The frequency of consanguineous marriages and their effects on offsprings in Tabriz city

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ABSTRACT

Objectives: Too much diversity and ever increasing number of genetic disorders appear as a big challenge in coming future. One of the main sources of genetic disorders is the consanguineous marriages which are, unfortunately very common in our society. In order to prepare ourselves to accept the challenge, the first step is to get complete information of their prevalence and their risk factors.

Patients and Methods: The study was made during 2003-2007 in Tabriz city of Iran. We selected 6000 families and a complete data was obtained on a questionnaire comprised of information regarding marital ages, number of pregnancies, type of delivery, ratio of consanguineous and non consanguineous marriages jobs of parents… etc and their effect on child malformations.

Results: Consanguineous marriages of all types were related with increased congenital malformations (with ratio 43/1000 for consanguineous marriages and for non consanguineous marriages 28/1000). Mother age less than 18 and more than 35 particularly was accompanied with increased malformations while education of mother came out to be inversely related to congenital malformation.

Conclusion: Increased stillbirths, consanguineous marriage and malformations, especially of musculoskeletal system require new planning on national level to control and aware people of the consequences of consanguineous marriages.

Key Words: Consanguineous marriage, congenital malformation, trisomy 21, genetic disease

INTRODUCTION

Congenital malformations are the malformations that are present at birth and may or may not have a genetic basis, some are very clear and present at birth, e.g. Amelia while others show themselves later in life e.g. different metabolic diseases and enzyme related disorders such as phenylketonuria (PKU). These have genetic basis while some others like congenital Rubella syndrome...
is not genetical. On the other hand, both kinds can result in defects called multifactorial disorders e.g. congenital heart disease. A single disorder seen in some members of the family (more than one) is called familial disorder. Familial disorders may or may not be hereditary e.g. congenital hypothyroidism due to low iodine in the environment, is repeated in the family.¹

Various causes of congenital malformations can be divided into 3 categories: Unknown, genetic and environmental. The cause of a majority of human malformations is unknown. A significant proportion of congenital malformations of unknown cause are likely to have an important genetic component. Malformations with an increased recurrent risk, such as cleft lip and palate, anencephaly, spina bifida, certain congenital heart diseases, pyloric stenosis, hypospadias, inguinal hernia, talipes equinovarus and congenital dislocation of the hip, fit in the category of multifactorial disease as well as in the category of polygenic inherited disease. The multifactorial/ threshold hypothesis postulates the modulation of a continuum of genetic characteristics by intrinsic and extrinsic (environmental) factors.

Spontaneous errors during development may account for some of the malformations that occur without apparent abnormalities of the genome or environmental influence. Occurring errors of development may indicate that we are far away from our goal of eliminating birth defects because a significant percentage of birth defects are attributable to the statistical probability of errors in the developmental process, similar to the concept of spontaneous mutation. It is estimated that the majority of all conceptions are lost before term, many within the first 3 weeks of development. The World Health Organization estimated that 15% of all clinically recognizable pregnancies end in a spontaneous abortion, 50% to 60% of which are attributable to chromosomal abnormalities. Finally, 3 to 6% of offspring are malformed, which represents the background risk for human maldevelopment². The chance that both parents are carriers of a mutant allele at the same locus is increased substantially if the parents are related and each of them have inherited the mutant allele from a single common ancestor, a situation called consanguinity.³

First cousin marriages are the most common reason for couples seeking genetic advice; these are legal in many western countries, but may be the subject of religious or social restrictions. In many Asian communities they are actively encouraged.¹ Consanguinity without known genetic disease in the family appears to cause an increase in mortality and malformation rate which is extremely marked in the children of incestuous mating, but which is of little significance when the relationship is more distant than that of first cousins. First cousin marriages, the most common counseling problem, seem to have an added risk of about 3 percent, so that a total risk of 5 percent for abnormality or death in early childhood, about double the general population risk, is a reasonable though approximate guide. It is possible but not certain that the risk is less for populations with a long tradition of cousin marriage; it is only recently that genetic disorders are being fully recognized and accurately diagnosed in these populations. By contrast, some immigrant groups of Asian origin in the UK show an unusually
high frequency of recessively inherited disorders, some extremely rare. This may well reflect increased consanguinity due to isolation and restriction of marriage partners. We performed this research in order to obtain the frequency of consanguineous marriages in our society and evaluate the consequences of such marriages related to congenital malformations in offspring and propose appropriate procedures for awaking people of the futurity.

