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Expression of PTHrP and PTHR (PTH/PTHrPr) mRNAs and polypeptides in bovine ovary and stimulation of bovine blastocyst development in vitro following PTHrP treatment during oocyte maturation.

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2	Expression of PTHrp and PTHR (PTH/PTHrP-r) mRNAs and Polypeptides in Bovine Ovary
3	and Stimulation of Bovine Blastocyst Development in vitro Following PTHrP Treatment
4	during Oocyte Maturation [*]
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18	

Abstract

20 Parathyroid hormone related protein (PTHrP) and its receptor have well established roles in the 21 development and regulation of many tissues, including bone and mammary gland. The objectives of 22 this study were: 1) to characterize the distribution of mRNAs encoding parathyroid hormone (PTH)-23 related protein (PTHrP) and receptor (PTHR) in bovine ovary; 2) to characterize the distribution of 24 PTHrP and PTHR polypeptides in bovine ovary; and 3) to examine the influences of PTHrP (1-141) 25 treatment during bovine oocyte maturation in vitro on blastocyst development. mRNAs encoding 26 PTHrP and PTHR were detected by *in situ* hybridization methods in oocytes, and granulosa cells in 27 all follicles from primordial to large antral. PTHrP and PTHR polypeptides displayed distinct 28 distribution patterns with PTHrP polypeptides primarily confined to oocytes from primordial to large 29 antral follicles. PTHrP polypeptides were detectable but at a reduced level in ovarian stroma and in 30 granulosa and thecal layers. PTHR polypeptides were detected in oocytes of all follicular stages but 31 were predominantly found in ovarian stroma, granulosa and theca follicular layers. Supplementation 32 of serum-free cSOFMaa oocyte maturation medium with PTHrP (1-141) resulted in a concentration-33 dependent increase in development to the blastocyst stage *in vitro*. The results suggest that granulosa 34 cells may be a primary site of PTHrP production and release. Oocytes from all follicular stages 35 stained strongly for PTHrP polypeptides and PTHrP enhanced development to the blastocyst stage in 36 vitro. 37 38 39

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Introduction

43 With the characterization of effective, defined and protein-free media to support the 44 maturation of mammalian oocytes *in vitro* it is now possible to examine the roles played by specific 45 hormone and growth factor modulators during oocyte maturation and upon subsequent early embryonic development (Saeki et al., 1991; Gardner, 1994; Eckert et al., 1995; Keskintepe and 46 47 Brackett, 1996; Hill et al., 1997; Avery et al., 1998; Krishner and Bavister, 1999; Watson et al., 48 2000). Parathyroid hormone related protein (PTHrP) was originally isolated from patients suffering 49 from humoral hypercalcemia of malignancy (Spiegel et. al., 1983; Strewler et. al., 1987; Stewart et. 50 al., 1987). As its name suggests, this molecule shares structural homology with parathyroid 51 hormone (PTH) (Stewart et. al., 1987; Horiuchi et. al. 1987), which enables PTHrP and PTH to 52 signal through a common PTH/PTHrP receptor (Rodan, et. al., 1988; Abou-Samra, et. al., 1992). 53 While PTH is predominantly (if not exclusively) produced by the parathyroid glands, PTHrP is 54 synthesized by many tissues including skin, brain, pancreas, adrenal glands, smooth muscle, heart, 55 lung, lactating breast, uterus, ovary and placenta (Urena et. al., 1993; Tian et. al., 1993; Lee et al., 56 1995; Weaver et. al., 1995; Downey et. al., 1997; Li et. al., 1995; Vasavada, et. al., 1998; Curtis et. 57 al., 1998; Ferguson et. al., 1998; Moseley and Gillespie, 1995; Gutmann et. al., 1993). PTHrP is 58 predominantly a paracrine/autocrine regulator of cellular growth and differentiation (Moseley and 59 Gillespie, 1995). The expression and distribution of gene products encoding PTHrP and the PTHR 60 have not been investigated within the bovine ovary. The objectives of the present study were: 1) to 61 characterize the distribution of mRNAs encoding parathyroid hormone (PTH)-related protein 62 (PTHrP) and receptor (PTHR) in the bovine ovary; 2) to characterize the distribution of PTHrP and 63 PTHR polypeptides in the bovine ovary; and 3) to examine the influences of PTHrP (1-141) 64 treatment during bovine oocyte maturation *in vitro* on blastocyst development.

