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ABSTRACT
Numerous articles have been published linking consanguineous marriage to an elevated prevalence of congenital heart disease, with ventricular septal defects and atrial septal defects the most commonly cited disorders. While initially persuasive, on closer examination many of these studies have fundamental shortcomings in their design and in the recruitment of study subjects and controls. Improved matching of cases and controls, to include recognition of the long-established community boundaries within which most marriages are contracted, and the assessment of consanguinity within specific levels and types of marital union would improve and help to focus the study outcomes. At the same time, major discrepancies between studies in their reported prevalence and types of congenital heart disease suggest an urgent need for greater standardization in the classification and reporting of these disorders.

Keywords: Congenital heart defects, consanguinity, endogamy, population stratification

INTRODUCTION
Marital unions between close biological kin are common in many populations, especially in South, Central and West Asia, and North and sub-Saharan Africa, and in migrant communities from those regions now resident in Europe, North America and Oceania.[1] From a medical genetics perspective, all marriages between couples related as second cousins or closer are regarded as consanguineous (derived from the Latin con sanguineus; i.e. sharing the same blood), and using this definition, it has been estimated that at least 10.4% of the current world population of 7.0 billion persons are consanguineous, with first cousin marriages especially popular.[1]

Despite the many investigations that have been conducted into the relationship between consanguinity and congenital heart disease (CHD), the precise nature and significance of the association remains unclear. At the most basic level it would be reasonable to assume that if a robust relationship could be demonstrated, it would be indicative of a causative mutation or the faulty expression of one or a number of otherwise rare recessive genes present in the parents in heterozygous form and adversely affecting embryonic and fetal development when homozygous. However, many of the more common CHDs appear to be genetically heterogeneous, whether diagnosed as isolated anomalies or accompanied by other heart defects.[2] If common predisposing genes are involved in the expression of CHD then, by definition, the likelihood of an association with consanguinity would be substantially reduced.[1]

The epidemiological data often are ambiguous, and although in many instances the results are suggestive of an association between consanguinity and CHD, they frequently fail to attain statistical significance. Several explanations can be advanced for this shortcoming, relating respectively to the diagnostic methods and disease criteria that are employed, the mode of recruitment of study subjects and the numbers and types of cases involved, the demographic and genetic structures of individual sub-populations and communities, and the methods of consanguinity assessment used, with the latter two factors especially relevant in the context of the Indian subcontinent.

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DIAGNOSTIC METHODS AND CRITERIA FOR CHD IDENTIFICATION

In most studies, CHD has been diagnosed by echocardiography and/or cardiac catheterization, with a birth prevalence of 4–8/1000 typically cited and approximately 90% of CHD reported as being multifactorial in origin. However, in a review by the American Heart Association Congenital Defects Committee endorsed by the American Academy of Pediatrics, an overall CHD incidence of 50/1000 births was described as conservative, on the grounds that a number of commonly occurring defects are routinely excluded from prevalence estimates. For example, bicuspid aortic valve, which affects 10–20/1000 people in the general population and may be associated with considerable morbidity and mortality in later life. Further, in studies on Israeli neonates, the incidence of ventricular septal defects (VSD) alone was 53/1000. The scale of the discrepancy in total CHD incidence between the generally accepted level of 4–8/1000 and 50+/1000 live births therefore raises significant concerns, and merits detailed attention and explanation.

As a first step it is important to determine exactly which types of heart defect are under investigation, the methods of ascertainment employed, the defects recorded and reported, and of equal importance which types of CHD were not identified or identifiable by the study techniques. With just a few exceptions, a consistent positive association has been reported between consanguinity and VSD and atrial septal defects (ASD), but both positive and negative associations between consanguinity and patent ductus arteriosus, atrioventricular septal defect, pulmonary atresia, tetralogy of Fallot and other CHDs have been recorded in different populations.

What is presently unclear is whether these inter-population differences are real and indicative of population-specific patterns of CHD, which in turn could be genetic and/or environmental in origin. Or if they also reflect variable diagnostic capacities, expertise and experience, the case definitions employed in the different study centers which have contributed to the literature, and whether ICD codes for malformations of the cardiac system were routinely applied, and disorders were subtyped in terms of embryological timing.

SELECTION AND COMPOSITION OF THE STUDY SAMPLE

A second area of concern relates to the composition of the study populations included in CHD investigations. While neonates only were recruited in a Lebanese study that reported a significant association between consanguinity and CHD, many other widely cited studies variously recruited symptomatic infants, including some with chromosomal anomalies such as Down syndrome, children of differing ages, and even adults in the third and fourth decades of life. Likewise, while some studies included control for the potential effects of sociodemographic and lifestyle variables, such as maternal age and education, socioeconomic status, and smoking, and clinical conditions including pre-eclampsia, maternal diabetes, and pregnancy-induced hypertension, other studies did not.

