B. 1993. *Human Cognitive Abilities: A Survey of Factor-Analytic Studies*. New York: Cambridge University Press.

Cattell, Raymond B. 1963. Theory of fluid and crystallized intelligence: A critical experiment. *Journal of Educational Psychology* 54: 1–22. [CrossRef]

Cheng, G., C. Huang, H. Deng, and H. Wang. 2012. Diabetes as a risk factor for dementia and mild cognitive impairment: A meta-analysis of longitudinal studies. *Internal Medicine Journal* 42: 484–91. [CrossRef]

Chinnery, Patrick F., and Douglass M. Turnbull. 2001. Epidemiology and treatment of mitochondrial disorders. *American Journal of Medical Genetics* 106: 94–101. [CrossRef]

Clark, Gregory. 2008. *A Farewell to Alms: A Brief Economic History of the World*. Princeton: Princeton University Press.

Coleman, Jonathan R. L., Julien Bryois, Hélène A. Gaspar, Philip R. Jansen, Jeanne E. Savage, Nathan Skene, Robert Plomin, Ana B. Muñoz-Manchado, Sten Linnarsson, Greg Crawford, and et al. 2019. Biological annotation of genetic loci associated with intelligence in a meta-analysis of 87,740 individuals. *Molecular Psychiatry* 24: 182–97. [CrossRef]

Cowan, Nelson. 2017. The many faces of working memory and short-term storage. *Psychonomic Bulletin & Review* 24: 1158–70. [CrossRef]

Currie, Thomas E., Peter Turchin, Edward Turner, and Sergey Gavrilets. 2020. Duration of agriculture and distance from the steppe predict the evolution of large-scale human societies in Afro-Eurasia. *Humanities and Social Sciences Communications* 7: 34. [CrossRef]

Darwin, Charles, and Alfred Wallace. 1858. On the tendency of species to form varieties, and on the perpetuation of varieties and species by natural means of selection. *Journal of the Linnean Society of London, Zoology* 3: 45–62. [CrossRef]

De Boeck, Paul, and Kristof Kovacs. 2020. The many faces of intelligence: A discussion of Geary’s mitochondrial functioning theory on general intelligence. *Journal of Intelligence* 8: 8. [CrossRef] [PubMed]

de Mello, Aline H., Ana B. Costa, Jessica D. G. Engel, and Gislaine T. Rezin. 2018. Mitochondrial dysfunction in obesity. *Life Sciences* 192: 26–32. [CrossRef]

Deary, Ian J., Alexander Weiss, and G. David Batty. 2010. Intelligence and personality as predictors of illness and death: How researchers in differential psychology and chronic disease epidemiology are collaborating to understand and address health inequalities. *Psychological Science in the Public Interest* 11: 53–79. [CrossRef]

Debatin, Tobias. 2020. Neuroenergetics and “General Intelligence”: A systems biology perspective. *Journal of Intelligence* 8: 31. [CrossRef]

Dicke, Ursula, and Gerhard Roth. 2016. Neuronal factors determining high intelligence. *Philosophical Transactions of the Royal Society B: Biological Sciences* 371: 20150180. [CrossRef]

Dickens, William T., and James R. Flynn. 2001. Heritability estimates versus large environmental effects: The IQ paradox resolved. *Psychological Review* 108: 346–69. [CrossRef]

Dye, Louise, Neil B. Boyle, Claire Champ, and Clare Lawton. 2017. The relationship between obesity and cognitive health and decline. *Proceedings of the Nutrition Society* 76: 443–54. [CrossRef]

Edwards, Lindsay M., Andrew J. Murray, Cameron J. Holloway, Emma E. Carter, Graham J. Kemp, Ion Codreanu, Helen Brooker, Damian J. Tyler, Peter A. Robbins, and Kieran Clarke. 2011. Short-term consumption of a high-fat diet impairs whole-body efficiency and cognitive function in sedentary men. *The FASEB Journal* 25: 1088–96. [CrossRef] [PubMed]

