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Estimating the Additive Heritability of Historiometric Eminence in a Super-Pedigree Comprised of Four Prominent Families

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Abstract

By merging analytical approaches from the fields of historiometrics and behavior genetics, a social pedigree-based estimate of the heritability of eminence is generated. Eminent individuals are identified using the *Pantheon* dataset. A single super-pedigree, comprised of four prominent and interrelated families (including the Wedgwood–Darwin, Arnold–Huxley, Keynes–Baha'u'lláh, and Benn–Rutherford pedigrees) is assembled, containing 30 eminent individuals out of 301 in total. Each eminent individual in the super-pedigree is assigned a relative measure of historical eminence (scaled from 1 to 100) with noneminent individuals assigned a score of 0. Utilizing a Bayesian pedigree-based heritability estimation procedure employing an informed prior, an additive heritability of eminence of .507 (95% CI [.434, .578]) was found. The finding that eminence is additively heritable is consistent with expectations from behavior-genetic studies of factors that are thought to underlie extraordinary accomplishment, which indicate that they are substantially additively heritable. Owing to the limited types of intermarriage present in the data, it was not possible to estimate the impact of nonadditive genetic contributions to heritability. Gene-by-environment interactions could not be estimated in the present analysis either; therefore, the finding that eminence is simply a function of additive genetic and nonshared environmental variance should be interpreted cautiously.

Keywords: Eminence; heritability; historiometrics; pedigree analysis

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Does eminence or extraordinary accomplishment owe more to ‘nature’ or ‘nurture’? This was the central question addressed by Francis Galton in his 1869 work *Hereditary Genius*. In doing so, Galton effectively created two fields, *historiometrics* (or *historiometry*) and *behavior genetics*. Historiometrics concerns itself with estimating the relative significance of a given cultural, political or scientific development, and the relative eminence of the individuals responsible for these developments, in addition to ascertaining their determinants (Simonton, 1984). This is accomplished using encyclopedias, biographies and other relatively neutral reference works as the basis for quantifying these properties (Simonton, 1984). Galton applied a crude form of historiometrics to the identification of extraordinary talent or eminence across a number of domains (science, industry, leadership and sports) within his own and in other notable British families — Galton believed that the assessment of individuals’ accomplishments by relevantly informed others offered the best basis for identifying eminence. Behavior genetics concerns itself with the measurement of the degree to which hereditary genetic and environmental factors (what Galton [1874] subsequently termed *nature* and *nurture*, respectively) influence variation with respect to a given behavioral or cognitive phenotype. In *Hereditary Genius*, Galton proposed

that concentrations of eminence within pedigrees could be employed to demonstrate the contribution of hereditary factors to variation in this trait.¹ In tracking the prevalence of eminent individuals across multigenerational pedigrees (including his own), Galton concluded that the elements of eminence were indeed hereditary. However, he was unable to provide either a quantitative estimate of their heritability or adequately control for environmental confounds, a point made by Augustin de Candolle (1873) who was a major critic of Galton’s approach.

Modern historiometrics (much of the groundwork for which was laid by Dean Keith Simonton) allows for relatively rigorous² quantification of the eminence of prominent individuals — this is achieved by determining the degree to which various neutral reference works converge with one another with respect to their coverage of prospectively eminent individuals and their accomplishments. For example, a scientist whose name appears in one list of field-specific significant figures, but not in others, would be less eminent than one whose name appears in multiple independently compiled lists. Similarly, a scientist who features in general reference works (such as encyclopedias), in addition to field-specific ones, is more eminent than the one who has field-specific, but not general, prominence. In the extreme, the most accomplished and influential individuals will be sufficiently well known that they are household names, becoming known simply by their surnames (e.g., Darwin, Einstein, Freud, Marx, Mozart; Murray, 2003).³ Historiometrically defined eminence is necessarily a *domain-general* phenomenon — encompassing extraordinary achievement across many different areas of accomplishment

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(e.g., STEM, politics, arts, literature, and sport). The historiometric approach is nevertheless also informative about the *domain-specific* dispositions of eminent individuals, since relevant biographic data are often rich in details that are reflective of the cognitive, conative, and situational factors that incline such individuals toward specific degrees and domains of accomplishment (Simonton, 1984, 2009, 2018).⁴

In terms of its broad determinants, Galton (1869) described eminence as the product of a ‘triple event, of ability, combined with zeal and with capacity for hard labor’ (p. 78). Johnson and Bouchard Jr. (2014) note that all of the elements comprising Galton’s ‘triple event’ are additively heritable; thus, it might be expected that historiometric eminence (presumably being a joint product of high levels of these elements) will itself be heritable. However, defining the phenotype of eminence for the purposes of behavior-genetic measurement models presents certain interesting challenges.

First, there is the problem of the sorts of historiometric thresholds that need to be met when identifying eminent individuals. Different researchers have different opinions on this matter. Charles Murray in his book *Human Accomplishment* (2003) employs a two-step procedure for identifying those whom he terms *significant figures* in the domains of science, technology, medicine, arts, literature, and philosophy (with such individuals being sourced from both Western and non-Western populations). Murray (2003) initially defines significant figures as those who are (a) featured in at minimum 50% of the various ‘qualified’ reference works that he consulted and (b) realized their accomplishments no later than 1950, so as to avoid ‘epochcentric’ or recency bias effects. Once this cutoff is achieved, such individuals can then be assigned an index score, which is a measure of their relative eminence based on that individual’s comparative prominence across the relevant reference works within a given inventory. This is essentially a function of the amount of text devoted to that individual across reference works (among which there is typically very high agreement), which is then normalized on a scale such that 1 is the lowest score and 100 is the highest score. This process resulted in the inclusion of 4002 significant figures in *Human Accomplishment*.

Yu and colleagues (2016) have generated an alternative global database of eminent individuals termed *Pantheon*, which is much broader than the one assembled by Murray, as it also includes politicians, economists, actors, and athletes among other categories, with no restrictions on year of accomplishment. As with Murray (2003), this dataset also employs a two-step inclusion criteria, with the first step being that for an individual to be included, they must have a *Wikipedia* page that has been translated into a minimum of (in the current version of the dataset) 15 languages. Earlier versions of the dataset had stricter inclusion rules (e.g., >25 languages in version 1.0; Yu et al., 2016). The relaxation of this inclusion rule led to an increase in the number of eminent individuals (what *Pantheon* terms *memorable people*) from 11,341 (in the earliest version of the dataset) to 88,506 (in the current version). The second step involves assigning each individual a historical popularity index (HPI) value, which is a measure of biographic prominence generated by combining information on the number of languages in which an eminent individual’s *Wikipedia* biography is available, with page views, and the amount of time that has elapsed since the individual’s birth. Epochcentric bias is controlled by adjusting the HPI values of individuals who have been known for <70 years. Additional weights are used to compensate for English-language bias. As with Murray’s (2003) index score, the HPI is also

normalized, such that 100 is the highest score and 1 is the lowest. The earliest version of *Pantheon* contained HPI values for 1570 individuals in the arts and sciences who overlapped with those included in Murray’s (2003) *Human Accomplishment*, which allowed Yu et al. (2016) to cross-validate their HPI values with respect to the latter’s index scores. They found that the two correlate modestly at .35 (95% C.I. [.31, .39]), which is consistent with the broader tendency noted by Murray (2003) for various neutral reference works to converge with one another in terms of their degree of coverage of eminent individuals.

