

A nonsense mutation in cGMP-dependent type II protein kinase (*PRKG2*) causes dwarfism in American Angus cattle

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Historically, dwarfism was the major genetic defect in U.S. beef cattle. Aggressive culling and sire testing were used to minimize its prevalence; however, neither of these practices can eliminate a recessive genetic defect. We assembled a 4-generation pedigree to identify the mutation underlying dwarfism in American Angus cattle. An adaptation of the Elston-Steward algorithm was used to overcome small pedigree size and missing genotypes. The dwarfism locus was fine-mapped to BTA6 between markers *AFR227* and *BM4311*. Four candidate genes were sequenced, revealing a nonsense mutation in exon 15 of cGMP-dependant type II protein kinase (*PRKG2*). This C/T transition introduced a stop codon (R678X) that truncated 85 C-terminal amino acids, including a large portion of the kinase domain. Of the 75 mutations discovered in this region, only this mutation was 100% concordant with the recessive pattern of inheritance in affected and carrier individuals (log of odds score = 6.63). Previous research has shown that *PRKG2* regulates SRY (sex-determining region Y) box 9 (*SOX9*)-mediated transcription of collagen 2 (*COL2*). We evaluated the ability of wild-type (WT) or R678X *PRKG2* to regulate *COL2* expression in cell culture. Real-time PCR results confirmed that *COL2* is overexpressed in cells that overexpressed R678X *PRKG2* as compared with WT *PRKG2*. Furthermore, *COL2* and *COL10* mRNA expression was increased in dwarf cattle compared with unaffected cattle. These experiments indicate that the R678X mutation is functional, resulting in a loss of *PRKG2* regulation of *COL2* and *COL10* mRNA expression. Therefore, we present *PRKG2* R678X as a causative mutation for dwarfism cattle.

achondroplastic dwarfism | BTA6 | cGKII | recessive genetic disease

Bovine dwarfism was widely observed from the late 1940s through the 1960s, making it one of the greatest problems in the history of the U.S. beef cattle industry (1). Even President Eisenhower showed great interest in the disease, because he raised Angus cattle on his ranch. Over the years, the disease has cost beef producers millions of dollars. Affected animals represent an economic loss, because of slow growth and lack of market value. Breeders have attempted to eliminate the disease by using test matings or removing parents in a herd known to produce affected offspring. These methods are extremely costly, and did not remove all of the carriers in the past. Yet by the 1970s, producers thought that dwarfism had been eliminated. Unfortunately, that was not the case, as 6 dwarf calves were discovered on several ranches across the United States in the spring of 2002. Today, there is no good estimate of the number of affected individuals in the United States. Rumors indicate that affected individuals are often not reported, and are removed from herds at birth to prevent negative publicity for the cattle breeder. Cattle breeders with suspect carriers have lost their livelihood due to the negative connotations associated with the damaging history of dwarfism in the beef industry. Development of a

definitive genetic test will finally allow producers to eliminate dwarfism while preserving genetically superior carrier animals with desirable genetic backgrounds in the Angus breed.

A multitude of dwarfism phenotypes occur in nature. Dwarfism occurs in many species, and several animal models exist. Notably, transgenic models in the mouse have been used to understand the implications of mutations within the fibroblast growth factor receptor 3 (*FGFR3*) gene. In humans, achondroplasia is caused primarily by a dominant mutation in *FGFR3* (2). Angus dwarfism is also characterized as a form of achondroplasia. Affected individuals have diminished endochondral ossification of growth plates, protrusions of the alar wing of the basisphenoid bone into the cranial cavity, abnormalities of the ventral vertebral bodies, and curvature of the vertebral processes (3). In American Angus, dwarfism appears to follow a recessive pattern of inheritance.

In cattle, few forms of dwarfism have been characterized at the molecular level (i.e., Dexter and Japanese Brown cattle) (4, 5). Disproportionate dwarfism in Japanese brown cattle is a recessive phenotype caused by insufficient endochondral ossification and abnormal arrangement of growth plate chondrocytes. Affected individuals have shortened stature, joint abnormalities, and ateliosis (5). Our group previously investigated the possibility that Angus dwarfism was caused by either of the 2 mutations in the *LIMBIN* gene that cause dwarfism in Japanese brown cattle. Analysis of the 2 mutations showed that Angus dwarf calves were wild-type for the *LIMBIN* mutations (3). We also investigated the possibility that dwarfism in the Angus and Dexter breeds could share a common genetic cause. Dexter dwarfism is a form of chondrodysplasia also caused by defective endochondral ossification due to 2 different mutations in aggrekan 1 (*ACAN*) (6). Homozygous mutants display a “bulldog” phenotype, including a retruded muzzle, cleft palate, abdominal hernia, shorted and misshaped long bones, and lethality at approximately 7 months of gestation. Individuals heterozygous

