Short communication

Racial effect on serum creatine-kinase: implications for estimation of heterozygosity risks for females at-risk for Duchenne dystrophy

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(Received 12 May 1988; revision received 1 October 1988; accepted 13 October 1988)

Key words: Creatine-kinase; Duchenne muscular dystrophy

Introduction

Duchenne muscular dystrophy (DMD), a lethal X-linked recessive genetic disorder of childhood, is the most common and severe of the muscular dystrophies. Until no effective treatment is found to cure or arrest the progression of the disease, prevention of new cases through detection of carriers and genetic counseling is fundamental.

About 50% to 80% of the heterozygotes for the DMD gene have increased creatine-kinase (CK) and pyruvate-kinase (PK) levels [1–4]. Studies on DNA polymorphisms have greatly improved identification of heterozygotes in informative families. However, techniques of DNA analysis are expensive and time consuming, and the results are not always absolute. Very often, they must be expressed in terms of probabilities, combined with results of serum enzymes determinations. Therefore, serum enzyme determinations will still be important first, as a screening test and second, for calculation of conditional probabilities (combined with DNA marker information through Bayesian methods) in heterozygosity risk estimations for females at risk for DMD [5,6].

Studies on serum CK in normal people from different racial background have shown that the black population has higher serum CK levels than the caucasoids or mongoloids [7–10]. In a recent study, we have compared serum CK and PK in healthy adult females of different black admixture. The preliminary results suggested that while PK did not differ among the different subgroups, CK was significantly higher in the black females than in the caucasoids [11].
On the other hand, results of serum enzymes in possible and obligate DMD carriers showed that the negroids have also higher serum CK levels than caucasoids [12]. These findings may have important implications in estimation of heterozygosity risks for females at-risk for DMD. In order to evaluate this hypothesis, we have extended this investigation and have determined serum CK and PK in 263 healthy female controls classified as caucasoids, negroids and mongoloids. The results were compared through discriminant analysis with serum enzyme activities from 82 caucasoids and 64 negroids DMD obligate carriers.

Subjects and methods

Healthy controls: A total of 263 adult healthy females who had no relatives with any neuromuscular disorders were included in the present study. The name, age, menses data, physical activity and racial group was registered for each individual.

According to ethnical origin, skin colour and facial features, the subjects were classified in three racial groups: caucasoids (C), negroids (N) and mongoloids (M). The negroid group was further subdivided in the following 4 subgroups: light mulattoes, medium mulattoes, dark mulattoes and blacks [13].

DMD carriers: Serum CK results from 146 obligate carriers classified as caucasoids (n = 82) or negroids (n = 64) were considered in the present study. A woman was classified as carrier when: she had a brother or other maternal relative and at least one affected son; two or more affected sons; two or more affected grandsons through different daughters.

Both enzymes were determined in all female controls (40 caucasoids, 50 mongoloids, 173 negroids) and DMD carriers.

Serum PK activity was determined in a Gilford spectrophotometer (model 2400) according to a procedure described elsewhere [14], always in fresh serum since it has been shown that PK activity decreases upon freezing [2,14]. The results are given in µmol/ml per h. Serum CK activity was performed with Sigma kits, in the same aliquots used for PK, with the colorimetric procedure 520-C, according to Hughes [15], as described in Sigma Bulletin [16]. This enzyme reaction is performed at 37 °C. Results are given in Sigma units/ml (S.u.). Usually the samples were assayed within 24 h or at most within 1 wk after collection in aliquots frozen at −20 °C, since no loss of enzyme activity has been observed in samples stored up to 2 mth [15,16].

For statistical analysis logarithmic transformation was performed [log(CK + 1), log(PK + 1)] with resulting distributions close to Gaussian ones. For comparing the enzyme activities of females from different racial subgroups, a variance analysis and Student's t test were applied according to each situation. A Tukey t test was used when a significant result was observed in the variance analysis.

The considered level of significance was 5%.

Results

Results of serum CK and PK determinations (transformed and untransformed data) in normal female controls from the three racial groups are summarized in Table I.
A significantly higher CK activity was observed in the negroid subgroups than in the caucasoid females ($t = 3.9; P < 0.001$). Furthermore, the CK activity varied and increased significantly with the degree of black admixture ($F = 5.08; P = 0.001$) but not with age ($F = 1.59; P = 0.21$). The mean CK activity in caucasoid females was significantly lower than in the groups of medium mulattoes, dark mulattoes and black females ($P < 0.05$), but was not significantly different from light mulattoes ($P > 0.05$).
The mean CK activity did not differ between mongoloids and caucasoids \((t = 1.48; \ P > 0.05)\); therefore, these two samples were joined for further analyses (group CM).

