A frameshift mutation in *GON4L* is associated with dwarfism in Fleckvieh cattle

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Abstract

Background

Low birth weight and postnatal growth restriction are the most evident symptoms of dwarfism. Accompanying skeletal aberrations may compromise the general condition and locomotion of affected individuals. Recently, several paternal half sibs with low birth weight and size have been noticed in the Fleckvieh cattle population.

Results

Affected calves were strikingly underweight at birth despite a normal gestation length and had craniofacial abnormalities such as elongated narrow heads and brachygnathia inferior. Despite a normal general condition, their growth remained restricted during rearing. We genotyped 27 affected and 10,454 unaffected animals at 44,672 SNP and performed association testing followed by homozygosity mapping to map the growth failure to a 1.85 Mb segment on bovine chromosome 3. Analysis of whole-genome re-sequencing data from one affected and 289 control animals revealed a 1bp deletion (g.15079217delC, rs723240647) in the coding region of the *GON4L* gene that segregated with the dwarfism-associated haplotype. We show that the deletion introduces intron retention and premature termination of translation, resulting in a severely truncated protein that lacks domains that are likely essential to normal protein function. The widespread use of an unnoticed carrier bull for artificial insemination has resulted in a tenfold increase in the frequency of the deleterious allele in the female population.

Conclusions

A frameshift mutation in *GON4L* is associated with autosomal recessive dwarfism in Fleckvieh cattle. The mutation has segregated in the population for more than fifty years without being recognized as a genetic disorder. However, the widespread use of an unnoticed carrier bull for artificial insemination caused a sudden accumulation of homozygous calves with dwarfism. Our findings provide the basis for genome-based mating strategies to avoid the

inadvertent mating of carrier animals and thereby prevent the birth of homozygous calves with impaired growth.

Background

Bovine stature is a prototypical complex trait that is controlled by a few loci with large effects and numerous loci with small effects. Genome-wide association studies using dense molecular markers revealed several quantitative trait loci (QTL) for growth-related traits in cattle [1],[2],[3]. The identified QTL account for a reasonable fraction of the phenotypic variation of bovine height [2],[4]. Sequence variants associated with mature height may also affect the size and weight of newborn calves [2],[3],[5],.

Birth size and weight varies between breeds, parities and the sex of the calf [6],[7]. The birth weight in Fleckvieh cattle typically ranges from 38 to 45 kg [8]. Calves with strikingly low birth weight and size despite normal gestation length are commonly referred to as "dwarfs".

Dwarfism (DW) has been observed in several cattle breeds including Fleckvieh [9],[10],[11]. Low birth size and postnatal growth restriction are the most apparent characteristics of DW. Undersized animals may be normally proportionated and have an undisturbed general condition (*i.e.*, proportional DW [12]). However, DW may also be accompanied by disproportionately shortened limbs and skeletal deformities (*i.e.*, disproportional DW, chondrodysplasia [13]). Depending on the severity of the structural aberrations, disproportionate DW may be fatal [14],[15].

Both autosomal recessive and dominant inheritance has been reported for bovine DW (*e.g.*, [12],[15]). Causative mutations for DW were identified in Angus (OMIA 001485-9913 [16]), Dexter (OMIA 001271-9913 [14]), Tyrolean Grey (OMIA 000187-9913 [13]), Holstein-Friesian (OMIA 001926-9913 [15]) and Japanese Brown cattle (OMIA 000187-9913 [17]). However, mutations causing DW have not yet been identified in Fleckvieh cattle.

Here we present the phenotypic and genetic characterization of autosomal recessive DW in Fleckvieh cattle. The use of genome-wide association testing, autozygosity mapping and massive re-sequencing data enabled us to identify a frameshift mutation in *GON4L* that is likely causal for the growth failure.

Methods

Animal ethics statement

Two animals were hospitalized at the animal clinic of Ludwigs-Maximilians-Universität München. Another two animals were pathologically examined at the Institute for Veterinary Disease Control (IVDC) of Austrian Agency for Health and Food Safety. One hospitalized calf was euthanized because of recurrent tympania with no prospect of improvement and subsequently necropsied. Tissue samples were collected during necropsy. All affected animals result from inadvertent carrier matings that happened in Fleckvieh farms. No ethical approval was required for this study.

