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EPITHELIAL INVOLVEMENT IN CHOLELITHIASIS IN GROUND SQUIRRELS

THE UNIVERSITY OF ALBERTA

BY 📜

, RAMJEET SINGH\_PEMSINGH

A THESIS SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH IN PARTIAL FULFILMENT OF THE OUIREMENTS FOR THE DEGREE: OF DOCTOR OF PHILOSOPHY

DEPARTMENT OF ANATOMY AND CELL BIOLOGY

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EDMONTON, ALBERTA

(FALL 1987)

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THE UNIVERSITY OF ALBERTA

FACULTY OF GRADUATE STUDIES AND RESEARCH

The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research for acceptance, a thesis entitled. Epithelial Involvement. in CholeLithiasis in Ground Squirrels submitted by. Ramjeet S. Pemsingh in partial fulfilment of the requirements for the degree of. Doctor of Philosophy in Anatomy and Cell Biology

(Supervisor)

(External examiner)



The search for an appropriate animal model to study cholelithiasis is ongoing. The present study was designed to determine whether the Richardson's ground squirrel (<u>Spermophilus richardsonii</u>) would be a suitable model in which to study cholesterol precipitation, the formation of biliary concrements, mucosal changes occurring prior to, during and after stone formation, and compare these findings with those in humans and in other models.

ABSTRACT

•Two hundred and thirty-eight ground squirrels of both sexes were divided into control and experimental groups. Each group contained at least four animals. The experimental groups were fed a 2% cholesterol-enriched rat chow diet for intervals of 6, 12, 18, 24 hours, 3, 5, 7 days, 2, 9 3, 10, 20 weeks, and 3 weeks on cholesterol-enriched diet followed by 3 weeks on normal rat chow diet.

The results indicated that in control animals, the gallbladder epithelium was composed of three types of cells: light, dark and edematous, the latter being extrule from the epithelial sheet. Normal columnar epithelial cells underwent mitoses. The basal lamina formed peg and socket interdigitations with the basal plasma membrane, and desmosomes were seen throughout the lateral membranes. Noradrenergic nerve fibers were seen beneath the lining epithelium and between muscle bundles. A basal level of both sial\_lated and sulphated mucins was exocytosed in membrane-bound granules by merocrine secretion.

In experimental animals, mucus hypersecretion and bile lithogenicity occurred as early as 18 hours and continued throughout the experimental period. Cholesterol monohydrate crystal precipitation occurred in 27 hours, microliths were formed in 2-3 weeks and mulberry stones occurred within 10 weeks. The larger stones contained 61-74% cholesterol. Mucosal changes observed include increased cellular proliferation in one week, cellular damage and extrusion, hypertrophy and hyperplasia in 10 weeks, plasma cell infiltration and Rokitansky-Aschoff sinuses in 20 weeks. Other mucosal changes observed were lipid accumulation intercellularly and intracellularly in 12 hours, deposition of cholesterol and neutral lipid supranuclearly and basally by 7 days and in the lamina propria and its macrophages by 20 weeks.

The present study demonstrated that the Richardson's ground squirrel is an excellent model to study the pathogenesis of cholesterol gallstone disease.

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### INTRODUCTION

Gallstones, although recognised for centuries, have commanded special attention over the past few decades. This has occurred because gallstone disease now represents a major health problem in Canada, affecting some 130,000 adults per year. It is the most common abdominal surgical problem in Canada with about 30,000 cholecystectomies per year performed at a cost of some \$280,000,000 (Shaffer, 1980; Vayda, 1973).

The presence of gallstones in the human population is higher because about 90% of the affected people with some form of gallbladder disease displayed minor or no symptoms (silent stones) (Bouchier, 1984).

The formation of gallstones depends on a number of factors. The presence of a lithogenic bile in the gallbladder is not sufficient to produce cholelithiasis. In addition the gallbladder mucosa (mucus hypersecretion), muscularis (stasis) and nucleating factor(s) are involved. The nucleating factor(s) have not been fully identified but there is increasing evidence that mucus is involved (LaMont et 30., 1984).

Human gallbladder tissue stained from

63)

cholecystectomies are generally in the end stage of the disease process and as such it is difficult to chronologically evaluate hepatobiliary physiology, gallstone formation or other disease of the gallbladder. This results in major gaps in our knowledge and the need for animal models to fill these gaps.

A number of animal models have been developed to study gallbladder disease (Malet, 1985; Holzbach, 1984; Hoffmann, 1984). Freston and Bouchier (1968) listed eight features that are optimal for an ideal model. (see literature review). A number of these models such as: dog, rabbit, mouse, hamster and guinea pig failed to fulfil these criteria either because of the diet used, type of stone formed, length of time needed for stone formation or pathological changes occurring in other organs besides the gallbladder.

The prairie dog model most closely fulfils the criteria of Freston and Bouchier (1968). The caloric distribution of the diet is similar to man, the type of stone formed, rapid development of stones, and an adequate amount of tissue and bile make this model ideal to study gallbladder disease.

The scarcity and protected status of the prairie dog

in Canada led Davison <u>et al.</u> (1982) and Fridhandler <u>et al.</u> (1983) to further developed Patton <u>et al.</u>'s (1961) ground squirrel model. The Richardson's ground squirrel, when fed a 1% cholesterol-enriched rat chow diet, secreted bile with an increased lithogenic index close to saturation levels within two weeks and defective gallbladder contractility before crystal precipitation (Fridhandler <u>et al.</u>, 1983). A thorough systematic investigation of this model in the cholelithiasic process however was lacking. This stuar was designed to:

(a) describe the morphology of the normal ground r squirrel gallbladder

(b) identify the chronology of crystal and stone ' formation

(c) describe the morphology and method of formation

of these biliary concrements (d) describe intraepithelial and mucosal changes occurring prior to, during and after\*stone formation.

