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(54) **SINGLE NUCLEOTIDE POLYMORPHISMS ASSOCIATED WITH BULL FERTILITY**

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CI2Q 1/6883 (2018.01)

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A01K 67/027 (2006.01)

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(52) **U.S. Cl.**
CPC *CI2Q 1/6883* (2013.01); *A01K 67/02* (2013.01); *A01K 67/027* (2013.01); *A61D 19/04* (2013.01); *A61K 35/54* (2013.01); *CI2Q 1/68* (2013.01); *CI2Q 1/6888* (2013.01); *A01K 2227/101* (2013.01); *A01K 2267/02* (2013.01); *CI2Q 2600/124* (2013.01); *CI2Q 2600/156* (2013.01); *CI2Q 2600/16* (2013.01)

(58) **Field of Classification Search**

None
See application file for complete search history.

(56) **References Cited**

PUBLICATIONS

Barendse et al. Mammalian Genome 8, 21-28 (1997) (Year: 1997).*

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(57) **ABSTRACT**

Single nucleotide polymorphic sites of the bovine MAP1B, PPP1R11, and DDX4 genes are associated with improved bull fertility as measured by e.g. sire conception rates. Nucleic acid molecules, arrays, kits, methods of genotyping and marker-assisted bovine breeding methods based on these SNPs are disclosed.

9 Claims, 6 Drawing Sheets

Specification includes a Sequence Listing.

FIG. 1

49681 tccagcctgt tctgtggatg tttgaacttg agaagtgggg tcttgcacca gaggaaacac
 49741 tgcttttgcg ctgttagagg atgggctcca tccgaatcat acccagtttt ttcccttgc
 49801 acttcctcat cttcccgatgg tttcatgtcg agtcagaatg taaggactgt ttagcttttgc
 49861 tgaggggcaa aaatgtttt ttgaactgaa caaggttaagg tttgaaccca ttcccttgc
 49921 ttcttggat acttcattt tcacitggag cacaaggcag gttggggaaag caggaggggg
 49981 gaagatgtt aatgtggata gagacagagg aaaaggcagg tgggggggtt gaaactgaacc
 50041 ccacttcctg cagccgttc ccagcgttgc ttgaaaagac tctgaaaaggaa gaataacgtc
 50101 tttaaatcaag agcaatagta ttatgttt tactataagt aatactttt tttgagccca
 50161 tatattttt accgggttag aatagatgtt agtttttcca gcagccatgtt ctatgttca
 50221 ggagttggat gtggggctggc aatagactgg ctgattacac ttttttagaaa taaacccctt
 50281 ttttggcgat ctcttctgtt gagaatgggtt cataaaaggcc tctgtggctg gttgttco
 50341 ggttgttgc ttatatacg ttatgttcc ttatgttgc ttatgttgc ttatgttgc
 50401 attataactc ttttgcagag ggcgttagggc ttttgcagttt ttttgcaggcaac atcacaaagg
 50461 ttttggaaagaa atctttaat taatgttgc ttttgcgttgc qttttttttt gttttttt
 50521 ctgaaatgtt tttatcttgc agntttttt ttttgcgttgc qttttttt
 50581 gcccggatgc cggccggatgc acacactgtt gttttttttt gttttttt
 50641 ctttgcgttgc ttttgcgttgc ctttgcgttgc ttttgcgttgc ttttgcgttgc
 50701 gttttttttt ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 50761 ctttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 50821 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 50881 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 50941 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51001 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51061 agatcttccaa accccaggcc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51121 agaaagggtt gggccaaata cggccggatgtt ttttgcgttgc ttttgcgttgc
 51181 gatctggccaa ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51241 aaaaaatctttt ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51301 acttccttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51361 aaaaaaaatgtt ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51421 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51481 acgtttttttt ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51541 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51601 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51661 [REDACTED] gggggatgtt ttttgcgttgc ttttgcgttgc ttttgcgttgc
 SNPs
 51721 acccttcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 SNP4
 51781 gggctccaa gaacaggaaa agaggatgtt ggggggggg gttttttttt ttttgcgttgc
 51841 tagtccctgtt gttttttttt ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51901 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 51961 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52021 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52081 gacacccat ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52141 aggaggaggcc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52201 agcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52261 aacttccttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52321 aacccat ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52381 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52441 ataaccattt ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52501 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52561 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52621 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52681 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52741 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52801 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52861 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52921 gatgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 52981 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 53041 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 53101 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 53161 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 53221 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 53281 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc
 53341 ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc ttttgcgttgc

85081 agtgctctgg ttaatcaacc agtgaggtoa gctgaccact gggtacccag tacagattga
 85141 gaaaagagca tccaagactc tacttattcc attctgacca cactgcctca ctaatacaga
 85201 ctcaacatct tgtrtttaggt cgaatattcag cttggcaaga gcaagcatcc ctgacttcat
 85261 gttcttagac ttatcaagg tctggcagtg atctgggtt ttgatgaaat tgggtttctg
 85321 atgatccagc ttcatatgt tgatcaaaa caatcacagg gattgaatt catatatitt
 85381 atttgcttta catagttact tgaaggattt agatcacagt tcacaaacat gtaaagcaaa
 85441 aaataagcaa cactttcttg atttattatg gaaaaattca gtathtagta ctttaggaag
 85501 tactagttac aggtacaagt ttttactttt aggcaacgtg aagcagcaat ttcaagactc
 85561 atatcagatt tccctctttt atttgcacat agaaaacaaa ctgaatttgt tcatgttag
 85621 aattttgtt qagccaccag ataataaaatg ttgatctaaa ggacttaaca gtgaccatgc
 85681 acttaggaga aaacatgaaa tcaattcaaa cagataaaaaa cccaaactgaa atttgcgtcc
 85741 aaactcatga aacttacat atagcccaca caattgtatt tatactttt tttttttttt
 85801 gtcatttaaa gataattttg agggaaatgt agttaattt gatttacattt gaggatgtt
 85861 tcccaacaga tttttttaaa agacaaaaact gcacattaa ttaattttaa aacaaacctg
 85921 ggtcaacttc cagtggttct atnagttrgg gtttctatga gcaatgttca tttgggtgtca
 85981 acgggagtga ttcaagggtgc aagtggaaac tgcaggcatt taaaaatatt agatgtatc
 86041 taactocaaa accctctqcta caagtcaaaa ttctttttt gatttacaca tgaatatgtt
 86101 taggactttt agtttagttt cattataatg gctggtaatf ctattcatga aatgatccag
 86161 tttatccaaa taccagttt gctgattttt actaccctt gcccctccaaa ataaaaataaa
 86221 accaggatcat agtgcattt qactgtggg tggcagtttatacatccc atggagaaaaq
 86281 gcaagagaat taattttagg ggtatgttgc agtattttaa ggggttac acgagtggc
 86341 tcaaaaccaga taacacattag cattttgggg gatgtttttaaaaatgtat gtttttttt
 86401 cagtgaaagcc ttaccatagc cattqaagcc agggcatctt tattttttttt gtttttttt
 86461 tctaatcatg tggccaggag gaagaaccac tgccttacaa tgcattttttt gtttttttt
 86521 caaccccttgc attagaacaa atcagaaagc caatttacat aacaaggtagt ccagaaactg
 86581 agattaatctt gaaaccttcat tttttttttt gttttttttt tttttttttt tttttttttt
 86641 acataccctac cccttaccc agtcccgat gttttttttt gttttttttt gttttttttt
 86701 totgaacccctt ctgtttgtca aagttttttt gttttttttt totgtttttt totgtttttt
 86761 attttaggaga ctgtttttt gttttttttt gttttttttt gttttttttt gttttttttt
 86821 aggtgttgc ggtttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 86881 gtgagggat aatatttgc tttttttttt gttttttttt gttttttttt gttttttttt
 86941 tactaagatg gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87001 ctattttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87061 gctgtttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 SNES

87121 agggaaacgc cccatttttcc attttttttt gttttttttt gttttttttt gttttttttt
 87181 agactgcccag tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87241 cccagagccg cttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87301 atttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87361 aagacgttca aaccccttgc tttttttttt gttttttttt gttttttttt gttttttttt
 87421 atcccccaacc atagcaatag tttttttttt gttttttttt gttttttttt gttttttttt
 87481 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87541 gatgtttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87601 cgggtttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87661 gaggcaccac ctggatccctt gttttttttt gttttttttt gttttttttt gttttttttt
 87721 gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87781 ggaaggccac tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87841 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87901 aagatgtttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 87961 aaaatcatgt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88021 cactcccaac accaaacat tttttttttt gttttttttt gttttttttt gttttttttt
 88081 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88141 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88201 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88261 acacacacac aaagacataa atctttttt gttttttttt gttttttttt gttttttttt
 88321 agaaggttttt cggttgcattt aataataaa gttttttttt gttttttttt gttttttttt
 88381 aatgtttttt acaatgtttt tttttttttt gttttttttt gttttttttt gttttttttt
 88441 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88501 gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88561 gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88621 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88681 ccacccatc tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88741 gatcccaactt catgtttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88801 gcaaggccac aacccatcatgg tttttttttt gttttttttt gttttttttt gttttttttt
 88861 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt
 88921 tttttttttt gttttttttt gttttttttt gttttttttt gttttttttt gttttttttt