65

Materials and methods

Detection of PTHrP and PTHR mRNAs in Bovine Ovary by in situ Hybridization 66 67 Bovine ovaries were collected at a local slaughterhouse and were quartered and placed 68 into 4% paraformaldehyde in phosphate buffered saline (PBS) pH 7.2 fixative for overnight 69 fixation. Ovarian pieces were then processed for routine paraffin embedding. In situ 70 hybridization (ISH) was carried out using biotin-labeled sense and antisense riboprobes and the 71 Genpoint CSA kit (Dako Diagnostics, Mississauga, ON, Canada) as described (Watson, et. al., 72 2000a; Watson, et. al., 2000b). Labeled riboprobes were prepared from plasmids containing 73 either a 2Kb cDNA corresponding to rat PTHR mRNA (Pausova, et.al., 1994; Amizuka, et.al., 74 1997) or a 0.7 Kb cDNA corresponding to rat PTHrP mRNA (Yasuda et.al., 1989) using T3 and 75 T7 RNA polymerases (Gibco, Burlington, ON, Canada) and a nucleotide labeling mix containing 76 biotin-16-UTP (Boerhinger-Mannheim, Burlington, ON, Canada). Tissue sections were 77 deparaffinized and rehydrated in a standard xylene and alcohol series and ISH performed 78 according to the manufacturer's directions. Briefly, slides were heated in Target Retrieval 79 Solution and proteinase K to undo protein cross-links caused by fixation. Endogenous peroxidase 80 activity was quenched in 0.3% hydrogen peroxide in methanol for 50 min at room temperature. 81 Sections were hybridized for 2 hours at 42°C with 5 ng/ml biotin-labeled riboprobe in the 82 supplied RNA hybridization buffer. After stringent washing at 55°C, sections were treated with 83 successive incubations in primary streptavidin-horse radish peroxidase, biotinyl-tyramide 84 solution and secondary streptavidin-horse radish peroxidase (Dako) for 15 minutes each at room 85 temperature. Color (golden-brown) was developed with the supplied diaminobenzidine diluted as directed (Dako) and sections counterstained with Carazzi's haematoxylin (blue) prior to
dehydration and mounting. We have employed this procedure using the same cDNAs to generate
riboprobes to localize PTHrP and PTHR mRNAs in rat tissues (including ovary) in previous
studies (Watson, et. al., 2000a; Watson, et. al., 2000b).

90

91 Immunocytochemical (ICC) Detection of PTHrP and PTHR polypeptides in Bovine Ovary

92 ICC was performed as described (Watson et. al., 1995; Watson et. al., 2000a; Watson et. al., 93 2000b). Briefly, tissue sections (as described above) were deparaffinized and rehydrated in a standard 94 xylene-ethanol series, post-fixed in 4% paraformaldehyde for 15 minutes and incubated for 10 95 minutes with 10 µg/ml proteinase K (Gibco, Burlington, ON, Canada) at room temperature to 96 retrieve fixation-concealed antigens. Sections were stabilized in 4% paraformaldehyde for a further 97 15 minutes before proceeding with the ICC procedure using the Vectastain ABC kit (Vector Labs, 98 Burlington, ON, Canada) following the manufacturer's directions. A polyclonal rabbit anti-rat PTHR 99 antibody was used: antibody PTH-IV raised against a portion of the first extracellular loop 100 (TLDEARLTEEELH; aa 249-262) (Babco, Berkeley, CA, USA) (Largo, Gomez-Garre, et. al., 1999). 101 The PTHrP antibody used was a mouse monoclonal raised against amino acids 34-53 of human 102 PTHrP (Oncogene Research Products, Cambridge, MA, USA). Both antisera are fully characterized 103 and their specificity for PTHrP and PTHR polypeptides is well-established (Watson et. al, 1999a; 104 Watson et. al., 1999b; Ferguson et. al., 1998, Largo et. al., 1999). Primary antibodies were diluted to 105 100 µg/ml in 1% BSA-PBS. As a control, some sections were treated with PTHR primary antibody 106 that had been pre-absorbed overnight at 4°C with the appropriate peptide. For the PTHrP antiserum, 107 control sections were treated with normal mouse serum instead of primary antibody. All sections

108 were counterstained with Carazzi's hematoxylin prior to dehydration and mounting.

109

110 Oocyte Collection, Insemination and Embryo Culture:

111 Cumulus oocyte complexes (COCs), were collected by razor blade slashing of slaughterhouse 112 ovaries within 4 h of removal from the animal (Wiemer et al., 1991; Watson et al., 1994). The COCs 113 were collected in oocyte collection medium (HEPES-buffered TCM-199 + 2% (v/v) NCS; Gibco 114 BRL, Burlington, ON, Canada) and then were washed 4 times in serum-free medium prior to 115 placement in oocyte maturation medium. Only denuded oocytes were discarded and a COC selection 116 strategy was not employed in this study. Following oocyte maturation (see below for specific 117 experimental conditions), oocytes were inseminated in vitro with frozen thawed bovine semen 118 (Semex Canada Inc., Guelph, ON, Canada) prepared using a "swim-up" method in Sperm TL 119 medium (HEPES-buffered modified Tyrodes solution as described in (Parrish et al., 1986). Matured 120 COCs were washed in sperm TL and placed in equilibrated fertilization drops (50 COCs/ 300 µl 121 drop) composed of bicarbonate-buffered modified Tyrodes solution under light paraffin oil (Parrish et al., 1986) BDH Inc., Toronto, ON, Canada). COCs and sperm (2.25x10⁵ motile spermatozoa/drop) 122 123 were incubated for 18h at 39°C under 5% CO₂ in air atmosphere before removal with a fine bore 124 glass pipette of the cumulus investment including all corona cells. Inseminated oocytes (40-50) were 125 placed into embryo culture consisting initially of 20 µl microdrops of citrate (0.5mM) and 126 polyvinylalcohol (PVA; 3 mg/ml) supplemented synthetic oviduct fluid medium (cSOFMaa) 127 (Keskintepe et al., 1995; Kestinkepe and Brackett, 1996; Watson et al., 2000) + 1X non-essential amino acids (NEA, Sigma-Aldrich Canada Ltd, Oakville, Ontario) and 1X essential amino acids 128 129 (MEM, Gibco, BRL) under paraffin oil in a humidified 5% CO₂ / 7% O₂ / 88% N₂ culture

atmosphere. Two days following initiation of culture, the microdrops were increased in volume by
addition of 20 µl of cSOFMaa medium. On days 5 and 7 of culture 20 µl of medium was removed
from each and replaced with 20 µl of fresh medium, thus keeping the microdrop volume constant for
the remainder of the eight day culture interval. Cleavage and blastocyst frequencies were assessed on
day 3 and 8 post insemination respectively.

