DEVELOPMENT OF SMALL BOWEL MOTILITY IN BEAGLE PUPPIES. 275 Frank H. Morriss, Steven B. Spedale, and Norman W. Weisbrodt, Univ. of Tx. Med. Sch. at Houston, Depts. of Pediatrics and Physiology, Houston.

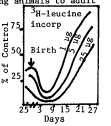
To investigate the neonatal development of upper small bowel (SB) peristalsis and the effect of pentagastrin (PG) on motility, we monitored duodenal intraluminal pressure manometrically in 11 beagle puppies from birth until weaning at 5 wk. The response rate the system employed was 150 mm Hg/sec. Electrodes were implanted on the SB serosal surface for subsequent monitoring of mycelectric activity during the first 2 mo. in 3 add'l puppies. During the first 3 wk there was an increase in duodenal contraction rate immediately after a liquid meal from 11+0.4/min to 16+1/min (p <0.001) followed by a slight decrease during weaning. Peak duodenal pressure increased from 17 to 38 mm Hg ing. Peak duodenal pressure increased from 17 to 38 mm Hg (p < 0.002), and subsequently decreased to 25 mm Hg with weaning. The duration of each contraction shortened during the first 3 wk from 3.4 to 2.4 sec (p < 0.001). Myoelectric monitoring revealed a gradual increase in slow wave activity from 14 to 20 cycles/min in both fasting and fed states, corresponding with the increased developmental changes are similar to those which occur in the esophageal sphincter (LES) and stomach during beagle lower suckling and weaning. However, in contrast to the lack of sensitivity to PG by the beagle LES and stomach during the first week, SB contractions were inhibited on day 1 with 0.03-8 ug/kg PG We speculate that the differential PG sensitivity of LES, stomach, and SB during the first week of life in the beagle is due to differential gastrin receptor number and/or affinity.

2766 POSSIBLE MECHANISM FOR NON-REJECTION OF THE DEVELOP-ING MAMMALIAN EMBRYO: THE ROLES OF UTEROGLOBULIN AND TRANSGLUTAMINASE. A. Mukherjee*, R. Ulane*, and A. Agrawal*. (SPON: J.D. Schulman) NICHD, NIH; Bethesda, MD 20205 The suppression of embryonic immunogenicity by the interaction of rabbit blastocysts with uteroglobin (UG) (a pregnancy specific protein) and transglutaminase (TG) (Factor XIIIa) has been inves-tigated in vitro. When incubated with maternal lymphocytes, washed, mitomycin-C inactivated blastomeres stimulated lymphocytes, suggesting recognition of embryonic antigens. When these blasto-meres are pretreated with pregnant uterine flushings (PUF), ³H-thymidine incorporation was dramatically reduced (1.0 x 10⁴ cpm/2 x 10⁶ lymphocytes). Pretreatment of blastomeres with UG alone, isolated to homogeneity from PUF, or in combination with alone, isolated to homogeneity from PUF, or in combination with TG caused significant dose dependent suppression of thymidine incorporation by the lymphocytes. Suppression to (100 cpm/ 2 x 10^6 cells was achieved by 250 ug/ml of UG alone; with added TG (3 u/ml) only 1.0 ug/ml of UG caused total suppression. Suppres-(3 u/ml) only 1.0 ug/ml of UG caused total suppression. Suppression was blocked by incubation of UG with its antiserum, or with TG plus anti-TG, prior to blastomere pretreatment. Inhibition of exogenous transglutaminase by neopentyl chloroethyl nitrosourea also greatly reduced the inhibition of thymidine incorporation by UG. The data clearly indicate that UG and TG suppress blastomere Immunogenicity, perhaps by a cross-linking mechanism between UG and blastomere antigens. We suggest that uteroglobin in conjunction with transglutaminase may inhibit immunorejection of embryos at about the time of implantation. at about the time of implantation.

CHANGES IN SENSITIVITY TO RICINUS COMMUNIS TOXIN 277 (RICIN) OF RABBIT JEJUNUM DURING THE SUCKLING PERIOD. A. Olson, R. Torres-Pinedo. University of Oklahoma

College of Medicine, Department of Pediatrics, Oklahoma City. Pinocytosis is the predominant route of penetration of macromolecules across the small bowel epithelium of newborn animals. We have developed a model system to study this process on jejunal mucosal explants using ricin. Ricin is a potent toxin which inhibits protein synthesis by a mechanism involving binding to galactose surface residues, internalization of its enzymatic compo-nent and inactivation of the 60 S ribosomal subunit. Protein synthesis (3H-leucine incorporation) in jejunal explants from fetal, suckling and adult rabbits was measured in organ culture following an initial exposure (30 min, 25°) to ricin (1-25 μ g). The rate of inhibition of protein synthesis was age dependent (Fig). High in the fetus, it increased further to a maximum at day 6 postnatally, then decreased rapidly in suckling an<u>imals to adult</u>

levels at weaning. The high sensitivity to ricin during the colostral period may be related to the following factors currently under study: 1) colostral-milk components stimulating a high rate of endocytosis, 2) transitional maturation changes in membrane properties and 3) changes in sensitivity at the target ribosomal site. The model presented is useful to study modifiers and events influencing mucosal permeability in the developing small intestine.