FIG. 4

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34261 ccagagtgtt ctatgaacta caaaaagggtgg gactgtgtga atcttggtca ttcacagat
34321 agataaaactg ggatgtctt gtctctgagt aggaacattg gagatatggg ggaagggaga
34381 agttgttagat taattacat acttgcataat cctgcattctg cttagggtga gatggtataa
34441 aaattatagt gctcagttct ggattatcta tagcagaca tggtaaaaata gcaacaatat
34501 ccacgaaaaa ccacagtggaa ctataaaaat tgctacaagt gtgcaaataat atttatgata
34561 gaacttttagt gtttggagct gcactagata catcatagtt ttgcgtcgaat cttggagata
34621 tggttttccc tttgcattata gatgttggc tcattgaata gatcattga tagcaggcc
34681 tccatgtgaa gctgagactt gctgttgatt tcactatagc tggatgatgaa ttgtgagggg
34741 cggtgggttag gaatttgggt gtgaatcagt tcagtcgctc aattygtgtt gacttttgtc
34801 gaccccatga attgcagcat gcccaggctc cctgtccatc accgactctt ggagtccatt
34861 caaaactcaag tccatcgagt cggtgatgcc atccaaccat ctcatcttctt gttgtccct
34921 tctccctctg ccccccaatcc ctcccaagcat cacagtctt tccaatgagt cagctcttgc
34981 catgaggttgg cccaaagttact ggagtttcag ctttagcattc attcccttcca aagaacaccc
35041 aggactgtatc tccctttagaa tggacttagtt ggatctctt ggacttccaa gggacttccaa
35101 gagtccttttca caacaccaca gttcaaaaacg atcaattctt cggcgtctcg ctttotttcac
35161 agtccaaactc tcacatccat atatgaccac tggaaaaacc atagcccttgc ctatatggac
35221 ctgttggc aaaaatagt ctgtgtttt cagatgtatc totaggttg tccataaatcc
35281 ctttccaaagg agtaagcgctc ttttaatttc acagctgcag tccatctgtc cagtgattt
35341 ggagcccgaga aaaataaaatg ctgcactgtt ttcactgtt tcccatcttca tttccatgt
35401 agtgatgggaa ccagatgcca tgatcttgc tttctgaatg ttgagcttta agccaaacttt
35461 ttcaactctcc tctttcaactt tcataaagag gctttttgtt tccctttcac tttctgcata
35521 agctgagttt ttaatggcaa ttgttaggggg ccctgcaatg atggggccgaa catttagtta
35581 agaaaatagac tgcgttttta acatgtgtt cttcccccct ttaacaagga gttttgcacac
35641 taatgttctt aaaaatcgatc tctttttgggtt ttctgcagaa cagtggtatctt ctttcttact
35701 attcagttttt ctttaatccat ttttaattca tattaatgtc cagcaatgaa aagccatgttgc
35761 cagtcgttttgc tgcgtgtatcc tgacttattg actagtgttag ttgttgcgtat aggggttcc
35821 ttgttgcgtat tttttttttatgtatccat tggatgttgc aaaaatgtgtt gttgcaccc
35881 ggtcaactcag taaaatcaga tttgtgtgtt aatagaacat tatctaaatgtt tatgttgc
35941 tatcatgtca tacaggaaca ggttgggtca tatgtcataa atgtatacag ctcaatgtatt

FIG. 5

FIG. 6

1**SINGLE NUCLEOTIDE POLYMORPHISMS
ASSOCIATED WITH BULL FERTILITY****CROSS-REFERENCE TO RELATED
APPLICATION**

This is a divisional application of U.S. application Ser. No. 14/754,729 filed on Jun. 30, 2015, claiming priority to U.S. patent application Ser. No. 13/798,181, filed on Mar. 13, 2013, the entire disclosure of which is incorporated herein by reference.

STATEMENT OF GOVERNMENT INTEREST

This invention was made with government support under 15 12-CRHF-0-6055 awarded by the USDA/NIFA. The government has certain rights in the invention.

FIELD OF THE INVENTION

The present invention relates to compositions and methods for testing and selecting cattle using molecular genetic methods by assaying for the presence of at least one genetic marker indicative of increased bull fertility. Specifically, genetic variations in the MAP1B and PPP1R11 genes are tested and used for selecting cattle animals with improved blastocyst or fertilization rates, or both.

BACKGROUND OF THE INVENTION

The dairy cattle genome has been significantly restructured over the past 30 years due to intensive breeding effort selecting for production traits, including high quality milk and high and sustained productivity. However, while those efforts led to dramatic improvement of productivity, there has been significant reproductive deterioration in high-producing dairy cows, which in turn has caused substantial economic loss in the dairy cattle industry (Lucy, 2007, Fertility in high-producing dairy cows: reasons for decline and corrective strategies for sustainable improvement. *Society of Reproduction and Fertility Supplement*. 64 237-254). Key factors contributing to decreasing fertility of dairy cow are low fertilization rates and decreased embryonic survival.

Fertility is a complex trait that comprises developmental stages such as combining sperm and egg to form a zygote, compaction of embryo cells to form a morula, establishment of the blastocyst, attachment of the embryo to the uterus, and fetal development (Amann and DeJarnette, 2012). This complexity makes accurate prediction of successful pregnancy difficult, as aberrant development of sperm, oocyte, embryo, or fetus all would lead to conception failure. Conception rate in dairy cattle is about 40%, and only 50% of the fertilized eggs produce viable embryos (Santos et al., 2004). The decline in reproductive performance in cattle over the past few decades (Dobson et al., 2007) has been ascribed primarily to fertilization failure and early embryonic loss (Santos et al., 2004).

Previous studies have shown that genetic makeup of an individual plays crucial roles in embryonic development and reproductive success (Weigel, 2006; Shook, 2006). Although a male and female parent each contributes half of its genetic material to the new zygote and both are necessary for embryo development, it is not obvious whether or not this contribution is equally important to pregnancy success. For example, it is well established that the paternal genome supports growth of extra-embryonic tissues while the maternal genome fosters development of the embryo proper

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(Barton et al., 1984). After fertilization, the development of an embryo is controlled by maternal genomic information that is accumulated during oogenesis (Telford et al., 1990). It is only at the 8-cell stage in the bovine embryo that the 10 embryonic genome activates and the embryo switches to transcribing its own RNA (Memili and First, 2000).

Despite that most breeding schemes in cattle are focused on the selection of elite bulls using progeny testing or genomic selection, and that some semen traits (e.g., sperm motility and percentage of abnormal sperm) show moderate to high heritabilities (Druet et al., 2009), most fertility studies in cattle have focused on the maternal contribution, and the paternal contribution to reproductive performance has not been thoroughly investigated, and only a few studies have been reported in the literature (Feugang et al., 2009; Khatib et al., 2010; Peñagaricano et al., 2012). Therefore, characterization of bull fertility markers is both feasible and highly desirable, and the deployment of these markers in 20 cattle breeding would lead to improved reproductive performance in cattle.

A recent comparative genomics study has characterized many genes involved in the control of spermatogenesis that were highly conserved from fly to human (Bonilla and Xu, 25 2008). Some of these genes were reported to be crucial for human fertility. However, it is not known whether or not these spermatogenesis genes play important roles in the fertility of bulls.

SUMMARY OF THE INVENTION

The present inventor carried out an association analysis between highly conserved spermatogenesis genes and sire conception rate (SCR) as a measure of bull fertility, with the 35 objective that significant polymorphisms associated with bull fertility can be used as genetic markers in breeding programs aimed at improving reproductive performance in cattle.

Specifically, an association analysis is performed between 40 highly conserved spermatogenesis genes and SCR in US Holstein populations as a measure of bull fertility. Sequence analysis revealed 24 single nucleotide polymorphisms (SNPs) in 9 genes in the bull population using the pooled DNA sequencing approach. These 9 genes were selected for 45 their high level of sequence conservation between flies and humans. Overall, the 24 SNPs were tested for association with SCR in a population of 1,988 bulls. Three SNPs located in the MAP1B gene, one SNP in the PPP1R11 gene and one SNP in the DDX4 gene showed significant associations with SCR. Nucleotide probes based upon these SNPs are found to 50 be useful for genetic testing of bull animals for improved fertilization rate.