### LITERATURE REVIEW

<u>ke of dietary cholesterol</u>

The source of cholesterol is twofold. As much as 70% is manufactured by the liver at a rate of 1000mg a day. The other source is dietary intake. Some foods have a high cholesterol content: beef kidney(90g) - 683mg; beef liver (90g) - 372mg; one egg- 275mg; doughnut(90g) - 10mg; french fries(90g) - 20mg (Langone, 1984).

Cholesterol, which is present in all diets chiefly as free alcohol, is hydrolyzed by pancreatic cholesterol esterase before it is absorbed. Bile and fat are necessary for cholesterol absorption. Bile, phospholipids, the aycerides (fat) and cholesterol form micelles which the tallon into the columnar epithelial cells of the small intestine by passive diffusion and active absorption in the terminal ileum. In the absence of bile and fat, no cholesterol is absorbed. Cholesterol fed in a fat-free diet may also be absorbed because endogenous fat, sufficient for micelle formation, is always present in the intestinal muceona, reaching a peak 3 hours after a meal in man, and then is slowly released into the lymph with a peak at 9 hours after the meal (Davenport, 1971).

Cholesterol in the intestinal epithelial cells comes from three cources: (1) bile(60%), (2) dietary(20%), (3) desquamated mucosal cells (20%). These pools of cholesterol in the mucosal epithelium do not mix. The mechanism of separation is unknown (Davenport; 1971; Turley and Dieteschy, 1982).

Lipid droplets in the lumen, emulsified by the action of bile salts, are broken down by pancreatic lipase. This process releases free fatty acids and monoglycerides that diffuse across the apical membrane and accumulate in the apical cytoplasm. Short chain fatty acids diffuse through the cell and enter the capillaries. Long chain, fatty acius and monoglycerides serve as substrates for triglyceride-synthesizing enzymes located in the smooth endoplasmic reticulum (SER). The resynthesized triglyceride, combined with cholesterol esters, phospholipid and a glycoprotein component (apoprotein) are synthesized in the rough endoplasmic reticulum (RER) to form a chydomicron. Chylomicrons are transported in vesicles, whose membrane is derived from the Golgi complex, to the lateral cell surfaces and are discharged by exocytosis into the intercellular spaces (Davenport, 1971; Turley and Dietschy, 1982; Davidson d Glickman, 1971).

Chylomicrons cannot pass through the bacement

membrane and therefore do not enter capillaries. Only short and medium size fatty acids (10%) enter the portal system and are taken to the liver. On the other hand, chylomicrons can enter lymphatic lacteals through the open channels between the interstitial spaces to the lymphatic lumen. The endothelial walls of the lacteal are thick but there is no enveloping basement membrane. The droplets page the endothelial separations. The cells forming the walls of lacteals contain many pinocytotic vesicles capable of carrying fat droplets across them. Once within the lacteals, the fat droplets are carried centrally by the tidal flow of the lymph caused by contraction of smooth muscle fibers of the villi or the gross movements of the mucosa. Onee delivered to the blood, chylomicrons do 🐋 recirculate in the lymph. In the capillaries of adipose and muscle tissue, the triglyceride bonds are cleaved by the enzyme lipoprotein lipase and the fatty acids are removed either to be stored as adipose tissue or used in muscle for oxidation to supply energy (Davenport; 1971; Davidson and Glickman, 1971).

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The remnant of the chylomicron containing the cholesteryl esters is removed from the circulation by, receptors at sites present only in the liver (Brown and Goldstein, 1984; 1986).

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Cholesterol in the liver

The cholesterol pool in the liver is fairly constant. It is maintained by either dietary absorption and synthesis ... or excretion Paumgartner and Paumgartner, 1984). Cholesterol in the liver is derived from three lifferent sources. (1) Dietary cholesterol or cholesterol synthesized in the intestinal mucosa that reached the liver. (2) Cholesterol that is synthesized in e: cahepatic tissues (skin, muscles) that meached the liver via high density lipoprotein (HDL) and very low density lipoprotein (VLDL). (3) Synthesis of cholesterol in the liver itself (Turley and Dietschy, 1982).

The loss of cholesterol from the liver pool is by: (1) loss directly from the body pool by sloughing of oily secretions and cells from the skin, through desquamation of cells from the stomach, small intestine and colon. (2) Cholesterol molecules may be metabolized to other product: such as testosterone and adrenocorticosteroids which may in turn be excreted from the body through urine or the gastrointestinal tract. (3) Synthesis of bile acids from cholesterol into bile. (4) Secretion of cholesterol into bile (supersaturation) (Turley and Dietschy, 1982). Bile is an aqueous solution continuously secreted at a variable rate by the erenchymal cells of the liver into bile canaliculi from which it flows to the hepatic duct. Bile secretion is under nervous, hormonal and chemical control. Stimulation of the vagus nerve, injection of secretin and administration of cholerectics increase bile flow (Van der Linden and Bergman, 1977; Paumgartner and Sauerbruch, 1983).

Bile is used as an excretory mechanism by the liver to rid itself of endugenous and exogenous products and maintain homeostasis by excretion of organic ions and cations, neutral compounds, heavy metals and bilirubin. Bilé also contains IgA from blood (vesicular transport), lysosomal enzymes (exocytosis) and Na<sup>+</sup> with water from intercellular spaces. Bile ductules and ducts alter the <sup>o</sup> composition and volume of canalicular bile by reabsorption and secretion of water and electrolytes (bicarbonate). ecretion of ductular bile is stimulated by secretin and other gastrointestinal hormones (Paumgartner and Paumgartner, 1984; Paumgartner and Sauerbruch, 1983).

