# Increased birth rank of homosexual males: disentangling the older brother effect and sexual antagonism hypothesis

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### **ABSTRACT**

Male homosexual orientation remains a Darwinian paradox, as there is no consensus on its evolutionary determinants. One intriguing feature of homosexual men is their higher male birth rank compared to heterosexual men. This can be explained by two non-exclusive mechanisms: an antagonistic effect (AE), implying that more fertile women have a higher chance of having a homosexual son, or an older brother effect (OBE), where each additional older brother increases the chances for a male embryo to develop a homosexual orientation due to an immunoreactivity process. However, there is no consensus on whether both OBE and AE are present in human populations, or if only one of these mechanisms is at play and its effect mimicking the signature of the other mechanism. An additional older sister effect (OSE) has also recently been proposed. To clarify this situation, we developed theoretical and statistical tools to study OBE and AE independently or in combination, taking into account all known sampling biases. These tools were applied on new individual data, and on various available published data (two individual datasets, all relevant aggregated data, and twin data). Support for OBE was apparent in aggregated data, with the OBE increasing linearly with fertility. The OBE was also supported in two individual datasets. An OSE seems to result from a sampling bias in presence of OBE, and is likely to be artefactual. AE was not supported, either in individual datasets, including the analysis of the extended maternal family, nor in twin data. The evolutionary implications of these findings are discussed.

### INTRODUCTION

Male homosexual orientation, i.e. preferential attraction of male subjects to same-sex partners for sexual intercourse and/or romantic relationships, is an evolutionary enigma because, in humans, preference for male-male relationships is partially heritable (Bailey et al., 2000; Långström et al., 2010), imposes a fertility cost (lower offspring number) (Bell and Weinberg, 1978; Iemmola and Camperio-Ciani, 2009; Rieger et al., 2012) and is relatively common in many societies (2%–6% in Western countries) for such a costly trait (Apostolou, 2020; Berman, 2003). The origin of male homosexual orientation has long been a matter of interest, and several evolutionary hypotheses were proposed, mostly involving kin selection or antagonistic pleiotropy (see Barthes et al., 2015, Gavrilets and Rice, 2006, and Apostolou, 2020 for reviews).

In this paper, we will examine some of the empirical evidence linked to the antagonistic pleiotropy hypothesis. This hypothesis explains the persistence male homosexuality by a sex-antagonistic effect (AE). Gavrilets and Rice (2006) have confirmed through modelling that this can maintain polymorphism for male sexual orientation under certain conditions. Several studies have indeed reported differences in fertility between families of homosexual and heterosexual men (Camperio-Ciani et al., 2009, 2004a; Iemmola Camperio-Ciani, 2009), and the authors proposed that families of homosexual men display a higher fertility on the maternal side, compared with the rest of the population, in accordance with the AE hypothesis.

Another line of research men d to decipher the proximal determinants of male homosexual orientation. Family determinants for the sexual orientation of men has been suspected for nearly a century, and two main directions have been explored (for a short review, see Wampold (2018). The first focused on birth order, and concluded that homosexual men had more older siblings than heterosexual men (e.g. Slater, 1962); the second focused on

sex-ratio, and concluded that homosexual men had more brothers than sisters (e.g. Lang, 1940). These two observations were reconciliated when it was proposed that homosexual men had more older brothers than heterosexual men (Blanchard and Bogaert, 1996a). This finding, commonly referred to as the older brother effect (OBE), has been replicated several times in Western (e.g., Blanchard, 2018a, 2018b; Blanchard and Bogaert, 2004; Bogaert and Skorska, 2011) and non-Western countries such as Turkey, Iran, Hong Kong, Samoa, and Indonesia (Blanchard, 2018c; Li and Wong, 2018; Nila et al., 2019). The underlying mechanism of the OBE is logical and prenatal, since homosexual orientation is influenced neither by the number of non-biological older brothers nor by the amount of time spent with biological or non-biological older brothers (Bogaert, 2006). The proposed explanation is a maternal immune reaction to successive male pregnancies, with each male foetus increasing the likelihood of an immune response from the mother. This maternal immune reaction would lead to an alteration of the typical development of sexually dimorphic brain structures relevant to the sexual orientation of the foetus (Bogaert and Skorska, 2011). Recently, possible molecular evidence of this specific immune reaction has been presented (Bogaert et al., 2018). Yet, these two lines of research leave many open questions.

Firstly, it is still unclear whether the OBE is universal. The OBE is not always found, even in some large samples from UK, Canada, or Australia (Bogaert, 1998; Kishida and Rahman, 2015; Rahman et al., 2008; Zjetsch et al., 2012, but see Blanchard and VanderLaan, 2015). It is thus possible that the OBE is only observed in some populations. Alternatively, the OBE could be restricted to subcategories of homosexual men, as suggested by Swift-Gallant et al. (2018). Additionally, the OBE is sometimes described from samples which are not comparable. For example, several eta-analyses (Blanchard, 2018c, 2018a, 2018b; Blanchard et al., 2020b, 2020a) attempting to demonstrate an OBE in homosexual men across multiple studies include data from transexuals, pedophiles, hebephiles, or gender-dysphoria

individuals. Similarly, other studies have focused on specific individual categories, such as sex offenders, psychoanalytic patients, individuals treated with feminizing hormones, clinically obsessional patients, or patients with paraphilic behaviours such as masochism, fetishism, and transvestism (e.g. Blanchard et al., 2012, 1998). As these different situations are drawn from highly non-representative populations (Zietsch, 2018), and are not necessarily the result of similar determinants as those for homosexuality, or could represent extreme values from a continuum, considering them could introduce some biases.

Secondly, an older sister effect (OSE) acting alongside the OBE has been described several times, e.g. in UK (King et al., 2005), Finland (Kangassalo et al., 2011), Samoa (VanderLaan and Vasey, 2011; Vasey and VanderLaan, 2007), Canada (Swift-Gallant et al., 2018), Netherlands (Ablaza et al., 2022), or in participants of a BBC internet survey (Blanchard and Lippa, 2020). Based on these findings, a recent meta-analysis proposed the presence of a pervasive OSE, in addition to the OBE (Blanchard et al., 2020a), even if this OSE is generally not as strong as the OBE. A further complication is that the OSE could in theory be a side-effect of an OBE: if the sex ratio is even, sampling individuals with more older brothers also means sampling individuals with more older sisters. If this were the case, homosexual men would generally have more older sisters than heterosexual men even if the only causal effect on the probability of being homosexual is the number of older brothers. It is thus unclear if explanations for the OBE should be sought independently or not from this additional sibling effect.

A third --and even more problematic-- question is whether the OBE and AE are both at play in human populations, or if they are different manifestations of the same biological phenomenon. Indeed, a higher fertility of mother of homosexual men implies that, when sampling homosexuals from a population, the mean birth order of homosexuals in gher than the mean birth order of heterosexuals. Conversely, if fertility varies within a population

independent of the occurrence of homosexuality, sampling high birth ranks (as is the case when sampling homosexuals in the presence of the OBE) will generate a sample from high-fertility mothers. The two phenomena (OBE and AE) thus lead to similar predictions of a higher birth rank of homosexuals and a higher fertility of families of homosexuals from population samples, and thus cannot be easily distinguished even if they rely on very different mechanisms: a plastic effect (maternal effect) in the case of OBE and a genetic effect in the case of the AE. This problem of causal attribution has been previously identified (e.g. VanderLaan and Vasey, 2011; Zietsch et al., 2008), and formally exposed (Khovanova, 2017), and three main methods have been proposed to study OBE while controlling for variation in female fecundity.

