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Isolation and characterisation of the bovine *Stearoyl-CoA desaturase* promoter and analysis of polymorphisms in the promoter region in dairy cows.

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The nucleotide sequence data reported in this paper have been submitted to GenBank and has been assigned the accession number AJ555480.

1 ABSTRACT

2

3 Conjugated linoleic acid (CLA) in milk arises through microbial biohydrogenation of 4 dietary polyunsaturated fatty acids (PUFA) in the rumen, and by the action of 5 mammary Stearoyl-CoA desaturase (Scd). A large variation (up to ten-fold) in the 6 concentration of this fatty acid in milk have been observed, even in cows receiving the 7 same diet. The reasons for this variation are not well understood. In this study, the 8 bovine core promoter region was isolated by a genome walking strategy from 9 genomic DNA GenomeWalker libraries and then cloned and characterised. This core 10 promoter sequence extended approximately 600bp upstream of the translation start 11 site. The presence of putative transcription factor binding sites conserved in bovine, 12 human, and mouse promoters were observed. Evidence that this promoter fragment 13 was functional in vivo was obtained from expression studies in a mammary cell line. 14 The promoter sequence of the scd gene was compared between cows selected for the 15 ability to produce high fatty acid methyl esters (FAME) (2.22-2.72) in their milk, with 16 the same promoter region of low FAME producing cows (0.81-1.12). However, such 17 comparisons of the sequences of the *scd* promoter region of cows producing high milk 18 CLA compared with low CLA revealed no polymorphisms in this promoter segment. 19 Furthermore, no sequence polymorphisms were observed between the *scd* promoter 20 region of Holstein Friesian, Montbeliarde, Normande, Norwegian Red, Charlois, 21 Limousin and Kerry breeds.

22

23 (Key words: Conjugated Linoleic acid, Stearoyl-CoA desaturase, Promoter,
24 Polymorphisms).
25

Abbreviation key: CLA = Conjugated Linoleic Acid, scd = Stearoyl-CoA desaturase, PCR = polymerase chain reaction, SNPs = Single Nucleotide Polymorphisms, PUFA = Polyunsaturated Fatty Acids, FAME = fatty acid methyl esters.

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38 INTRODUCTION

39

40 Conjugated Linoleic acid (CLA) is a collective term to describe one or more positional and geometric isomers of linoleic acid (cis-9, cis-12-C_{18:2}). The cis-9, 41 42 *trans*-11- $C_{18:2}$ isomer is the predominant one in the human diet as a result of microbial 43 biohydrogenation in the rumen. This isomer is produced in ruminants directly as an 44 intermediate during the microbial biohydrogenation of dietary linoleic acid, and 45 endogeneously from trans-vaccenic (trans-11-C_{18:1}) acid in mammary tissue by the 46 action of Stearoyl-CoA desaturase (Scd) (Griinari et al., 2000). Animal-fat containing 47 foods including dairy products, beef and lamb are rich sources of CLA (Chin et al., 48 1992; Fritsche and Steinhart, 1998; O'Shea et al., 2000).

49

50 CLA has attracted much attention in recent years, due to its many potential health 51 benefits. Studies have shown that CLA exhibits anti-carcinogenic activity in animal 52 models (Belury et al., 1995; Ha et al., 1990; Ip et al., 1996; Liew et al., 1995) and in 53 in vitro studies using a range of human cancer cell lines including mammary (Miller 54 et al., 2001; Park et al., 2000; Visonneau et al., 1997), prostate (Cesano et al., 1998; 55 Palombo et al., 2002) and colon, (Miller et al., 2001; Palombo et al., 2002). Other 56 properties of CLA include anti-atherogenic activity (Lee et al., 1994; Nicolosi et al., 57 1993), the ability to reduce the catabolic effects of immune stimulation (Cook et al., 58 1993; Millar et al., 1994), the ability to enhance growth promotion (Chin et al., 1994) 59 and the ability to reduce body fat (Pariza et al., 1996).

