Molecular Investigation on DNA-PKcs Gene and Identification of SCID Carriers among Iranian Arabian Horses Using a Test Based on PCR

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Abstract: Severe Combined Immunodeficiency Disease (SCID) is an autosomal recessive hereditary disease occurring among Arabian horses. The genetic defect responsible for this disease was recently identified as a 5-basepair deletion in the gene encoding DNA-Protein Kinase catalytic subunit (DNA-PKcs). Horses with one copy of the gene appear normal while horses with two copies of the gene manifest the disease. The only way to avoid the economic losses is early detection of SCID carriers. Because the real value of this test is that breeders can use it to ensure that they will never produce a SCID foal and that they can by breeding carriers to clear, preserve the unique and positive characteristics of exceptional carriers. And thereby breed out the undesirable gene over several generations.

Key words: SCID, DNA-PKcs, Iranian arab horse, genetic defect, carrier, preserve

INTRODUCTION

Severe Combined Immunodeficiency Disease (SCID) has been of grave concern to Arabian horse breeders for the past several decades (McGuire and Poppie, 1973). The first occurrence of SCID in Arabian foals was reported by McGuire and Poppie (1973). The paucity of mature T and B lymphocytes in affected Arabian foals is profound (Magnuson and Perryman, 1986). These foals produce no antibodies after infection or immunization and no evidence of leukemia has been detected SCID foals rarely survive up to 5 months of age before succumbing to infections caused by equine adenovirus, Pneumocystis carinii, Cryptosporidium parvum or many species of bacteria (Lunn et al., 1995). The disease is inherited as an autosomal recessive trait and was suggested that a defect in a single gene is responsible for equine SCID (Thompson et al., 1975; Perryman and Torbeck, 1980). The SCID gene was mapped to chromosome ECA9 (Bailey et al., 1997). Shin et al. (1997) described a 5-basepair deletion in the horse gene encoding DNA-Protein Kinase catalytic subunit (DNA-PKcs) is responsible for SCID in Arabian horses.

The mutation prevented translation of the 967 C-terminal amino acids, resulting in a nonstable protein. DNA-PKcs is related to the Phosphatidylinositol 3-Kinase (PI3K) family in which members function in a variety of roles such as signal transduction, control of cell cycle progression and maintenance of telomere length (Finnie et al., 1995). Variable-diversity-joining rearrangement is the mechanism by which gene segments (V, D and J) are joined to form the coding sequences of Ig and TCR variable regions (Weiler et al., 2000). Rearrangement involves two DNA cuts and relocations and is mediated by a lymphoid-specific endonuclease (the RAG 1 and RAG 2 proteins) and ubiquitous components of the DNA Double-strand Break Repair (DSBR) pathway (Shin et al., 1997).

The centrality of V (D) J recombination to the development of the immune system is illustrated in situations where the process is impaired (Malynn et al., 1988). At least, four factors are required for both V (D) J recombination and DSBR: the Ku heterodimer (Ku 86/Ku 70, XRCC5 and XRCC6), DNA-PKcs (XRCC7) and XRCC4 (Taccioli et al., 1994). Defective V (D) J recombination results in a complete block of B and T cell lymphopoiesis and the disease severe combined immune deficiency (Shin et al., 1997). Bosma et al. (1983) described a spontaneous mutation in C.B-17 mice resulting in defective V (D) J recombination and SCID. In SCID mice, the only step in V (D) J recombination that is impaired is the resolution of coding ends (Peterson et al., 1995).

It was demonstrated that SCID mice also have impaired DNA Double-strand Break Repair (DSBR). Like SCID mice, the immunologic defects in SCID foals can be corrected through transplantation of normal histocompatible bone marrow cells (Shin et al., 1997). This

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study intends to molecular investigation on DNA-PKcs gene and identification of SCID carriers among Iranian Arabian horses using a test based on PCR.