This first is a statistical control of fecundity: raw number of older brothers are not transformed, but sibling size is used as a control variable, for example as a dependent variable in a regression (e.g. Ablaza et al., 2022; Nila et al., 2019). Second, transforming the raw data using various *ad hoc* metrics controlling for family size: the general form of these metrics is (X + a)/(N + b), where X is the number of older brothers (or any other sibling category under study), N is the total number of siblings, and  $\{a, b\}$  are two scalars. Values of these scalars vary according to authors:  $\{-1, -1\}$  for Slater (1962),  $\{\frac{1}{2}, 1\}$  for Berglin (1980), and  $\{\frac{1}{4}, 1\}$  or  $\{\frac{1}{3}, -X+1\}$  for Blanchard (2014). These metrics have some drawbacks, e.g. Slater's index is not defined for only-children (N = 1), see Blanchard (2014) for further comments on these metrics. Other metrics have thus been subsequently proposed, based on ratio between the odds of observing an older brother for homosexuals, and the same odd for heterosexuals (OBOR, Blanchard, 2018c, 2018b), or based on the ratio of older brothers to older sisters, relative to the same ratio for heterosexuals (OR, Vilsmeier et al., 2021). Third, a data restriction: only families with a fixed number of child (i.e. 1 or 2) are considered (Khovanova, 2017). This can

lead, in populations displaying a relatively high fecundity, to discarding more than 60 % of the sampled individuals (Blanchard et al., 2020b).

Results using these methods indicated that OBE unlikely to be a sampling artefact, at least in some populations, although there is no consensus at a larger scale: several meta-analyses using an odds-ratio metric concluded that OBE is generally present (Blanchard, 2018c, 2018c, 2018b), while a meta-analysis using another odds-ratio metric concluded that "almost no variation in the number of older brothers in men is attributable to sexual orientation" (Vilsmeier et al., 2021). However, index ratios cannot be considered as reliable statistics to oddy OBE, as this approach assumes that samples of heterosexuals and homosexuals are adequately matched for potentially confounding variables affecting sibship size, such as age or social economic status, which is not always the case (see Price and Hare, 1969). Also, using an index ratio implicitly assumes that the expression of OBE is independent of the level of fertility, which remains to be shown.

To study female fecundity without the interference of birth rank, one possibility is to use additional family data: the fecundity of maternal aunts, for example, is considered to be independent of the birth rank of the sampled individuals (Camperio-Ciani et al., 2004a; Iemmola and Camperio-Ciani, 2009; King et al., 2005). However, this assumes that there is no correlation between the fecundity of the mother and her sisters, which is not the case (e.g. Anderton et al., 1987; Berent, 1953). Thus, even if an OBE sacting alone (i.e. no AE), homosexuals are sampled from families displaying a higher fecundity, and their maternal aunts are likely to display a higher fecundity as well (Zietsch et al., 2008). Another possible approach to control for birth rank is to consider only individuals with a specific birth rank, e.g. only first born or only second born (e.g. Blanchard et al., 2020b; Ciani and Pellizzari, 2012; Khovanova, 2017), but this results in a significant reduction of the available data and hence of inferential power.

Results using these methods are ambiguous: the presence of AE is claimed when additional family data are considered, such as maternal aunts (e.g. Camperio-Ciani et al., 2004b; Iemmola and Camperio-Ciani, 2009), but results are not controlled for birth rank. In a meta-analysis, when data are restricted to families with only one or only two sons, no AE is found (Blanchard et al., 2020b), although this analysis includes a paper retracted since then.

To sum up, there is still no consensus on whether both OBE and AE are present in human populations and contribute to the occurrence of male homosexuality, or if only one of these mechanisms exists and its effect mimics the signature of the other mechanism. If both mechanisms act together, we also need to estimate the relative contribution of each for the higher birth rank of homosexual men and higher fertility of their families. Last, we delete the older sisters and the probability of being homosexual, or if it is a by-product of the OBE. To clarify these questions, we developed methodological tools, allowing us to pursue three novel approaches.

Our main methodological development is a derivation of a population-level relationship between mean birth rank and mean fertility in a random population sample, without any OBE, OSE, or AE. This analytical relationship links the mean birth rank of individuals with the mean fertility of their population sample, allowing us to quantitatively estimate the role of OBE, independently of any fertility effect, when mean birth rank deviates from its value predicted by the population fertility estimate. We also determined analytically and through simulations the expected number of older sisters as a function of the number of older brothers to test if the OBE can indeed generate an OSE.

The three approaches were:

A) We checked if population samples from heterosexual and homosexual men gathered from the available literature followed the general expectation of mean birth rank given

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population fertility, or not. If population samples of homosexuals did not follow this general expectation but had higher birth rank given the mean population fertility, this would support an OBE effect.

B) We used two statistical frameworks to separate quantitatively the OBE and AE effects in family data from individual homosexual and heterosexual men sampled in France, Indonesia and Greece. The first framework was "classical" linear modelling (here generalized linear mixed model or GLMM) to estimate the fertility of the women on the mother's side (mothers and maternal aunts) after controlling for the birth rank of the focal subject (effectively comparing this familial fertility for first born, second born, etc.), allowing to test for AE after controlling for the OBE. If, for a given birth rank, homosexuals had more sibling or cousins than heterosexuals, the AE hypothesis would be supported. The second framework was to use Bayesian inference implemented as a hierarchical model in Nimble (de Valpine et al., 2017) to test whether the effects of the OBE, AE or both could be inferred from individual family data. Nimble generates simulated data based on different scenarios and provides quantitative support of the various scenarios from the empirical data. We evaluated several scenarios including solely or simultaneously the OBE and the AE favouring male homosexuality and women fertility. The OBE was modelled after determining the best-fit function from the empirical data.

C) In order to evaluate the presence of AE in an independent dataset, we exploited available twin data. Contrarily to monozygotic twining, dizygotic twining is a cue of higher fertility (Tong and Short, 1998): under the AE hypothesis, there should thus be more homosexual men among the dizygotic twins than among the monozygotic twins. We thus compared the proportion of homosexual men in all available monozygotic and dizygotic twin samples.

These different lines of reasoning allowed us for the first time to disentangle the effects of the birth rank (the OBE) and of antagonist genetic factors (the AE) affecting fertility and sexual orientation simultaneously in aggregated data and in three independent data sets, and to provide several tools that will be useful to address these questions in other human populations.

### MATERIAL and METHODS

# Aggregated family data

To find primary data on sexual orientation and family composition, we proceeded in two ways. First, we performed literature searches on accessible databases to find recent publications. Second, to find older data, we scanned cited literature. In addition, inspection of review articles ensured that no major older papers were overlooked (e.g. Blanchard, 2018b, 2018c, 2004, 1997; Blanchard et al., 2020a, 2020b, 2001; Bogaert and Skorska, 2011). Papers that present at least two samples of men, one homosexual and merosexual, along with the following information, were retained: number of individuals sampled, and the total number of each individual's older brothers, older sisters and siblings. Data used in several papers were retained only once, e.g. Rahman et al. (2008) used in Kishida and Rahman (2015). When some required information was not found in the publication, we attempted to contact the authors to obtain the missing information. When the size of a sibling category was given only as a proportion, this proportion was multiplied by the corresponding number of individuals and rounded to the nearest whole number, to obtain the expected number for this sibling category. When further computation was required (as in Blanchard, 2021), the sample was not considered. Data concerning sex-offenders, pedophiles, hebephiles, gender-dysphoria, psychoanalytic patients, individuals treated with feminizing hormones, hospital patients,

clinically obsessional patients, patients with paraphilic behaviours like masochism, fetishism, and transvestism, or transexuals, were not considered. Were also excluded samples concerned with children or adolescents (as the number of younger siblings may not be final), adopted individuals (as the biological sibling composition is generally not available) or twins (birth order is ambiguous, and dizygotic twining is a correct of higher fertility, Tong and Short, 1998). Retracted papers were not considered. The French sample collected for individual data (see below) was also considered here in its aggregated form. Bisexual individuals were pooled with homosexuals, and pairs of samples with at least 50 individuals for each sexual orientation were further retained. For each sample, the mean birth order among males was computed as OB/N + 1, where OB is the total number of older brothers, and N is the number of sampled individuals. Similarly, the mean birth order among females was computed as OS/N + 1, where OS is the total number of older sisters. The mean fertility was estimated as Sibs/N, where Sibs is the total number of older and younger siblings. The mean fertility for only males or only females was computed as half this overall fertility.