Elton, Sarah, Laura C. Bishop, and Bernard Wood. 2001. Comparative context of Plio-Pleistocene hominin brain evolution. *Journal of Human Evolution* 41: 1–27. [CrossRef] [PubMed]
Engle, Randall W. 2002. Working memory capacity as executive attention. *Current Directions in Psychological Science* 11: 19–23. [CrossRef]

Eppig, Christopher, Corey L. Fincher, and Randy Thornhill. 2010. Parasite prevalence and the worldwide distribution of cognitive ability. *Proceedings of the Royal Society B: Biological Sciences* 277: 3801–8. [CrossRef] [PubMed]

Fanta, Vaclav, Miroslav Šálek, Jan Zouhar, Petr Sklenicka, and David Storch. 2018. Equilibrium dynamics of European pre-industrial populations: The evidence of carrying capacity in human agricultural societies. *Proceedings of the Royal Society B: Biological Sciences* 285: 20172500. [CrossRef] [PubMed]

Finsterer, J. 2012. Cognition dysfunction in mitochondrial disorders. *Acta Neurologica Scandinavica* 126: 1–11. [CrossRef]

Flinn, Mark V., David C. Geary, and Carol V. Ward. 2005. Ecological dominance, social competition, and coalitional arms races: Why humans evolved extraordinary intelligence. *Evolution and Human Behavior* 26: 10–46. [CrossRef]

Flynn, James R. 1984. The mean IQ of Americans: Massive gains 1932 to 1978. *Psychological Bulletin* 95: 29–51. [CrossRef]

Foley, Robert. 1987. Hominid species and stone-tool assemblages: How are they related. *Antiquity* 61: 380–92. [CrossRef]

Foley, Robert, and Marta M. Lahr. 1997. Mode 3 technologies and the evolution of modern humans. *Cambridge Archaeology Journal* 7: 3–36. [CrossRef]

Geary, David C. 2005. *The Origin of Mind: Evolution of Brain, Cognition, and General Intelligence*. Washington, DC: American Psychological Association.

Geary, David C. 2015. *Evolution of Vulnerability: Implications for sex Differences in Health and Development*. San Diego: Elsevier Academic Press.

Geary, David C. 2017. Evolution of human sex-specific cognitive vulnerabilities. *The Quarterly Review of Biology* 92: 361–410. [CrossRef]

Geary, David C. 2018. Efficiency of mitochondrial functioning as the fundamental biological mechanism of general intelligence (g). *Psychological Review* 125: 1028–50. [CrossRef] [PubMed]

Gignac, Gilles E., and Timothy C. Bates. 2017. Brain volume and intelligence: The moderating role of intelligence measurement quality. *Intelligence* 64: 18–29. [CrossRef]

Gong, Yun, Jonathan Greenbaum, and Hong-Wen Deng. 2019. A statistical approach to fine-mapping for the identification of potential causal variants related to human intelligence. *Journal of Human Genetics* 64: 781–87. [CrossRef]

Havird, Justin C., and Daniel B. Sloan. 2016. The roles of mutation, selection, and expression in determining relative rates of evolution in mitochondrial versus nuclear genomes. *Molecular Biology and Evolution* 33: 3042–53. [CrossRef]

Hermann, Raphael, Andrea Dolfini, Rachel J. Crellin, Quanyu Wang, and Marion Uckelmann. 2020. Bronze age swordsmanship: New insights from experiments and wear analysis. *Journal of Archaeological Method and Theory*. [CrossRef]

Hill, Geoffre E. 2014. Cellular respiration: The nexus of stress, condition, and ornamentation. *Integrative and Comparative Biology* 54: 645–57. [CrossRef]

