

Myostatin is expressed in bovine ovarian follicles and modulates granulosal and thecal steroidogenesis

Article

Accepted Version

Cheewasopit, W., Laird, M., Glister, C. and Knight, P. G. (2018) Myostatin is expressed in bovine ovarian follicles and modulates granulosal and thecal steroidogenesis. Reproduction, 156 (4). pp. 375-386. ISSN 1741-7899 doi: https://doi.org/10.1530/REP-18-0114 Available at http://centaur.reading.ac.uk/79544/

It is advisable to refer to the publisher's version if you intend to cite from the work. See <u>Guidance on citing</u>.

To link to this article DOI: http://dx.doi.org/10.1530/REP-18-0114

Publisher: Society for Reproduction and Fertility

All outputs in CentAUR are protected by Intellectual Property Rights law, including copyright law. Copyright and IPR is retained by the creators or other copyright holders. Terms and conditions for use of this material are defined in the <u>End User Agreement</u>.

www.reading.ac.uk/centaur



CentAUR

Central Archive at the University of Reading

Reading's research outputs online

1	Myostatin is expressed in bovine ovarian follicles and
2	modulates granulosal and thecal steroidogenesis
3	Warakorn Cheewasopit ^{1,2} , Mhairi Laird ¹ , Claire Glister ¹ and Phil G Knight ¹
4	¹ School of Biological Sciences, Hopkins Building, University of Reading, Whiteknights,
5	Reading RG6 6UB, UK
6	² WC is now at Department of Biology, Ramkhamhaeng University, Bangkapi, Bangkok,
7	Thailand
8	
9	Correspondence: Phil G Knight, School of Biological Sciences, Hopkins Building,
10	University of Reading, Whiteknights, Reading RG6 6UB, UK
11	Email: p.g.knight@reading.ac.uk
12	
13	
14	Short title: myostatin and ovarian steroidogenesis
15	Keywords: GDF8, estrogen, androgen, ovary, cow

16 Abstract

17	Myostatin plays a negative role in skeletal muscle growth regulation but its potential
18	role in the ovary has received little attention. Here, we first examined relative
19	expression of myostatin (MSTN), myostatin receptors (ACVR1B, ACVR2B and
20	TGFBR1) and binding protein, follistatin (FST), in granulosa (GC) and theca (TC) cells
21	of developing bovine follicles. Secondly, using primary GC and TC cultures, we
22	investigated whether myostatin affects steroidogenesis and cell number. Thirdly, effects
23	of gonadotropins and other intrafollicular factors on MSTN expression in GC and TC
24	were examined. MSTN, ACVR1B, TGFBR1, ACVR2B and FST mRNA was detected
25	in both GC and TC at all follicle stages. Immunohistochemistry confirmed follicular
26	expression of myostatin protein. Interestingly, MSTN mRNA expression was lowest in
27	GC of large estrogen-active follicles while GC FST expression was maximal at this
28	stage. In GC, myostatin increased basal CYP19A1 expression and estradiol secretion
29	whilst decreasing basal and FSH-induced HSD3B1 expression and progesterone
30	secretion and increasing cell number. Myostatin also reduced IGF-induced progesterone
31	secretion. FSH and dihydrotestosterone had no effect on granulosal MSTN expression
32	whilst insulin-like growth factor and tumour necrosis factor-alpha suppressed MSTN
33	level. In TC, myostatin suppressed basal and LH-stimulated androgen secretion in a
34	follistatin-reversible manner and increased cell number, without affecting progesterone
35	secretion. LH reduced thecal MSTN expression whilst BMP6 had no effect.
36	Collectively, results indicate that, in addition to being potentially responsive to muscle-
37	derived myostatin from the circulation, myostatin may have an intra-ovarian
38	autocrine/paracrine role to modulate thecal and granulosal steroidogenesis and cell
39	proliferation/survival.

2

40 Introduction

41 Ovarian follicle development is dependent on the actions and interactions of systemic 42 and intra-ovarian regulatory signals. Whilst pituitary gonadotrophins (FSH, LH) are the 43 key endocrine signals driving follicle development, a complex array of locally-produced 44 growth factors also contribute to the modulation of follicular somatic cell proliferation 45 and differentiation, 'initial' and 'cyclic' follicle recruitment, steroidogenesis, dominant 46 follicle selection and ovulation (Campbell et al. 2003, Webb et al. 2003). Prominent 47 amongst these are various members of the transforming growth factor- β (TGF- β) 48 superfamily including growth and differentiation factor-9 (GDF9), anti-mullerian 49 hormone (AMH), inhibins, activins and several bone morphogenetic proteins (BMP) 50 including BMP2, BMP4, BMP6 and BMP7 (Shimasaki et al. 2004, Knight & Glister 51 2006). In the present study we examined the potential involvement of another TGF- β 52 superfamily member, myostatin (also known as GDF8) in regulating ovarian follicle 53 function.

54 Myostatin is well-recognised for its negative autocrine/paracrine role in skeletal muscle 55 development (Otto & Patel 2010, Schiaffino et al. 2013). Myostatin-null mice show a 56 pronounced increase in muscle mass due to muscle fibre hyperplasia and hypertrophy 57 (McPherron et al. 1997). Naturally occurring inactivating mutations in the myostatin 58 gene are also evident in several species including bovine (Kambadur et al. 1997), ovine 59 (Clop et al. 2006), canine (Mosher et al. 2007) and human (Schuelke et al. 2004) and 60 these also display a phenotype of substantially increased muscle mass. Conversely, 61 upregulation of myostatin is associated with pathological conditions characterised by 62 muscle wasting, notably sarcopenia and cachexia arising from late-stage cancer, chronic 63 kidney failure and congestive heart failure (Elkina et al. 2011, Elliott et al. 2012).

64	Apart from skeletal muscle, myostatin has also been implicated in the regulation of
65	cardiomyocyte and adipocyte function (review: (Elliott et al. 2012)), Moreover,
66	investigations into the expression and potential functional role(s) of myostatin in
67	reproductive organs including the human ovary have recently been reported (Chang et
68	al. 2015, Fang et al. 2015, Chang et al. 2016a, Chang et al. 2016b).
69	Myostatin signals through the activin receptor type 2B (ACTR2B), forming a signalling
70	complex with ACVR1B (ALK4) and/or TGFBR1(ALK5) that activates an intracellular
71	Smad 2/3-dependent signal transduction pathway. Myostatin receptor activation can
72	also signal in a Smad-independent manner via activation of MAPK and inhibition of
73	Akt pathways (Rebbapragada et al. 2003). Binding of myostatin to its signalling
74	receptors is modulated by follistatin (Amthor et al. 2004). Follistatin was initially
75	identified as a secreted activin-binding protein but has since been shown to bind several
76	other TGF-β ligands including BMP-2,-4,-6 and -7 (Fainsod <i>et al.</i> 1997, Iemura <i>et al.</i>
77	1998, Glister et al. 2004). Follistatin-null mice show decreased muscle mass (Matzuk et
78	al. 1995) likely arising from diminished antagonism of myostatin signalling. Conversely,
79	transgenic overexpression of follistatin promotes a hypermuscular phenotype
80	resembling that of myostatin-null mice (Lee & McPherron 2001).
81	Global microarray studies of the bovine ovary revealed that myostatin mRNA is
82	expressed in follicular granulosa (Skinner et al. 2008, Glister et al. 2014, Hatzirodos et
83	al. 2014b) and theca cells (Glister et al. 2013, Hatzirodos et al. 2014a) although studies
84	to confirm expression and explore the potential functional role(s) of myostatin in the
85	bovine ovary have not been reported. Myostatin mRNA expression has also been
86	documented in human reproductive tissues including ovary (Chang et al. 2015),
87	myometrium (Islam et al. 2014) and trophoblast (Peiris et al. 2014) and recent evidence