Accordingly, in one embodiment, the present invention provides an isolated oligo- or poly-nucleotide molecule consisting of

- 1) a nucleotide of Position 3066 of SEQ ID NO:1 and at least 12, but not more than 200 contiguous nucleotides of FIG. 1 adjacent to position 3066;
- 2) Position 3323 of SEQ ID NO:1 and at least 12, but not more than 200 contiguous nucleotides of FIG. 1 adjacent to position 3323;
- 3) Position 87071 of SEQ ID NO:2 and at least 12, but not more than 200 contiguous nucleotides of FIG. 3 adjacent to position 87071,
- 4) Position 112 of SEQ ID NO:3 and at least 12, but not more than 200 contiguous nucleotides of SEQ ID NO:3 adjacent to position 112,

- (5) Position 61646 of SEQ ID NO:4 and at least 12, but not more than 200 contiguous nucleotides of FIG. 6 adjacent to position 61646, and
- 6) Position 34239 of SEQ ID NO:5 and at least 12, but not more than 200 contiguous nucleotides of FIG. 5 adjacent to position 61646.

In one embodiment, the nucleotide molecule of the present invention comprises at least about 15 contiguous nucleotides adjacent to its respective position (hereinafter the "SNP position") of the respective figure. In one embodiment, the nucleic acid molecule of the present invention comprises at least about 20 contiguous nucleotides adjacent to the respective SNP position. In one embodiment, the oligonucleotide molecule of the present invention consists of not more than about 100 nucleotides. In one embodiment, the oligonucleotide molecule of the present invention consists of not more than about 50 nucleotides. In one embodiment, the SNP position of the nucleotide molecule of the present invention near or at the center of the molecule; alternatively, the SNP position is at the 3'-end of the oligonucleotide molecule.

Also provided herein is an array of nucleic acid molecules, comprising the isolated oligonucleotide molecule of the present invention, supported on a substrate. The substrate may be any suitable medium, known and readily available to one of ordinary skills in the art, and the array may be addressable.

The present invention further provides a kit comprising an isolated oligonucleotide molecule of the present invention, and a suitable container.

In another embodiment, the present invention provides a method for detecting single nucleotide polymorphism (SNP) in a gene listed in Table 1 below in a bovine cell, the method comprising optionally isolating an DNA from the bovine cell, determining the identity of a nucleotide on the gene of the cell at a SNP position identified in Table 1 below, and comparing the identity to the preferred nucleotide identity at a corresponding position in Table 1.

In one embodiment, the bovine cell may be an adult cell, an embryo cell, a sperm, an egg, a fertilized egg, or a zygote. The identity of the nucleotide may be determined by many methods known and readily available to those ordinarily skilled in the art, such as but not limited to sequencing a nucleic acid molecule comprising a suitable portion of the gene of the cell comprising a respective SNP position, or by hybridizing a suitable probe to a nucleic acid preparation from the cell, which probe may be suitably labeled e.g. fluorescently or radioactively.

The nucleic acid molecule may be isolated from the cell via a large variety of methods, known and readily available to an ordinarily skilled artisan, such as amplification by the polymerase chain reaction (PCR) of genomic DNA of the cell, or when appropriate, by RT-PCR of the mRNA of the cell.

In preferred embodiment, both copies of the gene in a diploid genome are genotyped according to the method of the present invention.

The identity of the nucleotide may be determined based on genotypes of the parent of the cell, genotypes of the daughter of the cell, or both, through genetic analysis methods well-known to those skilled in the art.

A method is further provided for determining whether an individual bovine animal is suitable as a gamete donor for natural mating, artificial insemination or in vitro fertilization, the method comprising detecting the SNP according to the above method of the present invention, and excluding as

gamete donor an individual which does not have the preferred allele identity at the respective SNP position as described in Table 1.

In one embodiment, the individual is excluded as a gamete donor if the individual, whose genotype is not homozygous of the preferred allele with regard to the respective SNP position.

The present invention additionally provides a method of selecting a bovine embryo for planting in a uterus, the 10 method comprising genotyping the embryo according to the present invention, while preserving the viability of the embryo, and excluding from planting an embryo which does not have the preferred allele identity at the respective SNP position as described in Table 1.

15 In another embodiment, the present invention further provides a method for selectively breeding cattle using a multiple ovulation and embryo transfer procedure (MOET), the method comprising superovulating a female animal, collecting eggs from said superovulated female, in vitro 20 fertilizing said eggs from a male animal which has at least one preferred allele identity at the respective SNP position as described in Table 1, implanting said fertilized eggs into other females allowing for an embryo to develop.

In another embodiment, the method for selectively breeding 25 cattle using MOET may further comprise a step of genotyping the male animal according to the method of the present invention.

In another embodiment, the method for selectively breeding cattle using MOET may further comprise a step of 30 genotyping the developing embryo, and allowing pregnancy to proceed only if the genotype of the embryo comprises at least one preferred allele identity at the respective SNP position as described in Table 1.

DESCRIPTION OF THE DRAWINGS

FIG. 1 depicts a partial genomic sequence of MAP1B, showing the locations of SNPs 1, 2 and 3 on the MAP1B gene (SEQ ID NO: 1) and the locations of the primers (1F and 1R, corresponding to positions 2859-2876, and positions 3346-3362, respectively) used to amplify the region comprising the SNP sites. The numbering of the sequences is according to that of GenBank Accession No. (Gene ID: 514739, updated on 12 Jul. 2012), incorporated herein by reference in its entirety.

FIG. 2 is the partial genomic sequence of the MAP1B gene (SEQ ID NO: 6) showing the locations of SNPs 4 and 6 on the MAP1B gene and the locations of the primers 2F and 2R (corresponding to positions 51447-51466, and positions 51931-51948, respectively) used to amplify the region comprising the SNP sites. The numbering of the sequence is according to that of GenBank Accession No. (Gene ID: 514739, updated on 12 Jul. 2012), incorporated herein by reference in its entirety.

55 FIG. 3 is the partial genomic sequence of the MAP1B gene (SEQ ID NO: 2) showing the location of SNP 5 on the MAP1B gene and the locations of the primers 3F and 3R (corresponding to positions 86634-86651, and positions 87260-87278, respectively) used to amplify the region comprising the SNP sites. The numbering of the sequence is per 60 that of GenBank Accession No. (Gene ID: 514739), incorporated herein by reference in its entirety.

65 FIG. 4. is the partial genomic sequence of the PPP1R11 gene (SEQ ID NO: 3) showing the locations of SNPs 1-4 on the PPP1R11 gene and the locations of the primers 1F and 1R (corresponding to positions -102 to -85, and positions 470-489, respectively) used to amplify the region compris-

ing the SNP sites. The numbering of the sequences is according to GenBank Accession No. (Gene ID: 504846), incorporated herein by reference in its entirety.

FIG. 5 is the partial genomic sequence of the DDX4 gene (SEQ ID NO: 5), showing the location of SNP 2 on the DDX4 gene and the locations of the primers 1F and 1R (corresponding to positions 34014-34031, and positions 34398-34417, respectively) used to amplify the region comprising the SNP site. The numbering of the sequences is according to that of GenBank Accession No. (Gene ID: 493725), incorporated herein by reference in its entirety.

FIG. 6 is the partial genomic sequence of the DDX4 (SEQ ID NO: 4) showing the location of SNP 1 on the DDX4 gene and the locations of the primers 2F and 2R (corresponding to positions 61531-61549, and positions 61867-61884, respectively) used to amplify the region comprising the SNP site. The number of the sequences is per that of GenBank Accession No. (Gene ID: 493725), incorporated herein by reference in its entirety.

DETAILED DESCRIPTION OF THE INVENTION

The present inventor has found that the spermatogenesis genes MAP1B, PPP1R11 and DDX4 showed significant associations with SCR. Table 1 below summarizes the SNPs demonstrated to be significantly associated with sire conception rates according to the present invention.

TABLE 1

Genetic markers significantly associated with sire conception rate

SNP ID	Location/Position	Nucleotide Identity of Preferred Allele	Nucleotide Identity of Polymorph in GenBank
SNP1.MAP1B	Position 3066 of FIG. 1	A	A
SNP3.MAP1B	Position 3323 of FIG. 1	T	T
SNP5.MAP1B	Position 87071 of FIG. 3	C	C
SNP1.PPP1R11	Position 112 of FIG. 4	G	T
SNP1.DDX4	Position 61646 of FIG. 6	A	A
SNP2.DDX4	Position 34239 of FIG. 5	G	G

Three SNPs in MAP1B, in low to moderate linkage disequilibrium (LD), were significantly associated with SCR. After correction for multiple testing, only one SNP in intron 5, SNP 5, showed the most significant association with SCR.

