

## Cloning and Sequence Analysis of Donkey Growth Hormone Gene

Wenjin Zhu, Yongmei Su and Jianhua Wu

Key Lab of Preventive Veterinary Medicine, College of Animal Science and Technology,  
Hebei Normal University of Science and Technology, Hebei Province,  
066600 Qinhuangdao, P.R. China

**Abstract:** The PCR primers were designed online according to result of gene homology comparison. Donkey *GH* gene DNA and cDNA sequence were cloned from liver and blood by RT-PCR and PCR, compared with *GH* gene sequences of different species by bioinformatics. The donkey *GH* gene sequence was 1928 bp including the 706 bp cDNA sequence with the complete CDS. By comparison for DNA and cDNA sequences, it was found that the *GH* gene sequence included 5 exons and 4 introns, encoding 216aa including signal protein of 26aa and matured protein of 190aa. Based on the analysis of the similarity of *GH* genes in different species on the level of cDNA, DNA and the deduced amino acid there was the most homology to the horse. The *GH* gene of donkey was conservative in the process of evolution and its promotor was not specific TATA box but was CATA box. The mutation of C-G in 1267 may affect the growth and development of donkeys and horses. All researches made an essential foundation for GH regulation of gene expression, evolution, polymorphism analysis in the future.

**Key words:** Donkey, growth hormone gene, cloning, sequence analysis, homology, China

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### INTRODUCTION

Growth Hormone (GH) is a single chain polypeptide hormone secreted by the anterior pituitary, it plays an important role during animal growth and development, studies have shown that *GH* gene consists of 5 exons and 4 introns in mammals and bird (De Noto *et al.*, 1981; Woychik *et al.*, 1982; Barta *et al.*, 1991; Buggiotti and Primmier, 2006). *GH* gene is the major gene controlling the level of GH secretion, regulating animal growth and development and other important physiological activities.

GH becomes one of hot spots in improving animal growth rate, growth performance and other aspects in recent years. Jorge AAM first reported the cloning of growth hormone gene sequences successfully in 1994 but study on donkey complete *GH* gene sequence and research has not been reported before. In the present study, the complete sequence of *GH* gene DNA and cDNA were cloned by PCR and RT-PCR and the sequence was analysed carefully in order to provide a theoretical basis for applied research.

### MATERIALS AND METHODS

#### Cloning of donkey *GH* gene cDNA

**Total RNA extraction:** The total RNA was extracted from the liver tissue of Guangling donkey based on the

instruction of total RNA extracted ultra light kit. The RNA was directly used for reverse transcription after it was detected through agarose gel electrophoresis.

**PCR primer design:** According to the growth hormone gene *cDNA* coding sequence of human (M13438.1), sheep (X12546.1, X15976.1), horse (U02929), pig (U19787) and cattle (AF034386) provided by NCBI, the upstream and downstream primers of donkey *GH* gene cDNA coding sequences was designed, respectively by Primer3 online after multiple sequence alignment. Forward primer was 5'-TGTGGACAGCTCACCCAAC-3'; reverse primer was 5'-GCACTGAGGAGGGTAACAG-3'.

**RT-PCR test:** RT-PCR was carried out based on RT-PCR kit (AMV) Ver. 3.0 instructions by using the total RNA extracted from liver as template, Oligo dT primer was used for reverse transcription and then the specific primers was designed for the PCR amplification, 3  $\mu$ L solution of the PCR was collected after the reaction for electrophoresis in 1% agarose. After detection, RT-PCR products were recovered and purified by rapid gel extraction kit produced by TaKaRa Biotechnology (Dalian) Co., Ltd. for further cloning.

**Cloning and sequencing of the target gene:** RT-PCR products were connected with vector pGEMR-T after

Table 1: PCR primer of donkey *GH* gene

Name	Primer sequence (5'-3')	The length of amplification (bp)	Annealing temp. (°C)
GH1	L: atgacgagcctggggacatg	267	68
	R: gtgcttacctgcagccatc		
GH2	L: caagagaggagcgggtacag	940	69
	R: atggtctcggagaagcagaa		
GH3	L: ttctgcttcccgagaccat	926	65
	R: gcactgaggagggttaaacg		
GH4	L: gggcagatcctaagcaaac	240	65
	R: ttattaggaaagatggtaggc		

recovered and then transformed into *E. coli* JM109, cultured on nutrient agar plate supplemented with AMP, IPTG and X-gal overnight. Positive colonies were picked for shaking culture and then plasmid was extracted for sequencing by Shanghai Biology Engineering Technology Ltd. (Beijing Sequencing Department) after colony PCR identification.