135

136 Experimental Design:

A PTHrP concentration experiment (0, 10⁻¹⁰ M, 1.59 ng/ml; 10⁻⁸ M, 15.9 ng/ml; and 10⁻⁶ 137 138 M, 159 ng/ml PTHrP) was conducted employing a randomized design that allocated equivalent 139 numbers of non-selected COCs (total number of 442 COCs representing 4 experimental 140 replicates with 25-30 oocytes allocated to each treatment group per replicate) to gonadotrophin 141 and estradiol 17- β -free cSOFMaa oocyte maturation medium employing a 5% CO₂ in air 142 atmosphere at 39°C for 22 h. Following maturation, oocyte pools were inseminated and zygotes 143 were placed into culture for assessment of frequency of development to the blastocyst stage as 144 described.

145

146 Statistical Analysis:

147 Culture data were analyzed using the SigmaStat (Jandel Scientific) software package.

148 One-way analysis of variance (ANOVA), followed by pairwise multiple comparisons

149 (Bonferronis Method), were used for analysis of differences in the means for two or more

150 populations. Differences of P# 0.05 were significant differences.

Results

152	Detection of PTHrP and PTHR mRNAs in Bovine Ovary by in situ Hybridization
153	PTHrP and PTHR mRNAs were localized in bovine ovarian tissue with biotinylated
154	antisense riboprobes. Experiments were applied to tissue sections obtained from three bovine
155	ovaries collected at slaughter and all in situ experiments were conducted a minimum of three
156	times. In total over 100 follicles including stages from primordial to large antral follicles were
157	examined. mRNAs encoding PTHrP were detected by in situ hybridization methods in oocytes,
158	and granulosa cells in all follicles from primordial to large antral (Fig. 1 B, D, F, E). The
159	granulosa cells of antral follicles displayed more intense signals for PTHrP mRNA (Fig 1E). A
160	positive signal for PTHrP mRNAs was observed in thecal layers of larger follicles and in ovarian
161	stroma in general (Figure 1 B,D,F, E). The signal observed for PTHrP sense controls (Fig. 1 A,
162	C) was very low and indicated the specificity of the antisense PTHrP riboprobe. mRNAs
163	encoding PTHR were detected in oocytes and granulosa cells in all follicles from primordial to
164	large antral (Fig2 B,D,F, E). A positive signal for PTHR mRNA was observed in thecal layers of
165	larger follicles and in ovarian stroma (Fig. 2 B,D, E,F). Sense PTHR mRNA controls displayed a
166	very low background signal in granulosa and thecal layers (Fig. 2 A,C).
167	
168	Immunocytochemical Detection of PTHrP and PTHR polypeptides in Bovine Ovary
169	PTHrP and PTHR immunoreactivity was studied in intact bovine ovarian tissue sections
170	to assess the distribution of these polypeptides within the bovine ovary. PTHrP and PTHR
171	polypeptides displayed distinct distribution patterns with PTHrP polypeptides primarily confined

to oocytes from primordial to large antral follicles (Fig. 3 B,D,F). PTHrP polypeptides were also

173	localized to a lesser extent in ovarian stroma and reduced signals were obtained in granulosa
174	cell and thecal layers (Fig. 3 B,D, E, F). PTHrP immuno-specificity was determined by
175	demonstrating a very low background signal in normal mouse serum-treated tissue sections (Fig.
176	3 A,C). PTHR polypeptides were detected in oocytes of all follicular stages (Fig 4, B, D, F).
177	However, the most intense signals for PTHR polypeptides were localized to ovarian stroma,
178	granulosa cell and thecal follicular layers (Fig 4 B,D,E, F). Pre-absorbed antibody controls
179	indicated a high level of specificity for the PTHR immunolocalization signal (Fig 4A, C).
180	
181	Influence of PTHrP during Oocyte Maturation on Development of Bovine Zygotes In vitro
182	cSOFMaa culture medium employed for bovine oocyte maturation was supplemented
183	with PTHrP and influences on development to the blastocyst stage were examined. No significant
184	differences in cleavage were observed among the four oocyte maturation treatment groups (p<
185	0.05; Fig. 5 and 6). Likewise, development to the 6-8 cell stage was not significantly influenced
186	by PTHrP treatment during oocyte maturation (Fig. 5 and 6). However, development to the
187	blastocyst stage as assessed by the proportion of total oocytes inseminated (Figure 5) or as a
188	proportion of cleaved zygotes progressing to the blastocyst stage was significantly enhanced
189	(p<0.05) by the addition of PTHrP to the oocyte maturation medium. The beneficial influence of
190	PTHrP during oocyte maturation on blastocyst formation arose following the 6-8 cell stage since
191	PTHrP treatment during oocyte maturation lead to a significant increase (p<0.05) in the
192	proportion of 6-8 cell zygotes progressing to the blastocyst stage (Figure 6).
193	