60

During biohydrogenation of linoleic acid in the rumen, CLA is produced. This CLA is largely a transient intermediate and is involved as an intermediate in the biohydrogenation pathway to produce stearic acid. A build up of *trans*-vaccenic acid

64 occurs (Harfoot and Hazelwood, 1988). Trans-vaccenic acid is also an intermediate 65 in the biohydrogenation of other PUFA (Griinari and Bauman, 1999). Several studies have shown that substantial amounts of trans-vaccenic acid (60-300g/day) reach the 66 67 duodenum of lactating cows (Kalscheur et al., 1997a, 1997b; Wonsil et al., 1994). 68 Infusion studies using trans-vaccenic acid post-ruminally resulted in elevated cis-9, 69 trans-11 CLA in milk-fat, indicating conversion in the mammary gland (Griinari and 70 Bauman, 1999). Several studies have found substantial Scd activity in both mammary 71 and adipose tissue of ruminant animals (Kinsella et al., 1972; Martin et al., 1999; St 72 John et al., 1991, Ward et al., 1998). Mammary Scd enzyme, which converts trans-73 vaccenic acid to cis-9, trans-11 CLA is believed to be responsible for the formation of 74 the majority of CLA in milk (Griinari and Bauman, 1999).

75

76 Animal diet is a major factor affecting the milk-fat content of CLA, with dietary supplements containing oils rich in PUFA, such as linoleic and linolenic acids, being 77 78 the most effective for CLA enrichment of milk. However, substantial variations in the 79 CLA content of milk-fat (ranging from 3 to 10-fold) of cows on the same dietary 80 treatment have been observed in a number of studies (Jiang et al., 1996; Stanton et al., 81 1997; Kelly et al., 1998a; 1998b; Lawless et al., 1998; 1999; Peterson et al., 2002; 82 Solomon et al., 2000; White et al., 2001). The reasons for this variation are not well 83 understood. Perhaps ruminally derived CLA is a more important contributor to milk 84 CLA in grass-fed cows. The variation in the CLA content of milk-fat may also be due 85 to a variation in rumen conditions leading to differences in the availability of CLA (or 86 CLA precursors) that escape from the rumen. It might also be caused by differences in 87 mammary Scd activity associated with either regulation of scd gene expression, 88 differences in structure of the enzyme due to gene polymorphisms, or differences in downstream factors that would affect interaction between enzyme and substrate (e.g.
phosphorylation) (Peterson *et al.*, 2002).

91

92 The promoter region of the human (Bene et al., 2001; Zhang et al., 2001), chicken 93 (Lefevre et al., 2001), and mouse (Ntambi et al., 1988; Kaestner et al., 1989; Mihara, 94 1990) scd genes have been isolated, cloned and characterized. It has been shown in 95 these studies that there is a conserved PUFA response region in all three, and that this 96 includes critical binding sites for Sterol Response Element Binding Protein (SREBP) 97 and Nuclear Factor-Y (NF-Y) transcription factors. Sequence comparison of the 98 human and mouse promoters indicated a second region of high homology including 99 the 5'UTR and basal/proximal promoter. It has been suggested that there are two 100 different transcription start sites in the human promoter, and that these may be 101 dependent on tissue-specific factors (Zhang et al., 2001).

102

103 In this study, the role that polymorphisms in the core promoter region of the bovine 104 scd gene might play in influencing regulation of the scd gene was investigated. The 105 bovine core promoter was therefore isolated from genomic DNA by a genome 106 walking approach using primers designed to a genomic database sequence for the 107 bovine Stearoyl-CoA desaturase gene (AF481915) that contained 162 bases of 108 5'UTR. Analysis of this promoter sequence identified a number of conserved 109 potential transcription factor binding sites based on comparison with the human and 110 mouse scd promoter regions and interrogation of the TRANSFAC 4.0 database of 111 transcription factor sequences (Wingender et al., 2000) with the MatInspector V2.2 112 program (Quandt et al., 1995). Importantly, and for the first time for bovine scd, this 113 core promoter was shown to drive transcription of the reporter gene luciferase in an *in*

vitro mammalian culture system, confirming its presumed function. A number of
animals were screened for possible polymorphisms in this promoter region. These
included cows producing high milk-fat CLA as a % of fatty acid methyl ester (%
FAME) content (2.22-2.72), compared with cows yielding a low milk-fat CLA as a %
of FAME content (0.81-1.12). Additionally, sequence comparisons of the *scd*promoter region from a number of different breeds were made.
A longer bovine scd promoter sequence (Acc No. AY241932) is now included in the

121 Genbank database but this was not available at the time of this study: the two

122 promoter sequences were submitted within a very short time of each other.