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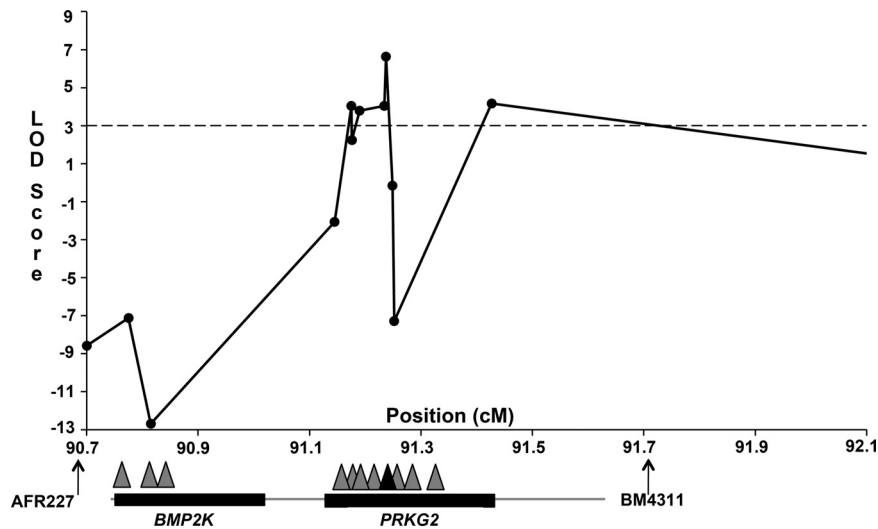


Fig. 1. The maximum LOD score for Angus dwarfism on BTA6 occurs at the *CT* nonsense mutation, indicated by a black triangle, in exon 15 of *PRKG2*. Microsatellite markers are indicated by arrows; SNP markers are indicated by triangles.

for mutations in the *ACAN* gene exhibit disproportionate dwarfism (4, 6). Angus dwarf samples were wild-type for Dexter dwarfism markers. Therefore, we concluded that Angus dwarfism was likely caused by a novel mutation.

Herein, we describe the fine mapping and analysis of positional candidate genes for dwarfism in American Angus cattle using a unique adaptation of the Elston-Steward algorithm to refine the position of the disease locus. We present strong evidence that a nonsense mutation within the kinase domain of *PRKG2* is the causative mutation for Angus dwarfism based on linkage mapping, cell culture, in vivo expression data, and similarities with existing mutant rodent models.

Results

Linkage Analysis Reveals a High Signal on BTA6. To find the Angus dwarfism locus, we began by genotyping an Angus pedigree consisting of 26 individuals, including 6 affected animals (Fig. S1). We prioritized genotyping of chromosomes that contained genes that have been shown to be associated with dwarfism in humans. After investigating several chromosomes with no statistical association, a significant log of odds (LOD) score of 4.53 was detected on BTA6 between 2 microsatellite markers, *AFR227* and *BM4311*. Because no other statistically significant linkage signals were detected, we focused on fine mapping BTA6. We added an additional 19 microsatellite marker genotypes within or closely flanking the critical region to verify the position of the dwarfism locus. Analysis of these new markers increased the size of the critical region from 0.8 to 2.8 cM, flanked by *AFR227* and *BMS511*. The size of the critical region expanded, because increased marker information was provided in the affected individuals by the new marker genotypes distal to *BM4311* (Fig. S2). However, the new critical region still encompassed the original region with approximately the same LOD score.