The Murray (2003) and Yu et al. (2016) databases could, by virtue of the differential breadth of their respective inclusion criteria, be said to yield *thin* versus *thick* measures of eminence, respectively.

The second problem concerns the rarity of eminence. Families containing one or more eminent twins are doubtlessly very rare, making the most straightforward behavior-genetic approaches to estimating the heritability of this phenotype untenable. But as Galton (1869) noted, historiometric eminence seemed to run in his own family and other prominent families. As pedigree data can be used to estimate heritabilities, such data can be employed to quantify the heritability of eminence among related eminent individuals defined based on historiometric criteria.

Methods

Eminence Measure

As eminent individuals are relatively rare, it was decided that using individuals sourced from the latest (2020) release of the thicker Yu et al. (2016) *Pantheon* dataset (<https://pantheon.world/data/datasets>) would be best, as this would allow for the largest number of eminent individuals to be identified within the super-pedigree described below, thus increasing the power available to estimate heritability. In assigning eminent individuals to the super-pedigree, each individual is assigned their respective their HPI value (as estimated in the current release of *Pantheon*⁵). Those who are noneminent are assigned values of 0, thus the model employed here is somewhat like a case-control design, in that a ‘liability threshold’ of biographic prominence has to be attained for an individual to get a nonzero degree of eminence; however, among the eminent, there is significant variation in terms of relative degree of eminence that provides the models used here with the variance necessary to establish stable heritability estimates. The possibility of using a strict case-control design to estimate heritability (i.e., where eminent individuals are assigned a value of 1 and noneminent individuals a value of 0) was also explored; however, the resultant model did not converge. This is consistent with the very low phenotypic variance associated with strict case-control designs that necessarily increases the error associated with parameter estimation. The use of a partial case-control model still has the benefit of bringing into sharp relief the most salient phenotypic difference, that is, whether the individual is eminent as measured by biographical prominence or not.

Super-Pedigree

Genealogical data were collected for a total of 301 individuals comprising a social (genealogical, nongenetic) super-pedigree involving four prominent and related British (and one British-Persian) families. The core of the super-pedigree was the Wedgwood–Darwin–Galton–Williams (henceforth Wedgwood–Darwin) pedigree, which includes three eminent Wedgwoods (Josiah

Table 1. Thirty eminent individuals sourced from *Pantheon* and contained within the super-pedigree, along with their birth and (where applicable) death years and accomplishment domains

Eminent individual (birth and death year)	Historical Popularity Index	Accomplishment domain
Josiah Wedgwood I (1730–1795)	66.56	Design
Thomas Wedgwood (1771–1805)	60.85	Invention
Emma Wedgwood (1808–1896)	65.66	Politics
Erasmus Darwin (1731–1802)	73.48	Biology
Robert Waring Darwin (1766–1848)	65.98	Medicine
Charles Robert Darwin (1809–1882)	93.29	Biology
Charles Galton Darwin (1887–1962)	63.51	Physics
George Howard Darwin (1845–1912)	70.6	Astronomy
Francis Darwin (1848–1925)	62.42	Biology
Francis Galton (1822–1911)	76.98	Statistics
Ralph Vaughan Williams (1872–1958)	69.33	Composing
Thomas Henry Huxley (1825–1895)	78.23	Biology
Aldous Huxley (1894–1963)	80.72	Writing
Andrew Huxley* (1917–2012)	69.51	Physics
Julian Huxley (1887–1975)	71.68	Biology
John Maler Collier (1850–1934)	65.78	Painting
Thomas Arnold the Elder (1798–1842)	65.24	Education§
Matthew Arnold (1822–1888)	65.44	Writing
John Maynard Keynes (1883–1946)	85.97	Economics
Alexander Skandar Keynes (1991–present)	53.96	Acting
Lydia Lopokova (1892–1981)	61.64	Dance†
Archibald Hill* (1886–1977)	69.77	Biology
Edgar Adrian* (1889–1977)	69.73	Medicine
Baha'u'lláh (1817–1892)	77.73	Religion
'Abdu'l-Bahá (1844–1921)	69.39	Religion
Shoghi Effendi (1897–1957)	66.68	Religion
Subh-i-Azal (1831–1912)	64.00	Religion‡
Anthony Wedgwood Benn (1925–2014)	57.87	Politics
Hilary James Benn (1953–present)	45.17	Politics
Margaret Rutherford (1892–1972)	68.41	Acting

Note: *The individual won a Nobel Prize. †*Pantheon* (as of July 8, 2021) incorrectly lists Lopokova as a painter, when she was an eminent ballerina. ‡*Pantheon* incorrectly lists Subh-i-Azal as a writer, when he was a religious leader. §*Pantheon* incorrectly lists Thomas Arnold as a philosopher when he was a prominent educator.

Wedgwood I, Thomas Wedgwood, and Emma Wedgwood⁶), six eminent Darwins (Erasmus Darwin, Robert Waring Darwin, Charles Robert Darwin, Charles Galton Darwin, George Howard Darwin, and Francis Darwin), in addition to Francis Galton and Ralph Vaughan Williams. The Wedgwood–Darwin pedigree employed here was compiled by Harry H. Laughlin in 1932, with the full data being republished in Berra et al. (2010). Another prominent pedigree is the Arnold–Huxley–Collier (henceforth Arnold–Huxley) pedigree, the members of which are related to the Wedgwood–Darwin pedigree via two instances of cross-marriage.⁷ This pedigree contains seven eminent individuals: Thomas Henry Huxley, Aldous Huxley, Julian Huxley, Andrew Huxley, Thomas Arnold the Elder, his son Matthew Arnold, and John Collier, who married two of Thomas Henry Huxley's daughters. Data sourced from *Wikipedia* on the Darwin–Wedgwood and Arnold–Huxley family trees (*Wikipedia*, 2021a,