123 MATERIALS AND METHODS

124

125 Construction of GenomeWalker (GW) libraries

126 To obtain the 5' flanking sequence of the scd gene, GenomeWalker (GW) libraries 127 were constructed from bovine total genomic DNA using a Universal GenomeWalker 128 kit (Clontech, UK) according to the manufacturer's instructions. High quality 129 genomic DNA was first extracted from cultured lymphoblast cells from a Holstein-130 Friesian animal. Briefly, a cell suspension (15 ml) was centrifuged at 1200 g for 5min 131 and the pellets mixed and incubated overnight at 37°C in 900 µl 0.2M EDTA 0.5% 132 sodium-n-lauroyl sarcosine and 25µl of proteinase K (20mg/ml). 10µl of RNase 133 (2mg/ml) was added to each tube and incubated at 37°C for 1.5 hr. The mix was split 134 in two, 200 µl phenol added, mixed and incubated for 30 min at 37°C in a rotary 135 mixer. Chloroform (200 µl) was added, tubes were shaken vigorously and incubated 136 at 37°C for 1 hr in a rotary mixer. Tubes were centrifuged at 16000 g for 15 min to 137 form two layers. The upper layer was transferred to a fresh tube and two volumes of 138 ice-cold 100% ethanol added. Tubes were inverted abruptly four times, and at this 139 point a DNA precipitate was obtained. The supernatant was decanted, 100 µl 70% 140 ethanol added and incubated overnight at 21°C in a rotary mixer. Ethanol was 141 decanted off and the pellet was allowed to air-dry. The DNA pellets were 142 resuspended in 50 µl 10mM Tris-HCl, pH8.0 containing 1mM EDTA (TE) buffer and 143 aliquots were checked for integrity on a 0.7% w/v agarose gel containing ethidium 144 bromide (EtBr) $(2\mu g/ml)$ gel.

The genomic DNA was digested with four restriction enzymes, *Dra*I, *Stu*I, *Eco*RV,
and *Pvu*II, to create four pools of DNA fragments. Adaptors were ligated onto both

148 ends of these fragments to create four GW libraries and these libraries were
149 designated DL1 (*DraI*), DL2 (*StuI*), DL3 (*Eco*RV), and DL4 (*PvuII*).

150

151 Isolation and characterisation of the *scd* promoter from GW libraries

To specifically isolate the scd promoter, genome walking primers pAKGW1 (5'-152 153 GCTCTCAGACACTGGGATCACTTTCTCGGG-3') (5'and pAKGW2 154 AACTGAGTGTAGACTAGTTCCTGAGCCTGC-3') were designed using Vector NTI software (InforMax, Inc.) to a genomic database sequence for the bovine scd 155 156 gene (AF481915) and synthesized by MWG Biotech (Ger). The primary PCR was 157 carried out using 1µl of the GenomeWalker libraries with the gene-specific primer, 158 pAKGW1, and adaptor primer, AP1 (from kit). The primary PCR products were 159 diluted 1:50 and used as template for nested PCR with a second gene-specific primer, 160 pAKGW2, and a second adaptor primer AP2 (again from kit). The reaction final 161 volume was 50 µl containing Taq DNA polymerase buffer (Invitrogen), 1.5mM 162 MgCl₂, 200µM dNTPs (Promega), 0.3µM each primer, and 1U Taq polymerase 163 (Invitrogen). Immediately prior to cycling 2.5 µl of DMSO was added to the mix. 164 The reaction was amplified for 35 cycles for both the first and second PCR. Cycling 165 was performed in a DNA Engine thermal cycler (MJ Research) and conditions were 166 95°C for 2min, followed by 35 cycles of 95°C for 30s, 60°C for 30s and 72°C for 3 167 min. This was followed by a 72°C final extension step for 7 min.