DISCUSSION

195 This study characterized the distribution of mRNAs and polypeptides encoding PTHrP 196 and the PTHR within bovine ovary and investigated the influence of PTHrP treatment during in 197 vitro maturation of bovine oocytes on development to the blastocyst stage. The results indicate 198 that the bovine ovary expresses both the mRNAs encoding PTHrP and PTHR and their 199 polypeptides. The granulosa layer may be a primary site of PTHrP production (high mRNA 200 signals) but the majority of this PTHrP does not appear to be stored within the granulosa layer 201 (low polypeptide signal). Why the granulosa highly expresses PTHrP mRNA but contains little 202 peptide is unclear. It may be that granulosa PTHrP mRNA is not translated well (or at all) or even 203 that the PTHrP peptide is secreted to the antral cavity. The answer to these possibilities awaits 204 further experimentation. Oocytes from all follicular stages stained intensely for PTHrP 205 polypeptides. The cognate receptor for PTHrP, the PTHR, was expressed in both oocytes and 206 granulosa cells of the bovine ovary suggesting that PTHrP may act in both an autocrine and 207 paracrine fashion to regulate bovine follicle development. PTHrP may also be an important 208 regulator of oocyte maturation since addition of PTHrP to bovine oocyte maturation medium 209 enhanced the development of fertilized bovine eggs to the blastocyst stage in vitro. 210 PTHrP and its receptor, the PTHR, are widely expressed in both skeletal and extraskeletal 211 tissues (Lee et al., 1995; Urena et. al., 1993; Tian et. al., 1993; Watson et. al, 2000a). Reports 212 suggest that the distribution of PTHrP and its receptor are nearly ubiquitous since this receptor 213 ligand duo are described in tissues as diverse as skin (Kaiser et. al., 1992), smooth muscle 214 (Moseley and Gillespie, 1995; Williams et. al., 1994; Yamamoto et. al., 1992; Watson, et. al.,

215 2000a), bone (Lee et. al., 1993; Kronenberg et. al., 1998), kidney (Lee et. al., 1996; Amizuka et.

216 al., 1997; Yang et. al., 1997; Watson, et. al., 2000a), uterus-placenta (Williams et. al., 1994; 217 Tucci and Beck, 1998; Watson, et. al., 2000b), brain (Weaver et. al., 1995), breast (Downey et. 218 al., 1997), intestine (Li et. al., 1995; Watson, et. al, 2000a), pancreas and cardiovascular system 219 (Vasavada, et. al., 1998). Few studies to date have investigated the expression or role of PTHrP 220 and the PTHR in the mammalian ovary. Ovarian expression of PTHrP was first demonstrated by 221 the humoral hypercalcemia of malignancy associated with ovarian small cell carcinomas and 222 PTHrP is now widely used as an ovarian tumour expression marker (Matias-Guiu et. al., 1994; 223 Inoue et. al., 1995). Northern blots were used to identify PTHR transcripts in rat ovary (Urena et. 224 al., 1993; Joun et. al., 1997) and PTHrP expression has been demonstrated in the ovary of the 225 developing frog (Danks et. al., 1997) and most recently in the porcine ovary (Garmey et al., 226 2000). Both the PTHR and its ligand, PTHrP, have been identified in rat ovary (Watson, et. al, 227 2000a). Only one study has identified PTHrP as a component of human ovarian follicular fluid 228 and demonstrated that the granulosa-luteal cells were the source of the PTHrP (Gutmann et. al., 229 1993). None of these studies have attempted to localize mRNAs encoding PTHrP and the PTHR 230 or their polypeptides as we have in the present study. In a recent study reporting nuclear 231 localization of the PTHR, we observed that transcripts encoding both the PTHR and PTHrP and 232 their polypeptides were present in rat ovary (Watson et. al., 2000a). 233 PTHrP is implicated as a regulator of embryo development (Behrendtsen et al., 1995; Lee 234 et al., 1995; Nowak et al., 1999). PTHrP and the PTHR are expressed during differentiation of 235 embryonal carcinoma or stem cells to primitive and parietal endoderm (van de Stolpe et. al., 236 1993) and PTHrP is essential to the outgrowth of parietal endoderm from isolated mouse embryo 237 inner cell mass (ICM) regardless of the substratum used for outgrowth (Behrendtsen et al., 1995). 238 In the developing mouse embryo, expression of PTHR transcripts was detected in parietal 239 endoderm from day 5.5 p.c. onwards. In the embryo proper, PTHR transcripts were highly 240 expressed at sites of epithelial-mesenchyme interaction in developing intestine, lung, kidney and 241 dermis starting at day 9.5 p.c. (Karperien, et. al., 1994). A comprehensive survey of PTHrP and 242 PTHR expression during rat fetal development found that, in extraskeletal tissues, PTHrP mRNA 243 was largely expressed in surface epithelia while PTHR mRNA was mainly localized to the 244 adjacent mesenchyme (Lee et al., 1995). This was true for tissues as diverse as lung, choroid plexus, teeth, heart and skin. These results are strongly suggestive of a paracrine role for PTHrP 245 246 during development. It has been proposed that PTHrP is a local, autocrine/paracrine regulator of 247 cell growth and differentiation (Tucci and Beck, 1998; Moseley and Gillespie, 1995). A 248 physiological role for PTHrP during mammalian endochondral bone formation and modeling is 249 well documented (Lee, et.al., 1996; Lanske et. al., 1996; Lanske and Kronenberg, 1998). In other 250 tissues PTHrP regulates smooth muscle relaxation, placental calcium transfer, breast 251 development and lactation, vascular smooth muscle tension and pancreatic β -cell growth (Mosely 252 and Gillespie, 1995; Vasavada et. al., 1998; Porter et. al., 1998). Very recently Nowak, et al., 253 (1999) reported that PTHrP and transforming growth factor β (TGF- β) both interact to promote 254 murine blastocyst outgrowth demonstrating that PTHrP may be an important regulator of early 255 developmental events.