The MAP1B gene belongs to the microtubule-associated protein family and is known to affect neuronal development such as axon growth (Tymanskyj et al., 2012), development of dendritic spine and synaptic maturation (Tortosa et al., 2011), and regulation of the interaction between microtubules and actin microfilaments for axonal development (Montenegro-Venegas et al., 2010). Recent reports on the expression of MAP1B in the male reproductive tract in both rat and human (Queiróz et al., 2006) and in testis of fruit fly and mouse (Bonilla and Xu, 2008) suggest important functions of this gene in the regulation of male fertility. The finding by the present inventor, disclosed herein, that the MAP1B gene is associated with SCR, supports the conclusion that MAP1B plays a role in male fertility across a wide range of species.

The present inventor also found that a SNP in the 5'UTR of PPP1R11 was associated with SCR in the bull population examined in this study. This is the first report of association between male fertility in cattle and PPP1R11, which is

consistent with previous reports on the roles of this gene in spermatogenesis in mouse and human. For example, the different isoforms of PPP1R11 (also known as TCTEX5) were found to be expressed in most mouse tissues with high expression in testis, epididymis, and in the head and tail regions of spermatozoa (Han et al., 2007). In a subsequent study, it was shown that mutations in the long transcript of PPP1R11 were associated with normal sperm function (Han et al., 2008). The authors concluded that PPP1R11 plays important roles in sperm motility and spermatogenesis. A recent study reported that an isoform of protein phosphatase 1 (PP1 γ 2), which has an essential role in spermatogenesis, forms a complex with PPP1R11 in the testis (Cheng et al., 2009). Given that PP1 γ 2 is regulated by PPP1R11, these results further support the idea that PPP1R11 has important functions in spermatogenesis.

The spermatogenesis genes investigated in this study were selected from a pool of genes whose expression is highly conserved in testis of both fruit fly and mouse (Bonilla and Xu, 2008). The protein sequence identities between cattle and human and between cattle and fly are 91% and 32%, respectively for MAP1B and 99% and 47%, respectively for PPP1R11. As such, the association of these genes with bull fertility testifies to the usefulness of the comparative genomics approach in selecting candidate male fertility genes.

To further explore involvement of male fertility genes identified in this study in female fertility, we tested the association of the SNPs in MAP1B, with fertilization and embryo survival rates using data from the IVF system. MAP1B genotypes of the cows, from which oocytes were extracted and used for fertilization and embryo culture, were significantly associated with differential fertilization rate and embryo survival rate. Recently, the expression of MAP1B was found to be downregulated in follicular cystic follicles compared to normal follicles, suggesting that alteration in MAP1B expression may be involved in reproduction failure in cattle (Choe et al., 2010). The instant disclosure again demonstrates the significance of both parental genomes to embryonic development and fertility.

Accordingly, the present invention provides nucleic acid-based genetic markers for identifying bovine animals, especially bulls, with superior fertility, specifically, sire conception rate as a measure of male fertility. In general, for use as markers, isolated oligonucleotide or polynucleotide molecules, or isolated nucleic acid fragments, preferably DNA fragments, as used. Such markers will be of at least 10 nucleotides (nt), preferably at least 11, 12, or 15 nt, usually at least 20 nt, often at least 50 nt. Such small DNA fragments are useful as primers for the polymerase chain reaction (PCR), and probes for hybridization screening, etc.

In one embodiment, the isolated nucleic acid molecule comprises not more than 1,500 nt, or not more than 1000 nt, or not more than 900 nt, or not more than 800 nt, or not more than 700 nt, or not more than 600 nt, or not more than 500 nt, or not more than 400 nt, or not more than 300 nt, or not more than 200 nt., or not more than 150 nt., or not more than 100 nt., or not more than 75 nt.

In the context of the present invention, the term "isolated" refers to a nucleic acid molecule purified to some degree from endogenous materials with which the nucleic acid molecule may naturally occur or exist. At the least, the term "isolated" refers to a nucleic acid molecule separated from chromatin or other protein or components of the genomic DNA. Preferably, the isolated oligonucleic acid molecule or polynucleic acid molecule of the present invention comprises a fragment that is shorter than that which is naturally occurring.

In the context of the present invention, the provided sequences also encompass the complementary sequence corresponding to any of the provided polymorphisms. Where appropriate, and in order to provide an unambiguous identification of the specific site of a polymorphism, the numbering of the original nucleic sequences in the GenBank may be used; alternatively, the numbering may simply refer to the specific sequence in the Sequence Listing accompanying this disclosure.

The term primer refers to a single-stranded oligonucleotide capable of acting as a point of initiation of template-directed DNA synthesis under appropriate conditions (i.e., in the presence of four different nucleoside triphosphates and an agent for polymerization, such as, DNA or RNA polymerase or reverse transcriptase) in an appropriate buffer and at a suitable temperature. The appropriate length of a primer depends on the intended use of the primer but typically ranges from 15 to 30 nucleotides. Short primer molecules generally require cooler temperatures to form sufficiently stable hybrid complexes with the template. A primer need not reflect the exact sequence of the template but must be sufficiently complementary to hybridize with a template. The term primer site, or priming site, refers to the area of the target DNA to which a primer hybridizes. The term primer pair means a set of primers including a 5' upstream primer that hybridizes with the 5' end of the DNA sequence to be amplified and a 3', downstream primer that hybridizes with the complement of the 3' end of the sequence to be amplified. One of these two primers is often referred to as the "forward primer," while the other the "reverse primer."

The term "probe" or "hybridization probe" denotes a defined nucleic acid segment (or nucleotide analog segment) which can be used to identify by hybridization a specific polynucleotide sequence present in samples, said nucleic acid segment comprising a nucleotide sequence complementary of the specific polynucleotide sequence to be identified. "Probes" or "hybridization probes" are nucleic acids capable of binding in a base-specific manner to a complementary strand of nucleic acid.

An objective of the present invention is to determine which embodiment of the polymorphisms a specific sample of DNA has. For example, it is desirable to determine whether the nucleotide at a particular position is A or G. An oligonucleotide probe can be used for such purpose. Preferably, the oligonucleotide probe will have a detectable label, and contains an A at the corresponding position. Experimental conditions can be chosen such that if the sample DNA contains an A, they hybridization signal can be detected because the probe hybridizes to the corresponding complementary DNA strand in the sample, while if the sample DNA contains a G, no hybridization signal is detected.

Similarly, PCR primers and conditions can be devised, whereby the oligonucleotide is used as one of the PCR primers, for analyzing nucleic acids for the presence of a specific sequence. These may be direct amplification of the genomic DNA, or RT-PCR amplification of the mRNA transcript of the gene of interest. The use of the polymerase chain reaction is described in Saiki et al. (1985) *Science* 230:1350-1354. Amplification may be used to determine whether a polymorphism is present, by using a primer that is specific for the polymorphism. Alternatively, various methods are known in the art that utilize oligonucleotide ligation as a means of detecting polymorphisms, for examples see Riley et al (1990) *Nucleic Acids Res.* 18:2887-2890; and Delahunty et al (1996) *Am. J. Hum. Genet.* 58:1239-1246. The detection method may also be based on

direct DNA sequencing, or hybridization, or a combination thereof. Where large amounts of DNA are available, genomic DNA is used directly. Alternatively, the region of interest is cloned into a suitable vector and grown in sufficient quantity for analysis. The nucleic acid may be amplified by PCR, to provide sufficient amounts for analysis.

Hybridization may be performed in solution, or such hybridization may be performed when either the oligonucleotide probe or the target polynucleotide is covalently or noncovalently affixed to a solid support. Attachment may be mediated, for example, by antibody-antigen interactions, poly-L-Lys, streptavidin or avidin-biotin, salt bridges, hydrophobic interactions, chemical linkages, UV cross-linking baking, etc. Oligonucleotides may be synthesized directly on the solid support or attached to the solid support subsequent to synthesis. Solid-supports suitable for use in detection methods of the invention include substrates made of silicon, glass, plastic, paper and the like, which may be formed, for example, into wells (as in 96-well plates), slides, sheets, membranes, fibers, chips, dishes, and beads. The solid support may be treated, coated or derivatized to facilitate the immobilization of the allele-specific oligonucleotide or target nucleic acid. For screening purposes, hybridization probes of the polymorphic sequences may be used where both forms are present, either in separate reactions, spatially separated on a solid phase matrix, or labeled such that they can be distinguished from each other.