### **Individual family data**

Sampling in France was performed between August 2006 until July 2016 in public areas, research institutes, and within social networks (mainly in the cities of Montpellier and Paris). A targeted sampling procedure was performed: when a beach mostly frequented by individuals with a homosexual preference was sampled, a nearby beach with no particular attendance bias was also sampled. Friends and acquaintances of both individuals reporting one or the other sexual preference were sampled. Upon agreement, a document describing the general purpose of the study and providing contact details of the person in charge (M.R.) was given to each participant. This document explicitly states that personal data will only be used for research purposes and that only global results —not individual data— will be published.

Written informed consent was obtained from all participants. The protocols used to recruit individuals and to collect data were approved the French National Committee of Information and Liberty (CNIL) through the CNRS (approval #1226659). Each individual was privately and anonymously interviewed and was requested to report his date of birth, his self-declared sexual preference, the sex and birth order of each of his full and half siblings or maternal side, the country of birth of his four grand-parents, the number of maternal aunts and corresponding number of cousins. Other personal and familial information have been collected to analysed elsewhere. Individuals below 18 years age of legal adulthood in France) were not considered. To reduce cultural heterogeneity, individuals with at least one grand-pal born outside Europe were not further considered. This resulted in 512 men who participate and to provide personal data (Table 1). Eighteen men declared a bisexual orientation and were grouped in the homosexual preference category. There was a total of 271 men declaring a homosexual preference (52.9%), and 241 men declaring a heterosexual preference (47.1%). Mean age was 33.8 years (range 18.3 - 75.4), with the homosexual preference group being slightly younger than the heterosexual group (32.0 vs 35.8 years, Wilcoxon test, W = 36838, P = 0.012).

Two other individual data sets were also considered: the Greek sample described by Apostolou (2020), and the Indonesian sample described in Nila et al. (2019). For the Greek data, only three categories of men were retained: "exclusively heterosexual", "bisexual", and "homosexual". Age information was missing 10 individuals (or 2.5 %), and was replaced by the mean age of the other individuals. For the Indonesian data, the number of maternal aunts and the corresponding number of cousins were also considered. Three individuals were removed due to incomplete data, resulting in a total sample size of N = 113. For both datasets, individual dates of birth were computed as the year of sampling (June 2018 for the Greek

sample) minus the age. For all data sets, the bisexual category was pooled with the homosexual category.

### Twin data

To find primary data on sexual orientation and twinning, we performed literature searches on accessible databases to find recent publications, and scanned the cited literature. Only papers based on samples of twins were retained (i.e. not primarily sampling homosexuals), for which the monozygotic/dizygotic condition was identified (Table S6). Data for same-sex dizygotic pairs and monozygotic pairs were compared for the percentage of homosexual men. Differences were tested using the Fisher exact test using a 2x2 contingency table (e.g. MZ/DZ vs homosexual/heterosexual).

# Statistical analysis of individual family data

To assess the presence of AE while controlling for OBE in the individual family data from France, Greece and Indonesia, three models were considered. First, a model with the number of cousins from the maternal aunts as the response variable (Model1), in order to assess the fertility of the maternal aunts. Second, a model with the number of siblings as the response variable (Model1) in order to assess the fertility of the mothers. For both models, the variable of interest was the sexual orientation of sampled men (qualitative variable), and the control variable was the male birth rank (qualitative variable) of sampled men. Considering male birth rank as a quantitative variable did not qualitatively change the results. The number of maternal aunts was also a control variable for Model2. As the number of cousins or siblings could be influenced by the age of the sampled men, this age was added as a confounding variable for both models. Generalized linear regression was performed, using a Poisson error structure. When the overdispersion parameter ( $\hat{c}$  = residual deviance/residual

degrees of freedom) was between 1 and 2, a quasiPoisson error structure was used instead. When  $\hat{c} > 2$ , a Gaussian error structure was used. The significance of each independent variable was calculated by removing it and comparing the resulting variation in deviance using the  $\chi^2$  test (for Poisson or quasiPoisson error structure) or F test (for Gaussian error structure), as done by the function Anova from the car R package.

In order to estimate the relative contribution of both OBE and AE on siblings data composition, we performed a Bayesian analysis using a hierarchical model, implemented in the nimble R package (de Valpine et al., 2017). The birth order effect, i.e. the probability p of displaying a homosexual preference according  $p=f(p_0,X)$  to the number of older siblings was modelled as where X is the number of older brothers (for modelling an older brother effect), or the number of older sisters (for modelling an older sister effect), and  $p_0$  is the probability of a firstborn displaying a homosexual preference, in the category considered. Various forms of the function f have been considered, notably logistic, geometric, linear, and polynomial (Table S1). These functions describing the OBE effect were compared by evaluating model fit to the sibling data using WAIC (Gelman et al., 2014; Watanabe, 2013): the mean of ten independent chains was used, each with a length of 20,000 samples and a burning phase of 10,000. To avoid the effects of small sample sizes for the number of older brothers or sisters in categories poorly sampled, we restricted the data to categories including at least 10 individuals. Fertility was assumed to follow a Poisson distribution with rate parameter  $\lambda$ . Three types of heterogeneity in  $\lambda$  were simultaneously considered. First, a temporal variation of  $\lambda$  during the last decades:  $\log(\lambda)$  was modelled as a linear function of the year of birth (yob, continuous variable, centred and scaled) of the sampled individuals,  $\lambda = e^{c_1 + c_2 \cdot yob}$ . Second, the possible presence of a subgroup displaying a higher fertility was modelled with parameter h (with  $h \ge 0$  ). For each individual, the probability that modified fertility rate parameter  $\lambda(1+h)$  applies followed a Bernoulli distribution with parameter  $\varphi$ . Third, the possible presence of a subgroup displaying a higher fertility and simultaneously a larger value of  $p_0$ . This antagonist effect was modelled as an increase in  $\lambda$  by a factor  $1+\beta$ , conjointly with an increase in  $p_0$  by a factor  $(1+\alpha)$  (with  $\alpha$  and  $\beta \ge 0$ ). For each individual, the probability that the AE applies followed a Bernoulli of parameter  $\varphi_{ae}$ . We implemented the model in a Bayesian framework by assigning uninformative prior distributions for all model parameters, and fitting the model using Markov chain Monte Carlo (MCMC) in the nimble R package. The birth order effect and the antagonist effect were simultaneously modelled in Nimble (de Valpine et al., 2017), and support for each effect, in the presence of the other one, was computed using reversible jump MCMC (RJMCMC, Green, 1995). Toggle samplers controlled the inclusion or exclusion of each effect according to RJMCMC transition probabilities. Two indicator variables controlled the presence or absence of the AE parameters ( $\alpha$  and  $\beta$ ), and one indicator variable dictated presence or absence of the OBE or OSE parameter  $(a_1)$ . RJMCMC was run 200,000 iterations, and the mean of the posterior distribution of the binary inclusion variables for each effect were used as an estimate of the support of the effect considered. When posterior samples of the binary inclusion variables appeared to have not converged to a stationary distribution, thus decreasing confidence in the posterior results, longer chains were applied (i.e. 2.106 iterations). Analyses were run in R 3.6.3 (R Core Team, 2020) using version 0.9.0 of the nimble package (de Valpine et al., 2020).