The characterization of optimized culture conditions for mammalian oocytes and early embryos is an important priority. Progress in the development of effective defined media for embryo culture has occurred rapidly in the last few years. Media supplements such as serum have, however, only recently been removed from standard culture protocols and little research has 260 investigated the role of ovarian factors on oocyte maturation and early development *in vitro* under 261 defined serum-free culture conditions (Saeki et al., 1991; Gardner, 1994; Eckert et al., 1995; Keskintepe and Brackett, 1996; Hill et al., 1997; Avery et al., 1998; Krishner and Bavister, 1999; 262 263 Watson et al., 2000). Research from our laboratories has characterized cSOFMaa medium for 264 bovine oocyte maturation (Watson et al., 2000). Our results indicate that supplementation of 265 cSOFMaa oocyte maturation medium with PTHrP may be of benefit for supporting enhanced 266 numbers of bovine zygotes through to the blastocyst stage. Future studies will explore the 267 mechanism underlying this positive influence and will determine whether PTHrP should become 268 a routine supplement to mammalian oocyte maturation and embryo culture media.

269

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278	References
279	Abou-Samra AB, Juppner H, Force T, Freeman MW, Kong XF, Schipani E, Urena P, Richards J
280	Bonventre JV, Potts Jr JT, Kronenberg HM and Segre GV (1992) Expression cloning of a
281	common receptor for parathyroid hormone and parathyroid hormone-related peptide from rat
282	osteoblast-like cells: a single receptor stimulates intracellular accumulation of both cAMP and
283	inositol triphosphates and increases intracellular free calcium. Proc Natl Acad Sci USA 89:2732-
284	2736
285	
286	Amizuka N, Lee HS, Kwan MY, Arazani A, Warshawsky H, Hendy GN, Ozawa H, White JH and
287	Goltzman D (1997) Cell-specific expression of the parathyroid hormone (PTH)/PTH-related
288	peptide receptor gene in kidney from kidney-specific and ubiquitous promoters. Endocrinology
289	138:469-481
290	
291	Avery B, Bavister BD and Greve T (1998) Development of bovine oocytes, in vitro matured in a
292	chemically defined protein-free medium, supplemented with different amino acid
293	formulations. Theriogenology 49:306.
294	
295	Behrendtsen O, Alexander CM and Werb Z (1995) Cooperative interactions between extracellular
296	matrix, integrins and parathyroid hormone-related peptide regulate parietal endoderm
297	differentiation in mouse embryos. Development 121:4137-4148
298	

301	Curtis NE, Thomas RJ, Gillespie MT, King RG, Rice GE and Wlodek ME (1998) Parathyroid
302	hormone-related protein (PTHrP) mRNA splicing and parathyroid hormone/PTHrP receptor
303	mRNA expression in human placenta and fetal membranes. J Mol Endo 21:225-234
304	
305	Danks JA, McHale JC, Martin TJ and Ingleton PM (1997) Parathyroid hormone-related protein in
306	tissues of the emerging frog (Rana temporaria): immunohistochemistry and in situ hybridisation.
307	J Anat 190 (Pt 2): 229-238
308	
309	Downey SE, Hoyland J, Freemont AJ, Knox F, Walls J and Bundred NJ (1997) Expression of the
310	receptor for parathyroid hormone-related protein in normal and malignant breast tissue. J Pathol
311	183:212-217
312	
313	Eckert J and Niemann H (1995) In vitro maturation, fertilization and culture to blastocyst of
314	bovine oocytes in protein-free media. Theriogenology 43:1211-1225.
315	
316	Ferguson II JE, Seaner RM, Bruns DE, Iezzoni JC and Bruns ME (1998) Expression and specific
317	immunolocalization of the human parathyroid hormone/parathyroid hormone-related protein
318	receptor in the uteroplacental unit. Amer J Ob Gyn 179: 321-329
319	
320	Gardner DK (1994) Mammalian embryo culture in the absence of serum or somatic cell support.
321	Cell Biology Inter 18:1163-1179.

324	Garmey JC, Schnorr JA, Bruns ME, Bruns DE, Seaner RM, Ferguson JE, Jayes FCL, Aquirre C, and
325	Veldhuis JD (2000) Expression of parathyroid hormone-related peptide (PTH-rp) and its receptor
326	in the procine ovary: regulation by transforming growth factor- β and possible paracrine effects of
327	granulose cell PTH-rp secretion on theca cells. Biol Reprod 62:334-339.
328	
329	Gutmann JN, Burtis WJ, Dreyer BE, Andrade-Gordon P, Penzias AS, Polan ML and Insogna KL
330	(1993) Human granulosa-luteal cells secrete parathyroid hormone-related protein in vivo and in
331	vitro. J Clin Endocrinol & Metab 76:1314-1318
332	
333	Hill JL, Wade MG, Nancarrow CD, Kelleher DL and Boland MP (1997) Influence of ovine
334	oviductal amino acid concentrations and an ovine oestrus associated glycoportein on
335	development and viability of bovine embryos. Mol Reprod Dev 47:164-169.
336	
337	Horiuchi N, Caulfield MP, Fisher JE, Goldman ME, McKee RL, Reagan JE, Levy JJ, Nutt RF, Rodan
338	SB, Scholfield TL, Clemens TL and Rosenblatt M (1987) Similarity of synthetic peptide from
339	human tumor to parathyroid hormone in vivo and in vitro. Science 238:1566-1568
340	
341	Inoue H, Kikuchi Y, Hirata J, Wada S, Seki K and Nagata I (1995) Dysgerminoma of the ovary with
342	hypercalcemia associated with elevated parathyroid hormone-related protein. Jap J Clin Oncol
343	25:113-117