Animals

Twenty-seven paternal half sibs (16 male, 11 female) with strikingly low birth weight and postnatal growth restriction were inspected by breeding consultants at the age of three weeks to 18 months. Ear tissue samples were collected by breeding consultants and DNA was prepared following standard DNA extraction protocols.

Genotyping, quality control and haplotype inference

Twenty-seven affected animals were genotyped with the Illumina BovineSNP50 v2 BeadChip interrogating genotypes at 54,609 SNP. The perindividual call-rate ranged from 98.96% to 99.60% with an average call-rate of 99.33%. In addition, genotypes of 10,454 unaffected Fleckvieh animals were

available from genotyping with the Illumina BovineSNP50 v1 BeadChip and the Illumina BovineHD BeadChip [18],[19]. The genotype data of cases and controls were combined and SNP that were present in both datasets were retained for further analyses. Following quality control (minor allele frequency above 0.5%, no deviation from the Hardy-Weinberg equilibrium (P>0.0001), per-SNP and per-individual call-rate higher than 95%), 10,481 animals (27 affected, 10,454 unaffected) and 44,672 SNP remained for association testing. The *Beagle* software [20] was used to impute sporadically missing genotypes and to infer haplotypes.

Haplotype-based association testing

A sliding window consisting of 25 contiguous SNP (corresponding to an average haplotype length of 1.42±0.43 Mb) was shifted along the genome in steps of 2 SNP. Within each sliding window, all haplotypes with a frequency above 0.5% (N=787,232) were tested for association with the affection status using Fisher exact tests of allelic association.

Generation of sequence data

Genomic DNA was prepared from a frozen semen sample of the supposed founder (DW_{het}) and from an ear tissue sample of one affected animal (DW_{hom}) following standard DNA extraction protocols. Paired-end libraries were prepared using the paired-end TruSeq DNA sample preparation kit (Illumina) and sequenced using the HiSeq 2500 instrument (Illumina). The resulting reads were aligned to the University of Maryland reference sequence of the bovine genome (UMD3.1 [21]) using the *BWA* software tool [22]. Individual files in SAM format were converted into BAM format using *SAMtools* [23]. Duplicate reads were marked with the MarkDuplicates command of *Picard Tools* [24]. We exploited sequence data from another 288 unaffected animals from nine cattle breeds (Gelbvieh, Nordic Finncattle, Fleckvieh, Holstein-Friesian, Brown-

Swiss, Original Braunvieh, Original Simmental, Red-Holstein, Ayrshire) that had been generated previously [25],[26].

Variant calling and imputation

Single nucleotide and short insertion and deletion polymorphisms were genotyped in DW_{hom}, DW_{het} and 288 control animals simultaneously using the multi-sample approach implemented in *mpileup* of *SAMtools* along with *BCFtools* [23]. *Beagle* phasing and imputation (see above) was used to improve the primary genotype calling by *SAMtools*. Larger insertions and deletions and structural rearrangements were identified in DW_{hom}, DW_{het} and 203 sequenced animals with an average genome fold coverage above 10x using the *Pindel* software package with default settings [27].

Identification of candidate causal variants

To identify mutations compatible with recessive inheritance, all polymorphic sites within the segment of extended homozygosity were filtered for variants that met three conditions: (i) DW_{hom} was homozygous for the alternate allele, (ii) DW_{het} was heterozygous and (iii) all control animals were homozygous for the reference allele. Candidate causal variants were annotated using the *Variant Effect Predictor* tool [28],[29]. Sequence variants of 1147 animals from Run4 of the 1000 bull genomes project [15] were analyzed to obtain the genotype distribution of candidate causal variants in a larger cohort.

Manual re-annotation of the bovine *GON4L*-gene

The genomic structure of *GON4L* (ENSBTAG00000020356) was predicted based on the University of Maryland (UMD3.1) bovine genome sequence assembly [21] and the Dana-Farber Cancer Institute bovine gene index release 12.0 [30] using the *GenomeThreader* software tool [31]. The *GenomeThreader*

output was viewed and edited using the *Apollo* sequence annotation editor [32].