Positional Candidate Gene Analysis: Discovery of a Nonsense Mutation in *PRKG2*. Four positional candidate genes were prioritized for resequencing based on their known function in bone development or relationship to genes known to be involved in bone development. They were bone morphogenetic protein 2 kinase (*BMP2K*), bone morphogenetic protein 3 (*BMP3*), fibroblast growth factor 5 (*FGF5*), and cGMP-dependant type II protein kinase (*PRKG2*). We sequenced the exons and some flanking intron sequence of each candidate gene in a WT, dwarf carrier,

and dwarf animal. Additionally, we sequenced one Hereford and one Brahman DNA sample as negative controls, which resulted in the identification of 75 single nucleotide polymorphisms (SNPs). We used the Brahman, Hereford, and unaffected Angus as control samples to determine if a given SNP followed the recessive pattern of inheritance for Angus dwarfism. Few heterozygous SNPs were found to be informative in *BMP3* or *FGF5* in Angus, but many heterozygous informative SNPs were detected in *BMP2K* and *PRKG2* (Tables S1–S4). Seven SNPs were detected within exons; however, only one was nonsynonymous. This mutation resulted in a C to T transition [Nt2032 (C-T)] that caused a nonsense mutation in exon 15 of *PRKG2*, herein referred to as the *R678X* marker. The T mutation was homozygous in the dwarf, heterozygous in carriers, and not present in the WT Angus, Hereford, or Brahman controls that we sequenced. Upon genotyping our pedigree, we found 100% concordance of the mutation with dwarf carrier and affected status. We genotyped 11 SNPs that surrounded the *R678X PRKG2* mutation and found a maximal LOD score of 6.63 for the marker interval, including the nonsense mutation (Fig. 1). The marker genotypes for the 6 affected individuals and 2 unaffected full siblings (sibs) show a relatively large segment of homozygosity in the dwarfs, but a much smaller one in the unaffected full sibs (Fig. S2). We genotyped additional breeds, including 8–10 randomly selected samples from Brahman, Hereford, Holstein, Limousin, and South Devon breeds, as well as crossbred cattle composed of Maine Anjou, Charolais, Simmental, and Angus ($n = 54$). However, we were unable to detect the *R678X PRKG2* mutation in additional breeds, indicating that this mutation may be Angus breed specific.

The *PRKG2 R678X* Mutation Causes a Loss of *COL2* Regulation in Cell Culture. Based on previous studies in the rat (7), we designed a cell culture experiment to determine the functionality of the *PRKG2 R678X* mutation found in bovine dwarfs. Human hepatoma (HUH-7) cells were transiently transfected with one of 4 treatments: (i) wild-type bovine (bt) *PRKG2* + human (h) *SOX9*; (ii) *R678X* bt*PRKG2* + h*SOX9*; (iii) h*SOX9* + empty vector; and (iv) empty vector (negative control). The *PRKG2* protein is known to regulate *SOX9* and downstream *COL2* mRNA levels. *PRKG2* downregulates *COL2* by activation of *SOX9* (7). Therefore, we hypothesized that the truncated dwarf *PRKG2* kinase domain would not be able to downregulate *COL2*. Cells transfected with the *R678X* bt*PRKG2* vector had a 38% increase in

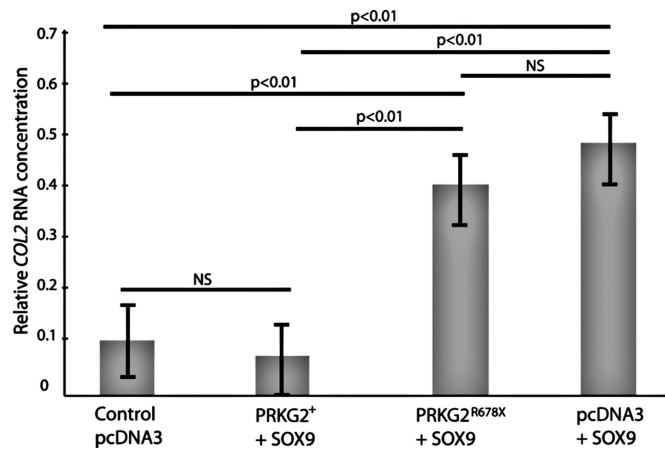


Fig. 2. Comparison of *COL2* mRNA expression levels between WT btPRKG2 and R678X btPRKG2 transfected HUH-7 cells. The lines above the bar graph denote the accompanying statistical difference for the 6 treatment comparisons. The *P* value directly above each line is associated with the difference between only the outermost 2 bars underneath the line. All *P* values are Bonferroni corrected to account for multiple testing. NS, not significant.