88	from studies on luteinized granulosa cells supports various functional roles. For instance,
89	treatment of human granulosa-lutein cells with myostatin down-regulated expression of
90	steroidogenic acute regulatory protein (STAR) and reduced progesterone secretion,
91	whilst increasing cytochrome P450 aromatase (CYP19A1) expression, FSHR
92	expression and estradiol secretion (Chang et al. 2015, Fang et al. 2015, Chang et al.
93	2016a). An anti-proliferative effect of myostatin on human granulosa-lutein cells was
94	also reported (Chang et al. 2016b). To our knowledge, there have been no reports on
95	effects of myostatin on non-luteinized granulosa cells, nor on theca cells from any
96	species.
97	Given the paucity of information on the ovarian expression and possible intraovarian
98	role(s) of myostatin, particularly in relation to actions on non-luteinized follicular cells,
99	the aims of the present study were to: (1) examine mRNA expression profiles for
100	myostatin, its signalling receptors and binding protein (follistatin; FST) in granulosa
101	(GC) and theca (TC) cells across different stages of bovine antral follicle development;
102	(2) use non-luteinized bovine GC and TC culture models to investigate whether
103	myostatin affects steroid production; (3) determine whether the effect of myostatin can
104	be attenuated by follistatin; (4) investigate whether thecal and granulosal expression of
105	myostatin mRNA is modulated by gonadotropins and several intrafollicular factors
106	implicated in the regulation of follicular steroidogenesis.

107

Page 6 of 39

Materials and Methods

109 Relative expression of myostatin, follistatin and myostatin receptor mRNAs in

- 110 *developing bovine antral follicles.*
- 111 Relative mRNA expression for myostatin (MSTN), myostatin receptors (ACVR2B,
- 112 ACVR1B and TGFBR1) and follistatin (FST) in theca and granulosa layers from
- bovine antral follicles was determined using RT-qPCR. Ovaries from randomly cycling
- 114 cattle were obtained from an abattoir (Anglo Beef Processors, Guildford, UK) and
- selected for follicle dissection as described previously (Glister et al 2001; 2004; 2010).
- Briefly, antral follicles of diameter 3-18mm were dissected out and sorted by size into
- 117 small (3-6mm; n = 30), medium (7-10mm; n = 43) and large (11-18mm; n = 37)
- 118 categories. For each follicle GC and TC layers were retrieved for RNA extraction and
- follicular fluid recovered for steroid hormone analysis. Follicles in the large (11-18mm)
- 120 category were subdivided retrospectively into large estrogen-active (LEA; E:P ratio >1)
- and large estrogen-inactive (LEI; E:P ratio <1) categories according to their
- 122 intrafollicular ratio of estrogen to progesterone (E:P ratio). Isolated GC and TC were
- homogenised in 0.5ml of Tri reagent (Sigma UK Ltd, Poole) and stored at -80° C for
- subsequent RNA purification. The number of GC and TC RNA extracts recruited to the
- study (n = 82 GC samples; n = 87 TC samples; see fig. 1 for n-values for individual
- 126 follicle categories) was lower than the number of extracts processed because samples
- 127 indicating >5% GC/TC cross contamination were rejected during an initial quality
- 128 control screen. This involved a RT-qPCR-based comparison of relative transcript
- abundance of four GC/TC-specific 'marker' transcripts (FSHR and CYP19A1 for GC,
- 130 CYP17A1 and INSL3 for TC) each normalized to β -actin transcript abundance (data not

131 shown).

132 Primary granulosa and theca cell culture models

133	Ovaries from randomly cycling cattle were collected from a local abattoir. As described
134	previously (Glister et al. 2001, Glister et al. 2005) GC and TC were isolated from 4-
135	6mm diameter follicles, plated out in either 96-well (75,000 cells/well; for steroid
136	secretion experiments) or 24-well (250,000 cells/well; for RNA extraction experiments)
137	plates and cultured for 7 days. To preserve a non-luteinized cellular phenotype
138	(Gutierrez et al. 1997, Campbell et al. 1998, Glister et al. 2001, Glister et al. 2005,
139	Sahmi et al. 2006) chemically-defined serum-free media was used throughout the
140	culture period. This consisted of McCoy's 5A modified medium supplemented with 1%
141	(v/v) antibiotic-antimycotic solution, 10 ng/ml bovine insulin, 2 mM L-glutamine, 10
142	mM Hepes, 5 μ g/ml apotransferrin, 5 ng/ml sodium selenite, 0.1% BSA. In the case of
143	GC cultures, media was also supplemented with 10^{-7} M androstenedione as aromatase
144	substrate (all media and supplements were purchased from Sigma). Media were
145	replenished and treatments added on days 2 and 4 (see below). Cultures were terminated
146	on day 7 when conditioned media were retained for hormone assays and viable cell
147	number was determined by neutral red uptake assay as described elsewhere (Glister et
148	al. 2001)
149	Effects of myostatin on granulosal and thecal steroid secretion and viable cell
150	number
151	P acombinant human myostatin (P & D Systems: 0.4% amino acid sequence homology
151	Recombinant numan myöstatin (Red) Systems, 9470 annio acid sequence nomology
152	with bovine myostatin) was added to wells to give final concentrations of 0.08, 0.4, 2,
153	10, 50 and 100ng/ml in the presence and absence of gonadotropin (FSH or LH). Highly
154	purified ovine FSH (oFSH 19SIAPP) and LH (oLH-S-16) were provided by the NHPP
155	(Torrance, CA, USA). In GC cultures, FSH was used at a final concentration of 0.3

156	ng/ml, shown previously to elicit optimal estradiol secretion (Glister et al. 2001, Glister
157	et al. 2004). GC were also treated with myostatin (100ng/ml) in the presence and
158	absence of LR3 IGF-1 analogue (Sigma;10 and 50 ng/ml) since IGF-1 is also a potent
159	stimulator of estradiol secretion (Gutierrez et al. 1997, Glister et al. 2001). In the case
160	of TC cultures, LH was used at a final concentration of 150 pg/ml, shown previously to
161	elicit maximal androstenedione secretion (Glister et al. 2005). Control wells received an
162	equivalent volume of culture medium as vehicle.
163	
164	Can follistatin neutralize the effect of myostatin on thecal androstenedione secretion?
165	To examine whether follistatin can neutralize the suppressive effects of myostatin on
166	thecal androgen secretion, TC were treated with myostatin (100ng/ml) in the

167 presence/absence of recombinant human follistatin-288 (R&D systems; 96% amino acid

sequence homology with bovine follistatin) at 0.25 and 1.25µg/ml. These

- 169 concentrations were shown previously to reverse the effects of 50 ng/ml activin and
- 170 BMP6 on bovine GC (Glister *et al.* 2004).
- 171

172 Effect of myostatin on granulosal expression of steroidogenic pathway components

- 173 To evaluate the effects of myostatin on expression of key transcripts involved in
- steroidogenesis (CYP11A1, HSD3B1, CYP19A1, FSHR) GC were cultured in 24-well
- 175 plates (250,000 cells/well) and exposed to fixed concentrations of myostatin (100
- 176 ng/ml) in the presence and absence of an optimal concentration of FSH (300 pg/ml). At
- the end of culture, media were removed and cell lysates were prepared for total RNA
- 178 extraction and RT-qPCR analysis.