Hirschfeld, Katherine. 2015. *Gangster States: Organized Crime, Kleptocracy and Political Collapse*. New York: Palgrave-MacMillan.
Hulgan, Todd, Asha R. Kallianpur, Yan Guo, Jill S. Barnholtz-Sloan, Haley Gittleman, Todd T. Brown, Ronald Ellis, Scott Letendre, Robert K. Heaton, David C. Samuels, and et al. 2019. Peripheral blood mitochondrial DNA copy number obtained from genome-wide genotype data is associated with neurocognitive impairment in persons with chronic HIV infection. *JAIDS Journal of Acquired Immune Deficiency Syndromes* 80: e95–102. [CrossRef] [PubMed]

Iozzo, Patricia, and Maria A. Guzzardi. 2019. Imaging of brain glucose uptake by PET in obesity and cognitive dysfunction: Life-course perspective. *Endocrine Connections* 8: R169–83. [CrossRef] [PubMed]

Jensen, Arthur R. 1998. *The g Factor: The Science of Mental Ability*. Westport: Praeger.

Johnson, Wendy, Andrew Carothers, and Ian J. Deary. 2008. Sex differences in variability in general intelligence: A new look at the old question. *Perspectives on Psychological Science* 3: 518–31. [CrossRef] [PubMed]

Jung, Rex E., and Richard J. Haier. 2007. The Parieto-Frontal Integration Theory (P-FIT) of intelligence: Converging neuroimaging evidence. *Behavioral and Brain Sciences* 30: 135–54. [CrossRef] [PubMed]

Kanazawa, Satoshi. 2004. General intelligence as a domain-specific adaptation. *Psychological Review* 111: 512–23. [CrossRef]

Kanazawa, Satoshi. 2008. Temperature and evolutionary novelty as forces behind the evolution of general intelligence. *Intelligence* 36: 99–108. [CrossRef]

Kaplan, Hillard, Kim Hill, Jane Lancaster, and A. Magdelana Hurtado. 2000. A theory of human life history evolution: Diet, intelligence, and longevity. *Evolutionary Anthropology* 9: 156–85. [CrossRef]

Koch, Rebecca E., Chloe C. Josefson, and Geoffrey E. Hill. 2017. Mitochondrial function, ornamentation, and immunocompetence. *Biological Reviews* 92: 1459–74. [CrossRef]

Kuczynski, B., W. Jagust, H. C. Chui, and B. Reed. 2009. An inverse association of cardiovascular risk and frontal lobe glucose metabolism. *Neurology* 72: 738–43. [CrossRef]

Lee, James J., Matt McGue, William G. Iacono, Andrew M. Michael, and Christopher F. Chabris. 2019. The causal influence of brain size on human intelligence: Evidence from within-family phenotypic associations and GWAS modeling. *Intelligence* 75: 48–58. [CrossRef]

Lee, Ji-Won, Ki D. Park, Jee-Aee Im, Moo Y. Kim, and Duk-Chul Lee. 2010. Mitochondrial DNA copy number in peripheral blood is associated with cognitive function in apparently healthy elderly women. *Clinica Chimica Acta* 411: 592–96. [CrossRef] [PubMed]

Leonard, William R., and Marcia L. Robertson. 1994. Evolutionary perspectives on human nutrition: The influence of brain and body size on diet and metabolism. *American Journal of Human Biology* 6: 77–88. [CrossRef] [PubMed]

Levine, Morgan E., Amal Harrati, and Eileen M. Crimmins. 2018. Predictors and implications of accelerated cognitive aging. *Biodemography and Social Biology* 64: 83–101. [CrossRef] [PubMed]

Longchamps, Ryan J., Christina A. Castellani, Stephanie Y. Yang, Charles E. Newcomb, Jason A. Sumpter, John Lane, Megan L. Grove, Eliseo Guallar, Nathan Pankratz, Kent D. Taylor, and et al. 2020. Evaluation of mitochondrial DNA copy number estimation techniques. *PLoS ONE* 15: e0228166. [CrossRef]

MacArthur, Robert H., and Edward O. Wilson. 1967. *The Theory of Island Biogeography*. Princeton: Princeton University Press.