COL2 mRNA compared with cells transfected with WT btPRKG2 (Bonferroni corrected $P < 0.01$; Fig. 2). There was no difference in *COL2* levels between the control (empty vector only) and WT PRKG2 + hSOX9 treatment (not significant, $P > 0.70$). The hSOX9 + empty vector control treatment exhibited the same *COL2* expression as the R678X btPRKG2 + hSOX9 treatment (not significant, $P > 0.30$) and increased *COL2* expression compared with btPRKG2 + hSOX9 (Bonferroni corrected $P < 0.01$) and the empty vector alone (Bonferroni corrected $P < 0.01$). These findings indicate that the R678X PRKG2 mutation severely reduced the ability of PRKG2 to regulate SOX9 to levels comparable to a complete loss of function of PRKG2 (vector + hSOX9 treatment alone).

Individuals Homozygous for the R678X Mutation Show Reduced Stature. More than 20 embryos were produced by carrier \times carrier and dwarf \times carrier matings; however, only 6 embryos survived to parturition. We blindly genotyped these 6 individuals using the R678X PRKG2 marker within 2 weeks of birth and successfully predicted dwarf status. Animals predicted to be dwarfs by the R678X PRKG2 marker had shorter metatarsals (reduced 4.5 cm; $P < 0.01$) and fused ulna/radius (reduced 7.5 cm; $P < 0.01$) bones. Although not significant, there were also trends toward significant differences between normal and dwarf metacarpals (reduced 2.75 cm; $P < 0.09$), femur (reduced 3.9 cm; $P < 0.17$), and tibia (reduced 4.9 cm; $P < 0.11$). No significant differences were observed between these animals for head length, calvarium (skull) width, or bone circumference for each of the major long bones previously listed ($P > 0.25$). Estimates of differences in stature and body length were also calculated for live dwarf and normal animals at ≈ 210 days of age. Differences in height of 15.8 cm ($P < 0.0001$) and 20.7 cm of spine length ($P < 0.0001$) were observed between animals declared normal versus those declared dwarf at birth by the R678X genotype. Based on height differences, gross differences in long bone length, and changes in vertebral length, we feel confident that both unaffected and dwarf animals were produced in these matings.

Individuals Homozygous for the R678X Mutation Have Altered COL2 and COL10 Gene Expression in Growth Plate Cartilage. Expression of *COL2* and *COL10* in growth plate cartilage was compared between 4 calves predicted to be dwarfs (PRKG2^{R678X/R678X}) and 2 calves predicted to be unaffected (PRKG2^{R678X/+}) at approx-

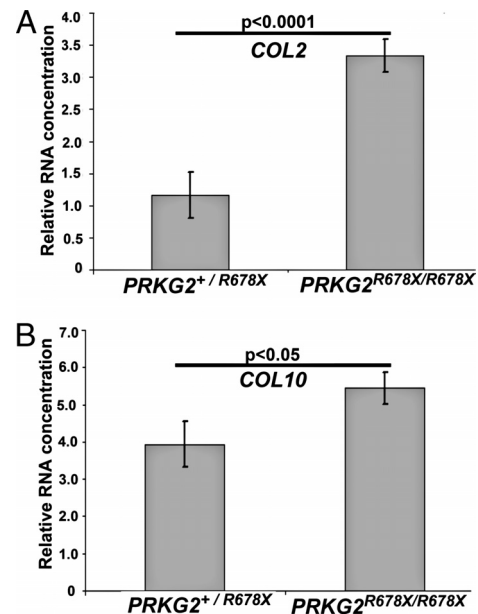


Fig. 3. In vivo gene expression of *COL2* and *COL10* was measured in bovine PRKG2^{R678X/R678X} (dwarf) and PRKG2^{+/R678X} (unaffected) tibia growth plates. (A) Difference in *COL2* expression in dwarf R678X PRKG2 mutants versus normal cattle growth plate. (B) Difference in *COL10* expression in dwarf R678X PRKG2 mutants versus normal cattle growth plate.

imately 7 months of age. Real-time PCR analysis confirmed a considerable increase in *COL2* mRNA in dwarf cattle compared with normal cattle ($P < 0.0001$; Fig. 3A). Real-time PCR of *COL10* indicated an increase in expression in dwarfs compared with normal individuals ($P < 0.05$; Fig. 3B). Expression levels of *COL2* and *COL10* were increased in dwarf (PRKG2^{R678X/R678X}) cattle, indicating expression levels comparable to PRKG2 null dwarf rats (7).