### Cloning of donkey *GH* gene DNA

**PCR primer design:** Four pairs of primer were designed according to the DNA sequence of Donkey (DQ009008), Chinese dwarf horse (DQ845298), Thoroughbred (DQ845297), pig (M17704), human (M13438.1). These primers were used to amplify different region of donkey *GH* gene (Table 1).

**Genomic DNA extraction:** The genomic DNA was extracted from the blood of Guangling donkey by applying convention method (Sambrook *et al.*, 1999).

**PCR amplification:** The system of PCR was 25 µL, annealing temperature is shown in Table 1. Purifying and sequencing of PCR products were done by Shanghai Biology Engineering Technology Ltd.

## RESULTS AND DISCUSSION

**DNA and cDNA sequence of donkey *GH* gene:** The length of 1928 bp donkey *GH* gene was amplified by PCR using the four pair primers (Fig. 1). The length of 706 bp donkey *GH* gene cDNA was obtained by PCR using the designed primer (shadow part in Fig. 1).

The sequence, 1928 bp DNA and 706 bp cDNA of donkey *GH* gene were compared with other animals in the NCBI database by BLAST, it was proved that it is true. The cDNA sequence is the same as 5 exons in DNA sequence, the DNA sequence included full-length of 5 exons and 4 introns. The 4 introns which lie between GT and AG and are 253, 214, 199 and 272 bp, respectively GT-AG is the signal of RNA splicing, it accords with GT-AG rule but the two initiation nucleotide of 5' terminal is GC in pig, it do not accord with the rule (Vize and Wells, 1987).

The 5 exons which are showed in italic type (Fig. 1) and are 10, 161, 117, 162 and 201 bp, respectively the length of exons in donkey *GH* gene is the same as majority mammals. But the first exon is 13 bp in cattle and sheep, the 3rd, 4th and 5th exon in human are 120, 165 and 198 bp, respectively. Length of donkey intron is equal to horse while it is largely different with other mammals. The open reading frame of donkey *GH* gene is 651 bp and initiation codon is ATG, stop codon is TAG. Two flanking regions of coding region are 5'UTR and 3'UTR. Three sites before poly (A) is AATAAAA which is the signal of synthetic polyadenylate.

**The protein of donkey *GH* gene:** According to Fig. 1, the open reading frame of donkey *GH* gene is 651 bp (23~673 bp) and encoding 216 amino acid precursor protein (Fig. 2). Precursor protein of donkey *GH* gene was analyzed online, it showed that the molecular weight was 24471.1, isoelectric point PI was 6.83, the molecular formula was C<sub>1098</sub>H<sub>1717</sub>N<sub>297</sub>O<sub>317</sub>S<sub>10</sub>. Signal peptide site was analyzed by SignalP online too, the most possible shearing site lies between 26aa and 27aa. The probability of signal peptide predicting is 1.0 and the probability of shearing site predicting is 0.917.

Membrane-spanning region of the protein was analyzed by TMPRED online. It has two strong membrane-spanning region and lie 1~22aa and 100~118aa residues, respectively. Majority signal peptide sequences are included in 1~22aa residues, the membrane-spanning region mainly lies in N-terminal through to be analyzed by TMHMM online.

**The homology of donkey and other animals in DNA, cDNA and the deduced amino acid sequence:** Using DNA man soft analysed the homology of DNA, cDNA and the deduced amino acid sequence of donkeys and other animals. Comparing the amino acid sequences of donkey with other animals, it suggests that precursor protein includes a signal peptide of 26aa and a mature peptide of 190aa, the result is the same as DeBao horse (Jiang *et al.*, 2009; Yan-Ru *et al.*, 2006).