Bile is composed of bile acids: cholic acid, chenodeoxycholic acid (primary), deoxycholic and

lithocholic acid (secondary), phospholipids (lecithin) and cholesterol. The primary bile acids are synthesised from cholesterol. Thus, cholesterol is a normal constituent of bile and biliary excretion is the only significant excretory route for this lipid. The mechanism of controlling the secretion of cholesterol in bile is not fully known (Van der Linden and Bergman, 1977; Turley and Dietschy, 1982).

Biliary cholesterol is derived from cholesterol synthesized in the liver, from extrahepatic tisgue and from dietary absorption. Under normal conditions, the secretion of cholesterol into bile is closely linked to phospholipids and bille acid secration. This coupling is brought about by the capacity of bile hoids to form micelles. These structures incorporate relatively large amounts of phospholipids . to form mixed micelles, which normally facilitate the complete solubilization of all cholesterol secreted into the bile Recently, it the shown that phospholipid vesicles existed in bile which contributed significantly to the sclubilization and transport of cholesterol (Holzbach, 1986). For reasons that are poorly understood, this coupling/reaction can become disrupted, so that more cholesterol enters the bile that can be solubilized by the hile acids and mospholipid present. The production of such a lithogenic bile is the initiating event in the

pat. Inesis of cholesterol gallstone disease. It can involve a variety of secretory defects ranging from excessive cholesterol secretion without any change in bile acid and phospholipid secretion rates to decreased bile acids and phospholipid secretion without any change in cholesterol ovtput (Goldstein and Brown, 1984; Turley and

Dietschy, 1982)

Mucus

Mucus also called mucin, mucoprotein, mucopolysaccharide and glycoprotein is the slimy secretion of epithelial membranes which contains water, salts and immunoglobulins (LaMont <u>et al</u>., 1984; Freston <u>et al</u>., 1969). The structure and composition of mucus from various organs and mammalian species appeared to vary only slightly. It is a high molecular weight substance with 75 carbohydrate, 20% peptide core and the remainder consisting of variable amount of sulphate, bound counter ions and water. Only 5 monosaccharides occur in mucus: (1) fucose, (2) galactose,•(3) sialic acid; (4) N-acetyl glucosamineand (5) N-acetyl galactosamine. These are branching side chains of 5 to 15 residued per chain linked to the peptide 'core'via an O-glycopsidic bond to either serine or threonine (LaMont <u>et al</u>., 1984; Allen, 1981; Smith and LaMont,

1985a).,

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Mucus secretions can be classified into 3 groups on the basis of their histochemical staining: (a) Neutral mucins which stain only with the PAS technique. (b) Sialic acid-containing mucin. Sialic acid is a generic name for various neuraminic acids which differ from each other in whether they are substituted with an acetyl or glycolly\_ residue on their amino acid group and the number of O-acetyl substitutions on the hydroxyl group. (c) Ester sulphate containing mucin. The acidic mucin stains with basic dyes like alcian blue and/or high iron diamine. The distinction between sialic acid and sulphate groups is achieved by avarying the salt concentration or the pH of the staining solution. The specificity of these techniques is increased by the use of enzymes to remove the sugar moietes; ie. neuraminadase selectively removes sialic acid (Reid et al., 1985; Pearse, 1968; Spicer, 1965). It is therefore possible to specifically characterize the various types of mucins secreted based on their staining properties.

Function of mucus in the G.I. tract

There are three main functions of mucus secreted by epithelial cells of the gastrointestimal tract.

(1) Protect the delicate mucosal epithelium of the G.L. tract from damage by food, fecal contents and vigorous

forces that accompany digestion.

(2) Provide slimy lubrication for the passage of solid material and a diffusion barrier to novious substances.

(3) Retain water and provide a perpetual aqueous, environment for the mucosal surface (Allen, 1981; Smith and LaMont, 1985b).

Role of mucus in gallstone formation

Mucin, a high molecular weight glycoprotein, is secreted by the gallbladder and biliary duct epithelium. It exists as a sol phase of monomers in the luminal secretion and gel phase of polymers adherent to the epithelial surface. A common feature in all animal modeles of gallstone formation has been mucus hypersecretion and /or concentration in all cholesterfol stones (LaMont et al., 1984). Mucus increases the wiscosity of bile. Gallbladder cin shares with other epithelial mucins the ability to bind lipids, bile pigments and calcium (Smith and Latont, 1983; Forstner and Forstner, 1975). It is the protein core of mucin which provides a hydrophobic domain that does the actual binding. The lipid binding properties of mucin may contribute to the nucleation process by providing a hydrophobic microenvironment in the mucu's gel which will be favourable to nucleation of cholesterol monohydrate

crystals either from the micellar phase or from a mesophase of redithin-cholesterol liquid crystal (LaMont <u>et al.</u>, 1984). Womack <u>et al.</u> (1963, 1971) demonstrated the presence of mucus at the center of human gallstones and showed histochemically that there was a thin layer of mucoid substance at the surface of the diseased gallbladder epithelium. Previous studies have shown that mucus hypersecretion actually precedes stone and crystal formation and is an an integral part of the nucleating process (Lee <u>et al.</u> 1981b; Freston <u>et al.</u>, 1969).

Mucin may also be involved in the pathogenesis of pigmented stones. Maki <u>et al</u>. (1971) reported the presence of sulphated glycoprotein in human pigmen. I stones. These concrements have a 3-20% glyprotein-mucin-bilikubin complex (Smith and LaMont, 1985a).