Validation of candidate causal variants

PCR primers were designed to scrutinise the rs723240647 polymorphism by classical Sanger sequencing. Genomic PCR products were sequenced using the BigDye® Terminator v1.1 Cycle Sequencing Kit (Life Technologies) on the ABI 3130x1 Genetic Analyzer (Life Technologies). Genotypes for rs723240647 and rs715250609 were obtained in 3,879 and 1,848 Fleckvieh animals, respectively, using KASPTM (LGC Genomics) genotyping assays.

Clinical examination of affected animals

Two affected calves were admitted to the animal clinic at the age of 57 and 93 days. Initial examination (including weighing) was performed upon admission. The younger calf suffered from recurrent tympania and was euthanized four days after admission because of no prospect of improvement and subsequently necropsied. Tissue samples were collected during necropsy. The older calf was hospitalized for a period of 400 days. Weight records were collected once a week.

RT-PCR

Total RNA from lymph nodes, thymus, lung, heart, pancreas, liver, kidney and spleen of the euthanized animal was extracted from tissue samples using Trizol (Invitrogen) according to the manufacturer's protocol with some modifications. After *DNasel* treatment (Ambion), RNA was quantified using a NanoDrop ND-1000 (PeqLab) spectrophotometer, and RNA integrity determined by RNA Nano6000 Labchip (Agilent Technologies).

Complementary DNA (cDNA) was synthesized using the SuperScript IV

transcriptase (Thermo Fisher Scientific). *GON4L* mRNA was examined by RT-PCR using primers 1F – GAGTCAAGCAGCTCAAACCC and 1R - AGCCAAGTCAGTTTCTCCATT, which hybridize to exons 20 and 21 and amplify a 348 bp product based on the mRNA reference sequence (NCBI accession number: XM_010802911) of the bovine *GON4L*. The shorter version of exon 21 was amplified using reverse primer 2R – CTCAGACTCACCCTCCTGACTC. RT-PCR was performed in 20 ml reaction volumes containing diluted first-strand cDNA equivalent to 50 ng input RNA. PCR products were loaded on 2% agarose gels.

Results

Phenotypic manifestation of dwarfism

Twenty-seven calves (16 males and 11 females) with strikingly low birth weight (~15kg) and size despite normal gestation length were noticed among the descendants of an artificial insemination bull. Four affected calves were clinically and pathologically examined. At age 61, 97, 101 and 143 days, they were underweight at 42, 79, 53 and 51 kg, respectively. The calves had multiple craniofacial aberrations (*i.e.*, brachygnatia inferior, elongated narrow heads, structural deformities of the muzzle) and spinal distortions. Wrinkled skin, areas with excessive skin and a disproportionately large head became evident during rearing (Figure 1). Although the general condition, feed intake and locomotion of the animals were normal, their growth remained restricted. The average weight gain of an affected animal during a hospitalisation period of 400 days was only 450 g per day, *i.e.* less than half the weight gain of healthy Fleckvieh bulls (Figure 1H). Since both sexes were affected and most dams had a common ancestor, an autosomal recessive mode of inheritance was assumed.