168

PCR products were directly cloned into the pCR2.1 vector (Invitrogen) and transformed using One Shot Top-10 chemically competent *E. coli* cells according to the manufacturer's instructions (Invitrogen). Colony PCR was performed to identify clones that potentially carried the *scd* promoter. Small-scale preparations of plasmid

DNA were made from these colonies using the procedure outlined by the
manufacturer (Sigma). Sequencing of three plasmids (DL1.2, DL1.4 and DL1.8) was
performed by MWG Biotech (Ger). The resulting sequences were analysed using
Vector NTI software (InforMax, Inc). Potential transcription factor binding sites were
identified using the MatInspector V2.2 program (Quandt *et al.*, 1995) by interrogation
of the TRANSFAC 4.0 database of transcription factor sequences (Wingender *et al.*,
2000).

180

181 Luciferase reporter vector construction

pCR2.1-based plasmids containing the putative *scd* promoter were restriction digested at 37°C to release the fragment and create compatible ends for sub-cloning into the pGL3-Basic (Promega) promoter-less expression vector. The pGL3-Basic vector was digested in the same manner. Ligation reactions were transformed into TAM Ultracomp chemically competent *E. coli* cells (Active Motif Europe), using instructions recommended by the manufacturer.

188

189 Cell culture

190 Chinese Hamster Ovary K1 (CHO-K1) cells obtained from ATCC (CCL-61) were 191 cultured in Dulbecco's Modified Eagles Medium F12 (Biowhittaker, UK) containing 192 10% (v/v) foetal bovine serum (Invitrogen). Human mammary MCF-7 cells (ATCC – 193 HTB22) were cultured in Eagles Minimum Essential Media (Invitrogen) containing 194 10% (v/v) foetal bovine serum (Invitrogen) and 1% (v/v) non-essential amino acids 195 Cells were routinely passaged every 3-4 days by washing with (Invitrogen). 196 phosphate buffered saline (PBS) and treating with 3ml EDTA-trypsin (Sigma) to 197 remove adherent cells. Cells were maintained in a humidified incubator at 37°C and

5% CO₂. Cells were grown to 80% confluency and both CHO-K1 and MCF-7 cells 198 199 were transiently co-transfected with luciferase reporter vectors using Fugene 6 200 transfection reagent (Roche Diagnostics). Six-well transfection plates were seeded at a concentration of 3 x 10^5 cells per well and incubated overnight at 37°C and 5% CO₂. 201 202 pGL3-Scd experimental constructs (1µg) were co-transfected with 25ng of pRL-SV40 203 plasmid (Promega) to control for transfection efficiency. Forty-eight hours after 204 transfection, media was removed from the wells, 300µl Passive Lysis Buffer 205 (Promega) was added followed by incubation at room temperature for 10min.

206

207 Luciferase assay

The Dual Luciferase Assay Kit (Promega) was used to measure both Renilla and Firefly luciferase expression by the reporter vectors on a Tecan Spectrafluor Plus luminometer using the Magellan software (Tecan). Mean Firefly luciferase activity values were corrected for variations in transfection efficiency using the corresponding mean Renilla luciferase figures. The corrected values were expressed as a percentage of the positive control value (pGL3-Control). Transfection values were a result of three independent transfections, with n=6, for both cell types.

215

216 Analysis of scd promoter polymorphism incidence

A dairy herd (n=75) had been on a ryegrass diet for a period of six months and their milk was analysed for fatty acid methyl esters content (% FAME) using the gas liquid chromatography (GLC) method described previously (Stanton *et al.*, 1997). FAME content was measured in milk samples from 59 animals taken at the evening milking on two occasions during the grazing season, in July and September. Nine of these 75 cows with consistently low and high milk CLA as a % of FAME values over both sampling times were selected for this study. The CLA isomer measured was the *cis*-9, *trans*-11 CLA isomer. High CLA animals produced greater than 2.0% FAME (2.222.72) in their milk, compared with low milk CLA producers of ~ 1.0% FAME (0.811.12). The CLA as a % of FAME ranged from 0.81 to 2.72 as quoted in Table I and
are the mean values of both sampling times.