**PATIENTS AND METHODS**

This study was a kind of social cross sectional investigation that was done during 2003-2007 in Tabriz city of Iran. An ethical approval form was prepared for the research.

The studies in other countries show that some congenital malformations have low incidence rate. Simple epidemiological studies in our country (Iran) also have been the proof performed of this. For involving all malformations it was necessary for the volume of the sample to be selected in a way that by considering P-value = 0.002, confidential results to be gained.

Since prevalence of Congenital Heart Diseases is approximately 1/125 to 1/250 and prevalence of Neural Tube Defects (NTDs) is 1/100 to 1/1500 and prevalence of cleft lip (without cleft plate disorders) is 1/250 to 1/600, so sample volume can be calculated by following formula:

\[
N = \frac{Z^2 \cdot pq}{d^2}
\]

\[
N = \frac{(1/100)^2 \cdot 0.002 \cdot 0.998}{(0.0002)^2} = 1917 \approx 2000
\]

If the average number of the children for each woman under study is considered to be two, for investigating the existence of diseases in these children, 100 families have to be interviewed and if the prevalence of consanguineous marriages is considered approximately 20%, for accessing to 1000 families with consanguineous marriages, 5000 families have to be interviewed. By considering all the aspects and for the results to be valuable and attributable to whole community and also considering at least 2 years period for spouses in order to bring a baby, the statistics experts advised that 6000 families to be interviewed in this research.

Samples were collected by random cluster sampling method and by consulting national cense published in 1993; according to which 287050 families with a total of 1211216 lived in Tabriz.

**RESULTS**

In this study average age for first marriage was 19.5±0.05 years for women and 26±5 years for men. At the time of study, average age for women and men was 35.5±8.4 and 43.4±9.2 years respectively. As for pregnancy experience, 14.5% of women under 20, 73.9% of women between 21-30 years and 11.6% of women above 35 had experienced pregnancy. Average number of pregnancies was 3.12 and average number of children per parents was 2.55. Out of 15301 alive born, 7980 were males and 7321 were females, with a female to male ratio of 1/1.09. Out of 5885 mothers, under study, 1425 mothers were totally illiterate while, 1595 had primary school education, 1040 middle school education, 1452 higher secondary school education and 370 mothers were university graduates. Out of 5695
fathers, enrolled in the study, 908 had primary school education, 1706 had middle school educated, 120 were high school educated, 1179 were higher secondary school educated and 776 were university graduates. Of the mothers 723 (12.28%) of them were working ladies outside the home and 439 (7.45%), did outside work at home and 4723 (80.25%) of them were housewives. Average period between marriage and first pregnancy was 14.14 months.

4696 (79.8%) of marriages were non consanguineous and 1189 (20.12%) were of consanguineous type (Graph 1).

The most common of these consanguineous marriages were third degree maternal first cousins marriage with a rate of 224 (36.07%) out of total 621 marriages (Graph 3).

Out of consanguineous marriages (1189 total), 621(52.22%) marriages were between third degree relatives, 337 (28.34%) between fourth degree relatives and 231(19.42%) between fifth degree relatives (Graph 2).

Number of mothers with single abortion was 1030 (17.5%) while 559 (9.49%) experienced repeated abortions (at least two abortions). There was no significant correlation between abortions and marriage types. Average neonatal mortality rate was 0.017 in non-consanguineous marriages and 0.21 in consanguineous marriages.

Out of 5885 mothers, 1294 of them had relative parents while 4591 had non-relative parents and the rate of infertility in the former was 643 (24%) and 311 (14%) for the later.