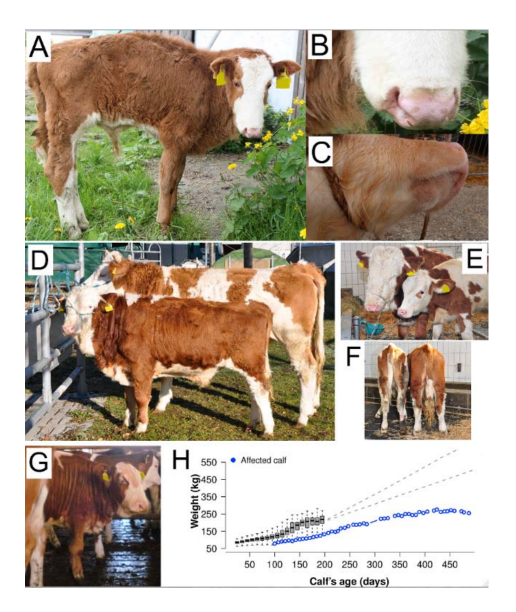


Figure 1 - Phenotypic manifestation of dwarfism in Fleckvieh cattle. A twenty-week old Fleckvieh calf with crooked back, structural abnormalities of the muzzle and brachygnathia inferior (A-C). A fifteen-month old Fleckvieh bull with dwarfism and a healthy coeval (D). Note the skin flaps in the neck area. The same animal with a six-month old healthy animal (E-F). Note the disproportionately large head of the affected animal (left) (E) although its height (right animal, height at withers: 112 cm) is similar to the nine-month younger healthy animal (left) (F). An approximately 18-month old animal with dwarfism with excessive skin in the neck area (G). The weight of one animal with dwarfism (blue) is compared to the weight of 74,422 Fleckvieh calves (grey boxes) (H). The upper dotted line represents the growth of Fleckvieh bulls observed in Geuder et al. [45]. The lower dotted line is a growth curve assuming an average weight gain of 1000 g/day, *i.e.*, a lower bound estimate for the growth of Fleckvieh bulls.

Dwarfism maps to chromosome 3

To identify the genomic region associated with DW, 27 affected and 10,454 unaffected animals were genotyped using a medium-density genotyping array. After quality control, 44,672 SNP were retained for genome-wide association testing. Haplotype-based association testing yielded a striking association with DW of a proximal segment of bovine chromosome 3 (Figure 2A). The most significant association signal (P=2.18 x 10⁻¹²⁴) resulted from two contiguous haplotypes located between 14,884,969 bp and 16,557,950 bp.

Autozygosity mapping revealed a common 1.85 Mb segment (14.88 Mb – 16.73 Mb) of extended homozygosity in 27 affected animals, corroborating recessive inheritance (Figure 2B). The common segment of extended homozygosity encompassed 71 transcripts/genes. However, none of them was previously associated with DW.

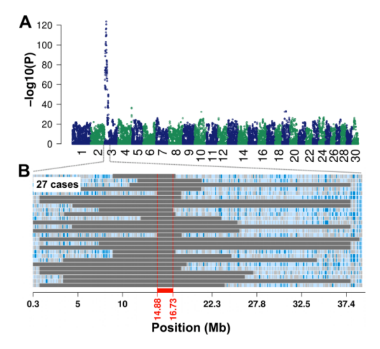


Figure 2 - Mapping of the genomic region associated with dwarfism. Association of 787,232 haplotypes with dwarfism in 27 affected and 10,454 unaffected animals (A). P-values were obtained by calculating Fisher exact tests of allelic association. Autozygosity mapping in 27 animals with dwarfism (B). Blue and pale blue represent homozygous genotypes (AA and BB), heterozygous genotypes (AB) are displayed in light grey. The solid grey bars represent segments of extended homozygosity in 27 animals with dwarfism. The red bar indicates the common segment of homozygosity.

Of 10,454 control animals, 81 were heterozygous and none was homozygous for the DW-associated haplotype, corresponding to a haplotype frequency of 0.38%. In the recent male breeding population (birth years 2000-2012), the frequency of the DW-associated haplotype was 0.25%. The haplotype frequency was considerably higher (2.6%) in the female population because of the widespread use of an unnoticed carrier bull for artificial insemination [33].

Haplotype and pedigree analysis enabled us to track the DW-associated haplotype back (up to twelve generations) to an artificial insemination bull (DW_{het}) born in 1959. DW_{het} was present in the maternal and paternal lineage of 21 affected animals. However, DW_{het} was not in the pedigree of six dams, possibly due to incomplete pedigree information and recording errors.

Identification of candidate causal variants for dwarfism

One affected animal (DW_{hom}) and DW_{het} were sequenced to an average read depth of 13x. To help identify the underlying mutation, we additionally exploited sequence data of 288 animals from nine breeds including 149 Fleckvieh animals. None of 149 sequenced control animals of the Fleckvieh population carried the DW-associated haplotype.