Previous studies have shown that it is difficult to measure muchs synthesis and secretion directly. Indirect measurements were done by estimating mucus gland hyperplasia or the viscosity of gallL'adder bile (LaMont <u>etfal</u>., 1984) Recent studies using organ culture explants from prairie dogs with radiolabelled glucosamine and autoradiography found that 80% or mode of this precursor was incorporated into synthesized mucus granules. This technique appears to provide a convenient and quantitative method to

measure the rate of myclus synthesis and secretion (lee et al., 1981b; LaMont et al., 1983).

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The stimulus for mucus secretion in the presence of lithogenic bile is unknown. However, bile duct ligation appears to prevent hypersecretion in the cholesterol-feil prairie dog suggesting that the stimulus is present in hepatic bile (Lee <u>et al.</u>, 1981b). Interestingly, gallbladder mucin release increased with age, an observation which may explain the increase in gallstone formation with age in experimental animals and man (LaMont et al., 1904).

### Factors régulating mucus secretion

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Mucusiis released by exocytosis of secretory granules in the apical portion of the gallbladder epithelial cell (Wahlin <u>et al.</u> 1074; Lee 1980).

(1) Cholecystokin<sup>4</sup> causes a rapid increase of mean release from its normal basal rate (Wahlin et al., 1996).

(2) Prostaglanding are known to stimulate mucin releasein the stomach and may also be involved in the release (t. )gallbladder mucin (LaMont et al., 1983).

(3) Arachidonic acid causes a 2 to 5 fold increase is mucin release in prairie dog gallbladder while indomethacin and aspirin block such a release by inhibiting the

synthesis of prostaglanding (LaMont et al., 1983; Lee et

<u>al.</u>, 1981a).

ς.

(4) Hydrocortisone and related steroid hormones inhibit gallbladder mucin release by inhibiting phospholipase  $A_2$  and thereby reducing the availability of arachidonate for prostaglandin synthesis.

(5) Lysolecithin, a membrane bound 2 itter ion, is a potent stimulator of gallbladder mucin release and hypersecretion (Neiderhiser et al., 1983).

Crystal and stone formation

Cholesterol gallstones are formed from cholesterol monohydrate crystals which are found in lithogenic bile (Small, 1980; Sedaghat and Grundy, 1980). There are tive stages in the formation of symptomatic gallstone disease. (1) There is a genetic trait or metabolic abnormality that leads to the production of a supersaturated bile; (2) a chemical stage involving the production of a supersaturated bile; (3) a physical stage when cholesterol monohydrate crystals precipitate out from supersaturated bile; (4) precipitated crystals aggregate to form macroscopic stones; (5) macroscopic stones cause symptoms by initiating cholecystitis or block the cystic duct.

Before cholesterol crystal precipitation can occur nucleation must take place. There are two types of nucleation: Homogenous nucleation takes place when crystals precipitate out from a highly supersaturated bile. Heterogenous nucleation can occur from a less supersaturated bile and is induced by other substances such as mucus (Small, 1980; Whiting and Watts, 1984; Levy et al., 1984).

Cholesterol monohydrate crystals are described as colorless, transparent and thin with parallel edges, often naving a notched corner and resembling pieces of broken windowpane (Juniper and Burson, 1957). They have also been described as being flat, plate-like parallelograms (Small, 1980), or rhomboidal (Osuga <u>et al.</u>, 1974; Ogata and Murata, 1971).

Not that many years ago no generally accepted theory of formation and growth of cholesterol gali : les existed. The major point of disagreement centered on the process of growth, whether it occurred from the nucleus outward (Naunyn, 1921), or from the periphery inward (Sweet, 1935; Tamura, 1943). Observations made on human stones (Ogata and Eurata, 1971), human stone development (Osuga et al., 1975) and development of experimentally-induced stones in the squirrel monkey (Osuga <u>et al.</u>, 1974) have clarified this question and confirmed the inficial theory put forward by Naunyn (1921) that growth occurred form the center

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outwards.

The morphology of human cholesterol gallstones was described in two studies using light and scanning electron microscopy (Ogata and Murata, 1971; Osuga <u>et al.</u>, 1975). The latter study proposed a chronological model for the development of macroscopic stones. Rhomboidal cholesterol monohydrate crystals form platy units either by layered growth or random aggregation. These plat, units are the basic building units for macroscopic stones which may be either spherical, multilobar or mulberry. The central area of macroscopic stones contained crystals in platy unit form as well as amorphous substance and is connected to the outer capsulo by radially arranged plates.

There has been only one study on an animal model to chronologically follow the development of cholesterol gallstones. The squirrel monkey when fed a diet of butter and cholesterol for nine months developed multilobar and mulberry gallstones. The pattern of stone development included the lamination or random aggregation of crystals into platy units. These platy units were the basic building blocks of microliths or macroscopic stones. An alternative route of stone development was the aggregation of wagch wheel shaped concrements into microliths and macrostones. ?

The basic sequence of crystal precipitation and stores formation was similar in man and the squirrel monkey. However, there were some noticeable differences such as: (1) fragmented crystals seen frequently in squirrel monkeys were only occasionally observed in man; (2) a tiny hole filled with air in the center of squirrel monkey's store was not seen in human concrements; (3) finally, the presence of microliths with a wagon wheel design dommon in the squirrel monkey gallbladder and hepatic bile was not a feature of human gallstores.