There Is No Evidence of Nonsense-Mediated Decay Due to the R678X Mutation. The R678X mutation occurs in the 15th exon of 18 exons, leading us to test the possibility of nonsense-mediated RNA decay. We tested PRKG2 mRNA expression by quantitative real-time PCR. No difference in PRKG2 expression was detected between R678X homozygous (dwarf) and R678X heterozygous (unaffected) individuals ($P > 0.20$). Because the R678X mutation is in the kinase domain, we would expect truncation of this domain to cause a loss of function even if the mRNA was produced. We wondered if the truncation could somehow occur near enough the end of the protein to not alter the structure of the protein. To predict the effect of the R678X mutation on PRKG2 protein structure, we used the SwissModel homology protein-modeling program to compare WT and R678X PRKG2 (8, 9). The modeling results indicate that a conformational change likely occurs (Fig. S3). Furthermore, a comparison of the primary amino acid sequence across 7 species suggests that the mutation resides in a conserved block of 11-aa residues (Fig. S4).

Discussion

We believe the use of the Elston-Steward algorithm—specifically, the ability to accommodate all possible genotypes for common ancestors in our small pedigree—was critical to our ability to quickly find the Angus dwarfism locus. The linkage mapping results strongly indicated that the R678X mutation in PRKG2 was the causative mutation for Angus dwarfism. Of the 76 mutations detected in the dwarfism-critical region, the R678X mutation was one of only 7 coding mutations, and the only

nonsynonymous coding mutation. Evaluation of dwarf and unaffected full-sib genotypes indicates a small region of homozygosity unique to affected individuals. Homozygosity in unaffected full sibs at markers flanking the *R678X* mutation caused the reduced LOD scores near the causative mutation as a result of analyzing 2-marker haplotypes (Fig. 1). Analysis of a larger haplotype does not exclude these flanking marker intervals; however, this limitation did not deter our ability to conclude which SNP was the causative mutation.

To evaluate the physiological effects in Angus dwarfs, we produced 6 dwarf calves by embryo transfer. Because only 6 of 20 individuals survived full term in pregnancy, this may indicate that embryo death may be an additional consequence in *PRKG2* mutants. This observation is consistent with research that indicates *PRKG2* may function in preovulatory follicles as a response to luteinizing hormone and progesterone (10). Analysis of bone lengths indicates that both dwarf and normal individuals were produced from our mating experiment. Analysis of such small data is difficult because of the limited degrees of freedom available. This experiment does not prove causality of the mutation, because there was likely considerable sharing of large chromosomal segments between these individuals and those in our linkage mapping pedigree due to shared common ancestors.

Rodent models show that *PRKG2* is a logical candidate gene for dwarfism in Angus cattle. Pfeifer et al. (11) created a knockout of *PRKG2* in the mouse, which resulted in mice that exhibited a dwarfism phenotype. These mice exhibited an unorganized growth plate with abnormal stacking of chondrocytes. More recently, Chikuda et al. (7) discovered a deletion in the coding region of *PRKG2* that caused dwarfism in the KMI rat. The loss of growth plate organization observed in these dwarf rats is very similar to what is observed in the Angus dwarf growth plate (Fig. S5).

To explain the role of *PRKG2* in growth plate organization, Chikuda et al. (7) showed that *PRKG2* regulates *SOX9* via phosphorylation. Phosphorylation of *SOX9* is required for nuclear translocation and transcriptional regulation of *COL2* to *COL10* within the growth plate. This is a key developmental checkpoint that allows proper transition of chondrocytes from a proliferative to hypertrophic growth phase. As a consequence, KMI mini-rats show increased *COL2* and decreased *COL10* mRNA expression levels, consistent with constitutive proliferative growth. Angus dwarfs also show a lack of regulation of *COL2* and *COL10* gene expression, as indicated by real-time PCR experiments. Functional changes in *COL2* mRNA levels in *R678X PRKG2* mutants are consistent with those observed in the KMI mini-rat. Results from cell culture indicate that the *R678X PRKG2* mutation was sufficient to change *COL2* mRNA expression. Control transfections using only *SOX9* did not significantly differ in *COL2* expression levels from *R678X* mutants, indicating that dwarf cattle would have *COL2* gene expression similar to a null *PRKG2* individual. Interestingly, the change in *COL10* expression in dwarf cattle was opposite of what was expected. Dwarf cattle show increased expression of *COL10*, unlike the rat *PRKG2* mutants. This could be due to differences in tissue collection, small sample size, temporal changes in expression, or species differences. Alternatively, differences in *COL10* expression could be explained, because bovine expression was measured in vivo, whereas expression measurements in the rat were carried out in cell culture. Because the *R678X* mutation occurred before the last exon, we considered nonsense-mediated decay as a cause of *PRKG2* loss of function. However, we observed no change in *PRKG2* expression in bovine growth plate cartilage. This would suggest that the change in *PRKG2* was caused at the protein level and not due to reduced *PRKG2* gene expression. It is possible that we were unable to detect a difference in expression due to small sample size or the lack of homozygous WT individuals in our assay. Because the mutation truncates a