In nonconsanguineous marriages, 110 (4.2%) of mothers had experienced single stillbirth and 50 (1.1%) of them had experienced more than one (at least two) stillbirths. In consanguineous marriages 43 (4.3%) of mothers had experienced single stillbirth and 33 (1.8%) of them had two cases of stillbirths. There was a significant statistical correlation (p<0.05) between the age of mother at the time of delivery and congenital defects in offspring (Ch- Square= 7.89, d.f=2).
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Congenital malformations were 1.7 times more in children of mothers with 35 years old or above compared to children of mothers between 18-35 years old. e.g. Down Syndrome (Trisomy 21) was 10 times more in mothers with 35 years old or above than mothers of 18-35 years old and was 2.1 times more in mothers with 15-18 years old (Chi-Square= 19.7999, d.f=4).

A significant correlation was seen between the type of marriage and congenital malformations; so that in case of non-consanguineous marriages congenital malformation rate was 28/1000 and for consanguineous marriages (third degree to 5th degree) it was 43/1000. Although malformations usually were seen on the all body organs, but skeletal system (bones and joints) malformations were the most common (Table 1).

**Table 1:** Prevalence of different congenital malformations.

<table>
<thead>
<tr>
<th>Organ</th>
<th>Risk of malformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bones and joints</td>
<td>0.0083</td>
</tr>
<tr>
<td>Central nervous system</td>
<td>0.008</td>
</tr>
<tr>
<td>Urogenital malformations</td>
<td>0.0038</td>
</tr>
<tr>
<td>Alimentary tract</td>
<td>0.0012</td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>0.005</td>
</tr>
<tr>
<td>Visual</td>
<td>0.0008</td>
</tr>
<tr>
<td>Acoustic</td>
<td>0.0003</td>
</tr>
<tr>
<td>Other</td>
<td>0.0006</td>
</tr>
<tr>
<td>Sum</td>
<td>0.028</td>
</tr>
</tbody>
</table>

There was a reverse correlation between mother education and congenital malformations (Chi-Square= 7.74, d.f=2) however, no significant correlation was found between father age and congenital malformations. As long as jobs of parents is concerned, no relation was observed between occupation and congenital malformations in offspring (Chi-Square= 0.602, d.f=2) (p>0.025).

Of 5885 females enrolled in the study 4914 (83.5%) considered consanguineous marriages as unfavorable and 971 (16.5%) contemplated them favorable. From those who disapproved consanguineous marriages, 4619 (94%) of them related it with increased rate of congenital malformations and those who approved such marriages, preferred them because of more acquaintance of spouse before marriage and also because of low expectations of partners and families.

**DISCUSSION**

The coefficient of inbreeding (F) is the probability that a homozygote has received both alleles at a locus from the same ancestral source; it is also the proportion of loci at which a person is homozygous or identical by descent.

In (Table 2) the coefficients of inbreeding for consanguineous marriages have been shown. If a person is inbred through more than one line of descent, the separate coefficients are summed to find his or her total coefficient of inbreeding.3
Table 2: Coefficients of inbreeding for the offspring of a number of consanguineous mating.

<table>
<thead>
<tr>
<th>Type</th>
<th>Degree of Relationship</th>
<th>Population of Genes in Common</th>
<th>Coefficient of Inbreeding of Child</th>
</tr>
</thead>
<tbody>
<tr>
<td>MZ twins</td>
<td>-</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>Parent-child</td>
<td>1&lt;sup&gt;st&lt;/sup&gt;</td>
<td>½</td>
<td>¼</td>
</tr>
<tr>
<td>Brother-sister (including dizygotic twins)</td>
<td>1&lt;sup&gt;st&lt;/sup&gt;</td>
<td>½</td>
<td>¼</td>
</tr>
<tr>
<td>Brother-half sister</td>
<td>2&lt;sup&gt;nd&lt;/sup&gt;</td>
<td>¼</td>
<td>1/8</td>
</tr>
<tr>
<td>Uncle-niece or aunt-nephew</td>
<td>2&lt;sup&gt;nd&lt;/sup&gt;</td>
<td>¼</td>
<td>1/8</td>
</tr>
<tr>
<td>Half uncle-niece</td>
<td>3&lt;sup&gt;rd&lt;/sup&gt;</td>
<td>1/8</td>
<td>1/16</td>
</tr>
<tr>
<td>First cousins</td>
<td>3&lt;sup&gt;rd&lt;/sup&gt;</td>
<td>1/8</td>
<td>1/16</td>
</tr>
<tr>
<td>Double first cousins</td>
<td>2&lt;sup&gt;nd&lt;/sup&gt;</td>
<td>¼</td>
<td>1/8</td>
</tr>
<tr>
<td>Half first cousins</td>
<td>4&lt;sup&gt;th&lt;/sup&gt;</td>
<td>1/16</td>
<td>1/32</td>
</tr>
<tr>
<td>First cousins once removed</td>
<td>4&lt;sup&gt;th&lt;/sup&gt;</td>
<td>1/16</td>
<td>1/32</td>
</tr>
<tr>
<td>Second cousins</td>
<td>5&lt;sup&gt;th&lt;/sup&gt;</td>
<td>1/32</td>
<td>1/64</td>
</tr>
</tbody>
</table>