Hybridization may also be performed with nucleic acid arrays and subarrays such as described in WO 95/11995. The arrays would contain a battery of allele-specific oligonucleotides representing each of the polymorphic sites. One or both polymorphic forms may be present in the array. Usually such an array will include at least 2 different polymorphic sequences, i.e. polymorphisms located at unique positions within the locus, and may include all of the provided polymorphisms. Arrays of interest may further comprise sequences, including polymorphisms, of other genetic sequences, particularly other sequences of interest. The oligonucleotide sequence on the array will usually be at least about 12 nt in length, may be the length of the provided polymorphic sequences, or may extend into the flanking regions to generate fragments of 100 to 200 nt in length. For examples of arrays, see Ramsay (1998) *Nat. Biotech.* 16:4044; Hacia et al. (1996) *Nature Genetics* 14:441-447; Lockhart et al. (1996) *Nature Biotechnol.* 14:1675-1680; and De Risi et al. (1996) *Nature Genetics* 14:457-460.

The identity of polymorphisms may also be determined using a mismatch detection technique, including but not limited to the RNase protection method using riboprobes (Winter et al., *Proc. Natl. Acad. Sci. USA* 82:7575, 1985; Meyers et al., *Science* 230:1242, 1985) and proteins which recognize nucleotide mismatches, such as the *E. coli* mutS protein (Modrich, *P. Ann. Rev. Genet.* 25:229-253, 1991). Alternatively, variant alleles can be identified by single strand conformation polymorphism (SSCP) analysis (Orita et al., *Genomics* 5:874-879, 1989; Humphries et al., in *Molecular Diagnosis of Genetic Diseases*, R. Elles, ed., pp. 321-340, 1996) or denaturing gradient gel electrophoresis (DGGE) (Wartell et al., *Nucl. Acids Res.* 18:2699-2706, 1990; Sheffield et al., *Proc. Natl. Acad. Sci. USA* 86:232-236, 1989).

A polymerase-mediated primer extension method may also be used to identify the polymorphism(s). Several such methods have been described in the patent and scientific literature and include the "Genetic Bit Analysis" method (WO92/15712) and the ligase/polymerase mediated genetic

bit analysis (U.S. Pat. No. 5,679,524). Related methods are disclosed in WO91/02087, WO90/09455, WO95/17676, U.S. Pat. Nos. 5,302,509, and 5,945,283. Extended primers containing a polymorphism may be detected by mass spectrometry as described in U.S. Pat. No. 5,605,798. Another primer extension method is allele-specific PCR (Ruao et al., Nucl. Acids Res. 17:8392, 1989; Ruao et al., Nucl. Acids Res. 19, 6877-6882, 1991; WO 93/22456; Turki et al., J. Clin. Invest. 95:1635-1641, 1995). In addition, multiple polymorphic sites may be investigated by simultaneously amplifying multiple regions of the nucleic acid using sets of allele-specific primers as described in Wallace et al. (WO 89/10414).

A detectable label may be included in an amplification reaction. Suitable labels include fluorochromes, e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein (JOE), 6-carboxy-X-rhodamine (ROX), 6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX), 5-carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA), radioactive labels, e.g. ³²P, ³⁵S, ³H; etc. The label may be a two stage system, where the amplified DNA is conjugated to biotin, haptens, etc. having a high affinity binding partner, e.g. avidin, specific antibodies, etc., where the binding partner is conjugated to a detectable label. The label may be conjugated to one or both of the primers. Alternatively, the pool of nucleotides used in the amplification is labeled, so as to incorporate the label into the amplification product.

It is readily recognized by those ordinarily skilled in the art that in order to maximize the signal to noise ratio, in probe hybridization detection procedure, the polymorphic site should at the center of the probe fragment used, whereby a mismatch has a maximum effect on destabilizing the hybrid molecule; and in a PCR detection procedure, the polymorphic site should be placed at the very 3'-end of the primer, whereby a mismatch has the maximum effect on preventing a chain elongation reaction by the DNA polymerase. The location of nucleotides in a polynucleotide with respect to the center of the polynucleotide are described herein in the following manner. When a polynucleotide has an odd number of nucleotides, the nucleotide at an equal distance from the 3' and 5' ends of the polynucleotide is considered to be "at the center" of the polynucleotide, and any nucleotide immediately adjacent to the nucleotide at the center, or the nucleotide at the center itself is considered to be "within 1 nucleotide of the center." With an odd number of nucleotides in a polynucleotide any of the five nucleotide positions in the middle of the polynucleotide would be considered to be within 2 nucleotides of the center, and so on. When a polynucleotide has an even number of nucleotides, there would be a bond and not a nucleotide at the center of the polynucleotide. Thus, either of the two central nucleotides would be considered to be "within 1 nucleotide of the center" and any of the four nucleotides in the middle of the polynucleotide would be considered to be "within 2 nucleotides of the center," and so on.

In some embodiments, a composition contains two or more differently labeled oligonucleotides for simultaneously probing the identity of nucleotides or nucleotide pairs at two or more polymorphic sites. It is also contemplated that primer compositions may contain two or more sets of allele-specific primer pairs to allow simultaneous targeting and amplification of two or more regions containing a polymorphic site.

Alternatively, the relevant portion of the gene of the sample of interest may be amplified via PCR and directly sequenced, and the sequence be compared to the wild type sequence shown in the figures. It is readily recognized that, other than those disclosed specifically herein, numerous primers can be devised to achieve the objectives. PCR and sequencing techniques are well known in the art and reagents and equipments are readily available commercially.

Alternatively, an invasive signal amplification assay, as described in e.g. U.S. Pat. No. 5,422,253 and Lyamichev et al., 2000, Biochemistry 39:9523-9532, both incorporated herein by reference in their entirety, may be used for detecting the SNP of interest. This assay takes advantage of enzymes such as the 5' nuclease activity of a DNA polymerase or the gene 6 product from bacteriophage T7 in their ability to cleave polynucleotide molecules by recognizing specific structures instead of specific sequences. A single-stranded target molecule is annealed to a pilot oligonucleotide such that the 5' end of the pilot forms a duplex with the target molecule. If the 3' end of the pilot oligonucleotide does not pair with the target, a 3' arm is formed. When exposed to a cleavage agent such as a DNA polymerase having a 5' nuclease activity or the gene 6 product from bacteriophage T7, the target molecule is cleaved in the 5' region, one nucleotide into the duplex adjacent to the unpaired region of the target. If a cut in a double-stranded molecule is required, the double-stranded molecule is denatured. Because this unpaired 3' arm can be as short as one nucleotide, this assay can be used for detecting a single-nucleotide difference, e.g. in the context of SNP detection. The pilot oligonucleotide is designed such that it pairs perfectly with one allele, but has a 3', single nucleotide mismatch with another allele. Cleavage only occurs if there is a mismatch between the target molecule and the pilot. To achieve signal amplification, the above invasive reaction is modified such that cleavage occurs on the pilot oligonucleotide. Two oligonucleotides are annealed in an adjacent manner to the target molecule. The resulting adjacent duplexes overlaps by at least one nucleotide to create an efficient substrate, called the overlapping substrate, for the 5' nucleases. The 5' end of the downstream oligonucleotide, also called the probe, contains an unpaired region termed the 5' arm (Lyamichev et al., 1993, *Science* 260:778-783.) or flap (Harrington and Lieber, 1994, *EMBO J* 13: 1235-1246) that is not required for the enzyme activity; however, very long arms can inhibit cleavage (Lyamichev et al., 1993, *Science* 260:778-783). Specific cleavage of the probe, termed invasive cleavage (Lyamichev et al., 1999, *Nat. Biotechnol.* 17 292-296; Kwiatkowski et al., 1999, *Mol. Diagn.* 4, 353-364.), occurs at the position defined by the 3' end of the upstream oligonucleotide, which displaces or "invades" the probe. If the overlap between the adjacent oligonucleotides is only one nucleotide, cleavage takes place between the first two base pairs of the probe, thus releasing its 5' arm and one nucleotide of the base paired region (Lyamichev et al., 1999, *Proc. Natl. Acad. Sci. USA*. 96: 6143-6148, and Kaiser et al., 1999, *J Biol. Chem.* 274: 21387-21394). If the upstream oligonucleotide and the probe are present in large molar excess over the target nucleic acid, and invasive cleavage is carried out near the melting temperature of the probe, a cut probe can rapidly dissociate, and an intact probe will anneal to the target more frequently than will a cut probe, thus initiating a new cycle of cleavage. This allows multiple probes to be cut for each target molecule under isothermal conditions, resulting in

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linear signal amplification with respect to target concentration and time (Lyamichev et al., 1999, *Nat. Biotechnol.* 17: 292-296).

DNA markers have several advantages; segregation is easy to measure and is unambiguous, and DNA markers are co-dominant, i.e., heterozygous and homozygous animals can be distinctively identified. Once a marker system is established selection decisions could be made very easily, since DNA markers can be assayed any time after a blood sample can be collected from the individual infant animal, or even earlier by testing embryos in vitro if very early embryos are collected. The use of marker assisted genetic selection will greatly facilitate and speed up cattle breeding problems. For example, a modification of the multiple ovulation and embryo transfer (MOET) procedure can be used with genetic marker technology. Specifically, females are superovulated, eggs are collected, in vitro fertilized using semen from superior males and implanted into other females allowing for use of the superior genetics of the female (as well as the male) without having to wait for her to give birth to one calf at a time. Developing blastomeres at the 4-8 cell stage may be assayed for presence of the marker, and selection decisions made accordingly.