On the other hand, the PK levels did not show any statistically significant difference between caucasoids and negroids \((F = 1.02; \ P = 0.41)\).

In order to evaluate the influence of black admixture on CK activity in the estimation of heterozygosity risks, conditional biochemical probabilities using a quadratic discriminant analysis [4] were determined considering the beneficial background from both carriers and controls. The estimated biochemical probabilities favoring the diagnosis of heterozygosity for caucasoid and negroid DMD carriers are shown in Fig. 1.

Discussion

Effect of black admixture on serum CK levels in healthy controls

Results from the present investigation confirm previous studies suggesting that serum CK activity is higher in healthy negroid females than in caucasoids [7–10] and are similar between caucasoids and mongoloids [9].

However, the gradual increase in the mean CK activity according to the degree of black admixture was apparently not reported previously in females, but only in healthy males [8].

The higher CK activity found in negroids than Caucasoids is still the subject of investigation. According to Meltzer [7] nutritional or physical activities differences are apparently not implicated in these results, and might be related to a greater muscle mass in the negroid group. More recently, Ama et al. [17] reported a higher muscle CK activity and a higher proportion of type 2a fibers in negroid than caucasoid young sedentary males.

It is important to observe that the females ascertained in the present study belong to the same social condition with similar nutritional habits and physical activities. Since the racial classification reflects genetic differences among the different negroid subgroups [13], the results on CK activity seem to confirm the genetic influence on serum enzyme levels already suggested by some authors. Indeed, comparative studies on serum CK in normal twins showed a higher concordance in monozygotic than in dizygotic twin [18,19].

The present results on PK activity confirm our previous suggestion that this enzyme is not influenced by race, as observed previously in a smaller sample of healthy controls [11].

Estimation of heterozygosity biochemical risks for females at risk for DMD according to racial background

Among serum enzymes, the creatine-kinase (CK) test is the most reliable and widely used. The concomitant use of serum PK may improve the capability of detecting heterozygotes females in about 10% [2,3].

Several factors have been reported to affect serum CK activity in DMD carriers such as age, pregnancy and physical exercise [5].
Recently Zatz et al [12] reported in females at risk for DMD the same effect which was found in healthy female controls, that is, a higher mean serum CK in negroids than in caucasoids but no significant difference in mean serum PK between these two populations. It was suggested that the CK difference was more likely due to the racial background rather than genetic heterogeneity in the DMD locus. This was supported also by a previous observation showing that the CK activity in caucasoid DMD boys did not differ from negroids [20].

Through DNA technology a great improvement in the identification of females at-risk for DMD has been achieved. However, to detect all females at-risk for DMD-BMD gene using only DNA probes is still difficult due to the extremely large size of this locus and its reported molecular heterogeneity [6]. The recent identification of dystrophin, described as the protein product of the Duchenne muscular dystrophy gene [21], may represent in the next future, a great step forward the detection of DMD carriers. However, if the DMD gene is expressed mainly in muscle [22], it would require a muscle biopsy, which is not an easy routine procedure. Therefore, results of simple tests, such as serum CK determinations, will still be important as screening and also as an adjunct for estimation of conditional probabilities.

The discriminant functions derived from serum enzyme activity show that the conditional biochemical probabilities favoring the diagnoses of heterozygosity for DMD varied for some CK values between caucasoids and negroids (Fig. 1). For example, if we consider a CK value of 11.0 S.u./ml, the biochemical probability for a female at-risk being a DMD carrier is 72% if she is a caucasoid and 54% is a negroid. These probabilities, combined with data from pedigree through Bayesian calculations will give a final estimate of heterozygosity risks for suspected carriers.

Therefore, based on the above observations it is suggested to take into account racial background for estimation of heterozygosity risks. Such considerations are not necessary for PK as its activity does not seem to vary according to the ethnical background in controls or DMD carriers.

Acknowledgements

The authors thank Dr. José Antonio Levy and Paulo Salum for referring patients and DMD families; Mrs. Marta Canovas and Mrs. Direynia B. Costa for technical assistance; to Dr. Paulo A. Otto and Dr. Fernando Carvalho for statistical analysis; to Dr. David Campion for support; to Debora Rapaport and Mariz Vainzof for reviewing the manuscript. This work was supported with grants from FAPESP, CNPq, CAPES and MDA from America.

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