### <u>Muche</u> changes

Even though there is no statistical evidence to link gallstones with carcinoma of the gallbladder, a causal relationship has been postulated (Dichl, 1980). The turnover rate of normal gallbladder epithelium is very slow (Bargmann, 1959; Dyban, 1973; Scott, 1978). In a number of studies of cholelithiasis utilizing animal models, an increased cell proliferation was observed before the presence of stones (Scott, 1978; Putz and Willems, 1981; Scott, 1976; Marsch-Zeigler and Palme, 1982; Lee and Forth, 1982). Scott (1978) examined the labelling indices of grabbits fed dihydrocholesterol diet, mice fed a cholesterol -cholic acid diet, and guinea pigs injected with lincomycle. showed labelling indices of 15-21; within the first week of

the cholelithiasic process. Increased mitotic and DNA synthetic activities also were observed in human lithiasic gallbladders (Lamote <u>et al</u>., 1983; Putz and Willems, 1978).

Kaye <u>et al</u>. (1966) examined cell replication if rabbit gallbladder and described proliferative zones in the valleys of mucosal folds. However, Mueller <u>et al</u>. (1972) demonstrated that valleys and folds were transient structures depending on the fullness of the gallbladder when fixed and Scott (1974) found no defined proliferative zones in the guinea pig gallbladder.

There have been three studies that systematically investigated the later stages of cell proliferation and accompanying mucosal changes, correlating them with stone formation (Marsch-Zeigler and Palme, 19 2; Lee and Scott) 1932; Lee <u>et al.</u>, 1986). In the mouse beiesterel-choic acid model, hyperplasia occurred within 6-3 days together with increased mitotic and fibroblast labelling indices ( Marsch-Zeigler and Palme, 1982; Lee and Scott, 1982). The latter authors also demonstrated muscle thickening and glandular metaplasia by 21-28 days and distinct Rokitansky Aschoff sinuses by 29-56 days. All these mucosal changes occurred long before the presence of stones. The gallbladder epithelium of rabbits fed a 15° oleic acid diet for 1-5 weeks showed interepithelial cell vacuoles and by
16 weeks Rokitansky-Aschoff sinuses were observed.

Scanning electron microscopy of pathological human gallbladders revealed defects in the epithelial sheet which were interpreted as empty goblet cells or the result of " chronic cholecystitis (Myllarniemi and Nickels, 1977) William and Smith, 1978).

Cholesterolosis lesions occur quite frequently in human gallbladders (Miettinen and Tilvis, 1985). There is only one study on the dog cholesterol-cholic model that examined the induced lesion (Holzbach et al., 1977). Both human and the dog studies used a single sampling interval.

Previous studies on the guinea pig gallbladder in <u>situ</u> using radiolabelled oldic acid, cholesterol and decithin, have demonstrated that this organ is capable of absorbing materials other than water and electrolytes (Neiderhiser <u>et al.</u>, 1071; Seiderhiser et al., 1073; Neiderhiser <u>et al.</u>, 1976; Hopwood <u>et al.</u>, 1983). Studies on human gallbladder cholesterologis have shown that lipid accumulation can occur either in epithelial cells, the lamina opria or both (Kouroumalis et al., 1984; Ross et <u>al.</u>, 1986; Koga, 1985; Nevalainen and Laito, 1972; Boyd, 1922; Illingueth, 1929; Hora and Schulz, 1970; English and Hopwood, 1985). Koga (1985) demonstrated a massive accumulation of lipids in the epithelial cells and also in macrophages found in the lamina propria of human gallbladders while Nevalainen and Laito (1972) showed lipid accumulation in epithelial cells only. In dogs fed cholesterol-cholic acid for nine months, lipid accumulation was seen only in the epithelial cells and not in macrophages (Holzbach <u>et al</u>., 1977).

Previous studies have proppsed that absorbed lipids reached the lymphatic or venous system via the intercellular spaces and passed to the lamina propria where they were engulfed by macropha (0.3 which then entered the general circulation (Niderhiser et al, 1976; Kouroumalis et al., 1984; Koğa, 1985; English and Hopwood, 1985).

Cells may also rid themselves of the absorbed lipid. by heterophagocytosis, which resulted in the formation of residual bodi (Nevalainen and Laito, 1972; Hora and Schultz, 1970), or esterification of the absorbed lipid to give discrete lipid droplets (Miettinen and Tilvis, 1985; Subbiah and Dicke, 1977; Tilvis <u>et al.</u>, 1982).

Characterization of lipid accumulation in the mucosa was done histochemically (English and Hopwood, 1985; Koga, 1985; Koga et al., 1975; Williamson, 1969; Okros, 1968):

Frozen sections stained with oil red O demonstrated neutral lipids while digitonin complexed experiments revealed free. cholesterol.

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Animal model. of gallstone disease

It is difficult to chronologically evaluate. hepatobiliary physiology or gallstone formation in humans because cholecystectomized gallbladders are generally in the end stage of the disease process. There is little if any, normal tissue. This results in major gaps in our understanding of the disease process and hence the use of animal models to fulfil this role.

The search for animal models to study galibladder disease is ongoing. There is no existing animal model that perfectly parallels human gallstone disease as cholelithfasis is multifactorial and involves: (a) biliary chole-(sterol supersaturation, (b) abnormality of biliary proteins, (c) relative gallbladder stasis, (d) hypersecretion of mucus and (e) other potential defects.

In all the models, induction of cholelithiapic requires manipulation of the exogenous factors is. cholesterol overload which in turn profoundly alters the normal metabolic state resulting in the supersaturation of bile.