179	
180	Do gonadotropins and other factors modulate MSTN expression by cultured GC and
181	TC?
182	GC (n>4 independent batches of cells) plated out in 24-well plates were cultured in the
183	presence/absence of FSH (300 pg/ml) and several other intrafollicular factors shown
184	previously to modulate steroidogenesis at the concentrations used here, including LR3
185	IGF-1 analogue at 10 ng/ml (Glister et al. 2001), TNFα at 10 ng/ml (Glister et al. 2014)
186	and DHT at 100nM (Wu et al. 2011, Hasegawa et al. 2017). RNA was harvested at the
187	end of culture for evaluation of relative gene expression by RT-qPCR. TC (n=9
188	independent batches of cells) plated out in 96-well plates were treated with LH (150
189	pg/ml) in the presence/absence of BMP6 (10 ng/ml) shown previously to suppress
190	thecal androgen production (Glister et al. 2005, Glister et al. 2013).
191	

192 *RNA isolation, cDNA synthesis and real-time PCR*

193 Total RNA was isolated using Tri-reagent as described previously (Glister *et al.* 2010).

194 cDNA was synthesized from 1µg of RNA using the AB High Capacity cDNA synthesis

kit (Thermo Fisher Scientific; used according to manufacturers protocol) in a 20µl

reaction primed with random hexamers. PCR primers (see table 1) were designed using

197 Primer-BLAST' (http://www.ncbi.nlm.nih.gov/tools/primer-blast) with BLAST

198 specificity checking against all known bovine (Bos taurus) transcripts to exclude

- 199 potential amplification of off-target sequences. Primer pairs were also validated using
- agarose gel electrophoresis to demonstrate amplification of a single product of the
- 201 predicted size. Melt curve analyses was included in each PCR assay to confirm the

Page 10 of 39

202	amplification of a single product in each sample. cDNA template log-dilution curves
203	were used to demonstrate satisfactory PCR efficiency and linearity. PCR assays were
204	carried out in a volume of 14µl containing 5µl cDNA template, 1µl each forward and
205	reverse primers (final concentration $0.36\mu M$) and $7\mu l$ QuantiTect SYBR Green QPCR
206	2x Master Mix (Qiagen, Crawley, W. Sussex, UK). Samples were processed on a
207	StepOne Plus thermal cycler (Applied Biosystems) with cycling conditions: 15min at
208	95°C (one cycle only) followed by 15s at 95°C and 1min at 60°C for 40 cycles. The
209	$\Delta\Delta$ Ct method (Livak & Schmittgen 2001) was used to compare the relative abundance
210	of each mRNA transcript. Ct values for each transcript in a given sample were first
211	normalized to the corresponding β -actin Ct value (i.e. Δ Ct value). In the case of theca
212	and granulosa tissue samples ΔCt values for each transcript in a given sample were then
213	normalized to the mean ΔCt value for that transcript in all tissue samples. Resultant
214	$\Delta\Delta$ Ct values were converted to fold-differences using the formula: fold-difference = 2 (-
215	$^{\Delta\Delta Ct).}$ In the case of cell culture experiments ΔCt values were normalized to the
216	corresponding ΔCt value for vehicle-treated control cells. $\Delta \Delta Ct$ values were then
217	converted to fold-differences using the formula: fold-difference = $2^{(-\Delta\Delta Ct)}$.
218	

219 Steroid hormone assays

- 220 Steroid concentrations were determined by competitive ELISA as described previously
- 221 (Glister et al. 2010, Glister et al. 2013, Glister et al. 2014). The progesterone assay had
- a detection limit of 20pg/ml and intra- and inter-assay CVs were 8% and 10%
- respectively. The androstenedione ELISA had a detection limit of 30 pg/ml and intra-
- and inter-assay CVs were 7% and 10% respectively. The estradiol ELISA had a
- detection limit of 15 pg/ml and intra- and inter-assay CVs were 6% and 9% respectively.

226

227 Immunohistochemistry

228	Bovine ovaries were dissected into segments and fixed in formalin for 48 hours, before
229	being dehydrated through an alcohol series, embedded in wax and sectioned (5 μ m) onto
230	Superfrost charged slides (VWR, Lutterworth, UK). Sections were dewaxed and
231	rehydrated prior to boiling in citrate buffer (10mM citric acid, pH6.0), blocking of
232	endogenous peroxidase (3% H_2O_2 in methanol) and blocking of nonspecific binding
233	with 20% normal goat serum (NGS, Vector Laboratories Ltd, Peterborough, UK). After
234	this, sections were incubated overnight at 4°C in rabbit antibody against GDF8 (1:200;
235	sc-28910, Santa Cruz) diluted in 2% NGS. Control sections were incubated with normal
236	rabbit serum (1:200) diluted in 2% NGS. Primary antibody binding was detected using
237	biotinylated goat anti-rabbit diluted 1:250 in 2% NGS and Vector Elite ABC reagents
238	(Vector), prepared as per manufacturers instructions. Visualization of bound antibodies
239	was achieved using 3,3'-diaminobenzidine tetrahydrochloride (DAB; Vector), prior to
240	slides being counterstained with haematoxylin, dehydrated through an alcohol series
241	and mounted with coverslips using DPX mounting medium. Sections were imaged
242	using a Zeiss Axioscop 2 microscope and AxioCam digital camera.

243

244 Statistical analysis

245 Steroid concentrations were log-transformed prior to statistical analysis to reduce

heterogeneity of variance. RT-qPCR data were analysed as $\Delta\Delta$ Ct values (i.e. \log_2

247 values) before conversion to fold-difference values for graphical presentation of relative

transcript abundance. ACTB was used as the normalization control and showed uniform

249	expression level across experimental groups being compared. Results were evaluated
250	using one- and/or two-way ANOVA and, where indicated, post-hoc pairwise
251	comparisons were made using Fisher's protected least significant difference (PLSD) test.
252	Results of cell culture experiments are based on a minimum of three replicate
253	experiments using independent batches of cells (see figure legends for numbers of
254	replicates)
255	
200	
256	Results
257	
258	Relative expression of myostatin, follistatin and myostatin receptors in theca and
259	granulosa layers
260	Myostatin
261	MSTN mRNA expression was found in both TC and GC of all antral follicles examined
262	and overall expression level was higher in TC than GC (Figure 1A). Interestingly, while
263	MSTN expression level in TC was uniform across antral follicle development,
264	expression in GC fell ~15-fold to a nadir in large estrogen active (LEA) follicles.
265	However, a higher expression level was maintained in GC of large estrogen inactive
266	(LEI) follicle. (Fig. 1A). Immunohistochemistry confirmed myostatin protein
267	expression in both TC and GC of antral follicles (Fig. 2). In addition myostatin
268	immunoreactivity was evident in preantral follicles and in vascular smooth muscle cells.
269	Both oocytes and granulosa cells of primordial, primary and secondary follicles
270	exhibited positive immunostaining for myostatin (Fig 2)

271	Follistatin
272	FST mRNA expression was found in both TC and GC at all stages of follicle
273	development examined with much higher expression levels in GC than TC (Fig. 1B).
274	Interestingly, the expression of FST in GC sharply increased in LEA follicles but
275	remained low in LEI follicles; this was opposite to what was observed for MSTN.
276	Myostatin receptors (ACVR2B, ACVR1B and TGFBR1)
277	ACVR1B, TGFBR1 and ACVR2B mRNA expression was found in both TC and GC at
278	all stages of follicle development examined. The expression of ACVR2B and ACVR1B
279	was generally higher in GC than TC while TGFBR1 expression levels were broadly
280	similar in the two cell types. No notable changes in cell-specific patterns of expression
281	of these receptors between each stages of follicle development were evident (Fig.
282	1C,D,E respectively).
283	
284	Effect of myostatin on basal and FSH-induced steroid secretion by GC
285	Myostatin promoted a marked increase in basal estradiol secretion by cultured GC (~12-

- fold; P<0.0001; Fig. 3A) but did not modulate the >30-fold increase in estradiol
- secretion elicited by FSH. Myostatin suppressed both basal (P<0.01) and FSH-induced
- 288 (P<0.001) progesterone secretion (Fig. 3B). In addition, myostatin promoted a modest
- though significant increase in cell number under basal conditions (~20% increase;
- 290 P<0.001), but not under FSH-stimulated conditions (Fig. 3C).