Malthus, T. Robert. 1798. *An Essay on the Principle of Population as it Affects the Future Improvement of Society with Remarks on the Speculations of Mr. Godwin, M. Condorcet, and Other Writers*. London: Printed for J. Johnson, in St. Paul’s church-yard.

Martin, Paul S. 1967. Prehistoric overkill. In *The Search for a Cause*. Edited by Paul S. Martin and H. E. Wright. New Haven: Yale University Press.

Martin, Paul S. 1973. The discovery of America: The first Americans may have swept the Western Hemisphere and decimated its fauna in 1000 years. *Science* 179: 969–74. [CrossRef]

Mattson, Mark P. 2012. Energy intake and exercise as determinants of brain health and vulnerability to injury and disease. *Cell Metabolism* 16: 706–22. [CrossRef]

Matzel, Louis D., Dylan W. Crawford, and Bruno Sauce. 2020. Déjà vu all over again: A unitary biological mechanism for intelligence is (probably) untenable. *Journal of Intelligence* 8: 24. [CrossRef]

McHenry, Henry M. 1994. Tempo and mode in human evolution. *Proceedings of the National Academy of Sciences of the United States of America* 91: 6780–86. [CrossRef]
Mengel-From, Jonas, Mikael Thinggaard, Christine Dalgård, Kirsten O. Kyvik, Kaare Christensen, and Lene Christiansen. 2014. Mitochondrial DNA copy number in peripheral blood cells declines with age and is associated with general health among elderly. *Human Genetics* 133: 1149–59. [CrossRef]

Montier, Laura L. C., Janice J. Deng, and Yidong Bai. 2009. Number matters: Control of mammalian mitochondrial DNA copy number. *Journal of Genetics and Genomics* 36: 125–31. [CrossRef]

Moore, H. L., A. P. Blain, D. M. Turnbull, and G. S. Gorman. 2020. Systematic review of cognitive deficits in adult mitochondrial disease. *European Journal of Neurology* 27: 3–17. [CrossRef] [PubMed]

Moore, Heather L., Thomas Kelly, Alexandra Bright, Robert H. Field, Andrew M. Schaefer, Alasdair P. Blain, Robert W. Taylor, Robert McFarland, Doug M. Turnbull, and Gráinne S. Gorman. 2019. Cognitive deficits in adult m. 3243A>G and m. 8344A>G-related mitochondrial disease: Importance of correcting for baseline intellectual ability. *Annals of Clinical and Translational Neurology* 6: 826–36. [CrossRef] [PubMed]

Napoli, Nicola, Krupa Shah, Debra L. Waters, David R. Sinacore, Clifford Qualls, and Dennis T. Villareal. 2014. Effect of weight loss, exercise, or both on cognition and quality of life in obese older adults. *The American Journal of Clinical Nutrition* 100: 189–98. [CrossRef] [PubMed]

Ngandu, Tiia, Jenni Lehtisalo, Alina Solomon, Esko Levalahti, Satu Ahtiluoto, Riitta Antikainen, Lars Bäckman, Tuomo Hänninen, Antti Jula, Tiina Laatikainen, and et al. 2015. A 2 year multidomain intervention of diet, exercise, cognitive training, and vascular risk monitoring versus control to prevent cognitive decline in at-risk elderly people (FINGER): A randomised controlled trial. *The Lancet* 385: 2255–63. [CrossRef]

O’Brien, Phillipe D., Lucy M. Hinder, Brian C. Callaghan, and Eva L. Feldman. 2017. Neurological consequences of obesity. *The Lancet Neurology* 16: 465–77. [CrossRef]

Pallier, Gerry, Richard D. Roberts, and Lazar Stankov. 2000. Biological versus psychometric intelligence: Halstead’s (1947) distinction revisited. *Archives of Clinical Neuropsychology* 15: 205–26. [CrossRef]

Pellerin, Luc, and J. Pierre. 1994. Magistretti. Glutamate uptake into astrocytes stimulates aerobic glycolysis: A mechanism coupling neuronal activity to glucose utilization. *Proceedings of the National Academy of Sciences of the United States of America* 91: 10625–29. [CrossRef]