Freston and Bauchier (1968) listed a number of characteristics and the ideal model of experimental cholelitaiasis against which all available models may be compared. The criteria were: (1) simple and reproducible method for induction; (2) a diet comparable to a typical human diet if dietary method of induction is used: (3) animals should be easy to procure and reasonably inexpensive; (4) reasonably rapid stone formation, forexample within several weeks rather than months; (F) gallstone composition similar to that of human, whether cholesterol or pigment stones are induced; (6) adequate amounts of tissue, bile, blood and adequately sized bile duct to allow for canulation to sample hepatic bild and enough stones for complete quantitative analyzis; and (7) other than the production of gallstones, there should be no other difference between experimental and control animals. A number of animal models have been developed to study both pigment and cholesterol gallstones; each one had an .identifiable deficiency (Freston and Bouchier, 1968; Gurl DenBesten, 1978; Hoffmann, 1984; Holzbach, 1984; Schoenfield, 1972; (Tepperman and Weiner, 1963; Van der Linden and Bergryn, 1977; Patton et al., 1961; Malet, 19\$5).

# <u>Pigment stones</u>

#### Mongrel dogs

A 1% cholesterol diet supplemented with 10% cusein, 50% sucrose, 26% starch, and 5% lard induced pigmented stones in dogs within one week and after 12 weeks on the diet all animals developed stones. The stones induced consisted of cholesterol, bilirubin and protein. Some calcium and bile acids were also contained within the concrements. The cost and size of the animals suggest that this model is not ideal for the study of cholelithiasis-(Malet, 1985; Van der Linden and Bergman, 1977).

## Hamster

Pigment stones consisting of calcium, phosphate, bile salts and pigment are developed in females and older hamsters. A number of diets including the addition of 30° ethanol to drinking water, lithogenic diets supplemented with thyroxine, and a 13 cholesterol-enriched diet induced pigment stone formation. These stones were basically similar to those in man (Malet, 1985; Van der Linden and Bergman, 1977).

Guinea pig

Guinea pigs of either sex injected with lincomycin subcutaneously for seven days develop pigmented stones. The composition of these stones included 25% calcium, 23% carbonate, 14% bilirubin and 7.7% cholesterol. This model provided more insight into gallbladder mucosal injury than actual stone formation. The stones formed contained a higher percentage of cholesterol than pigment stones (Malet, 1985).

Mice

Mice with hereditary new lytic anemia developed pigmented stones while ingesting a standard rat chow diet. The stones formed within 7-10 conthe in 77% of females but by 10-14 months only 56% of the males had developed stones. The concrements formed were similar to pigmented stones of man. This model may be suitable for studying the pathogenesis of pigment stones associated with anemia (Malet, 1985).

Cholesterol gallstones

## <u>Monkeys</u>

Several species of subhuman primates on long term,

high cholesterol diets, are known to be resistant to cholesterol gallstone formation, noteably the rhosus and African green monkey. Although chimpanzees and other species may show susceptability to cholesterol stone formation, cost of accquisition is prohibitive. Squirrel monkeys fed a diet of 25.5% butter and 0.5% cholesterol added to case in and sucrose developed gallstones within nine months in 83% of the animals (Osuga <u>et al.</u>, 1974). However, the long term feeding process (0 months) and accedent to primate, facilities often makes holding large numbers of animals for study at one time unfeasible (Holebach, 1984).

<u>Rat</u>

The rat, an animal which lacks a <u>sallhladder</u>, takes up cholesterol rapidly when fed a high cholesterol diet. The increased cholesterol level stimulates the enzyme cholesterol 7 Y-hydroxylase which converts cholesterol to bile acids. In this species, the administration of cholesterol does not induce the formation of supersaturated bile (Van der Linden and Bergman, 1977).

# <u>Mouse</u>

In the mouse model, both 18 cholesterol and 0.52 cholic acid is fed to the animals and gallstones are forced within two months. The cholic acid blocks the induction of bile acid synthesis and cholesterol increases in the cholesterol pool, resulting in more cholesterol secreted into the bile. The bile acid ingestion is a necessity for gallstone formation and hence is a disadvantage for comparison with human gallstones (Holzbach, 1984; Malet, 1985).

<u>Guine pigs</u>

Guinea pigs fed 0.5% cholesterol diet devoid of vitamin C developed cholesterol gallstones within five weeks. This major dietary deficiency, when compared to humans, suggested this model was not useful in the study of gallstone formation. However, due to the mechanism involved in stone formation, this model is useful to study the hepatic 7 Y-hydroxylase enzyme system (Malet, 1985).

Gerbils

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A cholesterol-cholic diet similar to that used in the meuse model also induced gallstone formation in this species. However, because of the diet used, the stone formed is not comparable to humans (Van der Linden and Bergman, 1977). Rabbits

Gallstones have been induced in rabbits by two different methods. One model involves reeding the animals a cholesterol analog, dihydroch esterol or cholestanol. The other uses a 40% protein rich diet connect with 20 office acid (Van der Linden and Bergman, 10% Lee et al., 1996). The stones formed were rich in glycodeoxycholic acid, a major bile adde in rabbits. Cholesterol constitutes only about one-fourth or less of the dry weight of these concrements. These stones differed significantly from humans which have a high cholesterol content of between the S5% (Malet, 1985).

Homsters .

Two of the most popular models utilize rödent species: the hamster and prairie dog. The hamster model has been in use for many years. Dam and Christensen (1952) demonstrated that the hamster is capable of forming cholesterol gallstones when fed a diet of refined sugar and no polyunbaturated fatty acids. However, this diet has little similarity to the average human diet. Bile from several animals has to be pooled for chemical analysis because of the small size of the animal gallbladder. They also developed diarrhea and exhibited growth retardation (Chang et al., 1973). Despite these drawbacks, the hamster model has, proved to be popular as acquisition and maintenance are relatively inexpensive and they form stones within 6-7 weeks (Van der Linden and Bergman, 1977). The work on this model has focussed more on cholesterol metabolism, prevention and/or dissolution of gallstones (Pearlman et al., 1979; Hoffmann, 1984).