Multi-sample variant calling within the 1.85 Mb segment of extended homozygosity revealed 11,475 single nucleotide and short insertion and deletion polymorphisms as well as 3,158 larger structural variants. These 14,633 polymorphic sites were filtered for variants that were compatible with recessive inheritance that is homozygous for the alternate allele in DW_{hom}, heterozygous in DW_{het} and homozygous for the reference allele in 288 control animals (assuming the mutation is recessive and specific for the Fleckvieh breed). This approach revealed ten candidate causal variants for DW (Table 1). Five of them were intergenic, four were located in introns of *KCNN3*, *ADAR* and *TDRD10*, and one variant was located in the *GON4L* coding region.

Table 1: Ten sequence variants compatible with recessive inheritance

Chr	Position (bp)	NCBI	Type	Ref	Alt	Affected	Effect
		reference ID				gene	
3	15,079,217	rs723240647	Indel	С	-	GON4L	p.E1430Kfs66
3	15,713,943*	rs524337907	SNP	G	С	-	- -
3	15,713,959*	rs719431247	SNP	G	Α	-	-
3	15,737,755*	rs723848297	SNP	С	Τ	-	-
3	15,737,992*	ss1457237026	Indel	Τ	-	-	-
3	15,738,245*	rs720131431	Indel	С	-	-	-
3	15,815,016*	rs723370534	SNP	G	Α	KCNN3	Intronic
3	15,924,914*	rs717718209	SNP	G	Α	KCNN3	Intronic
3	16,046,490*	rs720952332	SNP	Τ	С	ADAR	Intronic
3	16,131,785	rs715250609	SNP	Τ	С	TDRD10	Intronic

The chromosomal position (base pairs) of compatible variants was determined according to the UMD3.1 assembly of the bovine genome. The asterisks indicate eight variants that are polymorphic among 1005 animals from 28 breeds other than Fleckvieh that had been sequenced for the 1000 bull genomes project. Ref - reference allele, Alt - alternate allele.

Eight of ten compatible variants were excluded as being causative for DW because they segregated in 1005 animals from 28 breeds other than Fleckvieh that had been sequenced for the 1000 bull genomes project [15] (Table 1). In conclusion, only an intron variant in *TDRD10* (rs715250609) and a coding variant in *GON4L* (rs723240647) segregated with DW. The intron variant in *TDRD10* is unlikely to be deleterious to protein function because it is more than 4000 bp distant from the most proximal splice site. We therefore considered the coding variant in *GON4L* (gon-4-like) as the most likely causal mutation for DW.

A 1bp deletion in GON4L is associated with dwarfism

Bovine *GON4L* consists of 31 exons encoding 2239 amino acids. The variant compatible with recessive inheritance is a 1bp deletion (rs723240647, g.15079217delC, ENSBTAT00000027126:c.4285_4287delCCCinsCC) in exon 20 (Figure 3A). Sanger sequencing confirmed that DW_{hom} and DW_{het} were homozygous and heterozygous, respectively, for g.15079217delC. The deletion introduces a frameshift in translation predicted to alter the protein sequence from amino acid position 1430 onwards, and a premature translation termination codon at position 1496 (p.Glu1430LysfsX66). The mutant protein is

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predicted to be shortened by 745 amino acids (33%) and lack domains that are likely essential for normal protein function (Figure 3B).

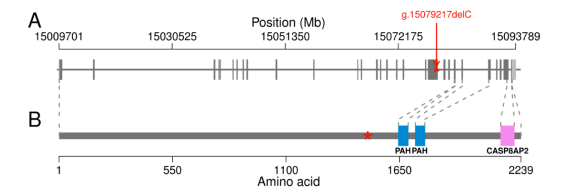


Figure 3 - A 1bp deletion in *GON4L* introduces a premature stop codon. Genomic structure of the bovine gon-4-like encoding gene *GON4L* (A). Grey and light grey boxes represent exons and untranslated regions, respectively. The red triangle represents a 1bp deletion (rs723240647, g.15079217delC) in exon 20 of *GON4L*. Gon-4-like consists of 2239 amino acids and contains highly conserved paired amphipathic helix (PAH) repeat and caspase 8-associated protein 2 myb-like (CASP8AP2) domains (B). The red star indicates the premature stop codon resulting from the 1bp deletion.