344

347	Joun H, Lanske B, Karperien M, Qian F, Defize L and Abou-Samra A (1997) Tissue-specific
348	transcription start sites and alternative splicing of the parathyroid hormone (PTH)/PTH-related
349	peptide (PTHrP) receptor gene: a new PTH/PTHrP receptor splice variant that lacks the signal
350	peptide. Endocrinology 138:1742-1749
351	
352	Kaiser SM, Laneuville P, Bernier SM, Rhim JS, Kremer R and Goltzman D (1992) Enhanced growth
353	of a human keratinocyte cell line induced by antisense RNA for parathyroid hormone-related
354	peptide. J Biol Chem 267:13623-13628
355	
356	Karperien M, Lanser P, de Laat SW, Boonstra J, and Defize LH (1996) Parathyroid hormone
357	related peptide mRNA expression during murine postimplantation development:evidence
358	for involvement in multiple differentiation processes. Int J Dev Biol 40:599-608.
359	
360	Keskintepe L and Brackett BG (1996) In vitro developmental competence of in vitro-matured
361	bovine oocytes fertilized and cultured in completely defined media. Biol Reprod 55:333-
362	339.
363	
364	Keskintepe L, Burnley CA and Brackett BG (1995) Production of viable bovine blastocyst in
365	defined in vitro conditions. Biol Reprod 52:1410-1417.

367	Krisher RL and Bavister BD (1999) Enhanced glycolysis after maturation of bovine oocytes in
368	vitro is associated with increased developmental competence. Mol Reprod Dev 53:19-26.
369	
370	Kronenberg HM, Lanske B, Kovacs CS, Chung U-I, Lee K, Segre GV, Schipani E and Juppner H
371	(1998) Functional analysis of the PTH/PTHrP network of lignads and receptors. Rec Prog Hor
372	Res 53:283-303
373	
374	Lanske B, Karaplis AC, Lee K, Luz A, Vortkamp A, Pirro A, Karperien M, Defize LHK, Ho C,
375	Mulligan RC, Abou-Samra AB, Juppner H, Segre GV and Kronenberg HM (1996) PTH/PTHrP
376	receptor in early development and indian hedgehog-regulated bone growth. Science 273: 663-666
377	
378	Lanske B and Kronenberg HM (1998) Parathyroid hormone-related peptide (PTHrP) and parathyroid
379	hormone (PTH)/PTHrP receptor. Crit Rev Eukaryotic Gene Express 8:297-320
380	
381	Largo R, Gomez-Garre D, Santos S, Penaranda C, Blanco J, Esbrit P and Egido J (1999) Renal
382	expression of parathyroid hormone-related protein (PTHrP) and PTH/PTHrP receptor in a rat
383	model of tubulointerstitial damage. Kid Inter 55:82-90
384	
385	Lee K, Brown D, Urena P, Ardaillou N, Ardaillou R, Deeds J and Segre GV (1996) Localization of
386	parathyroid hormon-parathyroid hormone-related peptide receptor mRNA in kidney. Amer J
387	Physiol 270:F186-F191

389	Lee K, Deeds JD, Bond AT, Juppner H, Abou-Samra A-B and Segre GV (1993) In situ localization of
390	PTH/PTHrP receptor mRNA in the bone of fetal and young rats. Bone 14:341-345
391	
392	
393	Lee K, Deeds JD and Segre GV (1995) Expression of parathyroid hormone-related peptide and its
394	receptor messenger ribonucleic acids during fetal development of rats. Endocrinology 136:453-
395	463
396	
397	Lee K, Lanske B, Karaplis AC, Deeds JD, Kohno H, Nissenson RA, Kronenberg HM and Segre GV
398	(1996) Parathyroid hormone-related peptide delays terminal differentiation of chondrocytes
399	during endochondral bone formation. Endocrinology 137:5109-5118
400	
401	Li H, Seitz PK, Thomas ML, Selvanayagam P, Rajaraman S and Cooper CW (1995) Widespread
402	expression of the parathyroid hormone-related peptide and PTH/PTHrP receptor genes in
403	intestinal epithelial cells. Lab Invest 73:864-870
404	
405	Matia-Guiu X, Prat J, Young RH, Capen CC, Rosol TJ, Delellis DA and Scully RE (1994) Human
406	parthyroid hormone-related protein in ovarian small cell carcinoma. An immunohistochemical
407	study. Cancer 73:1878-1881
408	
409	Moseley JM and Gillespie MT (1995) Parathyroid hormone-related protein. Crit Rev Clin Lab Sci

410 32:299-343

412	Nowak RA, Haimovici F, Biggers JD, and Erbach GT (1999) Transforming growth factor- β
413	stimulates mouse blastocyst outgrowth through a mechanism involving parathyroid
414	hormone-related protein. Biol Reprod 60: 85-93.
415	
416	Parrish JJ, Susko-Parrish JL, Leibfried-Rutledge ML, Crister ES, Eyestone WH and First NL
417	(1986) Bovine in vitro fertilization with frozen thawed sperm. Theriogenology 25: 591-600.
418	
419	Pausova Z, Bourdon J, Clayton D, Mattei M-G, Seldin MF, Janicic N, Riviere M, Szpirer J, Levan G,
420	Szpirer C, Goltzman D and Hendy GN (1994) Cloning of a parathyroid hormone/parathyroid
421	hormone-related peptide receptor (PTHR) cDNA from a rat osteosarcoma (UMR106) cell line:
422	chromosomal assignment of the gene in the human, mouse and rat genomes. Genomics 20:20-26.
423	
424	Porter SE, Sorenson RL, Dann P, Garcia-Ocana A, Stewart AF and Vasavada RC (1998) Progressive
425	pancreatic islet hyperplasia in the islet-targeted, parathyroid hormone-related protein-
426	overexpressing mouse. Endocrinology 139:3743-3751
427	
428	Rodan SB, Noda M, Wesolowski G, Rosenblatt M and Rodan GA (1988) Comparison of postereceptor
429	effects of 1-34 human hypercalcemia factor and 1-34 human parathyroid hormone in rat
430	osteosarcoma cells. J Clin Invest 81:924-927
431	