In one embodiment of the invention an assay is provided for detection of presence of a desirable genotype using the markers.

The term "genotype" as used herein refers to the identity of the alleles present in an individual or a sample. In the context of the present invention a genotype preferably refers to the description of the polymorphic alleles present in an individual or a sample. The term "genotyping" a sample or an individual for a polymorphic marker refers to determining the specific allele or the specific nucleotide carried by an individual at a polymorphic marker.

The present invention is suitable for identifying a bovine, including a young or adult bovine animal, an embryo, a semen sample, an egg, a fertilized egg, or a zygote, or other cell or tissue sample therefrom, to determine whether said bovine possesses the desired genotypes of the present invention, some of which are indicative of improved reproduction traits.

Further provided is a method for genotyping the bovine genes identified in Table 1, comprising determining for the two copies of the gene in a diploid genome present the identity of the nucleotide pair at the relevant SNP position (see below).

One embodiment of a genotyping method of the invention involves examining both copies of the gene, or a fragment thereof, to identify the nucleotide pair at the polymorphic site in the two copies to assign a genotype to the individual. In some embodiments, "examining a gene" may include examining one or more of: DNA containing the gene, mRNA transcripts thereof, or cDNA copies thereof. As will be readily understood by the skilled artisan, the two "copies" of a gene, mRNA or cDNA, or fragment thereof in an individual may be the same allele or may be different alleles. In another embodiment, a genotyping method of the invention comprises determining the identity of the nucleotide pair at the polymorphic site.

The present invention further provides a kit for genotyping a bovine sample, the kit comprising in a container a nucleic acid molecule, as described above, designed for detecting the polymorphism, and optionally at least another component for carrying out such detection. Preferably, a kit comprises at least two oligonucleotides packaged in the same or separate containers. The kit may also contain other components such as hybridization buffer (where the oligo-

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nucleotides are to be used as a probe) packaged in a separate container. Alternatively, where the oligonucleotides are to be used to amplify a target region, the kit may contain, preferably packaged in separate containers, a polymerase and a reaction buffer optimized for primer extension mediated by the polymerase, such as PCR.

In one embodiment the present invention provides a breeding method whereby genotyping as described above is conducted on a bovine animal, especially a bull, and based on the results, certain cattle are either selected or dropped out of the breeding program.

Through use of the linked marker loci, procedures termed "marker assisted selection" (MAS) may be used for genetic improvement within a breeding nucleus; or "marker assisted introgression" for transferring useful alleles from a resource population to a breeding nucleus (Soller 1990; Soller 1994).

A method is further provided for determining whether an individual bovine animal is suitable as a gamete donor for natural mating, artificial insemination or in vitro fertilization, the method comprising determining the identity of one or more SNPs according to the present invention using a method of the present invention, and excluding as gamete donor an individual which does not have the preferred allele identity at the respective SNP position as described in Table 1.

Specifically, an individual bovine animal, especially a bull, is excluded as a gamete donor, if its genome does not have at least:

- 1) adenine (A) at the position of its MAP1B gene corresponding to position 3066 of FIG. 1;
- 2) thymine (T) at the position of its MAP1B gene corresponding to position 3323 of FIG. 1;
- 3) cytosine (C) at the position of its MAP1B gene corresponding to position 87071 of FIG. 3;
- 4) guanine (G) at the position of its PPP1R11 gene corresponding to position 112 of FIG. 4;
- 5) adenine (A) at the position of its DDX4 gene corresponding to position 61646 of FIG. 6, or
- 6) G at the position of its DDX4 gene corresponding to position 34239 of FIG. 5.

In one embodiment, the individual is excluded as a gamete donor if the gene type of the individual is not homozygous of the preferred allele with regard to the respective SNP position.

The present invention additionally provides a method of selecting a bovine embryo for planting in a uterus, the method comprising genotyping the embryo according to the present invention, while preserving the viability of the embryo, and excluding from planting an embryo which does not have the preferred allele identity at the respective SNP position as described in Table 1.

In another embodiment, the present invention further provides a method for selectively breeding cattle using a multiple ovulation and embryo transfer procedure (MOET), the method comprising superovulating a female animal, collecting eggs from said superovulated female, in vitro fertilizing said eggs from a male animal which has at least one preferred allele identity at the respective SNP position as described in Table 1 and above, implanting said fertilized eggs into other females allowing for an embryo to develop.

In another embodiment, the method for selectively breeding cattle using MOET may further comprise a step of

genotyping the male animal according to the method of the present invention.

In another embodiment, the method for selectively breeding cattle using MOET may further comprise a step of genotyping the developing embryo, and allowing pregnancy to proceed only if the genotype of the embryo comprises at least one preferred allele identity at the respective SNP position as described in Table 1.

The following examples are intended to illustrate preferred embodiments of the invention and should not be interpreted to limit the scope of the invention as defined in the claims.

genotyping, genomic DNA was extracted from semen samples of 268 Holstein bulls (Genex Cooperative/CRI, Shawano, Wis.) using standard phenol/chloroform protocols. One DNA pool was constructed from 20 random semen samples with equal amounts of DNA. The DNA pool was amplified using primers designed in the 22 candidate genes to amplify 5' untranslated regions (UTRs), exons, introns, and 3' UTRs. The PCR products were sequenced, and SNPs were identified by visually inspecting sequence traces. PCR amplification and sequencing were performed as described in Khatib et al. (2008). Table 2 shows the primer sets used to amplify the nine candidate spermatogenesis genes found to be polymorphic in the bull population.

TABLE 2

Primers used to amplify the nine spermatogenic genes					
Gene	Forward Primer	SEQ ID NO:	Reverse Primer	SEQ ID NO:	Product Size
DCUN1D1	ATACCCTTAGGCAGTTAG	7	AATTGTAAACCCCTGAGAC	5	536
DDX4 (1)	AAACACGGAACAGAGGGT	9	AGGCAGGATTAGCAAGTATG	10	404
DDX4 (2)	AACCAAGTGGCTGGGATG	11	CAGACTCAAATGCGACAA	12	354
DNA11 (1)	CGGTAAGTGAGCAGCATC	13	ACTGAAGCCTTGCCTA	14	495
DNA11 (2)	CCCAGTGCTCCAATCCT	15	ATGGCTCATCTGTCTTCAGTA	16	413
DAN11 (3)	CGTGACTGGTTAGGAT	17	CTGGTGGCTGCTGTCTAT	18	602
GAPDHS (1)	CCAGGAAACGGCATCAC	19	ACACGCAGCAGGGCAACT	20	414
GAPDHS (2)	GTGAAGGCCAGGGACTATGA	21	ACATGAACAAGGGCTGCT	22	541
GSTM3	TTCTCTCCCTGCAAGTCGT	23	TGAGAACAGCTGCCATCATC	24	664
MAP1B (1)	CCATTTCTAAAGGCACAG	25	TTCCGCCATCTCCTTACA	26	504
MAP1B (2)	CTTATGGTCGTGATTATGAA	27	AAGGCTAACACTGCTGGT	28	502
MAP1B (3)	GGCTGTGACATACTTACC	29	CAGACCTCCCTACTTATT	30	645
PPP1R11 (1)	CACATTACGGCGGAACTA	31	ATCCCAAGCAGTATCACCTA	32	591
PPP1R11 (2)	ACCTGTTCTATCTCCTCCCA	33	GTCACCTACCCACCTTGC	34	543
SPATA20	TTGGAGAAGAAACCCACCA	35	CCTCACAAAGCAAGGCTAAGG	36	459
UBC	TCGCTCAGTCGTCTTAC	37	TCAACCAACGCCATAATGT	38	420

EXAMPLES

Materials and Methods

Associations of candidate fertility genes examined were carried out in two experiments. In the first part, single nucleotide polymorphisms (SNPs) in the spermatogenesis genes were tested for associations with sire conception rate (SCR) in a large bull population. In the second part, genes found significantly associated with SCR were tested for association with female fertility traits (fertilization and blastocyst rates). Male fertility genes that play roles in female fertility can be used to improve reproductive performance in cattle using genetic information from both males and females in breeding schemes.

Gene Selection, SNP Identification, and Genotyping for Bull Fertility

A total of 58 spermatogenesis genes, with conserved testicular expression from fly to human, were reported in Bonilla and Xu (2008). Of those genes, only 22 were annotated in the bovine genome. For SNP identification and

50 A total of 24 SNPs located in the nine spermatogenic genes were genotyped in the Genex population (268 animals) by MALDI-TOF MS (GeneSeek Inc., Lincoln, Nebr.).