kinase domain, the functional change is likely at the level of the *PRKG2* protein. Because severe coding mutations cause dwarfism in both transgenic and naturally occurring dwarf animals, it is logical that a similar mutation in cattle would cause dwarfism as well.

The function of *PRKG2* remains widely unknown, but it is clearly an important signal transduction pathway in numerous tissues (e.g., brain, growth plate cartilage, jejunum, kidney, lung, and pancreas). Its potential roles are diverse, ranging from kidney disease to cancer and chondrodysplasia (12, 13). Interestingly, the knockout of *PRKG2* shows no symptoms of infection when treated with STa toxin of *E. coli* (11). This may suggest that genetic variation in *PRKG2* is beneficial to circumvent bacterial infection. The role of *PRKG2* in transport of Na^+ , H^+ , HCO_3^- , Cl^- , and water have been well documented in the small intestine (11, 14, 15). Additionally, *PRKG2* has been shown to be highly expressed in the brain with a largely unknown function (<http://www.brain-map.org/>). Study of *PRKG2* knockouts indicated that *PRKG2* might regulate the circadian clock, behavior, and alcoholism (16–19). The use of *PRKG2* as a therapeutic treatment in medicine has considerable potential. It was previously suggested by Pfeifer et al. (11) that *PRKG2* could be used to enhance the process of bone healing. More recently, it has been suggested that the protein kinase G pathway may be important in developing treatments for diabetes (20). The role of *PRKG2* to regulate the cystic fibrosis transmembrane conductance regulator (*CFTR*) is particularly interesting. Analysis of the *PRKG2* pathway along with cAMP pathways may yield valuable information needed to create new treatments for cystic fibrosis patients. Angus dwarfs do exhibit labored, “raspy” breathing, but it is unclear if this is due to a lung phenotype or skeletal malformation. They do not exhibit additional phenotypes; however, many of referenced phenotypes are difficult to measure without examination at the molecular level.

The mutation for dwarfism in Angus is located on chromosome 6, along with numerous quantitative trait loci (QTLs) for production traits. Of particular interest are QTLs for growth rate, milk yield, and milk fat and protein that would be selected for optimal calf growth and maternal ability (see <http://bovineqtl.tamu.edu/>; http://www.vetsci.usyd.edu.au/reprogen/QTL_Map/; and <http://www.animalgenome.org/QTLdb/cattle.html>). Potential selection pressure for the QTL regions could have increased homozygosity of the region, which led to increased observance of the dwarfism allele. Additionally, the Angus breed went through a transition period where animals were selected to be short and stocky. It is possible that the occurrence of the mutation is a holdover from earlier years when smaller frame size was desired.

We present a strong argument that a nonsense mutation at *R678X* of *PRKG2* causes dwarfism in Angus cattle. First, considering a recessive gene action model, the mutation is 100% concordant with the pedigree, including both dwarfs and dwarf carriers. Second, the mutation causes a premature stop codon in a kinase domain. Third, knockout mouse and naturally occurring rat *PRKG2* mutants exhibit dwarfism with growth plate phenotypes similar to bovine dwarfs. Fourth, transfection of WT versus *R678X PRKG2* demonstrates that the *R678X* mutation causes loss of *PRKG2* function. Last, expression analysis from dwarf and normal cattle show that this mutation alters *COL2* and *COL10* expression levels. Because *COL2* and *COL10* expression level and timing are critical to proper growth plate development, it is likely that such a functional change will alter endochondral ossification. The immediate impact of this research is a genetic test for use by Angus producers to remove dwarfism. Continued research on knockouts and naturally occurring *PRKG2* mutants may unlock the doors to medical treatments for many debilitating genetic diseases. Additional functional analysis is underway

to characterize the impact of the *R678X PRKG2* mutation on bovine physiology.

Materials and Methods

Primer sequences for PCR and descriptions of general laboratory techniques are provided in *SI Text*.