2021b) were used to enhance each pedigree and identify the linkages necessary for merger of both into a common super-pedigree. Another prominent family is the Keynes–Lopokova–Hill–Adrian–Baha'u'lláh (henceforth Keynes–Baha'u'lláh) pedigree, which is connected to the Wedgwood–Darwin pedigree via one instance of cross-marriage.⁸ This pedigree contains seven eminent individuals, John Maynard Keynes, Alexander Skandar Keynes, Lydia Lopokova,⁹ Archibald Hill (who married John Maynard Keynes' sister, Margaret), Edgar Adrian, Baha'u'lláh,¹⁰ his half-brother Subh-i-Azal,¹¹ his eldest son 'Abd'u'l-Bahá, and his great-grandson Shoghi Effendi. As with the Arnold–Huxley pedigree, data permitting integration into the super-pedigree were obtained from the relevant significant figures' *Wikipedia* pages. The final constituent family in the super-pedigree is the Benn–Rutherford pedigree. This pedigree contains three closely related eminent individuals (Anthony Wedgwood Benn, Hilary James Benn, and Margaret

1 Robert Galton	101 Sarah Wedgwood	201 Joseph Fourier
2 John Galton I (Mother)	102 Sarah Wedgwood	202 John Fourier
3 John Galton I	103 John Bartlett	203 John Fourier
4 John Bruce	104 John Bartlett	204 John Fourier
5 Margaret	105 John B. Allen	205 John Fourier
6 Elizabeth Bruce	106 John B. Allen	206 Rosina Karlovna Douglas
7 John Frane	107 Samuel Galton III	207 John Fourier
8 John de Bussell (Mother)	108 Victoria Darwin	208 Sir John William Benn (Father)
9 John de Bussell	109 Robert Wang Darwin	209 John Fourier
10 William Darwin I	110 Susanna Wedgwood	210 Harriet Frances Keyes
11 Mary Huxley	111 William Wedgwood	211 John Fourier
12 William Darwin II	112 John Wedgwood	212 Susan John Keyes
13 John Wedgwood	113 Elizabeth Wedgwood	213 Anne Frances Ashmole
14 Margaret Buxton	114 John Ashmole	214 Anne Frances Ashmole
15 Thomas Wedgwood I	115 Elizabeth Allen	215 Randall Halse Keyes
16 Susan Wedgwood II	116 Francis Galton	216 Randall Halse Keyes
17 Thomas	117 Francis Galton	217 Randall Halse Keyes
18 John Galton II	118 Henrietta Emma Darwin	218 Simon Keyes
19 Robert Bates	119 George Darwin	219 George Darwin
20 Sarah Barr	120 Francis Darwin	220 Alexander Shanks Keyes
21 Charles Darwin	121 George Darwin	221 Alexander Shanks Keyes
22 Robert Barclay of Ury	122 George Darwin	222 Cecil Fife Huxley
23 Charles Midgton	123 George Darwin	223 George Darwin
24 David Barclay	124 George Darwin	224 George Darwin
25 Robert Ury	125 Emma Wedgwood III	225 Huxley Ashmole
26 Robert Barclay I	126 Susanna Wedgwood	226 George Darwin
27 Phyllis Gould	127 Henry Wedgwood	227 George Darwin
28 Priscilla	128 Hensleigh Wedgwood	228 George Darwin
29 Anne Earle	129 Anne Wedgwood	229 Randall
30 William Darwin III	130 Anne Wedgwood	230 Randall
31 Anne Paul	131 Francis Mackintosh	231 Henry Ethelred Ashmole
32 Robert Waring	132 William Darwin Darwin	232 George Darwin
33 Charles Darwin	133 Catherine E. Wedgwood	233 George Darwin
34 Thomas Wedgwood II	134 George Darwin	234 Alexander
35 Thomas Wedgwood II	135 George Darwin	235 George Darwin
36 Margaret	136 George Darwin	236 George Darwin
37 Susan Wedgwood II	137 Anne Wedgwood	237 George Darwin
38 Thomas Wedgwood (Aron Wedgwood lineage)	138 John Wedgwood	238 George Darwin
39 Susan Darwin	139 Robert Darwin (Mother)	239 George Darwin
40 Susan Darwin	140 Robert Darwin	240 George Darwin
41 Susan Darwin	141 Robert Darwin	241 George Darwin
42 Susan Darwin	142 Robert Darwin	242 George Darwin
43 Susan Darwin	143 Robert Darwin	243 George Darwin
44 Susan Darwin	144 Robert Darwin	244 George Darwin
45 Susan Darwin	145 Robert Darwin	245 George Darwin
46 Susan Darwin	146 Robert Darwin	246 George Darwin
47 Susan Darwin	147 Robert Darwin	247 George Darwin
48 Susan Darwin	148 Robert Darwin	248 George Darwin
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96 Susan Darwin	196 Robert Darwin	296 George Darwin
97 Susan Darwin	197 Robert Darwin	297 George Darwin
98 Susan Darwin	198 Robert Darwin	298 George Darwin
99 Susan Darwin	199 Robert Darwin	299 George Darwin
100 Susan Darwin	200 Robert Darwin	300 George Darwin