### **RESULTS**

### Methodological developments

1) Relationships between mean birth rank and mean fertility

Let us consider a population of N families with discrete generations, each family having a number of children drawn from a Poisson distribution of parameter  $\lambda$ . Among the  $\lambda N$  expected children,  $N(1-e^{-\lambda})$  are firstborn and  $N(1-e^{-\lambda}-\lambda e^{-\lambda})$  are second born. The probability of sampling an individual with birth order  $j \geq 1$  is (Appendix 1):

$$R_{j}(\lambda) = \frac{1 - e^{-\lambda} \sum_{k=0}^{j-1} \frac{\lambda^{k}}{k!}}{\lambda}$$
 (1)

The expected value  $\overline{R(\lambda)}$  of this sampling distribution is ( Appendix 1, Supplementary materials):

$$\overline{R(\lambda)} = \sum_{j=1}^{\infty} j.R_j(\lambda) = \frac{\lambda+2}{2}$$
 (2)

Simulation was used to verify equation (2) for  $\lambda$  values from 0.5 to 6 (Fig. S1). Simulation was further used to evaluate whether AE or OBE could modify this relationship. A total of 6000 families were generated, with fecundity drawn from a Poisson distribution of parameter  $\lambda$ , and with a 1:1 sex-ratio. The birth order of all siblings was recorded (male birth order among brothers, or female birth order among sisters). Sexual orientation for males was drawn from a binomial distribution of parameter p, with  $p = p_0 = 0.05$ . AE was modelled as an increase in  $\lambda$  by a factor  $1+\beta$ , conjointly with an increase in  $p_0$  by a factor  $p_0$  (with  $p_0$  and  $p_0$  decomposed included an extension of f7 (function f7'), where the parameter  $p_0$  varied linearly with fertility ( $p_0$   $p_0$ 

# 2) Estimating fecundity from sibling data

There are two known biases when fecundity is estimated from sibling data (Keyfitz and Caswell, 2005). First, mothers with no children cannot be sampled with this method, thus inflating fertility estimates. A zero-truncated sampling distribution is thus required. Second, the probability of sampling a member from a sibling class of any given size in the general population is proportionate to the sibling size (review in Berglin, 1980). A correction for this second bias was proposed in 1914, but "it is usual to find that authors are unacquainted with the trap" (Berglin, 1980). Both biases lead to an overestimate of fecundity, this overestimate being predominant for low fecundity values for the first bias (because the probability of sampling the zero-class becomes relatively high), and for high fecundity for the second bias (because the variance in sibling size increases). An unbiased estimate of population fecundity from sibling data, taking into account both sources of bias, and considering that fecundity follows a Poisson distribution, is given by the mean sibling size of the sampled individual (mean number of brothers and sisters, not including the sampled individuals), see Appendix 2, Supplementary materials. Simulation has been used to check the various corrections proposed, for  $\lambda$  values from 0.5 to 6 (Fig. S2). For aggregated data (population samples), the mean fecundity is given by the total sibship size divided by the number of sampled individuals (as

$$\sum_{n=1}^{\infty} P(\lambda) = P(n\lambda) \text{ , where } P(\lambda) \text{ is a Poisson distribution of parameter } \lambda).$$

# 3) Sampling distribution of number of older brothers or older sisters

When men are randomly sampled, in the absence of OBE, the sampling distribution of the number of their older brothers (ob) is given by  $Prob(ob = i) = R_{i+1}(\lambda/2)$ , where R is from Eq. (1), and  $\lambda/2$  is the fertility for males (i.e. half the overall fertility, assuming a balanced sex-ratio). The probability distribution of older sisters and older brothers should be the same, unless some male birth rank categories are over or under represented during the sampling (e.g.

when OBE is operating). Thus, in the absence of OBE, the probability distribution of older sisters Prob(os = i) could be also calculated considering that men of various male birth orders are sampled, giving (see Appendix 1b for derivation):

$$Prob(os=i) = OS_{i} = \sum_{j=0}^{\infty} R_{i+j+1}(\lambda) C_{i+j}^{j} \left(\frac{1}{2}\right)^{i+j}$$
(3)

# 4) OSE and OBE

When an OBE is absent, the sampling distribution of the number of older sisters is given by Eq. 3. When an OBE is present and homosexual men are sampled, some low male birth rank categories are under-represented (e.g. first born), and some high male birth rank categories are over-represented, changing the  $R_i(\lambda/2)$  and  $R_i(\lambda)$  values, thus affecting the sampling distribution of older sisters of Eq. 3. The sampling distribution of the number of older sisters in presence of an OBE is not easily tractable analytically, thus simulation was used to assess if an OSE is apparent when sampling occurs in presence of an OBE only. A total of 5000 families were generated, with mean fecundity 4 and 1:1 sex-ratio. An OBE was modelled by considering that the probability to be homosexual for i > 0 older brothers is increased by a constant *a* proportional to *i* (function f5, Table S1), or otherwise increased by a constant *a1* (function f7, Table S1), with  $a = a_1 = 0.2$ . From a random sample of 500 heterosexual and 500 homosexual men, the proportion of homosexual men is computed for each older brother or older sister category. The mean of 1000 replicates of this process was computed, with OBE modelled using functions f5 or f7, or without OBE as a control. A substantial older sister effect appears when randomly sampling heterosexual and homosexuals in the presence of OBE only (Figure 1, A and B). When OBE is not present, the same sampling process does not generate an OSE effect (Figure 1, C).

### **Empirical data analysis**

# Aggregated family data

A total of 23 pairs of samples of aggregated data were retained from the published literature, thus representing, with the conch dataset, a total of 43,362 homosexuals and 5,141,967 heterosexuals (Table 1). Mean fertility ranged on 1.0 to 6.4 for homosexual samples, and from 1.1 to 4.4 for heterosexual samples. The mean male birth rank was between 1.2 and 3.3 for homosexual samples, and between 1.3 and 2.2 for heterosexual samples. For heterosexuals, the relationship between mean male fertility (i.e. observed number of sons) and mean male birth rank was not different from the theoretical prediction (Figure 2): slope = 0.497 (SE = 0.04), not significantly different from the expected value of 0.5 ( $F_{1,21}$  = 0.007, P = 0.93), and intercept = 0.998 (SE =0.04), not significantly different from the expected value of 1 (t(22) = -0.04, P = 0.97). For homosexuals, this relationship displayed a slope of 0.72 (SE = 0.02), which is significantly higher than 0.5 ( $F_{1,22}$  = 85.5, P < 10<sup>-8</sup>).