Picard, Martin, and Doug M. Turnbull. 2013. Linking the metabolic state and mitochondrial DNA in chronic disease, health, and aging. *Diabetes* 62: 672–78. [CrossRef]

Picard, Martin, Caroline Trumpff, and Yan Burelle. 2019. Mitochondrial psychobiology: Foundations and applications. *Current Opinion in Behavioral Sciences* 28: 142–51. [CrossRef]

Picard, Martin, Robert-Paul Juster, and Bruce S. McEwen. 2014. Mitochondrial allostatic load puts The ‘gluc’ back in glucocorticoids. *Nature Reviews Endocrinology* 10: 303–10. [CrossRef]

Pietschnig, Jakob, and Martin Voracek. 2015. One century of global IQ gains: A formal meta-analysis of the Flynn effect (1909–2013). *Perspectives on Psychological Science* 10: 282–306. [CrossRef] [PubMed]

Pietschnig, Jakob, and Martin Voracek. 2015. One century of global IQ gains: A formal meta-analysis of the Flynn effect (1909–2013). *Perspectives on Psychological Science* 10: 282–306. [CrossRef] [PubMed]

Potts, Richard. 1998. Variability selection in hominid evolution. *Evolutionary Anthropology* 7: 81–96. [CrossRef]

Protzko, John. 2017. Raising IQ among school-aged children: Five meta-analyses and a review of randomized controlled trials. *Developmental Review* 46: 81–101. [CrossRef]

Pugazhenthi, Subbiah, Limei Qin, and P. Hemachandra Reddy. 2017. Common neurodegenerative pathways in obesity, diabetes, and Alzheimer’s disease. *Biochimica et Biophysica Acta (BBA)-Molecular Basis of Disease* 1863: 1037–45. [CrossRef] [PubMed]

Rae, Caroline, Alison L. Digney, Sally R. McEwan, and Timothy C. Bates. 2003. Oral creatine monohydrate supplementation improves brain performance: A double-blind, placebo–controlled, cross–over trial. *Proceedings of the Royal Society of London. Series B: Biological Sciences* 270: 2147–50. [CrossRef]

Richerson, Peter J., Robert Boyd, and Robert L. Bettinger. 2001. Was agriculture impossible during the Pleistocene but mandatory during the Holocene? A climate change hypothesis. *American Antiquity* 66: 387–411. [CrossRef]

Sauce, Bruno, and Louis D. Matzel. 2018. The paradox of intelligence: Heritability and malleability coexist in hidden gene-environment interplay. *Psychological Bulletin* 144: 26–47. [CrossRef]

Savi, Alexander O., Han L. J. van der Maas, Gunter K. Maris, and Maarten Marsman. 2020. Mitochondrial Functioning & general intelligence. *Journal of Intelligence* 8: 20. [CrossRef]

Schmidt, Reinhold, Helena Schmidt, J. David Curb, Kamal Masaki, Lon R. White, and Lenore J. Launer. 2002. Early inflammation and dementia: A 25-year follow-up of the Honolulu-Asia Aging Study. *Annals of Neurology* 52: 168–74. [CrossRef]
Schubert, Anna-Lena, and Dirk Hagermann. 2020. The evidence for Geary’s theory on the role of mitochondrial functioning in human intelligence is not entirely convincing. Journal of Intelligence 8: 29. [CrossRef]

Set, Kalloil K., Kuntal Sen, A. H. M. Huq, and Rajkumar Agarwal. 2019. Mitochondrial disorders of the nervous system: A review. Clinical Pediatrics 58: 381–94. [CrossRef] [PubMed]

Shemesh, Elad, Assaf Rudich, Ilana Harman-Boehm, and Tali Cukierman-Yaffe. 2012. Effect of intranasal insulin on cognitive function: A systematic review. The Journal of Clinical Endocrinology & Metabolism 97: 366–76. [CrossRef]