MATERIALS AND METHODS

About 120 whole blood samples were randomly collected among Iranian Arabian horses from Khuzestan and Kordan regions in the South-west and centre of Iran. Genomic DNA was extracted by the salting-out method (Miller et al., 1988) with some modifications. One set of allele-specific primers was designed to amplify only DNA containing the 5-basepair deletion. The primers have the following sequence:

F : 5′- TTC CTG TTG CAA AAG GAG GAG-3′
R : 5′-TTTGTG ATGATG TCA TCC-3′

Polymerase chain reaction was performed on Biometra PCR system. To produce 259 bp fragment of the DNA-PKcs gene the following PCR mix was composed: 1X PCR buffer; 5 mM MgCl2; 0.25 mM primers; dNTPs 200 μM; 1 unit of Taq polymerase; Genomic DNA 150 ng/reaction; dd H2O. The final volume was 15 μL. Samples were amplified under the following conditions: initial denaturation at 95°C for 2.5 min followed by 36 cycles of 95°C for 30 sec, annealing 60°C for 30 sec, extension 72°C for 45 sec, final extension for 5 min. The products were electrophoresed on 8% non-denaturing polyacrylamide gels at 80-v at overnight and bands visualized by rapid silver staining (Sarguinetti et al., 1994).

RESULTS AND DISCUSSION

The results showed total samples were homozygote for 259 bp band (Fig. 1) that is noncarrier. Some studies during 20 years ago indicated that the frequency of affected Arabian foals was around 3% but carrier frequency is about 28% and means that one out of every 3 or 4 adult Arabian horses is carrying the gene for this deadly disease (Poppie and McGuire, 1977; Studdert, 1978). Another way of stating is that 7-10% of all matings are between carriers and thus this proportion of matings is at risk to produce a foal that will soon die from SCID. Probable reasons for these results may be: Preliminary studies of the pedigrees of 21 carrier identified in this study implicate a popular stallion active in late 1920s (Bennico and Bailey, 1998). The pedigree of this stallion was limited to a single generation in the American Arabian horse registry. It may be there is not any progeny from this stallion within studied samples in Iran. The next studies samples were focused on infected flocks and therefore, it is expected high frequency for them. However, the sampling have been randomized. There are two aspects to examine when considering the reduction of SCID. The first is the removal of SCID from the Arab horse population as a whole and the second is the prevention of the conception of affected foals. The ultimate aim for the future would be the complete elimination of the SCID gene from the Arab horse breeding population. This would require the gradual withdrawal of all carrier horses from breeding. If the percentage of carriers in the population turns out to be high, this would take a number of years but would eventually remove the need to test any further, individuals for SCID. Realistically, preventing the breeding of carriers may in fact be undesirable, particularly if highly acclaimed animals prove to be carriers. The SCID defect is only one aspect of a carrier's genetic make-up and the desirable characteristics carried may outweigh this fault. One advantage which the new carrier test provides is that carriers can now be confidently used in breeding programmes with no fear of producing an affected foal. If carriers are only bred with normal horses, there is no possibility of generating a SCID foal. The progeny of all carriers will however, need to be tested to determine whether they are normal or are carriers.

CONCLUSION

The present study describes a PCR-based test for detection of the gene defect and the results from testing 120 randomly selected Arabian horses. There were not any carrier horse among tested individuals. But the researchers suggest further study on Arab horses throughout of Iran to finding any probable carrier.
RECOMMENDATIONS

The use of carriers in breeding should therefore carry no stigma but responsible breeders will now ensure that they do not pair a carrier with a carrier. In the long term, the aim for the breed should be to reduce the frequency of carrier-normal matings to lead to the eventual elimination of SCID whilst maintaining desirable genes. It should be pointed out that continued breeding of horses that are carriers of the SCID gene is indeed now possible without the worry of producing SCID foals. For example, carrier stallions that possess highly desirable traits can now be selectively bred to clear (homozygous normal) mares (and vice versa). The resulting foals would have an equal chance of being a carrier or clear of SCID but would definitely not be affected. The foals could be tested anytime after birth to determine their SCID genotype and future matings could be rationally planned. In this study, researchers suggest further study on Arab horses throughout of Iran to finding any probable carrier.

ACKNOWLEDGEMENT

This research was carried out at Animal Science Research Institute of Iran (Department of Biotechnology). Researchers thank N. Asadi, H. Emrani and S. Dordari for technical assistance.

REFERENCES