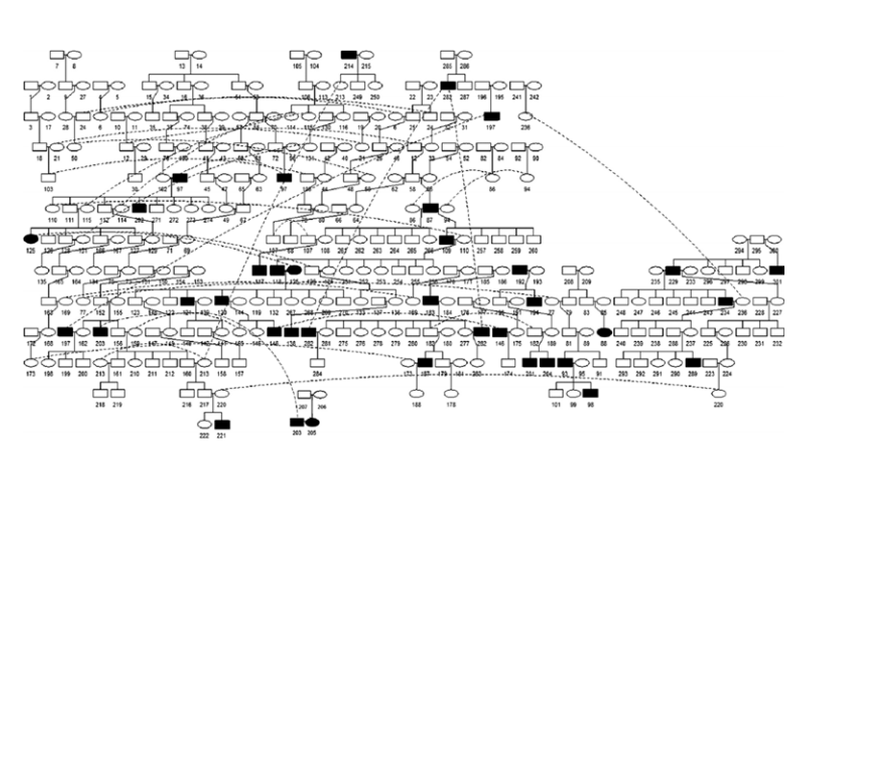


Figure 1. The super-pedigree. Shaded entries correspond to eminent individuals. Double lines indicate the presence of intermarriage (specifically marriage involving individuals related to one another to the degree of third cousin or closer). Dashed lines indicate the presence of the same individual in different parts of the super-pedigree. For example, Josiah Wedgwood III (126) is the husband of Caroline Darwin (124) and also the brother of Emma Wedgwood (125), Hensleigh Wedgwood (128), and Henry Wedgwood (127)

Rutherford), the first two whom are descended from the Wedgwood family via two distinct lines, both of which are collateral to the line of Josiah Wedgwood I. The precise nature of the connection between the Benn and Wedgwood families was ambiguous until recently, when the genealogist Maurice Frank was able to identify the relevant linkages (Frank, 2021). In total, there are 30 eminent individuals out of 301 (9.97%) comprising the super-pedigree. These individuals, along with their birth and (where applicable) death years, coupled with their (current as of this writing) HPI values and accomplishment domains (from their *Pantheon* listings) are shown in Table 1. The super-pedigree is presented in Figure 1 along with a table containing the names of all entrants.

Employing the thicker *Pantheon* database of eminent individuals yields both a more numerous and more diverse (in terms of accomplishment domains, nationality, and sex) sample of eminent individuals than does the thinner *Human Accomplishment* dataset. The latter contains only 8 eminent individuals, all of whom are male, British and whose accomplishments are restricted to science, mathematics, philosophy, literature, and the arts,¹² whereas the *Pantheon*-derived dataset contains data on 3 women (10% of the eminent sample) and 4 non-British (Persian) individuals (13.33% of the eminent sample), and spans 16 distinct accomplishment domains. Three of the individuals in the super-pedigree also won Nobel Prizes for scientific work (10% of the eminent sample).

Measurement Model

In addition to twin studies, geneticists have also adopted pedigree-based animal models to calculate heritability estimates. According

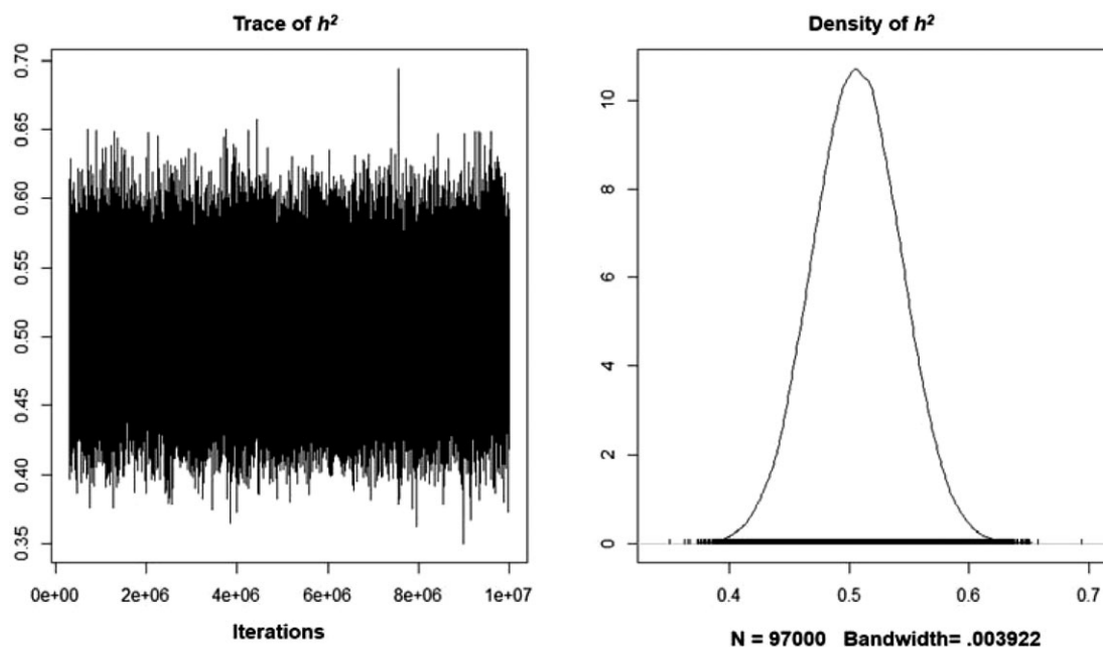
to Wilson et al. (2010), an animal model operates as a linear mixed-effects model (LMM) where the breeding value (the additive effect of a subject’s genotype upon a trait compared to the population’s average phenotype) is added to the statistical equation as a random effect. This term is included as a random effect because researchers do not know the breeding value for each individual in the sample. Moreover, per Wilson et al. (2010), random effects enable researchers to infer the effects’ distribution in a larger population. The implementation of this value as a random effect allows, via the computation of variance components, the estimation of the additive genetic variance (A) and the environmental variance (E). Thus, by computing these components, it is possible to calculate the trait’s narrow-sense heritability. Although animal models can be estimated as LMMs, geneticists have gravitated toward Bayesian estimation in the past decade.

In this study, several Markov Chain Monte Carlo (MCMC) generalized linear mixed models were computed with the R package *MCMCglmm* (Hadfield, 2015). This statistical procedure allows for the estimation of behavior-genetic variance components after accounting for the pedigree structure inherent in the data. Each model was computed using independent MCMC chains corresponding to 10,000,000 iterations, with sampling (thinning) occurring every 100 samples and a *burnin* period specified to disregard the first 300,000 iterations. The elimination of these initial iterations allowed the MCMC chain to explore and ascend through the likelihood landscape before sampling from the posterior distribution for the model’s parameters. Given the underlying skewedness of the variable’s residuals, a log-transformation was implemented to handle this issue. The model’s priors assumed that the G structure and the R structure were equivalent to the

Table 2. MCMCgImm evaluating the effects of sex on eminence

Model	Sex (95% CI)	Sex ^{pMCMC}	Mu	A	Vp	h^{2QG}	h^2 (95% CI)
Model 1: (A)	–	–	.211	2.516	4.961	.507	.510 (.434, .579)
Model 2: S+(A)	–.629(–1.098, –.153)	.0085	.554	2.469	4.867	.507	.507 (.434, .578)

Note. pMCMC: Significance test evaluating whether the parameter is different from zero. Mu: In variance components analysis, Mu is equal to the model's intercept. Although the pMCMC for Model 1's Mu was nonsignificant (.3010), the Mu value for Model 2 did reach statistical significance (.0220). A: additive variance, Vp: phenotypic variance. QG: heritability coefficients computed with the function QGparams in R.