291

292 Effects of myostatin on GC expression of steroidogenesis-related transcripts

293	The stimulatory action of myostatin on basal estradiol secretion was accompanied by a
294	~10-fold increase in CYP19A1 expression level (P<0.05; Fig. 3D). Concomitantly, a
295	reduction in CYP11A1 and HSD3B1 expression level was observed (P<0. 05; Fig. 3EF)
296	that mirrored the myostatin-induced decrease in progesterone secretion. Myostatin did
297	not affect FSHR expression (data not shown).
298	
299	Effect of myostatin on basal and IGF1-induced secretion of estradiol and
300	progesterone by GC
301	Fig.4 confirms the stimulatory effect of myostatin treatment (100 ng/ml) on basal
302	estradiol secretion by GC. However, myostatin did not modulate the stimulatory effect
303	of the LR3-IGF1 analogue on estradiol secretion or viable cell number. Myostatin
304	reduced both basal and IGF-induced progesterone secretion (P<0.05) but did not modify
305	the IGF-induced increase in viable cell number.
306	
307	Effects of FSH, LR3 IGF-1, TNF a and DHT on expression of MSTN mRNA by
308	cultured GC
309	Fig. 5 shows that treatment of cultured GC with FSH elicited a ~50-fold upregulation of
310	CYP19A1 expression (p<0.05) and estradiol secretion but did not affect MSTN
311	expression. Treatment with IGF-1 analogue also promoted a marked increase in
312	CYP19A1 expression (~10-fold; P<0.05) and estradiol secretion that was accompanied
313	by a 60% reduction in MSTN expression (P<0.05). Treatment with TNF α had no effect
314	on basal CYP19A1 expression but abolished FSH-induced upregulation of CYP19A1

- 315 expression and estradiol secretion. TNF α suppressed MSTN expression by ~80%
- 316 (P<0.05) under both basal and FSH-stimulated conditions. Treatment with DHT did not
- affect expression of either MSTN or CYP19A1.

318 *Effects of myostatin on thecal steroid secretion and viable cell number*

- 319 Myostatin suppressed androstenedione secretion in a dose-dependent manner (P<0.001)
- 320 with an IC₅₀ of \sim 10 ng/ml under LH-stimulated conditions (Fig. 6A). No effect of
- 321 myostatin on progesterone secretion was observed (Fig. 6B). Viable cell number was
- 322 increased (~25%; P<0.0001) by myostatin under both basal and LH-stimulated
- 323 conditions (Fig. 6C). LH increased both androstenedione and progesterone secretion but
- did not affect viable cell number.

325 Can follistatin neutralize the effect of myostatin on androstenedione secretion?

- 326 Treatment of cells with myostatin alone decreased androstenedione secretion by ~80%
- 327 (P<0.000; Fig. 7). Co-treatment with follistatin partially reversed this inhibitory action
- 328 (P<0.001). Treatment with follistatin alone tended to increase androstenedione secretion
- 329 but the effect was not statistically significant.
- 330

331 Effects of LH and BMP6 on MSTN mRNA expression by cultured TC

- Fig. 8 shows that treatment of cultured TC with LH elicited a 4-fold increase in
- 333 CYP17A1 expression and androstenedione secretion that was accompanied by a 40%
- 334 suppression of MSTN expression (p<0.05). Treatment with BMP6 profoundly
- suppressed basal and LH-induced CYP17A1 expression and androstenedione secretion.
- 336 Whilst BMP6 alone did not affect MSTN expression, it reversed the suppressive effect
- of LH on MSTN expression.

Page 16 of 39

338

339 **Discussion**

340

341	In this study, we first provide novel information on the spatio-temporal pattern of
342	mRNA expression of myostatin, its signalling receptors and the binding protein (FST),
343	at different stages of bovine antral follicles development. Expression of mRNA for
344	MSTN and its receptors was found in both GC and TC at all antral follicle stages
345	examined, consistent with and extending previous evidence from global microarray
346	studies (Skinner et al. 2008, Glister et al. 2013, Glister et al. 2014, Hatzirodos et al.
347	2014a, Hatzirodos et al. 2014b). Immunohistochemistry confirmed corresponding
348	expression of myostatin protein in follicular granulosa and theca interna layers of antral
349	follicles. Moreover, myostatin immunoreactivity was observed at earlier follicle stages
350	than those we analysed for mRNA expression, with positive staining in both oocytes
351	and GC of primordial, primary and secondary follicles and both GC and TC of late
352	preantral and early antral follicles. The inverse mRNA expression pattern of MSTN and
353	FST we observed in GC of large estrogen-active follicles is of interest since follistatin is
354	known to bind to and inhibit myostatin signaling (Lee & McPherron 2001, Amthor et al.
355	2004), a finding confirmed in this study by its ability to attenuate the effect of myostatin
356	on thecal androgen production. These results suggest, therefore, that GC-derived
357	myostatin and follistatin interact to regulate ovarian follicle physiology. In particular,
358	these observations suggest that autocrine/paracrine signalling by GC-derived myostatin
359	is attenuated in large healthy follicles (i.e. low myostatin/high follistatin), such as those
360	reaching the preovulatory stage of development. By contrast, at earlier antral follicle
361	stages (i.e. high myostatin/low follistatin), myostatin signalling via a Smad 2/3

362	dependent pathway may contribute to the suppression of thecal androgen production
363	whilst upregulating granulosal estradiol production and down-regulating progesterone
364	production. Thus, myostatin appears to act to prevent/delay premature follicle
365	maturation and luteinisation in a similar manner to that suggested previously for
366	activins and BMPs (Findlay et al. 2002, Knight & Glister 2006), both of which can
367	attenuate thecal androgen production, enhance granulosal estrogen output whilst
368	suppressing granulosal progesterone output.
369	The present results from experiments on non-luteinized ovarian cell models clearly
370	support the above with myostatin suppressing androgen secretion by theca cells. In the
371	case of granulosa cells, myostatin enhanced basal CYP19A1 expression and estradiol
372	secretion whilst suppressing CYP11A1 and HSD3B1 expression and secretion of
373	progesterone. In addition, treatment of human granulosa-lutein cells with myostatin was
374	recently reported to enhance FSH-induced upregulation of aromatase/estradiol
375	production, while inhibiting LH-induced upregulation of StAR/progesterone production
376	(Chang et al. 2016a). Moreover, the present study found that myostatin increased viable
377	cell number in both TC and GC cultures suggesting a positive effect on cell
378	proliferation and/or survival. This finding contrasts with a report that myostatin reduces
379	proliferation of human granulosa-lutein cells, evidently by upregulating connective
380	tissue growth factor expression (Chang et al. 2016b). The reason for this discrepancy is
381	not known but may reflect the effect of luteinisation, or a species difference.
382	An intrafollicular IGF system is firmly implicated in the autocrine/paracrine regulation
383	of follicle development, steroidogenesis and dominant follicle selection (Campbell et al.
384	1995, Glister et al. 2001, Silva & Price 2002, Webb et al. 2003). Like FSH, IGF-1 can
385	upregulate granulosal estradiol secretion; moreover, IGF-1 can augment follicular