Genotypes for rs723240647 were obtained in cases and controls using customized genotyping assays (Table 2). rs723240647 was highly significantly associated with DW (P=3.72 x 10⁻⁸⁸). Twenty-four calves with DW were homozygous for the deletion variant, while 3,855 unaffected animals were heterozygous and homozygous for the reference allele. One animal that carried the DW-associated haplotype was homozygous for the reference allele possibly indicating a swapped DNA sample, haplotype recombination or imperfect genotype phasing. The intron variant in *TDRD10* (rs715250609) was in almost complete linkage disequilibrium (r²=0.98) with rs723240647 (Table 2).

Table 2: Genotypes for two mutations segregating with the DW-associated haplotype

Haplotype	Genotyp	e at rs723	240647	Genotype at rs715250609			
status	C/C	C/del	del/del	T/T	T/C	C/C	
non-carrier	3581	-	-	1737	3	-	
carrier	1	82	-	-	53	-	
homozygous	-	-	24	-	-	24	
unknown	180	11	-	31	-	-	

Genotypes at the rs723240647 and rs715250609 polymorphisms were obtained in 3,879 and 1,848 Fleckvieh animals, respectively, using custom KASP genotyping assays. The haplotype status of the animals was determined using genotypes from the Illumina BovineSNP50 BeadChip.

The deletion in GON4L creates intron retention and mRNA degradation

The effect of the q.15079217delC variant on GON4L transcription was examined by RT-PCR using RNA extracted from several tissues of a homozygous animal. Using primers located in exons 20 and 21, we obtained two RT-PCR products of 348 bp and ~310 bp from a wild type and a mutant homozygous animal. The longer PCR fragment corresponded to the reference mRNA sequence (NM 001192626) of the bovine GON4L gene. The ~310 bp PCR fragment showed a superimposed sequence of 35 bp at the 5' end of exon 21, suggesting the presence of an alternative GON4L transcript. The intensity of the alternative cDNA fragment was higher in the mutant homozygous than in the wild type animal, possibly indicating degradation of the mutant transcript in the homozygous animal. We designed a reverse RT-PCR primer specific for the alternative exon 21, and obtained a unique 348 bp RT-PCR product from the wild type animal and two RT-PCR products of 313 bp and ~1500 bp from the mutant homozygous animal (Figure 4). DNA sequence analysis of the 348 bp wild type RT-PCR product revealed that it corresponded to the mRNA reference sequence of the bovine GON4L gene. Sequence analysis of the longer fragment in the mutant homozygous animal revealed retention of intron 20. The length of the longer PCR fragment was 1488 bp. The retention of intron 20 is predicted to introduce a frameshift resulting in a

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premature translation termination codon at position 1492. In conclusion, the animal homozygous for the g.15079217delC variant contained both the premature translation termination codons at positions 1496 and 1492.

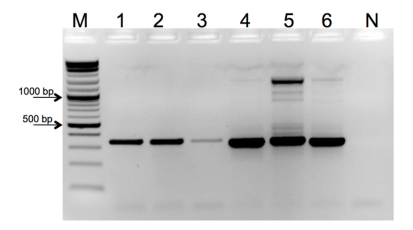


Figure 4 - Retention of intron 20 of the *GON4L* **gene in a Fleckvieh calf with dwarfism.** RT-PCR products were separated by 2% agarose gel electrophoresis. Lanes 1, 2, and 3 show 348 bp RT-PCR products from lung, lymph nodes, and liver of a wild type animal. Lines 4, 5, and 6 show 348 bp and 1488 bp RT-PCR products from lung, lymph nodes, and liver of a homozygous calf. M – size marker, N – negative RT-PCR control.