**Figure 2.** Trace and density plots representing the additive heritability estimate of eminence after controlling for sex

phenotypic variance divided by the number of V terms with the prior's matrices following an order of $N=1$. Thus, the observed phenotypic variance was divided between the informed priors (sourced from the meta-analysis of Polderman et al., 2015) corresponding to the model's genetic (.49) and residual effects (.51).

After generating each model, variance components were extracted and used to estimate the trait's narrow-sense heritability (h^2). These values were computed as the proportion between the additive genetic variance (A) and the total observed variance ($A + E$; where E is equivalent to the model's residual variance). Heritability coefficients were also estimated with the function QGparams associated with the package QGgImm (Villemereuil et al., 2016). In generating stable heritability values with pedigree data, the use of a minimum of 150 individuals is generally recommended (Perrier et al., 2018). The super-pedigree, therefore, exceeds this power requirement as it contains 301 individuals in total.

Concerning the estimation of fixed effects, the calculation of each parameter was accompanied by a significance test (pMCMC), evaluating whether the predictor was significantly different from zero. The models' deviance information criterion (DIC) values were also extracted for subsequent model comparisons. Similar to other model fit indicators (e.g., Akaike's information criteria [AIC], AIC second-order estimate [AICc], and Bayesian information criteria [BIC]), smaller (DIC) values indicate a better model fit. For a more accurate model appraisal, Δ DIC (the difference

between the best model's DIC and each of the competing model's DIC values) and DIC weights (the relative probability that a particular model features the best statistical fit) were also computed. All statistical analyses were conducted with the statistical packages MCMCgImm (Hadfield, 2015) and MuMin (Barton, 2020) in R v 4.0.1.

Results

MCMCgImm Variance Components, Heritability Estimation, and Model Comparison

The analysis did not identify significant autocorrelations for the various variance components for lags above 100 samples. Table 2 features the MCMCgImm results, including the narrow-sense heritability estimates based on the models' variance components. The $A + E$ model tested here determined that the additive genetic component explained a sizeable proportion of the systematic variance. The analysis also revealed that the model's intercept was significantly different from zero. Overall, the trait's narrow-sense heritability is .510 (95% CI [.434, .579]) and the unshared environmentality is $(1-h^2)$.490. A similar pattern was found for an alternative model containing sex as a fixed effect. The trait's narrow-sense heritability was slightly smaller (.507; 95% CI [.434, .578]). As reflected by Figure 2, the density function plot featured a normal distribution. In regard to the model's fixed effect, males, scored significantly higher on the eminence scale

Table 3. MCMCgIimm model comparison evaluating the effects of sex on eminence, ranked based on DIC weights

Model	Log-likelihood	DIC	Δ DIC	DIC weight
Model 2: S + (A)	-512.182	1146.6	.000	.970
Model 1: (A)	-515.953	1153.6	6.95	.030

Note: DIC, deviance information criterion. Smaller DIC values indicate a better fit. DIC weights reflect the probability that a specific model is the best model.

($p = .0085$). The best model was the one in which A was modeled as a random effect, and sex was included as a fixed effect. Table 3 features the results of a model comparison. The DIC weight of this model was .970, meaning that it had a 97% relative probability of being the best model.

Discussion

A and E behavior-genetic variance components were estimated using an informed prior, which was derived from the meta-analysis of heritability conducted by Polderman et al. (2015). Two separate models were estimated and then compared, one without and one with sex modeled as a fixed effect. The best-fitting model (with a 97% probability) was the one that included sex. Based on this model, the additive (A) heritability (h^2) was estimated to be .507 (95% CI [.434, .578]) meaning that 50.7% of the variance in eminence can be attributed to genetic variants with additive effects on phenotypic variance. The nonshared environmentality (E) variance is .493. This variance component captures all contributions to trait variance that are not directly attributable to the action of additive genetic variance and would include error. This finding is broadly consistent with the high pedigree-based heritability estimate of social status in a large English lineage presented by Clark (2021).

The most straightforward interpretation of these results is that variation in eminence, much like the extremes of more conventionally measured dimensions of individual differences (such as IQ), is mostly influenced by additive genetic variation, which in turn is consistent with existing knowledge of the basis of normal-range phenotypic variance in many traits believed to contribute in part to eminence (again, such as IQ; Shakesharf et al., 2015). Congruent with this, Roeling et al. (2016) estimated that the additive heritability of working in a creative profession (a potentially more normal-range manifestation of the sorts of phenotypes that contribute to 'eminence') is .70. Further, and interestingly, they found that the best-fitting model in their dataset was one estimating only A and E variances.

Visual inspection of the super-pedigree (Figure 1) yields clear indications of direct multiple generation-to-generation transmission of eminence. Erasmus Darwin was father to Robert Waring Darwin and grandfather to both Charles Robert Darwin (who was the son of Robert Waring Darwin) and Frances Galton. Josiah Wedgwood I was father to Thomas Wedgwood and grandfather to both Emma Wedgwood and Charles Robert Darwin. Josiah Wedgwood I and Erasmus Darwin were both great-grandfathers to George Howard Darwin, and Francis Darwin (both of whom were the children of Emma Wedgwood and Charles Robert Darwin), and great-great-grandfathers to Charles Galton Darwin (who was the son of George Howard Darwin). Anthony Wedgwood Benn was father to Hilary James Benn, Baha'u'llah was father to 'Abd'u'l-Bahá, who was grandfather to Shoghi Effendi, and Thomas Arnold the Elder was father to Matthew

Arnold. Thomas Henry Huxley was a grandfather to Aldous, Andrew, and Julian Huxley. More indirect instances of vertical transmission are also present in the data. Josiah Wedgwood I (along with Erasmus Darwin, through Robert Waring Darwin) was a great-great-grandfather to Ralph Vaughn Williams and great-great-great-great-great-grandfather (along with Erasmus Darwin) to Alexander Skandar Keynes through Charles Robert Darwin and Emma Wedgwood, who were his great-great-great-grandparents. Baha'u'llah was also great-great-great-grandfather to Alexander Skandar Keynes. One of the latter's great-grandfathers was Edgar Adrian. Thomas Arnold the Elder was great grandfather to both Aldous and Julian Huxley.