Page 18 of 39

386	responsiveness to FSH, providing a potential mechanism for selecting the dominant
387	follicle from the cyclically-recruited growing cohort (Campbell et al. 1995, Webb et al.
388	2003). It was therefore pertinent to investigate whether myostatin affected the GC
389	response to IGF-1 treatment. Although the results showed no effect on IGF-induced
390	estradiol production or cell number, myostatin increased basal estradiol production and
391	cell number whilst reducing basal and IGF-induced progesterone production. As such,
392	these observations further support the notion that myostatin has a role to delay
393	premature follicle maturation and luteinisation.
394	Whilst circulating or intrafollicular concentrations of myostatin in cattle have not been
395	reported to our knowledge, serum concentrations of 10-20 ng/ml in cynomolgus
396	monkey and human, ~24 ng/ml in rat and ~80 ng/ml in mouse have been documented
397	(Furihata et al. 2016, Hedayati et al. 2016, Palandra et al. 2016). A myostatin
398	concentration of ~3 ng/ml has been reported for human follicular fluid (Chen et al.
399	2012). Since myostatin suppressed thecal androgen production and granulosal
400	progesterone production in vitro with an IC50 value of ~ 10 ng/ml, it seems plausible
401	that levels reaching the well-vascularized theca interna from peripheral blood could be
402	sufficient to exert a regulatory action, regardless of the additional 'local' contribution
403	(perhaps considerable?) of TC and/or GC-derived myostatin. On the other hand, given
404	the greater diffusional barrier needed to reach the avascular granulosal layer, combined
405	with the somewhat higher myostatin concentration (~50 ng/ml) needed to upregulate
406	GC estradiol production, it is possible that GC are primarily responsive to locally
407	produced myostatin acting in an autocrine/paracrine manner. The establishment of a
408	bovine myostatin assay to allow comparison of endogenous concentrations in peripheral

409	blood and ovarian follicular fluid of cattle in different physiological states and in
410	follicles at different stages of development, would be useful in this regard.
411	As a first step towards investigating which endocrine and local paracrine and/or
412	autocrine signals regulate myostatin expression in bovine ovarian follicles, we found
413	that an LH-induced increase in thecal CYP17A1 expression and androstenedione
414	secretion was accompanied by reduced MSTN expression level, consistent with a
415	negative autocrine/paracrine action of myostatin on thecal androgen production, and
416	with in the findings of our myostatin dose-response study. Indeed, it is possible that the
417	stimulatory action of LH on thecal androgen production could be due, in part, to LH-
418	induced suppression of myostatin expression. The finding of a reduced MSTN mRNA
419	abundance in TC producing more androgen could reflect increased androgen receptor-
420	mediated signalling since raised androgen levels are also associated with decreased
421	MSTN expression in rat skeletal muscle tissue (Mendler et al. 2007). However, another
422	intraovarian growth factor, BMP6, shown here and elsewhere (Glister et al. 2005,
423	Glister et al. 2013) to greatly reduce thecal CYP17A1 expression and androstenedione
424	secretion, did not affect thecal MSTN expression, casting doubt on androgen having a
425	direct effect. Furthermore, treatment of cultured GC with the potent non-aromatisable
426	androgen DHT had no effect on MSTN expression, suggesting an absence of androgen
427	receptor-dependent regulation of granulosal MSTN expression. Consistent with
428	previous findings (Gutierrez et al. 1997, Glister et al. 2001) treatment of GC with FSH
429	and IGF analogue both promoted substantial increases in estradiol secretion but only
430	IGF analogue modulated MSTN expression, eliciting a ~60% reduction. This suggests
431	a possible interaction between IGF and myostatin signalling at the intrafollicular level
432	that warrants further investigation. In skeletal muscle IGF-1 is a prominent positive

Page 20 of 39

433	regulator of muscle cell proliferation and differentiation whilst myostatin opposes this
434	action (Valdes et al. 2013). Despite this, IGF signalling upregulates myostatin
435	expression in skeletal muscle tissue models, suggesting an inhibitory auto-regulatory
436	loop (Yang et al. 2007, Kurokawa et al. 2009, Valdes et al. 2013).
437	The pro-inflammatory cytokine, $TNF\alpha$, is also expressed at the intraovarian level and is
438	implicated in the regulation of follicle and luteal growth/regression and steroidogenesis
439	(Sheldon et al. 2014, Samir et al. 2017). Consistent with earlier findings (Glister et al.
440	2014) we showed that TNF α abolished FSH-induced upregulation of CYP19A1 and
441	estradiol secretion by GC. This was accompanied by a marked reduction in MSTN
442	expression reinforcing the view that myostatin has a positive role in granulosal estrogen
443	production. In skeletal muscle models, activation of the TNF α pathway suppresses
444	myogenesis but upregulates myostatin expression (Ono & Sakamoto 2017). Moreover,
445	IGF can reverse the TNF- α induced suppression of myogenesis (Zhao <i>et al.</i> 2015)
446	indicating interactions between positive (IGF1) and negative (myostatin, TNF- α)
447	regulators of myogenesis. Further studies are needed to decipher the regulatory signals
448	that contribute to the regulation of myostatin expression by ovarian follicular cells and
449	to place these in a physiological context.
450	With respect to myostatin-null mice, there are few, if any, references to their ovarian
451	phenotype and the potential impact of the mutation on gonadal function and fertility is
452	unknown to us. However, an in vivo study involving active immunization of female
453	mice against myostatin, showed that the number of developing ovarian follicles in their
454	female progeny was \sim 50% lower than that of control mice, with a similar diminution in
455	litter size (Liang et al. 2007). Double-muscled cattle with myostatin mutations,

456 reportedly show delayed puberty, reduced female fertility and a higher incidence of

457	dystocia and perinatal calf mortality/morbidity is associated with the large size of calves
458	(McPherron & Lee 1997). However, we are not aware of any studies examining whether
459	perturbations in ovarian follicle dynamics or steroidogenesis occur in double-muscled
460	cattle. Whilst information is currently lacking on the above, it is possible that the
461	physiological actions of myostatin in the ovary are functionally redundant owing to
462	compensatory effects of other TGF- β ligands (e.g activins) that can signal via the same,
463	or overlapping, receptors to elicit similar regulatory actions on theca and granulosa cells.
464	In summary, this study provides novel information on the expression of myostatin, its
465	signalling receptors and the binding protein, follistatin, in theca and granulosa cells of
466	developing bovine antral follicles. Myostatin expression in GC declined to a very low
467	level in large estrogen-active follicles in which expression of follistatin was maximal,
468	suggesting attenuation of GC-derived myostatin signalling at this stage. Since myostatin
469	suppressed thecal androgen production in a dose-dependent manner, an effect partially
470	rescued by follistatin, it is hypothesised that attenuation of myostatin signalling in large
471	antral follicles could facilitate thecal androgen production required as a substrate for
472	granulosal aromatase enzyme and estrogen synthesis. Paradoxically, however,
473	myostatin was found to promote CYP19A1 expression and estradiol production by
474	granulosa cells under 'basal' conditions whilst suppressing CYP11A1 and HSD3B1
475	expression and progesterone production (see Fig. 9). Taken together, this suggests a role
476	for myostatin in delaying follicle progression towards pre-ovulatory maturation and
477	luteinisation, in a manner similar to that suggested for granulosa-derived activin
478	(Findlay et al. 2002, Knight & Glister 2006). Further in-depth studies in other species,
479	including whole animal models, are required to confirm and extend these in vitro
480	observations based on bovine ovarian cell culture models. It is also speculated that