432	Saeki K, Hoshi M, Leibfried-Rutledge ML and First NL (1991) In vitro fertilization and
433	development of bovine oocytes matured in serum-free medium. Biol Reprod 44:256-260.
434	
435	Spiegel AM, Saxe AW, Deftos LJ and Brennan MF (1983) Purification of peptides with parathyroid
436	hormone-like bioactivity from human and rat malignancies associated with hypercalcemia. Hor
437	Metab Res 15: 299-304
438	Stewart AF, Wu T, Goumas D, Burtis WJ and Broadus AE (1987) N-terminal amino acid sequence of
439	two novel tumor-derived adenylate cyclase-stimulating proteins: identification of parathyroid
440	hormone-like and parathyroid hormone-unlike domains. Biochem Biophys Res Comm 146:672-
441	678
442	
443	Strewler GJ, Stern PH, Jacobs JW, Eveloff J, Klein RF, Leung SC, Rosenblatt M and Nissenson RA
444	(1987) Parathyroid hormone-like protein from human renal carcinoma cells: structural and
445	functional homology with parathyroid hormone. J Clin Invest 80:1803-1807
446	
447	Tian J, Smogorzewski M, Kedes L and Massry SG (1993) Parathyroid hormone-parathyroid hormone-
448	related protein receptor messenger RNA is present in many tissues besides the kidney. Amer J
449	Nephrol 13:210-213
450	
451	Tucci J and Beck F (1998) Expression of parathyroid hormon-related protein (PTHrP) and the
452	PTH/PTHrP receptor in the rat uterus during early pregnancy and following artificial deciduoma
453	induction. J Reprod Fertil 112:1-10

455	Urena P, Kong S-F, Abou-Samra A-B, Juppner H, Kronenberg HM, Potts Jr. JT and Segre GV (1993)
456	Parathyroid hormone (PTH)/PTH-related peptide receptor messenger ribonucleic acids are widely
457	distributed in rat tissues. Endocrinology 133:617-623
458	
459	
460	
461	van de Stolpe A, Karperien M, Lowik CW, Juppner H, Segre GV, Abou-Samra AB, de Laat SW and
462	Defize LH (1993) Parathyroid hormone-related peptide as an endogenous inducer of pariatal
463	endoderm differentiation. J Cell Biol 120: 235-243
464	
465	Vasavada RC, Garcia-Ocana A, Massfelder T, Dann P and Stewart AF (1998) Parathyroid hormone-
466	related protein in the pancreatic islet and the cardiovascular system. Rec Prog Hor Res 53:305-
467	340
468	
469	Watson AJ, Watson PH, Warnes D, Walker SK, Armstrong DT and Seamark RF (1994)
470	Preimplantation development of in vitro-matured and in vitro-fertilized ovine
471	zygotes:comparison between coculture on oviduct epithelial cell monolayers and culture
472	under low oxygen atmosphere. Biol Reprod 50:715-724.
473	
474	Watson P, Lazowski D, Han V, Fraher L, Steer B and Hodsman A (1995) Parathyroid hormone restores
475	bone mass and enhances osteoblast insulin-like growth factor I gene expression in

476	ovariectomized rats. Bone 16:357-365.
477	
478	Watson PH, Fraher LJ, Hendy GN, Chung U-I, Kisiel M, Natale BV and Hodsman AB (2000a) Nuclear
479	localization of the type 1 PTH/PTHrP receptor in rat tissues. J Bone Min Res 15 (6): 1033-1044.
480	
481	
482	
483	Watson PH, Fraher LJ, Natale BV, Kisiel M, Hendy GN and Hodsman AB (2000b) Nuclear
484	localization of the type 1 PTH/PTHrP receptor in MC3T3-E1 cells: association with the cell
485	cycle. Bone 26 (3): 221-225.
486	
487	Watson AJ, De Sousa P, Caveney A, Barcroft LC, Natale D, Urquhart J and Westhusin ME (2000)
488	Impact of bovine oocyte maturation media on oocyte transcripts levels, blastocyst development,
489	cell number and apoptosis. Biol Reprod 62:355-364.
490	
491	Weaver DR, Deeds JD, Lee K and Segre GV (1995) Localization of parathyroid hormone-related
492	peptide (PTHrP) and PTH/PTHrP receptor mRNAs in rat brain. Mol Brain Res 28:296-310.
493	
494	Wiemer KE, Watson AJ, Polanski V, McKenna AI, Fick GH and Schultz GA (1991) Effects of
495	maturation and co-culture treatments on the developmental capacity of early bovine embryos.
496	Mol Reprod Dev 30:330-338.
497	