There are alternative behavior-genetic models of extraordinary accomplishment that take into consideration nonadditive gene-by-gene interactions (genetic dominance and epistasis effects) and gene-by-environment interactions. An example of the former is *emergensis*, which posits that extreme and rare phenotypes may primarily result from the action of rare nonadditive gene combinations (Eysenck, 1995; Lykken et al., 1992; Jensen, 1997; Simonton, 1999), with the resultant phenotype being effectively larger than the sum of its parts owing to the multiplicative, as opposed to additive, ways in which the relevant genetic variants interact. An example of the latter is the *cosmopolitan hypothesis*. Cosmobia is an old concept from developmental biology which holds that rare physiological deformity can exhibit a degree of variation that does not necessarily handicap the individual exhibiting such deformities. For example, individuals can exhibit severe deformities, such as the presence of just one, or more than two, eyes (these are termed primary deformities), but within this range of phenotypes there are far less pathological, but nevertheless quite rare, secondary deformities, such as having two pupils per eye. While such cosmopolitan variation might stem from genetic differences among individuals, in some cases, it may purely be a function of the capacity for a particular genotype to produce substantially different phenotypes, given special environmental conditions experienced during development (Johnson & Bouchard Jr., 2014). Johnson and Bouchard Jr. (2014) have speculated that manifestations of extreme accomplishment (such as creative genius) might be another example of a cosmopolitan effect, with the possession of even extreme levels of the relevant dispositional traits being insufficient to bring about such a phenotype in the absence of certain unusual gene-by-environment correlations and resultant interactions. For discussion of models that similarly emphasize epigenetic and gene-by-environment interactions in the cultivation of extraordinary talent, see Papierno et al. (2005) and Simonton (1999).

There exist statistical packages that allow for various gene-by-gene interactions to be estimated based on pedigree data; however, the author of one such package (*Nadiv*; Matthew Wolak) informed the authors that the super-pedigree has insufficient varieties of consanguineous mating (intermarriage) necessary for the reliable estimation of nonadditive effects, based on the results of simulations that were run on behalf of the authors. Four clear instances of intermarriage (involving individuals related to one another at the degree of third cousin or closer) are present in the super-pedigree: (1) Josiah Wedgwood I married his third cousin Sarah Wedgwood, (2) Josiah Wedgwood III married his first cousin Caroline Sarah Darwin, (3) Robert Barclay II married his first cousin Lucy Barclay, and (4) Charles Robert Darwin married his first cousin Emma Wedgwood. For the estimation of nonadditive effects, much more elaborate patterns of intermarriage are

needed (such as those involving double-first cousins and half-cousins; Wolak, personal communication).

The cosmobia model could not be tested either, as it requires the estimation of gene-by-environment interactions, which cannot (currently) be done using the *MCMCglmm* software. Evidence of a substantial contribution to variance in eminence stemming from environmentality is nevertheless present, which might in part be reflective of factors such as the provisioning of ‘opportunities’ for the establishment of fortuitous gene-by-environment interactions via gene-by-environment correlation. Thus, it might be that eminence in this super-pedigree is a joint product of having high levels of additive genetic variants associated with the relevant dimensions of talent (e.g., high IQ, creativity, ambition, grit), in addition to having the rare opportunity to cultivate that talent to a truly extraordinary degree through access to unique environments (one such environment might be the home of a highly accomplished family with rich stores of knowledge, which may in turn encourage and support one to work extraordinarily hard in order to live up to expectations). The sex difference in degree of eminence (favoring males) might also be a partial function of the rigidity of historical sex roles and associated stereotypes; however, this finding is also in line with the greater male variability hypothesis (which is robustly supported with respect to a large variety of neuroanatomical and behavioral measures: Thöni & Volk, 2021; Thöni et al., 2021; Wierenga et al., 2020). It should be furthermore noted that evidence of a significant impact of vertical nongenetic (i.e., cultural) transmission on trait variance is scant, especially when extended twin studies are used to properly control vertical transmission effects for genetic confounds (see Eaves et al., 1999; Kandler et al., 2012; Martin et al., 1986; Swagerman et al., 2017). On this basis, it is not anticipated that these sorts of vertical transmission effects are (substantially) contributing to the patterns observed in these data.¹³

The super-pedigree is unusual in that it contains so many historiometrically identified (based on biographic prominence) eminent individuals, given their extreme rarity, even when established using thicker inclusion criteria, as per *Pantheon*.¹⁴ One possibility is that the members of this super-pedigree historically engaged in relatively high levels of intermarriage and assortative mating, which may have maintained at a high-frequency rarer genetic variants that predispose toward extraordinary accomplishment. Four instances of intermarriage involving individuals related to the degree of third cousin or closer have been noted in this super-pedigree, but it is possible that more cryptic forms of relatedness might also be present in this dataset (as an example of this, two separate Wedgwood lines, collateral to Josiah Wedgwood I, contributed to the Benn family via the marriage of William John Barker and Margaret Wedgwood who were fourth cousins). That many of the members of this super-pedigree may historically have been relatively highly interrelated is consistent with genome-wide estimates of linkage disequilibrium among representatively sampled individuals from the population of the USA, which indicate that general levels of inbreeding were higher among older individuals (Nalls et al., 2009). This would be consistent with historically more limited geographical dispersal among individuals (especially in the 19th and prior centuries) relative to their extended family members. Increased aggregate relatedness among partners would be expected to increase genomic linkage disequilibrium among offspring, raising trait heritability estimates. There may also be significant degrees of assortative mating in this super-pedigree, involving families who are similar with respect to specific talent-conferring genetic loci. Indeed, the merger

through marriage of the Arnold–Huxley, Keynes–Baha’u’llah, and Wedgwood–Darwin pedigrees could be evidence of this, as all three families were of equivalent social and intellectual standing (indeed Charles Robert Darwin and Thomas Henry Huxley were close scientific collaborators). Cryptic relatedness and assortative mating are difficult to estimate absent either genomic data or much more fine-grained and extensive social pedigree-based information.