481	muscle-derived myostatin conveyed to the ovary via the systemic circulation may
482	contribute to the regulation of follicle function. In a similar manner, testicular
483	steroidogenesis and gametogenesis may be influenced by circulating and/or locally-
484	produced myostatin although we are not aware of any studies, to date, examining this
485	possibility.
486	
487	Declaration of interests
488	The authors declare that there is no perceived conflict of interest that would prejudice
489	the impartiality of this scientific work
490	Funding
491	Supported by BBSRC (grant number BB/M001369 to PGK). WC was supported by a
492	postgraduate scholarship from the Thai Ministry of Science and Technology
493	Acknowledgements
494	We thank D Butlin and AD Simmonds for skilled technical assistance.
495	
496	References
497	
498	Amthor H, Nicholas G, McKinnell I, Kemp CF, Sharma M, Kambadur R & Patel
499	K 2004 Follistatin complexes Myostatin and antagonises Myostatin-
500	mediated inhibition of myogenesis. <i>Dev Biol</i> 270 19-30.
501	Campbell BK, Scaramuzzi RJ & Webb R 1995 Control of antral follicle
502	development and selection in sheep and cattle. J Reprod Fertil Suppl 49 335-
503	350.

504	Campbell BK, Baird DT & Webb R 1998 Effects of dose of LH on androgen
505	production and luteinization of ovine theca cells cultured in a serum-free
506	system. J Reprod Fertil 112 69-77.
507	Campbell BK, Souza C, Gong J, Webb R, Kendall N, Marsters P, Robinson G,
508	Mitchell A, Telfer EE & Baird DT 2003 Domestic ruminants as models for
509	the elucidation of the mechanisms controlling ovarian follicle development
510	in humans. <i>Reprod Suppl</i> 61 429-443.
511	Chang HM, Fang L, Cheng JC, Klausen C, Sun YP & Leung PC 2015 Growth
512	differentiation factor 8 down-regulates pentraxin 3 in human granulosa
513	cells. Mol Cell Endocrinol 404 82-90.
514	Chang HM, Fang L, Cheng JC, Taylor EL, Sun YP & Leung PC 2016a Effects of
515	growth differentiation factor 8 on steroidogenesis in human granulosa-
516	lutein cells. <i>Fertil Steril</i> 105 520-528.
517	Chang HM, Pan HH, Cheng JC, Zhu YM & Leung PCK 2016b Growth
518	differentiation factor 8 suppresses cell proliferation by up-regulating CTGF
519	expression in human granulosa cells. <i>Mol Cell Endocrinol</i> 422 9-17.
520	Chen MJ, Han DS, Yang JH, Yang YS, Ho HN & Yang WS 2012 Myostatin and its
521	association with abdominal obesity, androgen and follistatin levels in
522	women with polycystic ovary syndrome. <i>Hum Reprod</i> 27 2476-2483.
523	Clop A, Marcq F, Takeda H, Pirottin D, Tordoir X, Bibe B, Bouix J, Caiment F,
524	Elsen JM, Eychenne F et al. 2006 A mutation creating a potential
525	illegitimate microRNA target site in the myostatin gene affects muscularity
526	in sheep. <i>Nat Genet</i> 38 813-818.
527	Elkina Y, von Haehling S, Anker SD & Springer J 2011 The role of myostatin in
528	muscle wasting: an overview. J Cachexia Sarcopenia Muscle 2 143-151.
529	Elliott B, Renshaw D, Getting S & Mackenzie R 2012 The central role of
530	myostatin in skeletal muscle and whole body homeostasis. Acta Physiol
531	(<i>Oxf</i>) 205 324-340.
532	Fainsod A, Deissler K, Yelin R, Marom K, Epstein M, Pillemer G, Steinbeisser H
533	& Blum M 1997 The dorsalizing and neural inducing gene follistatin is an
534	antagonist of BMP-4. <i>Mech Dev</i> 63 39-50.
535	Fang L, Chang HM, Cheng JC, Yu Y, Leung PC & Sun YP 2015 Growth
536	Differentiation Factor-8 Decreases StAR Expression Through ALK5-
537	Mediated Smad3 and ERK1/2 Signaling Pathways in Luteinized Human
538	Granulosa Cells. Endocrinology 156 4684-4694.
539	Findlay JK, Drummond AE, Dyson ML, Baillie AJ, Robertson DM & Ethier JF
540	2002 Recruitment and development of the follicle; the roles of the
541	transforming growth factor-beta superfamily. <i>Mol Cell Endocrinol</i> 191 35-
542	
543	Furihata T, Kinugawa S, Fukushima A, Takada S, Homma T, Masaki Y, Abe T,
544	YOKOTA I, UDA K, UKITA K et al. 2016 Serum myostatin levels are
545	independently associated with skeletal muscle wasting in patients with
546	neart failure. Int J Caralol 220 483-487.
54/	Glister C, Lannetta DS, Groome NP & Knight PG 2001 Interactions between
548	follicie-stimulating hormone and growth factors in modulating secretion of
549	steroids and innibin-related peptides by nonluteinized bovine granulosa
550	cens. <i>Biol Reproa</i> 65 1020-1028.

551	Glister C, Kemp CF & Knight PG 2004 Bone morphogenetic protein (BMP) ligands
552	and receptors in bovine ovarian follicle cells: actions of BMP-4, -6 and -7 on
553	granulosa cells and differential modulation of Smad-1 phosphorylation by
554	follistatin. <i>Reproduction</i> 127 239-254.
555	Glister C, Richards SL & Knight PG 2005 Bone morphogenetic proteins (BMP) -4,
556	-6, and -7 potently suppress basal and luteinizing hormone-induced
557	androgen production by bovine theca interna cells in primary culture: could
558	ovarian hyperandrogenic dysfunction be caused by a defect in thecal BMP
559	signaling? Endocrinology 146 1883-1892.
560	Glister C, Satchell L & Knight PG 2010 Changes in expression of bone
561	morphogenetic proteins (BMPs), their receptors and inhibin co-receptor
562	betaglycan during boyine antral follicle development: inhibin can
563	antagonize the suppressive effect of BMPs on thecal androgen production.
564	<i>Reproduction</i> 140 699-712.
565	Glister C. Satchell L. Bathgate RA. Wade ID. Dai Y. Ivell R. Anand-Ivell R.
566	Rodgers RI & Knight PG 2013 Functional link between bone
567	morphogenetic proteins and insulin-like peptide 3 signaling in modulating
568	ovarian androgen production. <i>Proc Natl Acad Sci U S A</i> 110 E1426-1435.
569	Glister C. Hatzirodos N. Hummitzsch K. Knight PG & Rodgers RI 2014 The
570	global effect of follicle-stimulating hormone and tumour necrosis factor
571	alpha on gene expression in cultured bovine ovarian granulosa cells. BMC
572	Genomics 15 72.
573	Gutierrez CG, Campbell BK & Webb R 1997 Development of a long-term bovine
574	granulosa cell culture system: induction and maintenance of estradiol
575	production, response to follicle-stimulating hormone, and morphological
576	characteristics. <i>Biol Reprod</i> 56 608-616.
577	Hasegawa T, Kamada Y, Hosoya T, Fujita S, Nishiyama Y, Iwata N, Hiramatsu Y
578	& Otsuka F 2017 A regulatory role of androgen in ovarian steroidogenesis
579	by rat granulosa cells. <i>J Steroid Biochem Mol Biol</i> 172 160-165.
580	Hatzirodos N, Hummitzsch K, Irving-Rodgers HF & Rodgers RJ 2014a
581	Transcriptome profiling of the theca interna in transition from small to
582	large antral ovarian follicles. <i>PLoS One</i> 9 e97489.
583	Hatzirodos N, Irving-Rodgers HF, Hummitzsch K, Harland ML, Morris SE &
584	Rodgers RJ 2014b Transcriptome profiling of granulosa cells of bovine
585	ovarian follicles during growth from small to large antral sizes. <i>BMC</i>
586	<i>Genomics</i> 15 24.
587	Hedayati M, Nozhat Z & Hannani M 2016 Can the Serum Level of Myostatin be
588	Considered as an Informative Factor for Cachexia Prevention in Patients
589	with Medullary Thyroid Cancer? Asian Pac J Cancer Prev 17 119-123.
590	Iemura S, Yamamoto TS, Takagi C, Uchiyama H, Natsume T, Shimasaki S,
591	Sugino H & Ueno N 1998 Direct binding of follistatin to a complex of bone-
592	morphogenetic protein and its receptor inhibits ventral and epidermal cell
593	fates in early Xenopus embryo. <i>Proc Natl Acad Sci U S A</i> 95 9337-9342.
594	Islam MS, Catherino WH, Protic O, Janjusevic M, Gray PC, Giannubilo SR,
595	Ciavattini A, Lamanna P, Tranquilli AL, Petraglia F et al. 2014 Role of
596	activin-A and myostatin and their signaling pathway in human myometrial
597	and leiomyoma cell function. J Clin Endocrinol Metab 99 E775-785.