Shields, Grant S., Wesley G. Moons, and George M. Slavich. 2017. Inflammation, self-regulation, and health: An immunologic model of self-regulatory failure. Perspectives on Psychological Science 12: 588–612. [CrossRef]

Silzer, Talisa, Robert Barber, Jie Sun, Gita Pathak, Leigh Johnson, Sid O’Bryant, and Nicole Phillips. 2019. Circulating mitochondrial DNA: New indices of type 2 diabetes-related cognitive impairment in Mexican Americans. PloS ONE 14: e0213527. [CrossRef] [PubMed]

Singh-Manoux, Archana, Aline Dugravot, Eric Brunner, Meena Kumari, Martin Shipley, Alexis Elbaz, and Mika Kivimaki. 2014. Interleukin-6 and C-reactive protein as predictors of cognitive decline in late midlife. Neurology 83: 486–93. [CrossRef] [PubMed]

Smith, Patrick J., James A. Blumenthal, Benson M. Hoffman, Harris Cooper, Timothy A. Strauman, Kathleen Welsh-Bohmer, Jeffrey N. Browndyke, and Andrew Sherwood. 2010. Aerobic exercise and neurocognitive performance: A meta-analytic review of randomized controlled trials. Psychosomatic Medicine 72: 239–52. [CrossRef]

Spitznagel, Mary B., Misty Hawkins, Michael Alosco, Rachel Galioto, Sarah Garcia, Lindsay Miller, and John Gunstad. 2015. Neurocognitive effects of obesity and bariatric surgery. European Eating Disorders Review 23: 488–95. [CrossRef]

Spyridaki, Eirini C., Panagiotis Simos, Pavlina D. Avgoustinaki, Eirini Dermitzaki, Maria Venihaki, Achilles N. Bardos, and Andrew N. Margioris. 2014. The association between obesity and fluid intelligence impairment is mediated by chronic low-grade inflammation. British Journal of Nutrition 112: 1724–34. [CrossRef]

Spyridaki, Eirini C., Pavlina D. Avgoustinaki, and Andrew N. Margioris. 2016. Obesity, inflammation and cognition. Current Opinion in Behavioral Sciences 9: 169–75. [CrossRef]

Sripetchwandee, Jirapas, Nipon Chattipakorn, and Siriporn C. Chattipakorn. 2018. Links between obesity-induced brain insulin resistance, brain mitochondrial dysfunction, and dementia. Frontiers in Endocrinology 9: 496. [CrossRef]

Stankov, Lazar. 2020. “Turtles-all-the-way-down” from g to mitochondrial functioning. Journal of Intelligence 8: 23. [CrossRef] [PubMed]

Sternberg, Robert J. 2020. How mighty are the mitochondria in causing individual differences in intelligence?—Some questions for David Geary. Journal of Intelligence 8: 13. [CrossRef] [PubMed]

Stevenson, Anna J., Daniel L. McCartney, Robert F. Hillary, Paul Redmond, Adele M. Taylor, Qian Zhang, and et al. 2019. Childhood intelligence attenuates the association between biological ageing and health outcomes in later life. Translational Psychiatry 9: 1–8. [CrossRef] [PubMed]

Turchin, Peter. 2009. A theory for formation of large empires. Journal of Global History 4: 191–217. [CrossRef]

Turchin, Peter, Thomas E. Currie, Edward A. Turner, and Sergey Gavrilets. 2013. War, space, and the evolution of Old World complex societies. Proceedings of the National Academy of Sciences of the United States of America 110: 16384–389. [CrossRef]

Ujma, Péter Przemyslaw, and Kristof Kovacs. 2020. The mitochondrial theory of g is incompatible with genetic evidence and does not explain statistical phenomena. Journal of Intelligence 8: 27. [CrossRef]