228

229 Genomic DNA, for PCR amplification of the scd promoter fragment, was extracted 230 from whole blood (200µl) collected in heparinised tubes from the coccygeal vein from 231 these nine Holstein Friesian cows and in addition ten cows of different bovine breeds 232 using a Gentra capture column[™] (Gentra). Amplification of the bovine promoter was 233 performed using primers D9Dfor (5'-TGATGGGGTAGTGAGGAGC-3') and D9Drev (5'-GTTCCTGAGCCTGCTTTTGC-3') with 1µl of genomic DNA as 234 235 template (~200ng), in a final volume of 50µl, containing Taq DNA polymerase buffer (Invitrogen), 1.5mM MgCl₂, 200µM dNTPs (Promega), 0.3µM each primer, and 1U 236 237 Taq polymerase (Invitrogen). Cycling was performed in a DNA Engine thermal 238 cycler (MJ Research) and conditions were 95°C for 1min, followed by 35 cycles of 239 93°C for 1min, 58.5°C for 1min, 72°C for 2min and a final extension step of 72°C for 240 10min. PCR products were purified using a PCR purification kit (Qiagen). DNA was 241 eluted in 30µl of PCR grade water (Sigma).

243 **RESULTS and DISCUSSION**

244

245 Isolation of the bovine *scd* promoter

246 Isolation of the bovine *scd* promoter was achieved using a genome walking strategy, 247 which involved digestion of bovine genomic DNA with four restriction enzymes, 248 DraI, StuI, EcoRV, and PvuII, to create pools of short DNA fragments of varying 249 lengths. Adaptors were ligated onto the ends of these pools of DNA and PCR 250 employed to "walk" along the genome. Primers for PCR were designed to a bovine 251 genomic sequence for the scd gene (AF481915) that extends approximately 160bp 252 upstream of the translation initiation codon. PCR products were obtained for libraries 253 DL1 (DraI), DL2 (StuI) and DL3 (EcoRV), but not for the DL4 (PvuII) library. 254 Library DL1 yielded a mixture of fragments but as these were also the longest, these 255 fragments were purified as a mixed pool and cloned into the pCR2.1 vector. Colony 256 PCR was performed and three plasmids (DL1.2, DL1.4 and DL1.8) were sequenced. 257 BlastN analysis of the three sequences indicated that they were all fragments of the 258 bovine scd gene and the DL1.4 sequence extended upstream of the bovine genomic 259 database sequence, AF481915.

260

261 Comparison of bovine, human and mouse *scd* promoters

An alignment of the DL1.4 sequence was performed with a human promoter sequence (AF320307) (Zhang *et al.*, 2001) and a mouse promoter sequence (M21280) (Figure I). The bovine sequence displayed 67% identity with the human *scd* sequence and 59% homology with the murine promoter sequence *scd*1. The bovine sequence has been submitted to the Genbank database (Acc. No. AJ555480). Transcription start sites have been identified in both the mouse (Ntambi *et al.*, 1988) and human (Zhang *et al.*, 2001) genes at 28bp and 37bp respectively from the proximal TATA box in the bovine sequence. The bovine promoter sequence presented in Figure I is numbered relative to the human transcription start site, and extends to -407.

271

272 Potential transcription factor binding sites were compared with those predicted for the 273 human promoter (Zhang et al., 2001). Putative transcription-factor binding sites 274 identified using the MatInspector program (Quandt et al., 1995) included Nuclear 275 factor Y, Octamer binding factor 1, and muscle-specific Mt-binding sites. 276 Comparison of the mouse, human and bovine sequences identified a number of 277 conserved sequences in two particular regions of the promoters. The first area of 278 conservation occurs between -37 to -119, where there are two conserved TATA 279 sequences (5'-TTTAAAT-3' and 5'-TAAAA-3'), a fat-specific element (FSE) (5'-280 CTGAGGAAA-3') (-77 to -86), and binding sites for the transcription factors AP-1 (-281 64 to - 68), NF-1 (-102 to -109) and HNF4 (-116 to -119). Analysis of the human 282 SCD promoter has shown that for liver and hair follicles, there was a major 283 transcription initiation site 35 nucleotides downstream from the proximal TATA box 284 (Zhang et al., 2001). It was also shown that another transcription initiation site was 285 present 37 nucleotides downstream from the distal TATA box (Bene et al., 2001), and 286 it was suggested that the human SCD gene has different start sites that depend on 287 different tissue-specific factors (Zhang et al., 2001). The TATA sequence, 288 TTTAAAT, is somewhat unusual: this sequence, where the A in the second position 289 is replaced by a C, T, or G, has been shown to reduce the efficiency of a promoter in 290 in vitro studies (Conchino et al., 1983).