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<u>Prairie dog</u>

The herbivorous prairie dow\_is a model of unsurpassed popularity. Bronneman et al. (1952) were the first to realize the true potential of this species. These animals, fed 1.25% cholesterol-enriched diet, developed cholesterol gallstones within two weeks. The caloric distribution of the dict is similar to that of humans and the stones formed contained 75-85% cholesterol by weight. The amount of bile produced by each animal was sufficient for chemical analysis of cholesterol, phospholipids, and bile acids, and the animals remained maithy on the lithogenic diet Major problems as a model are that the animals are difficult. handle, seasonally unavailable and genetically hetero genous. Its major advantages are that they are comparatively inexpensive to acquire and maintain and their bile is comparable to that of humans insofar as it contains the same four major bile acids. The two primary acids: (a)

cholic acid and (b) chenodeoxycholic acid and two secondary acids: (a) deoxycholic acid and (b) "lithocholic acid (Van der Linden and Bergman, 1977; Chang <u>et al.</u>, 1973; Malet, 1985). In addition, the relative composition of the three main lipids are roughly comparable to those found in human bile (Van der Linden and Bergman, 1977; LaMont et al., 1984).

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The prairie dog model has been extensively used to provide information on a variety of parameters involved in gallstone formation: bile composition (DenBesten et al., 1974), gallbladder mucus hypersecretion (Lee et al., 1981); Doty <u>et al.</u>, 1983a), prevention of mucus hypersecretion by the administration of aspirin (Lee <u>et al.</u>, 1981a), inhibit -ion of bile stasis and stone formation with sphineterotomy and reversal with atropine (Hutton <u>et al.</u>, 1981a), i effects of bile acids and axozolings (Cohen <u>et al.</u>, 1984), , prevention of stone formation by hypersecventic acid (Singhal <u>et al.</u>, 1984), and the role of cyclic nucleotides and glycoproteins during stone formation (Zak et al.,

The scarcity and protected status of

1984).

Ground squirrels



Canada led Davison <u>et al</u>. (1982) and Fridhandler <u>et al</u>. (1983) to further develop Patton <u>et al's</u> (1961) ground squirrel model for cholelithiasis. The Richardson's ground squirrel, when fed a 1% commercially prepared cholesterolenriched rat chow diet, exhibited bile with an increased lithogenic index close to theoretical saturation values after two weeks. These authors did not observe the precipitation of crystals but described defective gallbladder contractility during this time interval. Prolonged exposure of animals to the enriched diet gave a lithogenic bile and resulted in either cholesterol •

Richardson's ground squirrels are plentiful in Alberta and are considered to be a pest by farmers. The classification of this species is:

Kingdom----- Animal Pholum----- Chordata

•young).

Order----- Rodentia (gnawing animals; enamel only on front incisors) Suborder----- Sciuromorphs (post orbital process, four cheek teeth above

and below). Squirrels, beavers, gophers, marmots,kangaroo rat, chipmunk,prairie dog and groundhog.

**Objectives** 

Genus-

Species

This model had been used previously to study bile lithogenicity and gallbladder stasis during the early stages of cholesterol formation (Patton <u>et al.</u>, 1061; Fridhandler <u>et al.</u>, 1983). A thorough systematic investigation of this model in the cholelithiasic process however was lacking. The current study was designed to : (a) describe the morphology of normal ground squirrel gallbladder

Spermophilus

richardsonii

- (b) identify the chronology of crystal and stone formation
- (c) describe the morphology of these biliary concrements
- (d) describe intra-epithelial and mucosal\_changes cocurring prior to, during and after stone formation.

#### MATERIALS AND METHODS

The bile and gallbladders of 238 Richardson's ground squirrels (Spermophilus richardsonii) of both sexes were studied. Ground squirrels weighing between 350-450 gm were trapped from the wild around the Edmonton, Alberta area and housed individually in plastic wired-topped cages at the Surgical Medical Research Institute (S.M.R.I.) Animal Holding Facility, University of Alberta. They were kept on a 12:12 hour (6 p.m.-6 a.m.) light:darkness photoperiod at a temperature of 23°C and humidity of 35-40%. The animals were maintained on a diet of rat chow (Wayne Lab-bloc, Allied Mills Inc., Chicago, Illinois) and water ad <u>libitum</u>. Animals were held for one month during which time they were checked for parasites, loss of hair, weight and diarrhea.

Animáls considered to be healthy were divided into control and experimental groups. Experimental animals were fed a 2% cholesterol-enriched rat chow diet (U.S. Biochemicals, Cleveland, OH) and water <u>ad libitum</u>. Control animals were maintained on the same commercial rat chow diet and water <u>ad libitum</u>. Each group consisted of four animals. Sampling intervals of 6, 12, and 18 hours; 1, 3, 5, and 7 days; 2, 3, 10, and 20 weeks; 3 weeks on on cholesterol diet followed by three week on normal diet were

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used. Animals in the shorter time intervals, 6 hc. through 1 day, were fed known quantities of food which were monitored before killing to chaure that they had actually eaten the cholesterol-enriched diet. These animals were not fasted overnight before killing.