598	Kambadur R, Sharma M, Smith TP & Bass JJ 1997 Mutations in myostatin
599	(GDF8) in double-muscled Belgian Blue and Piedmontese cattle. Genome Res
600	7 910-916.
601	Knight PG & Glister C 2006 TGF-beta superfamily members and ovarian follicle
602	development. <i>Reproduction</i> 132 191-206.
603	Kurokawa M, Sato F, Aramaki S, Soh T, Yamauchi N & Hattori MA 2009
604	Monitor of the myostatin autocrine action during differentiation of
605	embryonic chicken myoblasts into myotubes: effect of IGF-I. Mol Cell
606	<i>Biochem</i> 331 193-199.
607	Lee SJ & McPherron AC 2001 Regulation of myostatin activity and muscle growth.
608	Proc Natl Acad Sci U S A 98 9306-9311.
609	Liang YC, Yeh JY & Ou BR 2007 Effect of maternal myostatin antibody on
610	offspring growth performance and body composition in mice. <i>J Exp Biol</i> 210
611	477-483.
612	Livak KJ & Schmittgen TD 2001 Analysis of relative gene expression data using
613	real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. <i>Methods</i> 25
614	402-408.
615	Matzuk MM, Lu N, Vogel H, Sellheyer K, Roop DR & Bradley A 1995 Multiple
616	defects and perinatal death in mice deficient in follistatin. Nature 374 360-
617	363.
618	McPherron AC, Lawler AM & Lee SJ 1997 Regulation of skeletal muscle mass in
619	mice by a new TGF-beta superfamily member. <i>Nature</i> 387 83-90.
620	McPherron AC & Lee SJ 1997 Double muscling in cattle due to mutations in the
621	myostatin gene. <i>Proc Natl Acad Sci U S A</i> 94 12457-12461.
622	Mendler L, Baka Z, Kovacs-Simon A & Dux L 2007 Androgens negatively
623	regulate myostatin expression in an androgen-dependent skeletal muscle.
624	Biochem Biophys Res Commun 361 237-242.
625	Mosher DS, Quignon P, Bustamante CD, Sutter NB, Mellersh CS, Parker HG &
626	Ostrander EA 2007 A mutation in the myostatin gene increases muscle
627	mass and enhances racing performance in heterozygote dogs. PLoS Genet ${f 3}$
628	e79.
629	Ono Y & Sakamoto K 2017 Lipopolysaccharide inhibits myogenic differentiation
630	of C2C12 myoblasts through the Toll-like receptor 4-nuclear factor-kappaB
631	signaling pathway and myoblast-derived tumor necrosis factor-alpha. <i>PLoS</i>
632	<i>One</i> 12 e0182040.
633	Otto A & Patel K 2010 Signalling and the control of skeletal muscle size. <i>Exp Cell</i>
634	<i>Res</i> 316 3059-3066.
635	Palandra J, Quazi A, Fitz L, Rong H, Morris C & Neubert H 2016 Quantitative
636	measurements of GDF-8 using immunoaffinity LC-MS/MS. Proteomics Clin
637	<i>Appl</i> 10 597-604.
638	Peiris HN, Salomon C, Payton D, Ashman K, Vaswani K, Chan A, Rice GE &
639	Mitchell MD 2014 Myostatin is localized in extravillous trophoblast and up-
640	regulates migration. J Clin Endocrinol Metab 99 E2288-2297.
641	Rebbapragada A, Benchabane H, Wrana JL, Celeste AJ & Attisano L 2003
642	Myostatin signals through a transforming growth factor beta-like signaling
643	pathway to block adipogenesis. <i>Mol Cell Biol</i> 23 7230-7242.

644	Sahmi M. Nicola ES & Price CA 2006 Hormonal regulation of cytochrome P450
645	aromatase mRNA stability in non-luteinizing boyine granulosa cells in vitro.
646	<i>I Endocrinol</i> 190 107-115.
647	Samir M. Glister C. Mattar D. Laird M & Knight PG 2017 Follicular expression of
648	pro-inflammatory cytokines tumour necrosis factor-alpha (TNFalpha).
649	interleukin 6 (II.6) and their receptors in cattle: TNFalpha, II.6 and
650	macrophages suppress the cal and rogen production in vitro. <i>Reproduction</i>
651	154 35-49
652	Schiaffino S. Dvar KA. Ciciliot S. Blaauw B & Sandri M 2013 Mechanisms
653	regulating skeletal muscle growth and atronhy FERS 1280 4294-4314
654	Schuelke M. Wagner KR. Stolz LE. Hubner C. Riebel T. Komen W. Braun T.
655	Tohin IF & Lee SI 2004 Myostatin mutation associated with gross muscle
656	hypertronhy in a child <i>N Engl I Med</i> 350 2682-2688
657	Sheldon IM Cronin IG Healey GD Gabler C Heuwieser W Streyl D Bromfield
658	II Mivamoto A Fergani C & Dohson H 2014 Innate immunity and
659	inflammation of the boying female reproductive tract in health and disease
660	Reproduction 148 R41-51
661	Shimasaki S Moore RK Otsuka F & Frickson GF 2004 The hone mornhogenetic
662	protein system in mammalian reproduction <i>Endocr Rev</i> 25 72-101
663	Silva IM & Price CA 2002 Insulin and IGF-Lare necessary for FSH-induced
664	cytochrome P450 aromatase but not cytochrome P450 side-chain cleavage
665	gene expression in oestrogenic hovine granulosa cells in vitro <i>I Endocrinol</i>
666	174 499-507
667	Skinner MK Schmidt M Savenkova MI Sadler-Riggleman I & Nilsson FF 2008
668	Regulation of granulosa and theca cell transcriptomes during ovarian antral
669	follicle development. <i>Mol Reprod Dev</i> 75 1457-1472
670	Valdes IA Flores S Fuentes FN Osorio-Fuentealha C Jaimovich F & Molina A
671	2013 IGF-1 induces IP3 -dependent calcium signal involved in the
672	regulation of myostatin gene expression mediated by NFAT during
673	myoblast differentiation. <i>I Cell Physiol</i> 228 1452-1463.
674	Webh R. Nicholas B. Gong IG. Campbell BK. Gutierrez CG. Garverick HA &
675	Armstrong DG 2003 Mechanisms regulating follicular development and
676	selection of the dominant follicle. <i>Reprod Suppl</i> 61 71-90.
677	Wu YG. Bennett I. Talla D & Stocco C 2011 Testosterone, not 5alpha-
678	dihydrotestosterone, stimulates LRH-1 leading to FSH-independent
679	expression of Cvp19 and P450scc in granulosa cells. <i>Mol Endocrinol</i> 25 656-
680	668.
681	Yang W. Zhang Y. Li Y. Wu Z & Zhu D 2007 Myostatin induces cyclin D1
682	degradation to cause cell cycle arrest through a phosphatidylinositol 3-
683	kinase/AKT/GSK-3 beta pathway and is antagonized by insulin-like growth
684	factor 1. <i>I Biol Chem</i> 282 3799-3808.
685	Zhao Q, Yang ST, Wang JJ, Zhou J, Xing SS, Shen CC, Wang XX. Yue YX. Song I.
686	Chen M et al. 2015 TNF alpha inhibits myogenic differentiation of C2C12
687	cells through NF-kappaB activation and impairment of IGF-1 signaling
688	pathway. Biochem Biophys Res Commun 458 790-795.
689	