Discussion

A 1bp deletion in *GON4L* (g.15079217delC) was associated with DW in Fleckvieh cattle. The g.15079217delC variant introduces intron retention and premature translation termination resulting in a truncated protein. Compared to the wild type variant, the mutant GON4L protein is shortened by more than 30%. RNA analysis indicated that the mutant protein variant is less abundant, suggesting that it may be degraded *via* nonsense-mediated mRNA decay. If the truncated protein is (partially) retained, however, its function may be compromised because it lacks domains that are possibly essential for normal protein function. Loss-of-function variants in *Udu*, a gene that is highly homologous to *GON4L*, compromise cell cycle progression and response to

DNA damage and thereby disturb embryonic growth in *D. rerio* [34],[35],[36],[37]. In our study, the g.15079217delC variant was also associated with prenatal growth failure as evidenced by strikingly low birth weight of homozygous calves. The phenotypic manifestation of homozygosity for g.15079217delC, *i.e.*, pre- and postnatal growth restriction and craniofacial aberrations, resembles phenotypic patterns of human primordial DW resulting from DNA repair disorders [38],[39]. Such findings suggest that disturbed growth of homozygous animals might result from defective responses to DNA damage due to impaired GON4L function. However, the actual mechanism(s) and pathway(s) that cause the extremely low birth weight and postnatal growth restriction of homozygous animals have yet to be elucidated.

Congenital disorders that manifest as growth failure have been identified in several cattle breeds. Affected calves may be born underweight or fail to thrive during rearing [40],[41],[42],[26]. Homozygosity for g.15079217delC becomes evident at birth. Unlike mutations in *ACAN* and *COL2A1* that cause lethal disproportionate DW in cattle [14],[15], homozygosity for g.15079217delC is not fatal. Apart from large heads, affected animals were normally proportionate. Moreover, their general condition and locomotion was normal and their weight gain was constant, although considerably less than healthy animals. Thus, homozygosity for g.15079217delC is less detrimental than, *e.g.*, homozygosity for a mutation in *EVC2* which compromises both growth and locomotion of affected animals [13]. Nevertheless, homozygous animals are more likely to be culled at juvenile ages because of their reduced growth performance.

The g.15079217delC variant has segregated in the Fleckvieh population for more than fifty years, but due to its low frequency DW was rarely reported. Assuming a frequency of 0.2% of the deleterious allele, equal use of all bulls and 1,500,000 annual births in the German and Austrian Fleckvieh populations, one would expect only six homozygous calves with DW per year.

However, the widespread use of unnoticed carriers of rare recessive alleles in artificial insemination may cause a sudden accumulation of affected calves, as our study demonstrates. Twenty-seven calves with DW were descendants from a bull that was used for more than 290,000 inseminations. The heavy use of this carrier bull resulted in a more than tenfold increase in allele frequency in the female population [33]. Our findings now enable the rapid identification of carrier animals. The g.15079217delC variant was in almost complete linkage disequilibrium with the DW-associated haplotype, demonstrating a high sensitivity and specificity of haplotype-based identification of mutation carriers. Since all male breeding animals are routinely genotyped with dense genotyping arrays, carriers can be readily identified using haplotype information. Excluding carrier bulls from artificial insemination will prevent the emergence of homozygous animals and remove from the Fleckvieh population the rare DW-associated allele within a few generations. However, sophisticated strategies are required to simultaneously consider multiple deleterious alleles in genomic breeding programs while maintaining genetic diversity and high rates of genetic gain [43],[44].

Conclusions

A frameshift variant in *GON4L* was associated with autosomal recessive DW in Fleckvieh cattle. The deleterious allele persisted in the Fleckvieh population for more than fifty years at very low frequency without being recognized as a genetic disorder. However, the heavy use of an unnoticed carrier bull for artificial insemination resulted in an accumulation of homozygous calves with DW and a tenfold increase in frequency of the deleterious allele in the female population. Our results provide the basis for the rapid identification of carrier bulls and the implementation of genome-based mating strategies to avoid inadvertent carrier matings, thereby preventing the birth of homozygous calves with unsatisfactory growth performance.

List of abbreviations

DW: dwarfism; RT-PCR: reverse transcription polymerase chain reaction; SNP: single nucleotide polymorphism

Competing interests

The authors declare that they have no competing interests.

Authors' contribution

HP, and HS conceived the study; HP analyzed pedigree, genotyping and sequencing data; HS analyzed genotyping data and sampled affected animals; LM, MCL and GKS carried out clinical and pathological examinations; CW carried out the sequencing experiments; KF, SJ, ASch carried out the molecular genetic experiments; RF analyzed sequencing data; HP wrote the manuscript with the input from all authors. All authors read and approved the manuscript.

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