The mean birth rank of older sisters was between 1.2 and 3.1 for homosexual samples, and between 1.2 and 2.2 for heterosexual samples. For heterosexuals, the relationship between mean fertility for females (i.e. expected number of sisters) and mean birth rank of older sisters was not different from the theoretical prediction: slope = 0.55 (SE = 0.05), not significantly different from the expected value of 0.5 ( $F_{1,22}$  = 1.17, P = 0.30), and intercept = 0.94 (SE = 0.05), not significantly different from the expected value of 1 (t(22) = -1.1, P = 0.27). For homosexuals, this relationship displayed a slope of 0.67 (SE = 0.03), significantly higher than 0.5 ( $F_{1,22}$  = 23.8, P < 10<sup>-4</sup>).

Data simulation has been used to decipher which phenomenon could generate such a higher slope for the relationship between mean male fertility and mean male birth for homosexuals. When an AE alone was modelled, the resulting slope for homosexuals or heterosexuals were not higher (P > 0.5) that the theoretical value of 0.5 (Table S2). When an OBE alone was modelled, a significantly (P < 0.001) higher slope was observed when the

probability to be homosexual increased linearly with the number of older brothers (function f5, Table S1) or, for a threshold function, when the effect of having at least one older brother increased with the mean fertility (function f7') (Table S2). When AE and OBE were simultaneously modelled, results were globally similar to those with an OBE only. Thus, a slope larger than the expected value of 0.5 suggests the presence of an OBE. Using the aggregated family data, maximum likelihood estimate of the parameters were  $\hat{a} = 0.240$ , SE = 0.23 (function f5), and  $\mu = 0.152$ , SE = 0.14 (function f7').

# Individual family data

For the three datasets, the number of older brothers, older sisters, and siblings are given in Table 1, and descriptive statistics are given in Table 2. All datasets displayed a higher number of older brothers and older sisters for homosexual men (for the Indonesian sample, see Nila et al., 2019), and this difference was significant (Wilcoxon Mann-Whitney, P < 0.05) except for the French dataset (Older brother: P = 0.10; older sisters: P = 0.74). For the Greek and Indonesian samples, maternal fertility was significantly higher (Wilcoxon Mann-Whitney, P < 0.05) for homosexuals than heterosexuals. For the French sample, heterosexuals displayed a non-significantly (P = 0.49) higher maternal fertility. When controlled for birth rank, the fertility of mothers was not different in homosexuals compared to heterosexuals (Indonesia: P = 0.60; France: P = 0.32; Greece = 0.93, Table S3). The same result was found for the fertility of aunts: the number of cousins, controlled for the number of aunts, was not different between homosexuals and heterosexuals, for the same birth order (Indonesia: P = 0.82; France: P = 0.08, Table S4).

Bayesian inference was used to test whether an AE, or an OBE (or an OSE), or both, could be inferred from these individual family data. For each dataset, the various functions (Table S1) describing the variation of probability of a homosexual orientation according to the

number of older siblings (brother or sisters) were compared using WAIC. For an OBE, the function providing the minimum WAIC (or WAIC<sub>min</sub>,) were f7 (for France and Indonesia) and f5 (for Greece). For all datasets, these two functions provided a WAIC value lower than WAIC<sub>min</sub>+2 (Table S5). For an OSE, the minimum WAIC value resulted from function f7 (for Indonesia and Greece) and function f2 (for France). Only two functions (f4 and f7) provided a WAIC lower than WAIC<sub>min</sub>+2 for all datasets (Table S5). Thus function f7 was chosen to describe the older sibling effect for further modelling. This function fits two parameters, the probability, in the dataset, to sample a homosexual with no older siblings ( $p_0$ ), and the increase ( $a_1$ ) of this probability for one or more older sibling, thus describing a constant sibling effect starting with the first older sibling (Figure 3).

For each dataset, sibling data were fitted for an AE simultaneously with an OBE, or with an OSE. Fertility was fitted to take into account two possible sources of heterogeneity (variation of fertility with year of birth, and a subgroup of individuals with a different fertility), in addition to a possible AE. Means of the posterior distribution of the parameters are presented Table 3. In the Indonesian sample, the mean fertility was  $\lambda = e^{\epsilon_1} = 2.25$ , although some individuals (~19%) displayed a higher fertility of  $\lambda(1+h) = 7.72$ ; A decrease of the mean fertility with time was apparent, from 2.25 in 1983.2 (mean year of birth), to 1.66 ten years later (or 0.94 standard deviations later). An AE concerned 27.6% of individuals, mainly homosexuals ( $p_o(1+\alpha) = 0.84$ ), displaying a maternal fertility of  $\lambda(1+\beta) = 5.54$ . In the Greek sample, two groups of individuals were identified, one (54%) with a mean fertility of  $\lambda(1+h) = 2.96$ . No substantial temporal variation was detected. The AE concerned 25% of individuals, whose probability to be homosexual is  $p_o(1+\alpha) = 0.32$ , and displaying a fertility of  $\lambda(1+\beta) = 3.84$ . In the French sample, two groups of individuals were identified, one (55%) with a mean fertility of  $\lambda(1+h) = 3.28$ . A decrease of the

mean fertility with time was apparent, from (computed for the larger fraction  $\lambda(1+h)$ ) 3.28 in 1976.0 (mean year of birth), to 3.0 ten years later (or 0.76 standard deviations later). The AE concerned 49.2% of individuals, whose probability to be homosexual was  $p_o(1+\alpha) = 0.73$ , and displaying a fertility multiplied by  $(1+\beta) = 2.48$ .

Support values for OBE or OSE (in presence of AE), and for AE (in presence of OBE or OSE) were evaluated using RJMCMC. In the presence of an older sibling effect, either OBE or OSE, there was little support for an antagonist effect (Figure 4). The maximum support values were ~20% for the Indonesian dataset. In the presence of AE, there was a large support for an OBE or OSE, in the Indonesian and Greek datasets (all supports > 50%). For the French dataset, support was limited (~30%), or inexistent (<1%), for OBE, and OSE, respectively (Figure 4).

Twin data

A total of 5 samples of twins were retained from the published literature, representing a total of 1561 monozygote and 1065 dizygote twins (Table S6). In each study, homosexuals were not more frequent in the dizygotic twin sample compared to the monozygotic twin sample (Fisher exact test on 2x2 contingency table, P > 0.46). The same result was obtained after pooling the samples (Fisher exact test on a 2x2 contingency table, P = 0.44).

### **DISCUSSION**

Research on the biological determinants of male homosexual preference have long realized that the older brother effect (OBE) and the antagonist effect (AE) can both generate family data where male homosexual men have more siblings, more relatives and more older siblings than heterosexual men. Here, we developed several approaches to disentangle these two mechanisms from empirical population samples or family samples. By analysing three types of datasets with statistical tools correcting for known sampling biases, we were able for

the first time to separately test the actions of the birth rank and antagonist genetic factors on fertility and sexual orientation. We found unambiguous support for the OBE in aggregated population data from 24 independent samples, as well as in two individual datasets out of three. We showed that an apparent OSE can be generated by sampling bias in presence of an OBE, and conclude that the OSE reported in some previous studies is probably artefactual. We found no support for the AE in individual datasets including the extended maternal family, or in twin data. Levels of statistical support for OBE and/or AE, in the various datasets, are shown in Table 4.