Population. We constructed a 4-generation pedigree to determine the relationship between 6 individuals diagnosed as dwarfs (Fig. S1). This pedigree consists of 26 individuals that are related to 2 founders. Carrier and dwarf phenotypes were determined based on production of dwarf calves and diagnosis of dwarfism by veterinarians. Of the 26 individuals, DNA was available for only 23 individuals.

Identification and Analysis of Positional Candidate Genes. Very few bovine scaffold sequences were available for the Angus dwarfism locus in the spring of 2005. Traditional comparative genomic approaches were used to overcome this problem. We used the human-bovine comparative radiation hybrid map (<http://cagst.animal.uiuc.edu/RHmap2004/index.html>) to identify positional candidate genes (21). Human gene sequence was used to BLAST query bovine gene sequence from the preliminary bovine genome draft. Because this process required repeated BLAST searches and manual assembly of segmented exon and UTR sequences, we created an automated pipeline called BEAP (BLAST Extension and Alignment Program) to expedite sequence assembly and reduce sequence handling errors. We used BEAP to query bovine sequences and perform an *in silico* chromosome walk using CAP3 to assemble retrieved sequences into contigs (22). We identified 20 genes in cattle by BLAST analysis within or closely surrounding the critical region.

Statistical Analysis/Elston Stewart Methodology. We used an extension of the Elston-Stewart algorithm in a model-based linkage analysis to map the genomic location most likely to contain the dwarfism locus (23, 24). The method was modified to make it more computationally feasible when pedigrees contain multiple loops to common ancestors. The implementation is described in the following citations (24–26). The implementation of linkage mapping has been described previously (27). The model for linkage mapping assumes the disease allele follows a recessive pattern of inheritance that is present at a low frequency in the Angus population. Briefly, likelihood ratios were calculated for each marker interval assuming (i) the dwarf mutation is at the center of this interval (L1) and (ii) the dwarf mutation is in another linkage group (L2). The log base 10 of this likelihood ratio (L1/L2) resulted in the LOD score. Likelihood (L) can be expressed as

$$L \propto \Pr(y) = \sum_G \Pr(y | G) * \Pr(G),$$

where $\Pr(y)$ is a vector of dwarfism phenotypes, and $\Pr(G)$ is a vector of genotypes at the markers flanking the interval and the dwarf mutation. A LOD score greater than 3 was deemed significant. We used multiple disease allele frequencies in the model for the founder population to determine the influence of disease allele frequency on LOD score.

Matings and Data Generated from a Breeding Experiment. Affected and unaffected full-sib calves were produced using embryo transfer. Two cows, mother (carrier) and daughter (dwarf), were mated to a bull known to be a carrier of dwarfism. There were more than 20 embryos produced for implantation into surrogate mothers. Only 6 embryos survived to full term, 3 from each mating. The 6 calves were raised under the guidelines of the Iowa State University Institutional Animal Care and Use Committee protocols. These 6 calves were within the normal size range for newborn Angus calves and not distinguishable as dwarfs at birth. A DNA sample was collected from each calf and tested blindly to determine the genotype for the *PRKG2 R678X* marker. Calves were weighed, and height and bone measurements were taken monthly to approximate lengths for the head, femur, metacarpal, metatarsal, tibia, and spine. In addition, circumferences were measured for head, chest, metacarpal, knee, and pastern. All measurements were performed in triplicate to reduce measurement error. At approximately weaning age, the calves were killed to collect tissues for functional analysis of the *PRKG2 R678X* mutation and histological study. Growth plate cartilage was collected from the tibia of each individual. Length measurements were taken for the femur, head, tibia, fused ulna/radius, metacarpal, and metatarsal.

Statistical Analysis of Long Bone Length. Analysis of long bones and measurements from the 6 calves produced in the breeding experiment, described previously, using PROC GLM (SAS Institute). For consistency and to avoid

overfitting of the model, each model was fitted with the same factors, regardless of the independent variable (bone length). For long bone lengths, the model was

$$y_{ij} = g_i + e_{ij},$$

where g_i denotes dwarfism genotype (yes or no), and e is the random error term. For live measurements (spine and height lengths), the model was

$$y_{ijkl} = p_i + t_{ij} + d_k + e_{ijkl},$$

where s denotes sex, and Gr is the birth group (encompassing age, dam, and common environment).