While precursor protein of even-toed ungulate such as cattle, sheep and goat includes signal peptide of 27aa and a mature peptide of 190aa and primate includes a signal peptide of 26aa and a mature peptide of 190aa. The homology of the signal peptide sequence is distinctly lower than the mature peptide sequence.

Based on Table 2, the homology between donkey and horse is the highest according to DNA, cDNA or aa sequence. Compared donkey *GH* gene cDNA sequence with other animals, the homology successively is pig, cat, dog, cattle, sheep and goat. While the homology of aa

ATGACGAGCCTGGGGGACATGACCCAAGAGAGGAGCGGGTACAGGATGAGTGGGAGGAGTTCTAAATTAT  
 CCATCAGCAGAGGGCCCGTCAGTGGCCCATGCATAAATGTATAGAGAAAATAGGTGGGGCCGAGGGGAGA  
 GAGAAGGAGCCTGGGCATAAAAAGGGCCCGCAGGCGACCGATTCCAGGTTCCCTAGGACCCAGCTCCCCAAA  
 CTGCTCAGGGACCTGTGGACAGCTCACCCAACCTGCGATGCTGCTGAGTAAAGCACCTCTAAAATCCCCCTGGG  
 CTTGGAATCTACTGAGGGGTGGCAGGGGGCCCTGCAGATGGATGACAGCACTAACCCCTGGGCTTTGGGGCTC  
 CTGAATGTGAGCACAGACATTTGGTCAAGTTTGAATGCTCTCAGTCCCTGTGGGAAGGAGGGGGAGGAAA  
 GAAGTTTCCCTGAGGGAGGGAGAGGTCTGGCAGGAGACCCAGGCCTCTGCTCTCTGCCCGCCACCCTCCAT  
 GTGTTTCTCTAGGCCCTCGGACCTCCGTGCTCTGGCTTTCGCCCTGCTCTGCCCTGGCCTCAGGATGTGG  
 GCGCCTTCCCCGATGCCGTTGTCTAGCCTGTTTGCCAACGCTGTGCTCCGGGCCAGCACCTGCACCAACTGGC  
 TGCTGACACCTACAAAAGAGTTTGAAGCTTCCCAGGGATGGGTGCTCATGGAGGGTGGCAGGAAGGGGTGAAT  
 TCCCCACGAGGGACATAATGAGAAGAACTGACAAGTTCAGGGTTATTTTATCCAAGCGAAGATGCTCTCT  
 GGAGAGCATAAACTGAGGAGGGGTTCAAAAGAATCTCGGTGATGAGAACCGTGCACCAGCTTAGACCCCG  
 GGGGCGTCTTTCTCCCAGGAGCGCCTACATCCCCGAGGGACAGAGATACTCGATCCAGAACGCCAGGCT  
 GCCTTCTGCTTCCGAGACCATCCCCGCCCCACGGGCAAGGATGAGGCCAGCAGAGATCTGAGTGGCCCT  
 GCCAGGAAGAGGGGCTCCCTCTTTCTAAGAAGGCCGCCCTTCGCTCCCCGGGCCCTGGGGCCCTGTGTC  
 CCCTAGGTGGTGGAGCAGGGCAGAGGGTGTAGGGTGACGGGGTAAGGCCCTCGGGCAGAGCGGGCCCTAGA  
 GCGGCTGCCCTGCGCCTGCGCACCCACTGCGCCCTCCCCGAGCATGGAGCTGCTCCGCTTCTCGCTGCTG  
 CTCATCCAGTCTGGCTCGGGCCCGTGCAGTTCCTCAGCAGGGTCTTACCAACAGCCTGGTGTGGCACCTCGG  
 ACCGCGTCTATGAGAAGCTGAGGGACCTAGAGGAAGGCATCCAGGCCCTGATGCGGGGGGATGGCGTTCGG  
 GTCCCCCTCTCTGGGTCTGGAGGGCCCTCTGGCTTAGCTGAGGGGTGGGGCTTACATAGGCTGGGGAAGA  
 CAGATCTCGCCGCCCTCTTTGTAGCTGTCCAGCCCTGACCTAGGAAAAGATCTGACTCTTTCATTTCCCCTTTGA  
 ATCCCCCTGTCTTTCTAAGCCCGGAAGGGAAGGTGGAATGGAGGGGGAGGGGAGGGAGCAGCTTGC  
 AAGTCTCGCCCTCTTTCTCTTCTCTTCTGAGCTGGAAGACGGCAGCCCGGGCTGGGCAGATCC  
 TCAAGCAAACTACGACAAGTTTGACAAAACCTGCGCAGTGATGACCGACTGCTCAAGAAGTACGGGCTACTCC  
 TGCTCAAGAAGGACTGCACAAGGCTGAGACGTACCTGCGGGTATGAAGTGTGCGCGCTTCTGGAAAGCAGCT  
 GTGCCCTTAGTGTGGGCTTCTGTTACCCCTCCTCAGTGCCTCCCTTGATCTGGAGAGTGCCCTCTAGT  
 GCCTACCATCTTTCTAATAAAAGG