Sampling biases in presence of an older brother effect generate an artefactual older sister effect

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"The history of birth order studies is not a happy one". The wanderings mentioned by Price and Hare (1969), and regularly recalled since then (e.g. Berglin, 1980; Keyfitz and Caswell, 2005), correspond to several sampling biases, identified long since, but not always taken into account, while the generation of various ad hoc statistics to test various hypotheses has added to the confusion. For example, indexes proposed to study OSE have not considered sampling biases generated when OBE is present in the population. Thus, the report, in a recent meta-analysis, of a widespread OSE in addition to the OBE (Blanchard and Lippa, 2020), should be treated with caution, as we have shown how an apparent OSE is generated when only OBE is acting (fig. 1). The explanation is simple: individuals with more older brothers are more often sampled from larger sibship size (i.e. from a mother with a higher fertility), thus with also more older sisters, considering an even sex-ratio. This correlation between the number of older brothers and the number of older sisters has been widely acknowledged previously (e.g. Blanchard and VanderLaan, 2015), but not sufficiently considered. This sampling bias does not rule out the action of a genuine OSE in population data, but any claim

for an OSE, or for any additional sibling effect, should first control for the sampling bias generated by OBE. The sampling distribution of the number of older brothers or sisters in a sample of men (Eq. 1 and 3) is a first step to develop adequate statistics, although an explicit form of OBE should be introduced to define the sampling distribution of the number of older sisters, in presence of OBE. We thus conclude that there is, to date, no conclusive support for an OSE in empirical data. The only exception is perhaps from Ablaza et al. (2022), although their new regression method, using several highly correlated variables, requires a formal validation.

The older brother effect is well supported but its effects depend on mean population fertility

The generality of the OBE has generated much recent debate fuelled by analyses of aggregated data (Blanchard, 2018c; Blanchard et al., 2020a; Vilsmeier et al., 2021). Our reanalysis of all available and relevant aggregated data strongly supports the generality of the OBE, and excludes it being an artefact from AEs. Compared to previous studies, we believe we took several steps to clarify the situation. First, we filtered out dubious samples, such as samples not corresponding to adult homosexuality, such as pedophiles, or corresponding to non-representative populations, such as sex offenders, transexuals, psychoanalytic or hospital patients (Zietsch, 2018). Thus, our results can be safely associated with standard homosexual men. Second, we derived the relationship between mean birth rank and mean fertility in population samples, and showed that its slope was affected by the OBE but not by the AE: an OBE generates different slopes for homosexual and heterosexual samples while an AE moves the homosexual sample towards higher fertility but with the same slope as heterosexual men. We then showed that samples from heterosexual men fall on the expected line but that samples from homosexual men have a steeper slope: the increase in mean birth rank for a given increase in mean fecundity is higher for homosexuals than heterosexuals (Fig 1A),

which can thus be safely attributed to an OBE and not to any confounding fertility effect such as the AE. This also suggests that the expression of the OBE is fertility-dependent, and that a higher fertility triggers a stronger OBE. Thus, the probability of being homosexual must increase with fertility, and there are two ways to achieve this effect. First, the probability of being homosexual increases with male birth order, and mean male birth order increases with fertility. Second, the probability of being homosexual is constant for one or more older brothers, but this constant increases with mean fertility.

The fact that the expression of OBE is fertility-dependent has a consequence for the study of OBE. When indexes are used to control for fertility, such as those proposed by Slater (1962), Berglin (1980), Blanchard (2014; 2018b; 2018c), or Vilsmeier et al. (2021), the implicit assumption is that homosexual and heterosexual samples are compared independent of the mean fertility level, and the variation of those indexes as a function of the mean fertility is therefore not evaluated. When the mean fertility is low (e.g. male fecundity < 1), the mean rank of homosexuals is not very different from that of heterosexuals, and the OBE effect is not readily apparent (see Fig 2A). This probably explains the reports of an absence of OBE in single-sample studies from low fertility populations, e. g. from France (this study), and UK (Kishida and Rahman, 2015; Rahman et al., 2008). Thus, in a meta-analysis, including samples from populations with various fertility values, results will probably depend on the relative importance of samples from low or high-fertile populations, where OBE is differently expressed, unless the fertility-dependent expression of OBE is explicitly considered.

*The shape of the older brother effect remains elusive* 

At which rate additional older brothers increase the probability of homosexuality is not known. Five functions (f3 to f7, Table S1) were equally likely to fit the individual data in the three family datasets (Table S5), including a logistic, geometric, and linear functions. This

low resolution is explained by the paucity of individuals displaying a relatively high number of older brothers (e.g. ob > 3), thus precluding distinction of the various functions. We know of only one previous attempt to infer the shape of the OBE: Cantor et al. (2002) used the data from Blanchard and Bogaert (1996a) and Blanchard et al. (1998) to fit linear and quadratic functions, and concluded that quadratic terms (2<sup>nd</sup> or 3<sup>rd</sup> degree) were non-significant. A linear relationship between number of older brother and probability of being homosexual has been generally assumed since this seminal work. Here, various forms of piecewise linear functions are equally likely, such as a change in slope from the first older brother (f6), including a flat one (f7, i.e. an increase in probability of being homosexual when having one older brother and no additional effect for additional older brothers), although logistic or geometric functions cannot be discarded. Large samples from high fertility populations would be necessary to achieve a reasonable number of individuals with 3 or more older brothers, allowing discrimination between the various functional forms.

# Proximate and ultimate mechanisms of the older brother effect

The main candidate for a proximate mechanism for the OBE is a maternal immune response to male-specific antigens (Blanchard and Bogaert, 1996b; Bogaert and Skorska, 2011), for which possible molecular evidence has been recently presented paert et al., 2018). Based on current knowledge, this OBE is only found in humans and not in any other species of mammals, even those closely related. This suggests that the OBE is not a mere constraint of the gestation in primates, and thus the interference of male birth order with sexual orientation requires an evolutionary explanation. Under this hypothesis, the OBE would be an adaptive plastic manipulation of the phenotype of male offspring by the mother. Nila et al. (2019) have proposed that the OBE could decrease competitive ability of later-born sons, which would reduce the cost of sibling competition in males. Such a mechanism could

be selected for in patrilocal societies, but probably not in matrilocal ones, where males usually migrate, thus reducing local competition. It should be then interesting to evaluate if an OBE is present or not in matrilocal societies. More generally, the relative contribution of adaptive responses and developmental constraints in shaping the OBE, and the selective pressures generating the OBE, remain in urgent need of investigation.

*No support for antagonistic pleiotropy through female fertility* 

After controlling for the confounding effect of the OBE on fertility in families of heterosexuals and homosexuals, we have found no direct association between higher maternal fertility and male homosexual orientation, i.e. no support for genetic factors increasing fertility of females and increasing at the same time the probability that any given son is homosexual. The larger sibship size displayed by homosexual men is indeed best explained by their high male birth rank (or OBE). More fertile women are more likely to produce homosexual sons because they are more likely to produce sons with a high birth rank (thus with several older brothers), and not because they have a higher propensity to produce homosexual sons at any given birth rank, compared to lower fertility women. Sampling homosexual men, randomly relatively to their birth rank, will thus result in individuals with a higher number of older brothers due to OBE, but also with a larger number of siblings. These highly fertile mothers are likely to have sisters also displaying high fertility, due to correlation of fertility among sisters (Anderton et al., 1987; Berent, 1953). Controlling for male birth rank showed that mother or aunt fertility does not differ between samples of homosexuals and heterosexuals. In addition, AE was neither supported in the analysis of individual datasets, including one displaying a high fertility (Fig. 3), nor in the analysis of twin data (Table S6).