498	Williams ED, Leaver DD, Danks JA, Moseley JM and Martin TJ (1994) Effect of parathyroid
499	hormone-related protein (PTHrP) on the contractility of the myometrium and localisation of
500	PTHrP in the uterus of pregnant rats. J Reprod Fertil 102: 209-214.
501	
502	Yamamoto M, Harm SC, Grasser WA and Thied MA (1992) Parathyroid hormone-related protein in
503	the rat urinary bladder: a smooth muscle relaxant produced locally in response to mechanical
504	stretch. Proc Natl Acad Sci (USA) 89:5326-5330.
505	
506	Yang T, Hassan S, Huang YG, Smart AM, Briggs JP and Schnermann JB (1997) Expression of PTHrP,
507	PTH/PTHrP receptor, and Ca(2+)-sensing receptor mRNAs along the rat nephron. Amer J
508	Physiol 272:F751-F758
509	
510	Yasuda T, Banville D, Rabbani SA, Hendy GN and Goltzman D (1989) Rat parathyroid hormone-like
511	peptide: comparison with the human homologue and expression in malignant and normal tissue.
512	Mol Endocrinol 3:518-525
513	

Figure 1: Detection of PTHrP mRNA in bovine ovary. Representative photomicrographs display A) sense control, B) detection of PTHrP mRNAs in primordial follicles included oocytes and pre-granulosa cells (arrowheads), C) sense control, D) PTHrP mRNAs in oocyte (o) and granulosa(g) layer of a secondary follicle and stromal (s) tissue, E) PTHrP mRNAs in granulosa and theca (t) cell layers of antral follicles (a), F) PTHrP mRNAs in oocyte, granulosa cells, and theca of antral follicles. Bar = 50μ M.

521 Figure 2: Detection of PTHR mRNAs in bovine ovary. Representative photomicrographs 522 display A) sense control, B) detection of PTHR mRNAs in primordial follicles included oocytes 523 and pre-granulosa cells (arrowheads), C) sense control, D) PTHR mRNAs in oocyte (o) and 524 granulosa (g) layer of primordial and secondary follicles and stromal (s) tissue, E) PTHR mRNAs 525 in granulosa and theca (t) layers of antral (a) follicles, F) PTHR mRNAs in oocyte, granulosa 526 cells, and theca of early antral follicle. Bar = $50 \mu M$ 527 528 Figure 3: Detection of PTHrP polypeptides in bovine ovary The photomicrographs represent 529 A) mouse normal serum control, B) detection of PTHrP polypeptides in primordial follicles included oocytes and pre-granulosa cells (arrowheads), C) control, D) PTHrP polypeptides in 530 531 oocyte (o) of secondary follicle, E) PTHrP polypeptides in theca (t) layers of antral follicles, F) 532 PTHrP polypeptides in oocyte of early antral (a) follicle and in ovarian stroma (s). g, granulosa

533 Bar = 50 μ M.

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538	Figure 4: Detection of PTHR polypeptide in bovine ovary The photomicrographs represent A)
539	pre-absorbed control, B) detection of PTHR polypeptides in primordial follicles included oocytes
540	and pre-granulosa cells (arrowheads) and ovarian stroma (s), C) control, D) PTHR polypeptides
541	in oocyte (o) and granulosa (g) cells of secondary follicle and stromal tissue, E) PTHR
542	polypeptides in granulosa and theca (t) layers of antral (a) follicles and stroma, F) PTHR
543	polypeptides in oocyte, granulose cells and theca of antral follicle. Bar = $50 \ \mu M$
544 545	
546	Figure 5: Influence of PTHrP during oocyte maturation on blastocyst development.
547	Proportion of inseminated oocytes cleaving (cleavage), or reaching the 6-8 cell stage, 6-8/insem,
548	or morula and blastocyst stage of development, mor/bl/insem and blastocysts blast/insem
549	displayed by oocytes matured in 0, 1.59, 15.9, and 159 ng/ml (0, 0.1, 1 and 10 nM, respectively)
550	of human PTHrP (1-141)-supplemented cSOFMaa. Cleavage, development to the 6-8 cell and
551	morulae /blastocyst stages did not vary significantly among the treatment groups. The proportion
552	of zygotes that progressed to the blastocyst stage varied significantly among the treatments. Bars
553	represent the mean \pm SEM of n=4 replicates. Values with different superscript letters are
554	significantly different (p<0.05).
555	

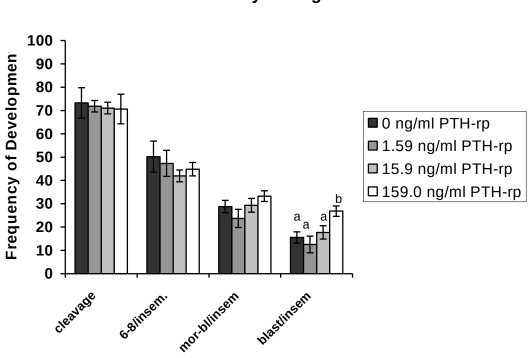


Figure 5: Influence of PTHrP During Bovine Oocyte Maturation on Development to the Blastocyst Stage

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559	Figure 6: Influence of PTHrP during oocyte maturation on progression of 6-8 cell embryos
560	to the blastocyst stage. Proportion of 6-8 cell stage embryos (over cleaved embryos, 6-8/clvd),
561	morulaeblastocysts (over cleaved embryos, mor-blast/cleaved) blastocysts (over cleaved
562	embryos, blast/clvd) and 6-8 cell embryos progressing to the blastocyst stage, (blast/6-8)
563	displayed by oocytes matured in 0, 1.59, 15.9, and 159 ng/ml (0, 0.1, 1 and 10 nM, respectively)
564	of murine PTHrP (1-141)-supplemented cSOFMaa. Development to the 6-8 cell and morula-
565	blastocyst stages did not vary significantly among the treatment groups. However the proportion
566	of blastocysts and 6-8 cell embryos that progressed to the blastocyst stage varied significantly.
567	Bars represent the mean \pm SEM of n=4 replicates. Values with different superscript letters are
568	significantly different (p<0.05).