Age-independent case selection makes rigorous inter-study comparisons well-nigh impossible, as defects that would, or at least could, have been diagnosed shortly after birth may have spontaneously corrected early in life. For example, in the Israeli study in which no association was found between consanguinity and the incidence of VSD, the defects detected were asymptomatic and 89% closed spontaneously within the first 10 months of life. This outcome was confirmed in an associated study on preterm neonates which reported 87.5% spontaneous closure of VSD by age 6–11 months.

Many of the more seriously affected cases of CHD diagnosed in developing countries die in childhood, and unless assessed in early infancy they would not be included in prevalence statistics. By comparison, adults who are not diagnosed with CHD until their 20s and 30s but are included in prevalence statistics presumably do not have an especially life-threatening condition. Their inclusion may thus convey an erroneous impression as to the nature and clinical severity of the underlying cardiac lesions in the population. With these factors in mind, the unqualified use of mixed-age study groups in the investigation of CHD and the calculation of prevalence rates is questionable. More specifically, in the present context, is it appropriate to use the results obtained with such heterogeneous patient groups to assess the influence of consanguinity on CHD prevalence and clinical outcomes?

POPULATION STRATIFICATION IN CASE-CONTROL AND ASSOCIATION STUDIES

To ensure scientific and clinical relevance, case-control and association studies rely on the close matching of affected persons with other individuals who do not exhibit symptoms and have no family history of the disorder under investigation. As previously noted, for this reason cases and controls are routinely compared and matched in terms of age and gender, and with respect to variables such as residence and socioeconomic status. While this type of matching might be sufficient when largely environmental disorders are under investigation, when a genetic predisposition or cause is suspected it is of obvious importance that matching in terms of
inheritance also needs to be ensured.

This requirement is especially important in South Asia due to the very marked patterns and levels of population stratification. For example, in India, there are numerous co-resident ethnicities and religions, some 50,000–60,000 caste and non-caste communities, and up to 30% uncle-niece and first cousin marriages reported in the southern states of Andhra Pradesh, Karnataka and Tamil Nadu, and in southern Maharashtra.[19-22] Similarly, in Pakistan, the population is subdivided into ethnic communities and biraderi (literally translated as “brotherhoods”), that is, traditional social and occupational groupings inherited through the male lineage, and with first cousin marriage rates of 40–50+% consistently reported in regional and national surveys.[23-25] In both countries, the long history of marriage within strict community boundaries means that significant genetic differentiation predictably would have occurred, with some disease mutations unique to specific communities. Ongoing studies have suggested that on average each human carries 50–100 genetic variants that have been implicated in inherited disorders.[26] Ample opportunities would therefore have existed for major inter-community genetic differentiation during the estimated 3000 years since the initial establishment of the caste system.

It is the routine failure to control for population genetic subdivisions brought about by community endogamy that is of greatest significance and concern in association studies, including the investigation of CHD. Besides the widely recognized geographical, ethnic and religious divisions, the genetic effects of caste and sub-caste endogamy have been clearly demonstrated in India,[22,27,28] and an equivalent role is played by biraderi subdivisions in the Muslim populations of India and Pakistan.[29-31] Yet, in the selection of cases and controls, little attempt appears to have been made to control for these differences, despite the clear role of community-specific founder mutations in many genetic disorders,[1] for example, β-thalassemia in the populations of the subcontinent.[32,33]

This problematic situation was demonstrated in a bibliographic exercise based on information compiled from PubMed, using the terms association study, case-control study and clinical trial as keywords and matching of these terms with caste or biraderi and consanguinity.[34] Of the 948 association studies published on Indian populations, only 4.2% mentioned caste, while for 2030 case-control studies the comparable figure was 1.9%, and for 1381 clinical trials it was 0.5%. The data for Pakistan were essentially the same, with just 6.6% of association studies, 1.9% of case-control studies and 0.6% of clinical trials indicating caste or biraderi affiliations. The combined data therefore suggest non-recognition or inadvertent avoidance of the central roles played by caste and biraderi membership in determining the genetic structure of South Asian populations.

Failure to account for the probability of inter-community genetic differences means that the results obtained with case-control and gene association studies may at least in part represent flawed recruitment protocols and thus inappropriate investigative design, which could explain the notable lack of reproducibility in many gene association studies. However, another factor to be considered is that because of the highly endogamous nature of caste and biraderi lineages, even children described as “non-consanguineous” because their parents were unrelated prior to marriage may in fact be homozygous at a significant proportion of their gene loci.