The antagonist pleiotropy hypothesis proposes that the reproductive cost of homosexual men is at least counter-balanced by a reproductive advantage of relatives, and

that both effects are driven by the same genetic factors. If the advantage is greater than the cost, then those genetic factors increase in frequency, and the frequency of homosexual men increases in the population up to the point where the cost is too high (if the fitness loss of males is sufficiently large, leading to protected polymorphism, see Gavrilets and Rice, 2006). It was first proposed that maternal female relatives were concerned (e.g. mothers and aunts), and expressed a higher fecundity (Camperio-Ciani et al., 2004b). The higher fecundity of maternal female relatives was subsequently also found in several independent datasets (Iemmola and Camperio-Ciani, 2009; Rahman et al., 2008; Vasey and VanderLaan, 2007) However, the OBE was not controlled for in those analyses, possibly leading to artefactual results due to the sampling biases we have described above. Unfortunately, the original data used by Camperio-Ciani and collaborators to test for the AE are no longer available (Camperio-Ciani pers. comm., February 2020), and could not be reanalysed. Until additional data or additional analyses are presented, we suggest that there is currently no evidence for antagonist pleiotropic factors favouring female fertility at the cost of increasing the frequency of homosexual sons.

Zietsch et al. (2008) were the first to suggest that the fitness advantage of genetic factors increasing the occurrence of homosexuality could be expressed by heterosexual relatives of both sexes, in the form of a higher number of sex-partners. The idea is that genes predisposing to homosexual orientation may also increase mating success in heterosexuals. This hypothesis has recently received empirical support from genomic evidence for such pleiotropic genes (Zietsch et al., 2021), where the pleiotropic advantage associated with male or female homosexuality seemed to be restricted to a mating advantage of heterosexual male relatives (Table S2 of Zietsch et al., 2021). Reproductive output is notoriously difficult to measure for males, due to variable mating strategies and extra-pair copulation, although paternity uncertainty turned out to be low in some human populations (Larmuseau et al.,

2016, 2016; Larmuseau et al., 2019). Further studies are required to confirm the presence of such an intra-sex antagonist effect. Interestingly, the cross-sex genetic correlation for male homosexuals (i.e. between male homosexuals and number of children for female relatives) in Zietsch et al. (2021) was non-significant, consistent with the absence of AE for female fertility that we report here.

### Limits and future directions

This study has several limitations, although none of which call our results into question. First, the aggregated data mainly remes on published studies, thus generating a potential publication bias (excess of publication with significant OBE). However, the new method of analysis proposed here should not be very sensitive to such publication bias, as it mainly relies on samples displaying different fertilities (in order to estimate the slope of the increase of mean birth rank as a function of mean fertility). A larger proportion of studies from high-fertility population would nevertheless strengthen the results. Second, only three individual datasets were analysed, thus restricting the generally of the results concerning the joint analysis of OBE and AE. Finally, modelling assumed that maternal fertility in human populations follows a Poisson distribution, which is not always the case (e.g. Austerlitz and Heyer, 1998; Hruschka and Burger, 2016). Even if several sources of additional fertility variability have been explicitly incorporated, alternative probability distributions, such as the negative binomial, should be also considered.

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Table 1. Aggregated data collected from published studies. For each pair of homosexual and heterosexual male samples, the number of focal individuals (N), their total number of older brothers (OB), older sisters (OS), and siblings (older and younger) are shown, as well as the origin of the sample and the reference.

Reference	Sample origin	Sexual orientation	N	ОВ	os	sibs
Zucker and	USA. White volunteers	Homosexual	569	286	256	1104
Blanchard, 1994	from earlier studies <sup>a</sup>	Heterosexual	281	123	100	528
Blanchard and	Canada : white volunteers <sup>b</sup>	Heterosexual	434	209	206	977
Bogaert, 1996a	Gunda : winte volunteers	Homosexual	302	213	182	735
Blanchard and Bogaert, 1996b	USA: white individuals	Homosexual	799	556	470	1814
Doguett, 19900	from Kinsey database <sup>a</sup>	Heterosexual	3807	2223	2052	8667
Blanchard et al.,	USA, UK : volunteers,	Heterosexual	225	73	96	357
1998	from earlier studies <sup>a</sup>	Homosexual	385	205	164	728
Bogaert, 1998	USA: non-white individuals from Kinsey	Heterosexual	594	630	534	2370
Dogaett, 1990	database.	Homosexual	229	237	182	810
Ellis and	USA, Canada : volunteers <sup>a</sup>	Homosexual	175	117	85	374
Blanchard, 2001		Heterosexual	971	494	482	1892
Bogaert, 2005	UK, USA: individuals from prob. sample of households from earlier	Homosexual	79	72	50	204
	studies	Heterosexual	2721	1870	1758	7249
Blanchard et al., 2006	Canada : volunteers from previous studies (sample « Bogaert (Other) ») <sup>a</sup>	Heterosexual	148	75	67	242
		Homosexual	267	219	174	655
Frisch and Hviid, 2006	Denmark: national cohort <sup>d</sup>	Homosexual	1890	699	594	2704
		Heterosexual	429181	147704	117529	651577
Blanchard and	UK & other : BBC internet	Homosexual	79519	35580	35368	143134
Lippa, 2007	survey	Heterosexual	8279	4387	4235	15326
Vasey and VanderLaan, 2007	Samoa : volunteers	Homosexual	83	188	173	533
		Heterosexual	114	140	142	497
Bogaert, 2010	UK : probability sample from previous study	Homosexual	132	90	88	331
		Heterosexual	5472	3174	3119	12148
Schwartz et al.,	USA & Canada:	Heterosexual	873	486	446	1892
2010	volunteers	Homosexual	677	539	445	1891
VanderLaan and Vasey, 2011	Samoa : volunteers	Homosexual	133	255	226	747
		Heterosexual	208	179	212	903
Kishida and	UK : volunteers	Homosexual	905	570	534	1891

Rahman, 2015						
		Heterosexual	999	559	529	2117
Currin et al.,	USA: internet volunteers <sup>a</sup>	Homosexual	118	61	57	261
2015	COTT CAMELINE COMMITTEE	Heterosexual	500	285	245	1080
Skorska and Bogaert, 2017	USA: national representative cohort of adolescents, sampled when	Homosexual	225	68	36	289
- 6	adult <sup>d</sup>	Heterosexual	6562	1722	1480	8675
Xu and Zheng,	China: internet volunteers	Homosexual	481	118	226	484
2017		Heterosexual	392	108	164	438
Swift-Gallant et al., 2018	Canada, USA, UK, Australia, New-Zealand : volunteers	Homosexual	243	141	122	467
		Heterosexual	91	50	39	191
Nila et al., 2019	Indonesia : volunteers	Heterosexual	62	46	71	210
		Homosexual	113	120	140	432
Apostolou, 2020	Greece: internet volunteers <sup>b</sup>	Heterosexual	593	206	182	850
•		Homosexual	221	107	87	336
Gómez	Mexico : volunteers	Homosexual	244	284	278	855
Jiménez et al., 2020	wiexico . voiunteers	Heterosexual	194	145	135	527
Ablaza et al.,	Netherlands : marriages, population registers <sup>b</sup>	Homosexual	26542	18890	16145	59984
2022		Heterosexual	4607785	2795390	2687875	10874373
This study	France : volunteers	Heterosexual	241	89	100	405
	Trance . volunteers	Homosexual	271	128	116	425

a. Data available in Blanchard (2018c). b. Data provided upon request. c. Data reconstructed taking into account missing values. d. Data available in Blanchard (2018b).

**Table 2.** Descriptive statistics for the three individual samples. The Greek and Indonesia samples are from Apostolou (2020) and Nila et al. (2019), respectively. Numbers of OB, OS, and siblings are in Table 1.