Imputation of SNPs for Validation of Significant SNPs Found in Genex Population in a Larger Bull Population Obtained from the USDA

Solely for the purpose of validating the conclusions drawn 55 from the results using the Genex population, and to increase the sample size and improve the statistical power of the study, the 24 SNPs identified in the 9 candidate genes and genotyped in 268 bulls (from here forward, the reference population) were imputed in a total of 1,720 bulls (from here forward, the imputed population) so that a final dataset of 1,988 bulls with genotypic data was generated for subsequent statistical analyses. Bulls in the reference and imputed populations have been previously genotyped with the Illumina BovineSNP50 Bead Chip, and hence shared SNP were used to infer the genotypes of the unshared SNP in the imputed population. Genotypes of the reference (n=268) and the imputed populations (n=1,720) for the 50K SNP Chip

were provided by Genex Cooperative/CRI (Shawano, Wis.) and the Animal Improvement Programs Laboratory of the United States Department of Agriculture (AIPL, USDA; Beltsville, Md.), respectively. SNP with minor allele frequencies below 5% were removed. After data editing, 38,265 SNP spanning the entire bovine genome were available in both populations for the imputation process.

Imputation of SNPs was carried out for each candidate gene separately. In each case, a total of 100 SNP on each side of the gene were used to infer the genotypes of the ungenotyped SNP. Imputation was performed using the population-based haplotype clustering algorithm of Scheet and Stephens (2006), which was implemented via the fastPHASE version 1.2 software using the default settings for all parameters (University of Washington TechTransfer Digital Ventures Program, Seattle, Wash.).

Phenotypic Data for Bull Fertility

The 1,988 bulls genotyped with the 50K SNP Chip were evaluated for sire conception rate (SCR), a phenotypic evaluation of bull fertility provided to dairy producers by AIPL-USDA as described in Peñagaricano et al. (2012). Briefly, SCR is the expected difference in conception rate of a sire compared with the mean of all other evaluated sires (Kuhn and Hutchison, 2008; Kuhn et al., 2008).

In this study, SCR values ranged from -10.66% to +6.80%, and the number of breedings per bull ranged from 303 to 111,402. SCR data were obtained from seven consecutive evaluations provided by AIPL-USDA between August 2008 and December 2010. For bulls with multiple evaluations, the most recent SCR evaluation was used in the analysis.

Statistical Analysis for Bull Fertility

The association between each SNP and SCR was evaluated using the following mixed linear model,

$$\text{SCR}_{ijkl} = \mu + \text{EVAL}_j + \beta \text{SNP}_k + \text{sire}_i + e_{ijkl}$$

where μ is the general mean, EVAL_j is the fixed effect of the j^{th} AIPL-USDA SCR evaluation ($j=1, 2, \dots, 7$), SNP_k is the number of copies of one allele of the SNP (corresponding to 0, 1 or 2 copies) carried by the i^{th} animal ($i=1, 2, \dots, 1988$), β is the regression coefficient for the SNP considered (also known as the allele substitution effect), sire_i represents the random additive genetic effect of the i^{th} sire ($i=1, 2, \dots, 246$) of the i^{th} animal, and e_{ijkl} represents the random residual for each observation. To detect possible deviations from the additive model, associations between genotype and SCR were evaluated using SNP as a categorical variable.

Random effects were assumed to follow the multivariate normal distribution,

$$\begin{pmatrix} s \\ e \end{pmatrix} \left| \sigma_s^2, \sigma_e^2 \right. \sim N \left[0, \begin{pmatrix} A\sigma_s^2 & 0 \\ 0 & W^{-1}\sigma_e^2 \end{pmatrix} \right]$$

where s and e are the vectors of sire and residual effects, respectively: σ_s^2 and σ_e^2 are the sire and residual effect variances, respectively; A represents the matrix of additive relationships between sires in the pedigree ($1,558 \times 1,558$) and W is a diagonal matrix of order 1,988 with its elements representing reliabilities of SCR values. The A matrix was calculated based on a five-generation pedigree of sires downloaded from AIPL-USDA. The association between each SNP and SCR was tested using a likelihood ratio test by comparing to a reduced model without the SNP effect.

Phenotypic and Genotypic Data for Cow Fertility

The most significant SNP for SCR (rs109423562 located in MAP1B) was further investigated for association analysis with fertilization and blastocyst rates—the main cow fertility traits—using an IVF system. The procedures of in vitro fertilization and subsequent embryo culture were described in Khatib et al. (2008). To generate fertilization and blastocyst rate data, a total of 6,282 in vitro fertilizations were performed, and a total of 4,207 embryos were produced using oocytes from 359 ovaries collected from 359 Holstein cows and semen samples from 12 Holstein bulls. For 74 ovaries, oocytes were fertilized by two different bulls each. Fertilization rate was calculated as the number of cleaved embryos at Day 2 post-fertilization divided by the total number of fertilized oocytes collected from one ovary. Blastocyst rate was calculated as the number of embryos that reached the blastocyst stage (Day 8) and appeared normal out of the total number of embryos produced.

The 359 ovaries were genotyped for SNP rs109423562 (G/A) using PCR-RFLP. A 171 bp fragment was amplified using the primers 5'-GCAGCTCTTTAGGAGTGT-TAGCGTCTGAT-3' (SEQ ID NO: 39) (forward) and 5'-CT-CACAGAGGGCATTGACA-3' (SEQ ID NO: 40) (reverse). The PCR product was then digested by the restriction enzyme Hinfl and electrophoresed on a 2.0% agarose gel. Allele G was cut while allele A was uncut.

Statistical Analysis for Cow Fertility

Association between SNP rs109423562 (G/A) in MAP1B and fertilization and blastocyst rates were analyzed using the following mixed linear model,

$$y_{ijk} = \mu + \text{ovary}_i + \text{sire}_j + \text{SNP}_{ijk} + e_{ijk}$$

where y_{ijk} represents the fertilization or blastocyst rate of oocyte k from ovary i fertilized with semen from bull j , μ represents a general mean for the trait considered, ovary_i represents the random effect of the individual ovary from which oocytes were harvested, sire_j represents the random effect of the sire used in the fertilization, SNP_{ijk} represents the fixed effect of the ovary genotype for the SNP considered, and e_{ijk} represents the residuals, assumed normal, independent and identically distributed with mean 0 and variance $I\sigma_e^2$. Ovaries and bulls were assumed uncorrelated with variance structures $I\sigma_o^2$ and $I\sigma_s^2$, respectively. Association between the SNP and fertilization or blastocyst rate was tested again using a likelihood ratio test by comparing with a reduced model without the SNP effect. All the statistical analyses were performed using the pedigreemm package (Vazquez et al. 2010) of the R language/environment (R Development Core Team 2009).

Results

SNP Identification and Association of Candidate Genes with SCR

Sequencing analysis revealed 24 SNPs in 9 spermatogenesis genes (DCUN1D1, DDX4, GAPDHS, 55 GSTM3, MAP1B, PPP1R11, SPATA20 and UBC) in the bull population using the pooled DNA sequencing approach. All 24 SNPs located in 9 candidate genes were tested for association with sire conception rate first in Genex population and then in a larger population of 1,988 bulls for validation.

Association Analysis in Genex Population

SNPs in the genes MAP1B, PPP1R11, and DDX4 are associated with SCR

For MAP1B gene, a SNP C/T in intron 5 at position 65 9331992 (University of Maryland bovine version 3.1; UMD3.1) showed significant association with sire conception rate. Primers used to amplify the gene and SNP location

are shown in FIG. 1. Frequency of allele T was 25% and frequency of allele C was 75%. The allele substitution effect was -0.15 ± 0.01 ($p\text{-value}=0.01$) so that allele C is favorable for SCR.

For PPP1R11, one SNP (T/G) in the 5'UTR region at position 28710268 (UMD3.1) was significantly associated with SCR in the Genex population, with allele G associated with increased SCR. SEQ ID NO:6 shows SNP location and primers used to identify the SNP.

For DDX4, two SNPs located at positions 23382814 (SNP1; A/G) and 23410221 (SNP2; G/A) were identified in the gene. For SNP locations see FIG. 3. The two SNPs were in almost complete linkage disequilibrium, so they have the same allele frequencies in the bull population examined. Genotype AA of SNP1 has 0.783 units of SCR versus -0.214 SCR units for GG genotype ($P\text{-value}=0.05$). Similarly, GG genotype of SNP2 has 0.749 units of SCR versus -0.494 for AA genotype.