Cloning, Cell Culture, Transfection, Real-Time PCR and Analysis. We cloned the full-length bovine *PRKG2* cDNA into pGEMT-easy vector (Promega). The insert was then shuttled to the pcDNA3 expression vector (Invitrogen) by digestion with XbaI and Apal restriction enzymes (New England Biolabs) followed by ligation with T4 ligase (Promega). This bovine *PRKG2* vector was then mutated to contain the *PRKG2* exon 15 *R678X* mutation by site-directed mutagenesis using the QuikChange II kit (Stratagene). The resulting vectors were verified using restriction fragment length polymorphism (RFLP) and DNA sequence analysis. Cell culture was performed as described (7). Briefly, HUH-7 cells were transiently transfected using Lipofectamine 2000 (Invitrogen) and treated with 8Br-cGMP (Sigma) 4 h posttransfection. Cells were transfected with the following vector combinations (treatments): (1) WT bovine (bt) *PRKG2* + human (h) *SOX9*; (2) *R678X* bt*PRKG2* + h*SOX9*; (3) h*SOX9* + empty vector; and (4) empty vector (negative control). Treatments 1, 2, and 4 were replicated 4 times in 2 separate experiments, for a total of 8 replicates. Treatment 3 was replicated 4 times in a single experiment along with the other treatments. Cells were harvested 72 h posttransfection and total RNA was isolated with the Qiagen RNeasy Kit (Qiagen) according to manufacturer's protocol. RNA was reverse transcribed with SuperScript III (Invitrogen) for downstream amplification of cDNA in a real-time PCR that used Syber Green chemistry (Bio-Rad). *COL2* mRNA levels were evaluated using the primers F- CTTAGGC-CCGAGAGAGAAGG, R- ACTCAGGGTGGCAGAGTTTC, and the housekeeping gene *GAPDH*: F- TGATGACATCAAGAAGGTGGTGAAG, R- CCTTGAGGCCATGTAGCCAT. Real-time PCR was replicated 3 times for each treatment for both *COL2* and *GAPDH* expression.

In Vivo Real-Time PCR Analysis of Bovine Growth Plate Cartilage. Expression levels were measured in 4 *R678X* homozygote and 2 *R678X* carriers produced from the previously described breeding experiment. Total RNA from growth plate cartilage was analyzed by replicating each sample 3 times for the target and housekeeping gene in 2 separate plate replicates (i.e., 6 total replicates of both the target and housekeeping gene) using the same *COL2* primers, *COL10* primers F- AAT CTG AAA TGC AAG GTG CT, R- AAG ACT CAA ATA GTC ATT AAA GCA AA, *PRKG2* primers F- TGGTTTAAATTGGGAGGGACT, R- AAGCTTTGTCCAGCCTGA, and β -actin as the housekeeping gene, F- AA GGA CTC GTA CGT GGG GGA TGA, R- AA GGA CTC GTA CGT GGG GGA TGA. All genes were amplified using the same thermocycling protocol: (1) 95 °C for 10 min, (2) 95 °C for 15 s, (3) 60 °C for 30 s, (4) repeat steps 2 and 3, 40×, (5) 60 °C + 0.5 °C/cycle for 70 cycles, (6) end.

Statistical Analysis of Real-Time PCR. Standard curves for the reference gene were made and evaluated for all genes analyzed. Real-time PCR expression data were analyzed first using MyiQ software to normalize the data (Bio-Rad). The normalized expression data were then statistically analyzed using the natural log of the starting quantity of cDNA for transfection data and the ddCt method (28) for cattle expression data in PROC GLM (SAS Institute). The model used to analyze the cell culture transfection data were

$$y_{ijkl} = p_i + t_{ij} + d_k + e_{ijkl},$$

where y is the ln starting value of *COL2* mRNA adjusted for ln starting value of housekeeping gene mRNA (ln starting number *COL2* – ln starting number *GAPDH*), p is the real-time plate replicate, t is treatment 1–4 as described in Results, d is the experiment date, and e represents error. The P values calculated from the cell culture data were Bonferroni corrected to adjust for multiple testing (29). The model used to analyze the expression data from cattle growth plate was

$$y_{ijkl} = p_i + g_{ij} + e_{ijk},$$

where y is the ddCt, target gene (either COL2 or COL10) – housekeeping gene (β -actin), for the dwarf subtracted from the target gene – housekeeping gene, for a unaffected individual; p is the real-time plate replicate; g is the genotype, and e represents error.

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