buffer. Thiocarbamyl-nitro-BT was used as the reduction indicator. Incubation was carried out at 15°C for 2 hrs and the tissues rinsed in buffer and post-fixed in 1 p. 100 osmium tetroxide in buffer for 1 hr. They were processed for electron-microscopy and thin sections stained with lead citrate. HSD enzymes were localised by precipitation of electron-dense osmium compounds.

DICHTOMY OF RESPONSE TO HCG BETWEEN THE GRAAFAIAN FOLLICLE AND OOCYTE IN PIGS. — R. H. F. HUNTER. University of Edinburgh (Scotland).

Recent experiments on the morphology and steroidogenic potential of the Graafian follicle, particularly those conducted in vitro, give little indication of the functional status of the oocyte. In studies on maturation of the follicle and oocyte in pigs, we have observed that if ovulation is induced with an injection of HCG in the early follicular phase of the oestrous cycle, many primary oocytes are ovulated. These precociously liberated oocytes do not exhibit a zona reaction, but invariably become highly polyspermic when confronted by capacitated spermatozoa. This dichotomy of response to HCG between the follicle and its oocyte will be discussed.


Studies on ultrastructural events during fertilization in mammals have been largely confined to laboratory species, although we have published an extensive report on the domestic pig. Using a system that induces varying degrees of polyspermic penetration, ranging from dispermy to > 20 vitelline sperm, we have examined the formation of pronuclei and the fate of the male elements. EM features of these eggs will be presented, with emphasis on decondensation of the nuclear chromatin and apparent anomalies of the male pronuclei. Observations will also be included on structural abnormalities in the midpiece of spermatozoa from the fertile boar used in these studies.


The storage of cow eggs at room temperature (19-21°C) and after cooling to low temperature (0-7.5°C) was examined in two experiments. Morulae (Day 5) and 8-celled eggs (Day 3) were stored in PBS (Dulbecco Phosphate Buffer) or TCM 199 (Tissue Culture Medium 199) for 1-2 hr or 6-7 hr at 19-21°C. After storage, eggs were transferred to the ligated rabbit oviduct for 48 hr (Day 5) or 96 hr (Day 3). Storage in PBS resulted in a higher proportion of apparently normal embryos than storage in TCM 199, particularly for Day 3 eggs. Cooling of morulae (Day 5 and Day 6) to temperatures of 0°, 2°, 5° and 7.5° for 2 min, 30 min and 24 hr resulted in a high proportion of degenerate eggs. These findings will be discussed in relation to the recent attempts to freeze cow eggs.