Fig. 1: The sequence of donkey GH gene

MAAGPRTSVLLAFALLCLPWPQDVGAFPAMPLSSLFANAVLRAQHLHQLAADTYKEFERAYIPEGQRYSIQNAQ  
 AAFCFSETIPAPTKGDEAQRSDMELLRFLLLLIQSWLGPVQFLSRVFTNSLVFGTSDRVYEKLRDLEIGIQALMRE  
 LEDGSPRAGQILKQTYDKFDNLRSDALLKNYGLLSCFKKDLHKAETYLRLVMKCRRFVSSCAF

Fig. 2: The encoded amino acid sequence of donkey GH gene

Table 2: The homologous comparison of DNA cDNA and amino acid of GH gene in donkey and different species

Species	DNA	cDNA	AA
Debao horse 67	98.8	99.5	99.5
Debao horse 46	98.3	98.9	98.6
Chinese dwarf horse	98.5	99.2	98.1
Braise horse	98.6	99.4	99.1
Thoroughbred	98.4	99.1	98.6
Mongolian horse	98.2	99.2	98.6
Cat	-	92.8	98.1
Dog	-	92.8	97.7
Pig	79.4	94.9	97.2
Cattle	68.3	91.1	89.8
Sheep	70.9	90.9	88.9
Goat	-	90.6	88.9
Human	68.3	79.6	68.4

successively is cat, dog, pig, sheep, goat and cattl. The homology of donkey between primate is the lowest. This conclusion is consistent with morphological classification which prove that conservation of donkey GH gene is high. The further analysis of the homology of equine GH gene shows that the highest homology of aa sequence is DeBao and BaiSe horse then is Thoroughbred, Mongolian horse and it is the same trend in DNA sequence of GH gene.

In the equine aa sequence, the 117st location is Phe in donkey while it is Leu in all horse. This result is caused

by the diversity of GH gene DNA sequence in 1267st site. Base of GH gene DNA sequence in 1267st site change C for G in equine which results in that aa change Leu for Phe in 117st site. This may be the reason causes the growth and development difference between donkeys and horses.

**Promoter of donkey GH gene:** Promoter is a specific DNA sequence which combines RNA polymerase and start to transcription.

Various promoters have the same or similar sequence called consensus sequence which determines the transcription activity of promoter. TATA box is a location found when researched transcription mechanism in prokaryote. In subsequent researchs, the location was discovered in majority advanced animals (Sun *et al.*, 1990).

**CONCLUSION**

The 5'UTR promoter sequence of donkey GH gene is predict by NNPP soft. Promoter sequence begin at the 149st and end at 199st of DNA sequence and it's length is 50 bp (agcctgggcataaaaaggccccgagcaccgattccagg

ttcctagga). The site of predicted transcription beginning of the promoter is at 189st and it's accuracy is 1.00. In predicted sequence, TATA frame is CATAAA (CATA box) rather than TATAAA (TATA box) in majority mammals.

This suggests that transcription mechanism in donkey is different with other animals, the combination mechanism of RNA polymerase to template is different too. If want to express a certain gene of donkey in other mammals' cells or individuals, the transcription initiation factor of donor must be removed.

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