690	
691	Table 1 List of primers used for real-time PCR
692	
693	
694	
695	Figure Legends
696	Fig. 1. Relative abundance of mRNA transcripts for (A) MSTN, (B) FST, (C)
697	ACVR1B, (D) TGFBR1 and (E) ACVR2B in theca and granulosa layers of small (3-
698	6mm), medium (7-10mm) and large (11-18mm) bovine antral follicles. Large follicles
699	are subdivided into estrogen active (E:P ratio >1) and estrogen-inactive (E:P ratio <1)
700	categories referred to as LEA and LEI follicles, respectively. Intrafollicular E:P ratios
701	for each follicle category are shown in panel F. Numbers in parenthesis in panel A are
702	n-values for each group. Values are mean \pm SEM and summarized two-way ANOVA
703	results are shown. Within each cell type means without a common letter are
704	significantly different (P<0.05).
705	
706	Fig. 2 Immunohistochemical staining of bovine ovary sections showing myostatin
707	immunoreactivity (brown) in oocyte and granulosa cells of primordial (pF) and primary
708	(PrF) follicles (A), secondary (SF) follicles (B,C) and in thecal (T) and granulosal (G)
709	layers of antral follicles (AF) (D,E). Myostatin immunoreactivity was also evident in
710	vascular smooth muscle cells (bv) (E). No staining was observed in control sections
711	treated with normal rabbit serum instead of primary antibody (F).
712	
713	Fig. 3 Effect of myostatin on basal and FSH-induced secretion of (A) estradiol and (B)
714	progesterone by bovine granulosa cells, and on (C) viable cell number; Panels (D-F)
715	show the effect of myostatin \pm FSH on expression of CYP19A1, CYP11A1 and
716	HSD3B1 mRNA, respectively. Values are means \pm sem (n = 5 independent cultures).

717	Results of 2-way ANOVA are summarized; *P<0.01, **P<0.01 ***P<0.001 compared
718	to respective control with zero myostatin (panels A, B, C). In panels D-F means without
719	a common letter are significantly different (P<0.05).
720	
721	
722	Fig. 4 Effect of myostatin on basal and LR3 IGF-1-induced secretion of (A) estradiol
723	and (B) progesterone by bovine granulosa cells and on (C) viable cell number. Values
724	are means \pm SEM (n = 3 independent cultures). Means without a common letter are
725	significantly different (P<0.05).
726	
727	Fig. 5 Effect of different treatments known to modulate GC steroidogenesis on
728	granulosal expression of (A) MSTN and (B) CYP19A1 and on (C) secretion of estradiol
729	Values are means \pm SEM (n = 4 independent cultures): Means without a common letter
730	are significantly different ($p<0.05$).
731	
732	Fig. 6 The effects of myostatin on basal and LH-induced secretion of (A)
733	androstenedione and (B) progesterone by bovine theca cells. Panel (C) shows effects on
734	viable cell number. Values are mean \pm SEM (n=12 independent cultures); Two-way
735	ANOVA p-values are shown
736	
737	Fig. 7 Ability of follistatin to antagonize myostatin-induced suppression of thecal
738	androstenedione secretion. Values are means \pm SEM (n=6 independent cultures)
720	
/39	
740	Fig. 8 Effect of LH and BMP6 on thecal expression of (A) MSTN and (B)
741	CYP17A1and on (C) secretion of androstenedione. Values are means \pm SEM (n = 8
742	independent cultures); means without a common letter are significantly different
743	(p<0.05).

744

- Fig. 9 Schematic diagram illustrating potential involvement of systemic and/or locally
- produced myostatin in the modulation of thecal and granulosal steroidogenesis.

Target	Accession number	Forward primer 5' to 3'	Reverse primer 5' to 3'	Amplicon size (bp)
LUCCD	NING 174201 1			02
LHCGK	NM_1/4381.1	ATIGUTUAGIUGATGUUAGAUU	AAAAAGCCAGCCGCGCTGC	92
STAR	NM_174189	TTTTTTCCTGGGTCCTGACAGCGTC	ACAACCTGATCCTTGGGTTCTGCACC	103
CYP11A1	NM_176644	CAGTGTCCCTCTGCTCAACGTCC	TTATTGAAAATTGTGTCCCATGCGG	99
HSD3B1	NM_174343.2	GCCACCTAGTGACTCTTTCCAACAGCG	TGGTTTTCTGCTTGGCTTCCTCCC	111
FSHR	NM_174061.1	GCCAGCCTCACCTACCCCAGC	AATTGGATGAAGGTCAGAGGTTTGCC	75
CYP17A1	NM_174304	GACAAAGGCACAGACGTTGTGGTCA	TGATCTGCAAGACGAGACTGGCATG	301
CYP19A1	NM_174365	TCTGTCCCCACTGAATCCTCCTGG	GGGTTTCATGGTGCTGTGTGGC	102
MSTN	NM_001001525.2	GTTCGATGTCCAGAGAGATGCCAGC	ACTTGCGTTAGAAGATCAGACTCCGTGG	114
ACTB	NM_173979.3	ATCACCATCGGCAATGAGCGGTTC	CGGATGTCGACGTCACACTTCATGA	128

 Table 1: List of primers used for quantitative RT-PCR

Fig 1









Fig 5

MSTN 1.2 Α а а 1 а ANOVA Relative transcript abundance Ī .8 P <0.001 b .6 .4 b b .2 T 0 FSH CON DHT IGF TNF TNF+FSH CYP19A1 ANOVA а 100 В P < 0.001 а T 10 b b b b 1 T T .1 DHT FSH CON IGF TNF **TNF+FSH** Estradiol secretion С а Estradiol (ng/ml) а Ι ANOVA T 10 P < 0.001 1 b bc bc С T .1 CON FSH DHT IGF TNF **TNF+FSH**







Fig. 9