In our study no significant correlation observed between mother activity (employed or housewives) with congenital malformations, so in our community type of activity and environment does not act as risk factor. However, in the study performed by Brandt and Neilsen in 1990, mother’s jobs were related to congenital malformations where movie making jobs seemed to be the most affective cause for congenital malformations. Similarly in another case control study a striking correlation was seen between neural tube defects (NTDs) in first trimester and mother being in contact with organic solvents at home or outside.

Influence of father’s profession on anencephaly was studied by Suarez and Breuder in 1990 in Texas City. They showed that children whose fathers were in contact with organic solvents had 1.55 times more risk of congenital malformations. But in our study after grouping fathers according to their different occupations, no statistical correlation was seen between types of jobs and congenital malformations in their children (Chi-Square= 0.602 d.f=2) (P>0.025).

Similarly occupation of consanguine parents had no important consequences on their children malformations. In our study Congenital malformations were significantly more in children of mothers with 35 years old or above compared to children of mothers between 18-35 years old. For example the known Down syndrome (Trisomy 21) was 10 times more in mothers with 35 years old or above than mothers of 18-35 years old and was 1-2 times more in mothers with 15-18 years old. These findings are compatible with other references.

It has been seen a direct correlation between the age of mothers and congenital malformations, particularly pregnancy after 35 years old is accompanied by high prevalence of congenital malformations, which is the main risk factor for Down syndrome and also mother’s age below 18 years old involves the risk of increased congenital malformations, especially numerical chromosomal abnormalities.
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Congenital defects rate came out to be 28 in every 1000 births in the world with average of 25 in 1000 births, but it differs from place to place because of different social customs in marriages, nutritional habits, hygienic conditions and genetic potentials and may be more or less of this average value. Bone and joints malformations are the most prevalent malformations in this study, while in authenticated references central nervous system malformations are seen the most (Table 3). This requires new researches and more investigations for planning and finding ways to decrease musculoskeletal malformation in Tabriz city.

**Table 3: Prevalence of congenital malformations in authenticated references.**

<table>
<thead>
<tr>
<th>Organ</th>
<th>Risk of malformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Central nervous system</td>
<td>0.007</td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>0.004</td>
</tr>
<tr>
<td>Kidney and urinary tract</td>
<td>0.0015</td>
</tr>
<tr>
<td>Limbs</td>
<td>0.0035</td>
</tr>
<tr>
<td>Other</td>
<td>0.0035</td>
</tr>
<tr>
<td>Sum</td>
<td>0.025</td>
</tr>
</tbody>
</table>

In our study neural tube defects rate was a little less than reported elsewhere. This increased rate in other countries may be related to alcoholic consumption in mothers which is not consumed in our country.

In our country visual and acoustic malformations are more common than developed and non Muslim countries and it is because of high rate of consanguineous marriages in our country. As 80% of congenital blindness and 70% of congenital deafness follow autosomal recessive hereditary pattern which express themselves in homozygous conditions resulted by consanguineous marriages.