There are some limitations in these data that need to be addressed. Chief among these is the social, as opposed to genetic, nature of the super-pedigree. It has been noted that ‘[h]eritability estimates ... [are] slightly higher’ (Perrier et al., 2018, p. 863) when estimated using genetic, as opposed to social pedigree data. This discrepancy is attributed by Perrier et al. (2018) ‘to incorrect pedigree links, including extra-pair paternity, and to lower information content’ (Perrier et al., 2018, p. 838). Therefore, the estimates of the additive heritability of eminence presented here are likely to be *lower* than would have been the case had corresponding genetic data for the super-pedigree been available, owing to error in identifying linkages, making these estimates likely somewhat conservative.

Another limitation concerns the use of the thicker *Pantheon* dataset, as opposed to a thinner, more selective dataset such as that of *Human Accomplishment*. It could be argued that some of the individuals counted as eminent might have had their historiometric bibliographic prominence inflated for relatively mundane reasons (such as in the case of those who are the offspring of someone eminent), or due to epochcentric effects (although the HPI values used here do attempt to correct for ‘recency bias’). The majority of the individuals that were included as ‘eminent’ are, however, clearly independently and (by world historical standards) significantly accomplished and therefore are eminent in their own right (such individuals include several unambiguous creative geniuses, three Nobelists, a major 18th century industrialist, and the founder of a significant world religion). Another advantage (beyond greater model power) of using the thicker *Pantheon* dataset is the breadth of domains covered. There are likely to be highly general genetic factors that predispose toward extraordinary achievement across multiple domains of accomplishment (including things as far afield as athletics and science), such as those contributing to individual differences in grit and achievement orientation. In the case of the present result, it is likely that the heritability estimated here for eminence is capturing the heritability of the relevant interactions among all of those dispositional factors in Galton’s ‘triple event’, therefore making the phenotype of eminence consistent with what has come to be termed the *First Law of Behavior Genetics*, specifically ‘all human behavioral traits are heritable’ (Turkheimer, 2000, p.160). Finally these findings potentially bring closure to the Galton–de Candolle debate, as eminence appears to be equal parts nature and nurture.

Conflict of interest. We the authors declare that there are no conflicts of interest.

Notes

- 1 In subsequent work, Galton (1874, 1875) proposed that comparisons involving identical and fraternal twins could be used in the estimation of the relative impacts of nature and nurture on a given trait.
- 2 ‘Relatively’ rigorous compared to most research on eminence, which is typically qualitative and impressionistic.
- 3 It should be noted that historiometric appraisal of eminence is by no means a ‘perfect science’. There will obviously be cases of individuals who do not get

recorded by history, but who would nevertheless qualify as eminent (e.g., whoever first devised a means of reliably starting controlled fires, whoever first invented the wheel). Knowledge of the life and works of some individuals might have been suppressed by virtue of them having lived in anti-intellectual or intolerant cultures. The importance of this limitation should not be exaggerated; however, as for historiometric methods to be useful, it is merely necessary that relevant neutral historical data on the lives and accomplishments of a reasonably representative sample of eminent individuals exist (Murray, 2003; Simonton, 1984).

4 This was usefully exploited by Cox (1926) in estimating the historical IQs of various creative geniuses — for another example, see Simonton's (2006) estimates of the historical IQs and other qualities of US presidents.

5 Note that these values fluctuate slightly owing to continuous updating. The values reported here are correct as of July 8, 2021.

6 In addition to being known as the wife of Charles Robert Darwin, Emma Wedgwood was also known as a strident advocate of political reform (see: Loy & Loy, 2010).

7 1. Richenda Pease, who was a granddaughter of Josiah Wedgwood IV, married Andrew Huxley. 2. A great-granddaughter of Thomas Henry Huxley, Angela Darwin (née Huxley), married George Pember Darwin, a great-grandson of Charles Robert Darwin.

8 John Maynard Keynes' brother Geoffrey married Elizabeth Darwin, a granddaughter of Charles Robert Darwin, with whom he had two children, Richard and Quentin Keynes.

9 In addition to being the wife of John Maynard Keynes, Lydia Lopokova was also a prominent ballerina.

10 Who founded the Bahá'í faith.

11 Who founded the Bayání faith

12 The possibility of using the *Human Accomplishment* dataset as a replication sample (as per Yu et al., 2016) was considered and rejected on the basis that it contained too few significant figures to yield a 'convincing' estimate of the heritability of eminence. Moreover, the index values estimated by Murray (2003) only allow for comparisons to be made within specific inventories, unlike the *Pantheon* HPI values, which are intended to allow global ranking among eminent individuals. Therefore, even though there is significant apparent agreement between the two sets of values (Yu et al., 2016), the possibility of confounding stemming from comparing 'apples and oranges' when using Murray's (2003) index scores exists.

13 Although one potential instance of purely culturally transmitted eminence is the succession to the leadership of the Bahá'í faith which was transmitted to the descendants of Baha'u'lláh through decrees enacted via wills.

14 Of the approximately 107 billion people who have ever lived, only 88,506, or 0.00008% have appeared in the *Pantheon* dataset as of this writing.