This phenomenon relates to the deeper ancestral heritage of humans and has been clearly demonstrated in studies in Asian, European and North American populations.[35-38] Even families known not to have contracted consanguineous marriages for at least 5–10 generations exhibit numerous regions of the genome in which there are long uninterrupted runs of homozygosity that are indicative of close kin unions in much earlier generations.[36,37]

**ASSESSING THE INFLUENCE OF CONSANGUINITY IN CASE–CONTROL AND ASSOCIATION STUDIES**

The situation with regard to non-recognition of the possible influence of intra-familial marriage in case-control and association studies is actually more extreme, with in India consanguinity mentioned in just 1.0%, 0.2% and <0.1% of association studies, case-control studies, and clinical trials, respectively, possibly due to the strict avoidance of unions between biological relatives in many communities. But the fact that only 4.6% of association studies, 4.0% of case-control studies and no clinical trials conducted in Pakistan discussed the possible genetic effects of consanguinity in a country where 40–50+% of all marriages are between first cousins can be highly misleading, especially for clinicians and researchers who may be unfamiliar with the high prevalence of this form of marriage and its consequences in terms of increased genomic homozygosity.[34]

Even when consanguinity is included as a variable, as has been the case in many studies of CHD, the evaluation very frequently is reduced to a simple “consanguineous” versus “non-consanguineous” dichotomy. In an uncle-niece or a double first cousin marriage, the couple is assumed to have inherited 1/4 of their genes from a common ancestor, whereas in first cousin unions the assumption is that the couple has inherited 1/8 of their genes from a common ancestor, and for a second cousin...
couple the comparable proportion is 1/32. This means that on average the progeny of an uncle–niece or a double first cousin marriage will be expected to have inherited identical gene copies at 1/8 of all their loci, defined as a coefficient of inbreeding $F = 0.125$. It follows that for first cousin progeny, $F = 0.0625$, that is, 1/16 loci predictably are homozygous, whereas for second cousins, $F = 0.0156$, that is, 1/64 of loci are homozygous.$^{39}$

It is unclear and unhelpful why in investigating the impact of consanguinity on the prevalence and types of CHD, researchers very frequently have opted to coalesce three quite distinct levels of consanguinity ($F = 0.0156, 0.0625$ and 0.125), representing an eightfold average difference in genomic homozygosity, into a single ambiguous and ill-defined “consanguineous” category. Failure to recruit adequate numbers of cases in each specific category of consanguineous union is not a sufficient excuse for the adoption of this practice in order to attain statistical significance.

It is also important to recognize that the preferred types of consanguineous marriages vary between communities with, for example, uncle–niece marriages commonly contracted in Dravidian Hindu communities in South India, whereas first cousins are by far the most common form of consanguineous marriage in Muslim populations.$^{40}$ Also, since in many communities consanguinity is a long-established tradition, the effect of uninterrupted intra-familial marriages contracted in successive generations would have resulted in much higher cumulative coefficients of inbreeding than would be predicted on the basis of consanguineous marriage in a parental generation only.$^{41}$

Perhaps because of these complex and variable factors, it was not unexpected that a study conducted on consanguineous progeny in Bangalore, the capital of Karnataka where uncle–niece and first cousin marriages remain popular, was unable to identify a single gene of major effect in a clinically heterogeneous cohort of patients with CHD.$^{42}$ As suggested by the authors, the statistical power of follow-up studies could be significantly increased by investigating consanguineous families in which multiple children have been diagnosed with the same form of CHD. However, a complementary approach which would enable more genetically meaningful comparisons to be drawn would be to match cases and controls according to their community of origin, rather than simply on the basis of parental consanguinity.$^{41,42}$

**DISCUSSION**

Obviously, there must be some practical limitation in the degree to which cases and controls can be matched, and control for potential genetic variables is currently much more intractable than for environmental or sociodemographic factors. At the same time, it should be incumbent on authors to avoid claims for their findings that are based on unproven and/or questionable study designs, particularly when major variables of genetic significance, such as caste/biraderi affiliation and the specific levels of consanguinity of individual cases and controls, have often been effectively ignored or inadequately controlled.

In India, the selection of cases and controls matched solely on the basis of personal religion is inadequate because most members of the Muslim, Christian and Buddhist communities are the descendants of converts from Hinduism in the recent or relatively recent past, and so they would be expected to share significant portions of their genomes with sectors of the majority Hindu community.$^{43}$ Given the approximately 120 generations since the establishment of the caste system, the restricted numbers of founder individuals in each caste, the effects of subsequent caste sub-divisions, and the action of genetic drift, the opportunity for genetic differentiation and the accumulation of caste-specific mutations would have been equivalently large.

From a global perspective, more appropriate subject recruitment based on social and demographic realities should lead to improved investigative outcomes for CHD and other common disorders. This conclusion applies not only to South Asia but also to the Middle East, Africa and many other regions of the world where clan and tribal affiliations should be integral factors in the recruitment of cases and controls. At the same time, to facilitate credible inter-population comparisons, the standardization of diagnostic methods and particularly disease classificatory systems needs to be urgently addressed, together with improved statistical analysis and reporting of findings. Once these basic changes have been effected, it should be possible to more accurately assess the impact of consanguineous marriage on the health of populations in general, and more specifically on the prevalence and types of CHD at family and community levels.

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