Association of Spermatogenesis Genes in the Combined Genex and USDA Populations

Three SNPs located in MAP1B and one SNP in PPP1R11 showed significant associations with SCR (Table 3). The SNP with the most significant association with SCR is SNP5.MAP1B located in intron 5 with an allele substitution effect of -0.24 and a $P\text{-value}$ of 0.001. The other two significant SNPs were SNP1.MAP1B and SNP3.MAP1B, both located in intron 1 with an allele substitution effect of 0.15 and $P\text{-values}$ of 0.025 and 0.039, respectively. After Bonferroni correction for multiple testing, only SNP5.MAP1B remained significant ($P\text{-value}=0.024$). Pairwise linkage disequilibrium (LD) tests of MAP1B SNPs showed a moderate LD ($r^2=0.38$) between SNP1.MAP1B and SNP3.MAP1B and SNP5.MAP1B. The LD between SNP3.MAP1B and SNP5.MAP1B was relatively low ($r^2=0.14$).

SNP1.PPP1R11 located in the 5'UTR region of PPP1R11 showed significant association with SCR with an allele substitution effect of 0.15 and a $P\text{-value}$ of 0.046.

TABLE 3

Genetic markers significantly associated with sire conception rate				
SNP ID	Gene	Genotype (N)	Allele substitution effect \pm SE	P-value
SNP1.MAP1B	M4P1B	GG (463)	0.15 ± 0.01	0.025
		GA (987)		
		AA (538)		
SNP3.MAP1B	M4P1B	CC (972)	0.15 ± 0.01	0.039
		CT (847)		
		TT (169)		
SNP5.MAP1B	MAP1B	CC (1081)	-0.24 ± 0.01	0.001
		TC (763)		
		TT (144)		
SNP1.PPP1R11	PPP1R11	TT (1033)	0.15 ± 0.01	0.046
		TG (821)		
		GG (134)		

Association of MAP1B with Fertilization Rate and Embryo Survival Rate

SNP5.MAP1B located in intron 5 of MAP1B showed the most significant association with sire conception rate in the bull population analyzed above. To characterize its impact on female fertility, we tested the association of this SNP with fertilization rate and blastocyst rate in the IVF system. SNP5.MAP1B showed significant associations with both fertilization rate ($P\text{-value}=0.027$) and blastocyst rate ($P\text{-value}=0.029$) (Table 4). Oocytes collected from genotype CT cows showed the lowest fertilization rate (59.9%) com-

pared with that from CC (66.4%) and TT (66.3%) cows (Table 4). For blastocyst rate, the CT genotype again showed the lowest rate (27.1%) while homozygous CC and TT individuals showed blastocyst rates of 31.0% and 41.8%, respectively (Table 4).

TABLE 4

Association between SNPs of MAP1B gene and fertilization rate and blastocyst rate				
Genotype (N)	Fertilization rate		Blastocyst rate	
	Estimate \pm SE	P-value	Estimate \pm SE	P-value
CC (321)	0.664 ± 0.03	0.027	0.310 ± 0.02	0.029
CT (84)	0.599 ± 0.03		0.271 ± 0.03	
TT (28)	0.663 ± 0.04		0.418 ± 0.05	

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<223> OTHER INFORMATION: r is g or a

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 <223> OTHER INFORMATION: y is c or t

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acttcttcat cttcccgtgg tttcatgtcg agtcagaatg taaggactgt ttagcttttg	180
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tttcttgtat acttccattt tcacttttagt cacaaggcag gttggggaaag caggaggggg	300
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39

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<210> SEQ ID NO 7
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<400> SEQUENCE: 7

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<210> SEQ ID NO 8
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Reverse Primer DCUN1D1

<400> SEQUENCE: 8

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aattgttaaac cctgagac 18
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<210> SEQ ID NO 9
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
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<400> SEQUENCE: 9

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<400> SEQUENCE: 11

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<210> SEQ ID NO 12
<211> LENGTH: 18
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<213> ORGANISM: Artificial Sequence
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<400> SEQUENCE: 12

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cagactcaaa tgcgacaa	18
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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
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<400> SEQUENCE: 13

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<210> SEQ ID NO 14
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<212> TYPE: DNA
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<223> OTHER INFORMATION: Reverse Primer DNAI1(2)

<400> SEQUENCE: 16

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<210> SEQ ID NO 17
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<212> TYPE: DNA
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<220> FEATURE:
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<400> SEQUENCE: 17

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18

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18

<210> SEQ ID NO 19
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<400> SEQUENCE: 19

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18

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<400> SEQUENCE: 21

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 <212> TYPE: DNA
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<400> SEQUENCE: 22

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<400> SEQUENCE: 23

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<210> SEQ ID NO 24
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<212> TYPE: DNA
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<400> SEQUENCE: 24

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<210> SEQ ID NO 25
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<212> TYPE: DNA
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<220> FEATURE:
<223> OTHER INFORMATION: Forward Primer MAP1B(1)

<400> SEQUENCE: 25

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<223> OTHER INFORMATION: Reverse Primer MAP1B(1)

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<210> SEQ ID NO 28
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<400> SEQUENCE: 28

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18

<210> SEQ ID NO 29
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<220> FEATURE:
<223> OTHER INFORMATION: Forward Primer MAP1B(3)

<400> SEQUENCE: 29

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<400> SEQUENCE: 30

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19

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<220> FEATURE:
<223> OTHER INFORMATION: Forward Primer PPP1R11(1)

<400> SEQUENCE: 31

cacattacgg cggaaacta

18

<210> SEQ ID NO 32
<211> LENGTH: 20
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<223> OTHER INFORMATION: Reverse Primer PPP1R11(1)

<400> SEQUENCE: 32

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<210> SEQ ID NO 33
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<223> OTHER INFORMATION: Forward Primer PPP1R11(2)

<400> SEQUENCE: 33

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<210> SEQ ID NO 34
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<220> FEATURE:
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<400> SEQUENCE: 34

gtcacctacc caccttg

18

<210> SEQ ID NO 35
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<223> OTHER INFORMATION: Forward Primer SPATA20

<400> SEQUENCE: 35

ttggagaaga aaccaccag

20

<210> SEQ ID NO 36
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<212> TYPE: DNA
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<223> OTHER INFORMATION: Reverse Primer SPATA20

<400> SEQUENCE: 36

cctcacaagg aaggctaagg

20

<210> SEQ ID NO 37

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<211> LENGTH: 19
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<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Forward Primer UBC

<400> SEQUENCE: 37

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19

<210> SEQ ID NO 38
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<212> TYPE: DNA
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<223> OTHER INFORMATION: Reverse Primer UBC

<400> SEQUENCE: 38

tcaaccaacg cctaatgt

18

<210> SEQ ID NO 39
<211> LENGTH: 30
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Forward Primer PCR-RFLP

<400> SEQUENCE: 39

gcagctcttt taggagtgtt agcgctctgat

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<210> SEQ ID NO 40
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<220> FEATURE:
<223> OTHER INFORMATION: Reverse Primer PCR-RFLP

<400> SEQUENCE: 40

ctcacagagg gcatttgaca

20

40

What is claimed is:

1. A method for implanting an embryo comprising:
in vitro fertilizing cattle eggs to obtain fertilized eggs,
culturing fertilized eggs into developing embryos
detecting the identity of a nucleotide of a MAP1B gene of
an embryo at a position corresponding to position 1986
of SEQ ID NO: 1, wherein the MAP1B gene comprises
the nucleotide sequence of SEQ ID NO: 1, and
identifying an embryo that has an adenine at said position,
and
implanting said identified embryo into a suitable female
bovine.
2. The method of claim 1, wherein the identity of both
copies of the gene in the embryo is determined.
3. A method according to claim 1, wherein the nucleotide
is detected by sequencing the gene or a relevant fragment
thereof.
4. A method according to claim 3, wherein the gene or a
relevant fragment thereof is isolated from the animal's
nucleic acid sample via amplification by a polymerase chain
reaction.
5. A method of selecting a bull bovine animal as a breeder,
the method comprising:

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obtaining a sample of the animal's nucleic acid, wherein
the nucleic acid comprises at least a partial MAP1B
gene comprising SEQ ID NO: 1,
detecting an adenine nucleotide at a position of the
MAP1B gene of the animal corresponding to position
1986 of SEQ ID NO: 1,
selecting the bull bovine animal that has an adenine at said
position, and
using bovine cell or tissue from the selected bull animal
in a breeding procedure.

6. The method according to claim 5, wherein the bovine
cell or tissue is sperm.

7. A method according to claim 5, wherein the nucleotide
is detected by sequencing the MAP1B gene or a relevant
fragment thereof.

8. A method according to claim 7, wherein the gene or a
relevant fragment thereof is isolated from the animal's
nucleic acid sample via amplification by a polymerase chain
reaction.

9. The method of claim 5, wherein the identity of both
copies of the gene in the bovine bull animal is determined.

* * * * *

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