292 The second conserved region occurs at -313 to -390 and has been designated the 293 PUFA response region. Putative binding sites for transcription factors NF-Y (-313 to 294 -317), NF-1 (-333 to -336), NF-Y/NF-1 (-351 to -361), SREBP (-366 to -376), and 295 SP-1 (-386 to -390) are conserved between the three promoter sequences. The PUFA 296 response region present in the bovine sequence is an element previously shown to 297 mediate the down-regulation of mouse scd expression in response to PUFA (Waters et 298 al., 1997). The Sterol Response Element binding protein (SREBP) has been shown to 299 activate scd genes in the mouse (Tabor et al., 1999; Shimomura et al., 1998), and it 300 has also been shown that PUFA negatively regulate SREBP mRNA and protein 301 activation (Xu et al., 1999; Yahagi et al., 1999). This bovine promoter fragment also 302 contains the CCAAT box at -356 to -360 previously shown to be critical for 303 transcriptional activation in the human promoter (Zhang et al., 2001). This CCAAT 304 element is also required for full activation of mouse scd1 and scd2 promoters and 305 binds the NF-Y transcription factor (Tabor et al., 1999).

306

307 Transcription studies in CHO-K1 and MCF-7 cells

308 To determine whether this 407bp putative promoter sequence was sufficient to direct 309 transcription, luciferase reporter gene constructs containing the putative promoter 310 cloned into the promoter site of the promoter-less expression vector pGL3-Basic were 311 constructed. A plasmid (pRL-SV40) providing constitutive expression of Renilla 312 luciferase was co-transfected to serve as transfection efficiency control. Two cell 313 types were transfected, a human mammary cell line, MCF-7, and a Chinese hamster 314 ovary cell line, CHO-K1. The CHO-K1 cell line is most commonly used to express 315 mammalian genes in vitro and should contain the general transcription factors necessary for activating transcription of most genes. The human mammary cell line 316

317 contains more tissue-specific transcription factors, and is a more suitable system for318 studying regulation of a gene expressed in the mammary gland.

319

The putative promoter showed promoter activity in both CHO-K1 and MCF-7 cell lines (Figure II). Promoter activity was greatest in the mammary MCF-7 cell line being 6-fold higher in MCF-7 cells than in CHO-K1 cells. These results indicate that the 407bp region upstream of the proposed transcription start site is sufficient to direct transcription.

325

326 **Promoter polymorphism screen**

327 Animals receiving an identical dietary treatment of ryegrass for a 6-month period 328 were seen to have milk CLA as a % of FAME varying from 0.81 to 2.72. The sample 329 size selected was initially 59 Holstein Friesian cows, from which milk was analysed 330 for CLA content and subsequently 9 cows were selected and divided into two groups 331 of high and low CLA producers. There were no differences in the milk yields, feed 332 intake and milk fat content of the high v's low CLA animals. However, we did 333 observe that blood glucose and non-essential fatty acids (NEFA) differed significantly 334 between high and low CLA cows, with glucose being higher in the blood of the low 335 CLA group whereas NEFA was lower in the low CLA group compared to the high 336 CLA group. To investigate whether polymorphisms in the bovine *scd* promoter could 337 explain the considerably large inter-cow variation in milk fat CLA content, the 338 nucleotide sequences of the promoter region of nine cows (Table I) were analysed to 339 identify sequence polymorphisms which may be related to elevation of milk fat CLA.

In an attempt to also include an increased genetically diverse element, the promoter
region of the *scd* gene was amplified and sequenced from ten animals of different
breeds (high genetic merit Holstein Friesian, low genetic merit Holstein Friesian,
Montbeliarde, Normande, Norwegian Red, Irish Friesian, Dutch Freisian, Charlois,
Limousin, Kerry). No information regarding the milk CLA status of these animals
was available.