Congenital blindness and deafness are malformations that do not affect life span too much, but worsen quality of the person’s life. The studies in different countries of the world show that the main risk factor of increase in these malformations is consanguineous marriage. In a study in the north of India, it has been found that the IQ of children in consanguineous marriages is significantly lower than the children in control group that their parents are not consanguine. A study in 1993 in Glasgow of England on 205 blind kids has shown that 44% of them were due to consanguineous marriages. Also investigation in 1991, in Russia on 2848 blind individuals has revealed that 1108 of them were due to consanguineous marriages. In 1991, in America, from 1000 blind children, 42% were children as a result of consanguineous marriages.

In 1996, in Shahid Moradi school of Tabriz, out of 94 blind students 51 (54%) were the consequences of consanguineous marriages. There were 122 congenitally blinds in our study and 32 of them had consanguine parents. Considering this fact that 20.2% of the mothers had consanguineous marriages, if we did not consider consanguineous marriages as the risk factor for increase of blindness in offspring we would expect 24 blind children, while this number has increased to 38 blind children because of the effects of defective and recessive genes in consanguineous marriages. On the whole, 25 in 1000 of the children who were born from non consanguine parents had congenital malformations.
whereas this rate was 43 in 1000 for children whose parents were consanguine. In this study there was no significant correlation between the father education and congenital malformations in offspring (P>0.30), but it was not the same for mothers education and congenital malformation rate (P<0.05).

The malformation ratio in the children of uneducated or elementary educated mothers to the children of the mothers who had secondary school education was 1.1 to 1. This ratio was 1.7 to 1 for mothers with university education. By correlating mother’s education, mother’s age and type of marriage the following formulas were achieved:

\[
\text{Congenital malformation} = 1 + \text{mother education} + \text{type of marriage}.
\]

\[
\text{Congenital malformation} = 1 + \text{mother education} + \text{mother age}.
\]

Above relations show that age, educations of the mothers and marriage type have independent effects on congenital malformations.

The incidence of malformations in males and females was the same but type of malformations was different in two genders (P<0.005). Skeletal and joint malformations such as club foot and congenital dislocation of hip bone, were more common in female children than males (1.3 to 0.8) while urogenital malformation were seen more in male than females. The Urogenital malformations not being appearing in female children at the time of birth might be the cause of such difference which needs more researches to be done by urologists.

Simultaneous studying of type of marriage showed the following formula:

\[
\text{Congenital malformation} = 1 + \text{type of marriage} + \text{age}.
\]

The above model shows that type of marriage and age had no counter effects on congenital malformations. But studying the type of malformations show that neonates of mothers above 35 years old at the time of delivery and with consanguineous marriage were in high risk of congenital malformations compared to those neonates whose mothers had the ages between 18 to 35 years old and their parents were non consanguine. The records of mothers deliveries show that although congenital malformations rate increased with each pregnancy (being 2.5% in first, 2.8% in second, 3% in third and 3.6% in fourth pregnancy and so on), but these differences were not statistically significant. The positive history of abortions had significant correlation with congenital malformations (P<0.05), since mothers with no abortion, one abortion and more than one in the past had 2.7%, 3.7% and 4.8% of children with congenital malformations, respectively.

Also the mothers whose parents were non consanguine had less stillbirth cases compared to mothers whose parents were consanguine. Average stillbirth rate was 0.071 ± 0.035 in the former and 0.12 ± 0.045 in the later. We have no standards for comparing the later group with international statistics but the rate of stillbirths in the former is compatible with the international statistics. It seems that factors like acquaintance of spouses, less economical problem and interference of grandparents in consanguineous marriages and on the
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other hand cultural differences, prevalence of perversity specially possibility of drug addiction in the young in non Consanguineous marriages are the most important factors bringing tendency towards consanguineous marriages.

However, the cooperation of the government and related official institutes for solving the economical problems of the young, warning the young and their families about risks of consanguineous marriages, helping them by providing genetic counseling and required facilities, congenital disease screening projects, preventing Consanguineous marriages seriously and preparing educational informative programs can be effective steps in preventing consequences of the consanguineous marriages.

REFERENCES