THE MATERNAL AND FETAL CATECHOLAMINE (CAT) RESPONSE 278 278 TO HYPOXIA IN SHEEP. Sue M. Palmer, Gary K. Oaks, <u>Robert Lam, Calvin J. Hobel, and Delbert A. Fisher</u>. UCLA School of Medicine, Harbor-UCLA Medical Center, Departments of Obstetrics-Gynecology and Pediatrics, Torrance, CA.

The plasma CAT responses to hypoxia were investigated in six pregnant ewes at 133-140 days gestation. Hypoxia was induced by exposure of the ewe to $6-9\% 0_2 - 3\% CO_2/nitrogen$ for 30 min. Mat-ernal and fetal arterial blood samples were obtained at 5 min in-tervals before and during the hypoxic stress for measurement of cervals before and during the hypoxic stress for measurement of epinephrine (E), norepinephrine (NE) and blood gases. Analyses of maternal CAT responses revealed a significant elevation of mean NE from 191 to 685 pg/ml (p<0.05) and E from 129 to 311 pg/ ml (p<0.05). The levels of plasma NE correlated with duration of hypoxia (r=0.93, p<0.05) based on mean NE values at 10, 20 and 30 min). Fetal CAT responses correlated (r=0.54, p<0.05) with sever-ity of maternal hypoxia as follows:

mater	nal nypoxia	as tollows:	
	%Mat paO ₂	Fet. E	Fet. NE
	100 (C)	26	179
	>51	43	265
	40-50	7,332*	2,101*
	30-40	30,333*	33,190*
$pa0_2 =$	pa0 ₂ during	hypoxia/pa0 ₂ control X	100.
		cant maternal F and NF	

% Mat Conclusions: 1) Significant maternal E and NE secretion occurs with hypoxia. 2) The levels of maternal NE correlate linearly with time of hypoxic exposure. 3) Fetal E and NE secretion increase with maternal hypoxia. 4) There is a direct correlation between fetal E or NE levels and counsity of between fetal E or NE levels and severity of maternal hypoxia.

SYMPATHO-ADRENAL RESPONSE TO UMBILICAL CORD CUTTING 279 279 (UCC). James F. Padbury, Emmanuel S. Diakomanolis, Calvin J. Hobel, Alvin Perelman and Delbert A. Fisher UCLA School of Medicine, Harbor-UCLA Medical Center, Departments of Pediatrics and Obstetrics-Gynecology, Torrance, CA. Catecholamine (CAT) release has been observed in near term fe-Fisher,

tal sheep in response to a variety of stimuli including hypoxia, maternal hypovolemia or exercise and parturition. In earlier studies of fetal sympatho-adrenal activity in response to parturi-tion it has not been possible to differentiate the effects of la-bor, intrapartum asphyxia, delivery and UCC on CAT release. We have used the acutely exteriorized near term fetal lamb to study the effects of delivery and UCC on CAT release. Results show that delivery alone evokes an elevation of newborn plasma CAT which is brief and followed by return to basal values by 30 min. Subse-quent UCC evokes a marked release of norepinephrine (NE) and epquent UCC evokes a marked release of norepinephrine (NE) and ep-inephrine (E) (peak plasma levels 32,000 pg/ml and 35,000 pg/ml), maximal at 5 min. and persisting over the 4 hour study period. There is a concomitant rise in plasma free fatty acids (FFA) and reversal of post-delivery hypothermia. The magnitude of the CAT surge is inversely proportional to the degree of acidosis; a blunted FFA response and slower correction of hypothermia were observed in more acidotic animals despite higher CAT levels. Conclusions: 1) UCC is an important stimulus for fetal CAT re-lease. 2) UCC stimulated neurosympathetic activity is of suf-ficient magnitude to influence many metabolic and cardiovascular functions. 3) Acidosis stimulates CAT release in the near term ovine fetus. 4) Acidosis obtunds neonatal chemical thermogenesis.

THE EFFECTS OF POSTNATAL AGE ON THE WHOLE BODY PROTEIN METABOLISM AND SKELETAL MUSCLE PROTEIN BREAK-280 DOWN OF PREMATURE INFANTS. <u>P. Pencharz, M. Masson,</u> <u>F. Desgranges, A. Papageorgiou.</u> (Spon: T. Heim) The Hospital for Sick Children, Toronto; and The Jewish General Hospital, Montreal, Quebec, Canada. Rates of whole body nitrogen flux (Q), protein synthesis (S),

and breakdown (C), and skeletal muscle protein breakdown were measured in 24 growing premature infants. The infants were each studied twice. The first study (A) was conducted once the in-fants were clinically stable and ingesting an oral intake of at least 120 kcal/kg/d. The second study (B) was started 2 weeks later.

There was a marked increase in rates of skeletal muscle pro-tein breakdown, from 0.93 ± 0.21 g to 2.9 ± 0.26 g/kg/d (p<0.01). During Study A, very low birth weight (<1500 g) infants had sig-nificantly higher rates than larger infants (1.19 cf 0.66 g/kg/ d); however, these differences were no longer present during Study B; both groups showing a marked increase in muscle break-down rates (2.95 cf 3.03 g/kg/d). There were no differences in Q, S or C with increasing postnatal age. It appears that skele-tal muscle protein metabolism increases postnatally from about 8% to 24% of whole body protein turnover with a concommitant decrease in other tissues. Skeletal muscle has been estimated to constitute about 22% of total body protein in infants. Thus, these changes may reflect an increase in skeletal muscle protein turnover from a low rate to one that is in proportion to skeletal muscle's contribution to whole body protein.