347

348 The scd promoter region sequences were aligned for these nineteen animals (nine 349 animals showing high and low milk CLA as a % of FAME and ten animals of varying 350 bovine breeds) and compared to search for any polymorphisms in the area (Data not 351 shown). This alignment showed the total absence of any polymorphic sites between 352 the bovine scd promoters of 19 (nine high and low milk CLA and ten animals of 353 different breeds). This high conservation may be significant in indicating that the 354 regulation of this gene is under extremely rigid control by transcription factors, and a 355 high conservation is necessary for full activation.

356

357 CONCLUSION

358

359 In conclusion we have isolated and partially characterised the bovine *scd* promoter. 360 This promoter fragment showed 59% and 67% similarity to both the mouse and 361 human sequences, respectively. Differences in the CLA content of milk from these 362 animals were not a consequence of polymorphisms within the core promoter of this gene. The *scd* promoter sequence exhibits remarkable sequence conservation not just 363 364 across high and low CLA yielding Holstein Friesians but also across ten different 365 breeds. In contrast considerable differences were seen between human and mouse 366 sequences which included variation in non-binding regions and binding regions for

AP-1, SP-1, and NF-KB transcription factors. Consequently the observed variation in
the levels of milk CLA produced in the Holstein Friesian animals may be explained
by differences in ruminant synthesis of CLA or CLA precursors, polymorphisms in
the coding sequences of the bovine *scd* gene, or differences in the regulatory proteins
themselves, an area that requires further investigation.

372

373 ACKNOWLEDGEMENTS:

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- line, MCF-7, was a gift from Dr. C Curran, University College Hospital, Galway.

TABLES and FIGURES

377 Table I: High and low CLA animals.

Animal	Breed	% FAME
330	Holstein Friesian	0.81 +/- 0.49
385	Holstein Friesian	0.88 +/- 0.49
387	Holstein Friesian	1.04 +/- 0.49
44	Holstein Friesian	1.12 +/- 0.49
728	Holstein Friesian	2.22 +/- 0.49
67	Holstein Friesian	2.27 +/- 0.49
249	Holstein Friesian	2.32 +/- 0.49
376	Holstein Friesian	2.34 +/- 0.49
225	Holstein Friesian	2.72 +/- 0.49

Figure I: Alignment of Human and Mouse Stearoyl-CoA desaturase promoters with DL1.4 (Bovine) sequence.