		Indonesia		Greece			France		
	Homo	Hetero	All	Homo	Hetero	All	Homo	Hetero	All
N	113	62	175	221	593	814	271	241	512
Mean age (SD)	31.6 (9.4)	30.0 (12.6)	31.0 (10.6)	29.8 (8.5)	35.1 (11.9)	33.7 (11.3)	32.0 (10.0)	35.8 (13.4)	33.8 (11.8)
Mean year of birth	1983.4	1985.0	1984.0	1988.7	1983.4	1984.8	1978.5	1973.1	1976.0
Mean OB	1.06	0.74	0.95	0.42	0.35	0.38	0.47	0.37	0.44
Mean OS	1.24	1.15	1.21	0.39	0.31	0.33	0.43	0.41	0.42
Mean sibs Aunts	3.82 221	3.39 124	3.67 345	1.52 -	1.43 -	1.46 -	1.56 394	1.68 288	1.62 682
Cousins	535	316	851	-	-	-	794	497	1291

**Table 3.** Parameters estimates when AE and OBE are simultaneously considered. Mean and standard deviation (SE) of the parameter posterior distribution. The proportion of individuals concerned (Ind. Conc.) is indicated for parameter values dependent on a latent variable. The variation in time is described by the intercept ( $c_1$ ) and a slope ( $c_2$ ). See text for interpretation of the parameters.

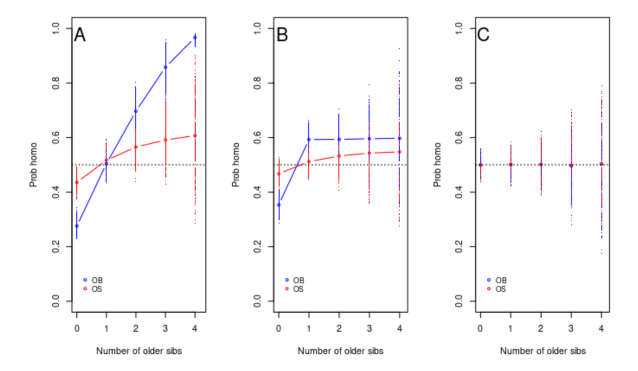
	Indonesia		Greece		France		
	Mear	Mean (SE)		Mean (SE)		Mean (SE)	
Fertility							
Variation in time	:						
c1	0.810	(0.506)	-0.105	(0.588)	-0.271	(0.801)	
c2	-0.321	(0.044)	0.015	(0.032)	-0.117	(0.034)	
Hyperfertility:							
h	2.435	(1.553)	2.286	(2.285)	3.297	(2.212)	
Ind. Conc.	0.190	(0.294)	0.460	0.426	0.446	(0.404)	
FFE:							
α	0.685	(0.584)	0.492	(0.506)	0.853	(0.589)	
β	1.464	(1.350)	3.193	(1.931)	1.483	(2.272)	
Ind. Conc.	0.276	(0.311)	0.249	(0.391)	0.492	(0.411)	
OBE		. ,		. ,		. ,	
p0	0.498	(0.098)	0.215	(0.048)	0.392	(0.110)	
a1	0.155	(0.311)	0.104	(0.035)	0.082	(0.042)	

**Table 4.** Results of the various tests to detect the presence of OBE or AE, in the different datasets considered. A dash indicates that the hypothesis could not be tested.

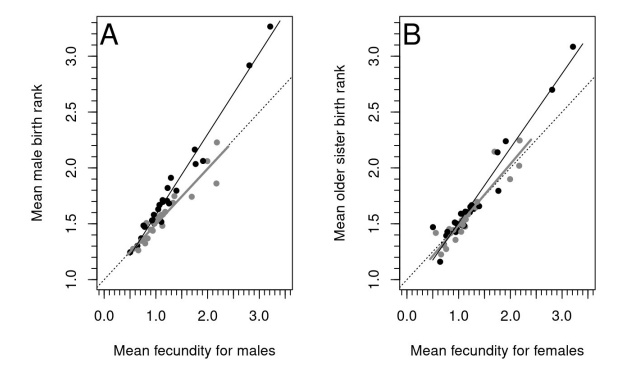
	Hypothesis tested			
Type of data	OBE	AE		
Aggregated data	Yes	-		
Individual data				
Indonesia:				
Mother fertility	-	No		
Aunts fertility	-	No		
Sibs	Yes	No		
Greece:				
Mother fertility	-	No		
Sibs	Yes	No		
France :				
Mother fertility	-	No		
Aunts fertility	-	No		
Sibs	No	No		
Twin data	-	No		

## **FIGURES**

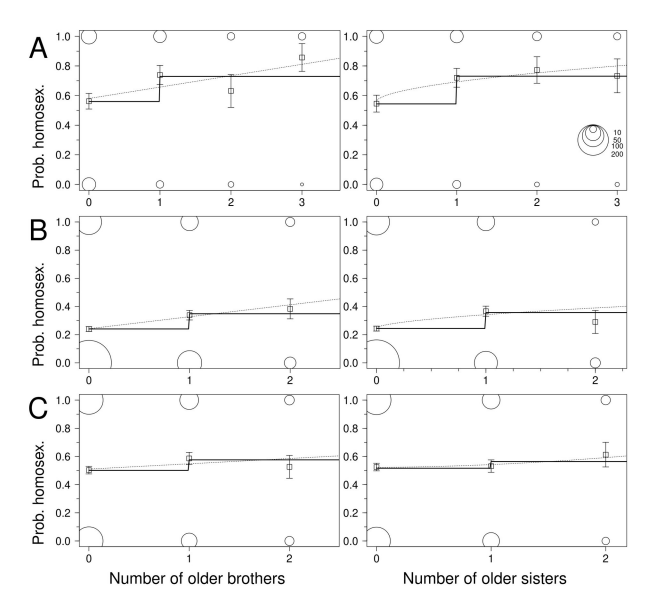
**Figure 1**. Apparent OSE when sampling men from a population with only an OBE. A) OBE generated with function f5 (cf Table S1). B) OBE generated with function f7. C) Control (no OBE). For each of the 1000 replicates, the proportion of homosexual men is computed for each older brother (blue) or older sister (red) category, and is represented as a dot. The mean for these replicates, for each category of older siblingss, is depicted as an empty circle with the corresponding colour. The dotted black line represents the expected curve when sexual orientation is independent of male birth rank.



**Figure 2.** Mean birth rank in relation to fecundity. A. Birth rank among males. B. Birth rank among older sisters. Each sample is represented as a dark dot for homosexuals, and as a grey dot for heterosexuals. The solid lines are regression lines for homosexual samples (black), and heterosexual samples (grey). The dotted line represents the theoretical expectation between mean birth rank and mean fecundity (i.e. Equation 2).



**Figure 3.** Modelling the older sibling effect. Data for the three datasets are presented (A: Indonesia; B: Greece; C: France), for each type of older sibs (left: older brothers; right: older sisters). The solid lines correspond to function 7. The dotted lines correspond to function 5 (left panels) or function 4 (right panels). For each older sibling category, circle areas are proportional to sample size, according to the reference top-right. The frequency of homosexuals is depicted as a square, with the corresponding +/- SE range. Data with elevated number of older siblings (A: more than 3; B-C: more than 2) are not represented.



**Figure** 4. Support for an antagonist effect (AE) or an older sibling effect, conjointly modelled. Coordinates of each point represents the support for AE in presence of an older sibling effect (x-axis), and the support for an older sibling effect in presence of an AE (y-axis). The older sibling effect is either an older brother effect (OBE, full symbol), or an older sister effect (OSE, empty symbol). Point shapes vary according to datasets (circle, square, and triangle for Indonesia, Greece, and France, respectively).