Veronese, Nicola, Silvia Facchini, Brendon Stubbs, Claudio Luchini, Marco Solmi, Enzo Manzato, Giuseppe Sergi, Stefania Maggi, Theodore Cosco, and Luigi Fontana. 2017. Weight loss is associated with improvements in cognitive function among overweight and obese people: A systematic review and meta-analysis. Neuroscience & Biobehavioral Reviews 72: 87–94. [CrossRef]
Volkow, Nora D., Gene-Jack Wang, Frank Telang, Joanna S. Fowler, Rita Z. Goldstein, Nelly Alia-Klein, and et al. 2009. Inverse association between BMI and prefrontal metabolic activity in healthy adults. *Obesity* 17: 60–65. [CrossRef]

Vrba, Elizabeth S. 1995. The fossil record of African antelopes (Mammalia, Bovidae) in relation to human evolution and paleoclimate. In *Paleoclimate and Evolution, with Emphasis on Human Origins*. Edited by Elizabeth S. Vrba, George H. Denton, Timothy C. Partridge and Lloyd H. Burckle. New Haven: Yale University Press.

Wachsmuth, Manja, Alexander Huebner, Mingkun Li, Burkhard Madea, and Mark Stoneking. 2016. Age-related and heteroplasmy-related variation in human mtDNA copy number. *PLoS Genetics* 12: e1005939. [CrossRef] [PubMed]

Warne, Russell T., and Cassidy Burningham. 2019. Spearman’s g found in 31 non-Western nations: Strong evidence that g is a universal phenomenon. *Psychological Bulletin* 145: 237–72. [CrossRef]

Whitley, Elise, Catharine R. Gale, Ian J. Deary, Mika Kivimaki, and G. David Batty. 2011. Association of maternal and paternal IQ with offspring conduct, emotional, and attention problem scores: transgenerational evidence from the 1958 British birth cohort study. *Archives of General Psychiatry* 68: 1032–38. [CrossRef] [PubMed]

Willette, Auriel A., Barbara B. Bendlin, Erika J. Starks, Alex C. Birdsill, Sterling C. Johnson, Bradley T. Christian, Ozioma C. Okonkwo, Asenath La Rue, Bruce P. Hermann, Rebecca L. Koscik, and et al. 2015. Association of insulin resistance with cerebral glucose uptake in late middle-aged adults at risk for Alzheimer disease. *JAMA Neurology* 72: 1013–20. [CrossRef]

Wingo, Aliza P., Eric B. Dammer, Michael S. Breen, Benjamin A. Logsdon, Duc M. Duong, Juan C. Troncosco, Madhav Thambisetty, Thomas G. Beach, Geidy E. Serrano, Eric M. Reiman, and et al. 2019. Large-scale proteomic analysis of human brain identifies proteins associated with cognitive trajectory in advanced age. *Nature Communications* 10: 1–14. [CrossRef] [PubMed]

Yaffe, Kristene, Alka Kanaya, Karla Lindquist, Eleanor M. Simonsick, Tamara Harris, Ronald I. Shorr, and et al. 2004. The metabolic syndrome, inflammation, and risk of cognitive decline. *JAMA* 292: 2237–42. [CrossRef]

Yang, Yingkai, Grant S. Shields, Cheng Guo, and Yanling Liu. 2018. Executive function performance in obesity and overweight individuals: A meta-analysis and review. *Neuroscience & Biobehavioral Reviews* 84: 225–44. [CrossRef]

Zeng, Tian C., Alan J. Aw, and Marcus W. Feldman. 2018. Cultural hitchhiking and competition between patrilineal kin groups explain the post-Neolithic Y-chromosome bottleneck. *Nature Communications* 9: 2077. [CrossRef]

Zerjal, Tatiana, Yali Xue, Giorgio Bertorelle, R. Spencer Wells, Weidong Bao, Suling Zhu, Raheel Qamar, Qasim Ayub, Aisha Mohyuddin, Songbin Fu, and et al. 2003. The genetic legacy of the Mongols. *American Journal of Human Genetics* 72: 717–21. [CrossRef]

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