References

- Bartoń, K. (2020). MuMIn: Multi-model inference. R package version 1.43.6. <https://CRAN.R-project.org/package=MuMIn>
- Berra, T. M., Alvarez, G., & Shannon, K. (2010). The Galton–Darwin–Wedgwood pedigree of HH Laughlin. *Biological Journal of the Linnean Society*, 101, 228–241.
- Clark, G. (2021). *For whom the bell curve tolls: A lineage of 400,000 English individuals 1750–2020 shows genetics determines most social outcomes*. Manuscript in preparation.
- Cox, C. M. (1926). *Genetic studies of genius. II. The early mental traits of three hundred geniuses*. Stanford University Press.
- de Candolle, A. (1873). *Histoire des sciences et des savantes depuis deux siècles d'après l'opinion des principales académies ou sociétés scientifiques*. Geneva.
- de Villemeureuil, P., Schielzeth, H., Nakagawa, S., & Morrissey, M. (2016). General methods for evolutionary quantitative genetic inference from generalized mixed models. *Genetics*, 204, 1281–1294.
- Eaves, L., Heath, A., Martin, N., Maes, H., Neale, M., Kendler, K., Kirk, K., & Corey, L. (1999). Comparing the biological and cultural inheritance of personality and social attitudes in the Virginia 30,000 study of twins and their relatives. *Twin Research & Human Genetics*, 2, 62–80.
- Eysenck, H. J. (1995). *Genius: The natural history of creativity*. Cambridge University Press.
- Frank, M. (2021). Anthony Neil Wedgwood Benn: Family tree. <https://gw.geneanet.org/swintonbarterer?iz=0&lang=en&n=benn&oc=0&p=anthony+neil+wedgwood&type=tree>
- Galton, F. (1869). *Hereditary genius: An inquiry into its laws and consequences*. Macmillan.
- Galton, F. (1874). On men of science, their nature and their nurture. *Proceedings of the Royal Institution of Great Britain*, 7, 227–239.
- Galton, F. (1875). The history of twins, as a criterion of the relative powers of nature and nurture. *Journal of the Anthropological Institute*, 5, 391–406.
- Hadfield, J. (2015). Package 'MCMCglmm'. <http://cran.r-project.org>
- Jensen, A. R. (1997). The puzzle of nongenetic variance. In R. J. Sternberg & E. L. Grigorenko (Eds.), *Intelligence, heredity, and environment* (pp. 42–88). Cambridge University Press.
- Johnson, W., & Bouchard Jr., T. J. (2014). Genetics of intellectual and personality traits associated with creative genius: Could geniuses be cosmopolitan dragon-kings? In D. K. Simonton (Ed.), *The Wiley handbook of genius* (pp. 269–296). Wiley-Blackwell.
- Kandler, C., Bleidorn, W., & Riemann, R. (2012). Left or right? Sources of political orientation: The roles of genetic factors, cultural transmission, assortative mating, and personality. *Journal of Personality & Social Psychology*, 102, 633–645.
- Loy, J. D., & Loy, K. M. (2010). *Emma Darwin: A Victorian life*. University of Florida Press.
- Lykken, D. T., McGue, M., Tellegen, A., & Bouchard Jr., T. J. (1992). Emergence. Genetic traits that may not run in families. *American Psychologist*, 47, 1565–1577.
- Martin, N. G., Eaves, L. J., Heath, A. C., Jardine, R., Feingold, L. M., & Eysenck, H. J. (1986). Transmission of social attitudes. *Proceedings of the National Academy of Sciences USA*, 83, 4364–4368.
- Murray, C. (2003). *Human accomplishment: The pursuit of excellence in the arts and sciences, 800 BC to 195*. Harper Collins.
- Nalls, M. A., Sanchez-Simon, J., Gibbs, J. R., Paisan-Ruiz, C., Bras, J.T., Tanaka, T., Matarin, M., Scholz, S., Weitz, C., Harris, T. B., Ferrucci, L., Hardy, J., & Singleton, A. B. (2009). Measures of autozygosity in decline: Globalization, urbanization, and its implications for medical genetics. *PLoS Genetics*, 5, e1000415.
- Papierno, P. B., Ceci, S. J., Makel, M. C., & Williams, W. M. (2005). The nature and nurture of talent: A bioecological perspective on the ontogeny of exceptional abilities. *Journal for the Education of the Gifted*, 28, 312–332.
- Perrier, C., Delahaie, B., & Charmantier, A. (2018). Heritability estimates from genome wide relatedness matrices in wild populations: Application to a passerine, using a small sample size. *Molecular Ecology Resources*, 18, 838–853.
- Polderman, T. J. C., Benyamin, B., De Leeuw, C. A., Sullivan, P. F., Van Bochoven, A., Visscher, P. M., & Posthuma, D. (2015). Meta-analysis of the heritability of human traits based on fifty years of twin studies. *Nature Genetics*, 47, 702–709.
- Roeling, P., Willemsen, G., & Boomsma, D. I. (2016). Heritability of working in a creative profession. *Behavior Genetics*, 47, 298–304.
- Shakesharp, N. G., Trzaskowski, M., McMillan, A., Krapohl, E., Simpson, M. A., Reichenberg, A., Cederlöf, M., Larsson, H., Lichtenstein, P., & Plomin, R. (2015). Thinking positively: The genetics of high intelligence. *Intelligence*, 48, 123–132.
- Simonton, D. K. (1984). *Genius, creativity and leadership: Historiometric inquiries*. Harvard University Press.
- Simonton, D. K. (1999). Talent and its development: An emergent and epigenetic model. *Psychological Review*, 106, 435–457.
- Simonton, D. K. (2006). Presidential IQ, openness, intellectual brilliance, and leadership: Estimates and correlations for 42 U.S. chief executives. *Political Psychology*, 27, 511–526.
- Simonton, D. K. (2009). Varieties of (scientific) creativity: A hierarchical model of domain-specific disposition, development, and achievement. *Perspectives on Psychological Science*, 4, 441–452.

- Simonton, D. K.** (2018). *The genius checklist: Nine paradoxical tips on how you can become a creative genius*. MIT Press.
- Swagerman, S. C., Van Bergen, E., Dolan, C., de Geus, E. J., Koenis, M. M., Pol, H. E. H., & Boomsma, D. I.** (2017). Genetic transmission of reading ability. *Brain & Language*, 172, 3–8.
- Thöni, C., & Volk, S.** (2021). Converging evidence for greater male variability in time, risk, and social preferences. *Proceedings of the National Academy of Sciences USA*, 118, e2026112118.
- Thöni, C., Volk, S., & Cortina, J. M.** (2021). Greater male variability in co-operation: Meta-analytic evidence for an evolutionary perspective. *Psychological Science*, 32, 50–63.
- Turkheimer, E.** (2000). Three laws of behavior genetics and what they mean. *Current Directions in Psychological Science*, 9, 160–164.
- Wierenga, L. M., Doucet, G. E., Dimi, D., Agartz, I., Aghajani, M., Akudjedu, T. N., Albajes-Eizagirre, A., Alnæs, D., Alpert, K. I., Andreassen, O. A., Anticevic, A., Asherson, P., Banaschewski, T., Bargallo, N., Baumeister, S., Baur-Streubel, R., Bertolino, A., Bonvino, A., & Tamnes, C. K.** (2020). Greater male than female variability in regional brain structure across the lifespan. *Human Brain Mapping*. Advance online publication. doi: 10.1002/hbm.25204.
- Wikipedia.** (2021a). Darwin-Wedgwood family. https://en.wikipedia.org/wiki/Darwin%E2%80%93Wedgwood_family
- Wikipedia.** (2021b). Arnold-Huxley family tree. https://en.wikipedia.org/wiki/Talk%3ADarwin%E2%80%93Wedgwood_family#/media/File:Huxley-Arnold_family_tree.svg
- Wilson, A. J., Reale, D., Clements, M. N., Morrissey, M. M., Postma, E., Walling, C. A., Kruuk, L. E. B., & Nussey, D. H.** (2010). An ecologist's guide to the animal model. *Journal of Animal Ecology*, 79, 13–26.
- Yu, A. Z., Ronen, S., Hu, K., Lu, T., & Hidalgo, C. A.** (2016). Pantheon 1.0, a manually verified database of globally famous biographies. *Scientific Data*, 3, 150057.