408		-407 -390
409	Human	-GGGGGAGCGAGGAGCTG GCGGC AGAG GGAA C AGCAGATTGCG CCGA GCCAATGGCAA CG
410	Mouse	GGAGAGACGGAGAAGCTA GAGGC AGAG GGAA C AGCAGATTGCG CCTA GCCAATGGAAA AG
411	Bovine	-GGGGTAGTGAGGAGCTCCGCGGCAGAGAGGGAACATCAGATTGCGCCGAGCCAATGGCAACG
412		* * * *** **** * *********************
413		SP-1 KFAI SKEBF INF-1/INF-1/KFAI
414		-313
415	Human	GCAGGACGAGGT GGCA CCAAATTCCCTTCGG CCAAT GACGAGCCGGAGTTTACAGAAGCC
416	Mouse	GCAGGACAAGGT GGCA CCAAATTCTCTTTGG CCAAT GACAAGACGGGCTTCACAGGAGGC
417	Bovine	GCAGGACGAGGT GGCA CCAAATTCCCCTTCGG CCAAT GACGCGCCAGAGTCTACAGAAGCC
418	DOVINC	****** ********************************
110		NF-1 NF-Y/AP-1
120		
$\frac{120}{121}$	Humon	
421	Hullian	
422	Mouse	ACATTAGCATTTATCCCCAGGCAGGGGGTTGGAGCAGCGCCCCTGTTGATGCCTTCAGC
423	Bovine	-CATTAGCATTTCCCCAGGGGCAGGGG-CAGAGGCAGGGGCTGCGGCGGCCAA
424		********* ** ******* * ***** * *****
425		
426		
427	Human	GTGTCGGTGTCGGCAGCATCCCCGGCGCCCTGCTGCGGTCGCCGCGAGCC
428	Mouse	ATCCCGGCGCCTCCAAGGTCTACTCTGGAATCTACTTGGCTTTCTTT
429	Bovine	GCCGCGGTGTGTGTGCAGCATCCAGTTCTTGCTTCTTCGGCCCCCAG-C
430		*** * * * * * * *
431		
432		
433	Human	TCGGCCTCTGTCTCCTCCCCCTC
434	Mouse	CCGCCCTCTCTCTCCCCCCCCCCCCCCCCCCCCCCCCCC
435	Bovine	ACGCCTCGGCGCTCTGTCTCCTC
436		** * *** * ***
437		
438		-119
439	Human	
440	Mouse	ĊĊŢĊĊĊŢĊĂĊĊŢĊŎĊĠĊĊŢĠĊĊŢŢĊĊŢŢĊĠĊŢĂĠĊŢĂŢĊŢĊŢĠĊĠĊŢ ĊŢŢŢ ŎĊĊŢŢŢ ŢĠ
441	Bowine	
442	DOVINC	** * ** * *** ** * ** * * * * *** ****
443		HNF4
$\Lambda\Lambda\Lambda$		
115	Humon	
116	Hullian	
440	Mouse	
447	Bovine	TTGGCAACGAATAAAAGAGGTCTGAGGAAATACGGGGACACAGTCA-CCCCCTGCCAGCGC
448		NF_1 ΤΔΤΔ FSF ΔP_1
449		-37
450		•
451	Human	TAGCC TTTAAAT TCCCGGCTCGGG-GACCTCCACGCACCGCGGCTAGCGCCGAC A ACCAG
452	Mouse	TACCC TTTAAAA TCCCAGCCCAGGAGATCTGTGCACAGCCAGACCGGGCTGAAC A CCCAT
453	Bovine	TAGCC <u>TTTAAAT</u> CCCCCAGCATAGCAGGTCGGGTCCGGAC <u>A</u> CCGGT
454		** ***** *** *** *** * * * * * * * *
455		TATA
456		+1
457	Human	CTAGCGTGCAAGGCGCCGCGGCTCAGCGC-GTACCGGCGGGCTTCGAAACCGCA
458	Mouse	CCC G AGAGTCAGGAGGGCAGGTTTCCAAGCGCAGTTCCGCCACTCGCCTACACCAAC
459	Bovine	$\texttt{CCA} \underline{\texttt{G}} \texttt{CGCGCACCGTGCAGCGGAAGGTCCCCGAGCGCAGCGCTGCGGATC-CCCACGCAAA-}$
460		* * * * * * * * * * * * * * * * *
461		
462		
462 463	Human	GTCCTCCGGCGACCCCGAACTCCGCTCCGGAGCCTCAGCCCCCTGGAAAGTG
462 463 464	Human Mouse	GTCCTCCGGCGACCCCGAACTCCGCTCCGGAGCCTCAGCCCCCTGGAAAGTG GGGCTCCGGAACCGAAGTCCACGCTCGA-TCTCAGCACTG-GGAAAGTG
462 463 464 465	Human Mouse Bovine	GTCCTCCGGCGACCCCGAACTCCGCTCCGGAGCCTCAGCCCCCTGGAAAGTG GGGCTCCGGAACCGAAGTCCACGCTCGA-TCTCAGCACTG-GGAAAGTG



466 Figure II: Transient transfection of mammalian cell lines with promoter
467 constructs.



488 **FIGURE LEGENDS**

489

Figure I: Sequence alignment of the DL1.4 sequence with database entries for the 490 491 human (AF320307) and murine (M21280), scd1 promoters. Numbering is indicated 492 above sequence. Transcription factor binding sites are underlined and bold and these 493 sites are based on TRANSFAC 4.0 database search results and comparison of human 494 and mouse scd gene promoters. The critical CCAAT box from human studies is 495 overlined. Numbering is relative to the human transcription start site and is indicated 496 by +1. The transcription start site in the mouse promoter is indicated by the symbol \bullet . 497 Abbreviations: RFX-1=X-box-binding regulatory factor 1, SREBP=Sterol response 498 element binding protein, NF-Nuclear factor. 499 500 Figure II: Transient transfection of mammalian cells with the putative bovine scd

promoter sequence. (a) Luciferase assay following transient co-transfection of CHOK1 cells with promoter construct. (b) Luciferase assay following transient cotransfection of MCF-7 cells with promoter construct. Luciferase activities are given
as a ratio of Firefly to Renilla (pRL-SV40) values

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