Animals were fasted overnight and killed by cervical dislocation between 0900 and 1000 hours. The majority of bile was withdrawn from the gallbladder by a loc syringe and stored at -20°C for assay later. I do do the will was examined by polarizing microscopy to level presence of crystals or microliths. Bicchemical assay of the bile to determine cholesterol, bile acids and phospholipids was performed by the Biochemistr — nit of S.M.R.I. From these parameters the lithogenic index of bile was calculated (Carey, 1978; Thomas and Hoffmann, 1973).

The same volume of fixative, 2.50 gluteral whyle in Millonig's buffer pH 7.2 was injected into the gallbladder. The organ was then excised and immersed in the same fixative for one hour at room temperature, split along iter longitudinal axis and rinsed of its luminal contents.

The luminal rinsings were examined by direct light and polarizing microscopy, photographed and stored in 16 neutral buffered formalin. Luminal rinsings from gall-

bladders that did not contain macroscopically visible ' stones were filtered through a 0.22µm Millex 'GS filter ' (Millipore Corp., Bedford, MA) which was subsequently air-dried, mounted on a stub and sputter-coated with gold. Luminal rinsings that contained macroscopic stones were Piltered through a Whatman (England) =4 filter paper and left to air-dry. The dried stones were transferred onto stubs counted with electron conducting paint and sputter . coated with gold. Crystals and stones isolated by either technique were examined using a Philips 505 SEM. Larger stones were fractured after surface observations were recorded, recoated and the internal architecture examined. Macroscopically visible stones of some animals were washed in distilled water to remove any bile and debris and then dried in a vacuum desiccator until a constant weight was achieved. Stones were dissolved in a mixture of ethanol/ether (3:2 vol/vol) at 40°C for 72 hours in the dark with vigorous mixing. The mixture was then centrifuged at 1500g (Smith and LaMont, 1985a). The colorless supernatant was decanted and stored while the residue was ₀ washed again and recentrifuged. On average, the ratio of stone weight to dissolution solvent was 15mg/ml. The supernatant and washings were combined and assayed for cholesterol content using the method of Carr and Drekter :0 (1956). Analysis was done by the Biochemistry Unit of S.M.R.I.

The gallbladder of the ground squirrel is small and the various techniques listed below Were not all performed on each organ. Similar experiments were repeated and those used as determined by procedure. One longitudinal strip of the organ<sup>w</sup>as diced into 1mm<sup>2</sup> pieces for transmission electron microscopy and immersed in the same fixative for two hours at room temperature, postfixed in 1% osmium tetroxide for one hour, blocked stained in saturated aqueous wanyl acetate for one hour, dehydrated in an ascending series of ethanol and propylene oxide and embedded in LX-112 resin (Appendix I). Three of these blocks were randomly selected for electron microscopy. Ultrathin sections of 600-900A<sup>0</sup> were cut on a Reichert Ultracut microtome; mounted on uncoated 200 mesh copper grids, stained with saturated a polous uranyl acetate for 10 minutes and lead citrate (Reynol ., 1963) for 2 minutes and examined with a Philips 410 electron microscope at an accelerating voltage at 80kv. Micrographs were taken on Kodak Fine Grain Positive 35mm film.

For morphometry of mucus secretory granules, sections from coded blocks where columnar epithelial displayed a nucleus, basal and apical regions were randomly delected and photographed at the same magnification of 4400 on 35mm fine grain positive film and the negativer were enlarged 2 times when printed. Eight to ten cells were photographed

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from each block. From the 24-30 prints from each animal; ten were randomly selected for planimetry. For each group of animals (n=4), 40 prints were digitized. Morphometric study was performed using a Nikon Microplan II semiautomatic image analyser to determine the area occupied by the mucus secretory granules and cytoplasm. The volume density (Vd) of the mucus granules was expressed as a percentage in which the "otal area of granules per cell was divided by the total cytoplasmic area multiplied by 100. Data was expressed as a percentage of means and standard error of the mean (SEM). Analysis of data was performed using the one-tailed unpaired Student's t test (Wahlin et al., 1976; Pradal et al., 1984; Weibel and Elias, 1967). The section thickness and compression factor (Holme's effect) were ignored because they did not affect the, statistical treatment of the data (Wahlin et al., 1976; Pradal <u>et al.</u>, 1984).

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Another longitudinal strip of the gallbladder was . immersed in the 2.5% gluteraldehyde in Millonig's buffer pH 7.2 for four hours at room temperature, postfixed in 1% . osmium tetroxider for one hour, block-stained in saturated uranyl acetate for one hour, pinned on a cork with no tension and dehydrated in an ascending series of acetone. The tissue was critical-point dried in liquid carbon a dioxide using a Seevac critical-point-drier and mounted on

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aluminum stubs using colloidal silver paint (Appendix III). These dried specimens were sputter coated with gold using an Edwards \$150B sputter coater, examined and photographed with a Fhilips 505 scanning electron microscope using Kodak Plus-X Pan 35mm<sup>4</sup> film.

Another longitudinal strip of the organ was processed for light microscopy. The specimen was refixed in 10 buffered formalin for 24 hours, washed overnight in running water, and dehydrated in an ascending series of ethanol to 95%. Each strip of the specimen was further divided longitudinally into two parts. One part was dehydrated to 100% ethanol, cleared in xylene and embedded in paraffin (Luna, 1968); while the other was processed through to embedding in glycol methacrylate (Polysciences Inc. duta sheet #123,=1982). Paraffin, methacrylate and histochemical slides were examined with a Leitz Orthoplan photomicroscope and selected areas photographed using Kodak Panatomic-X film (Appendix II).

To establish a mitotic index (MI), groups of animals (n=8) from sampling intervals of 1,2 10 and 20 weeks were killed, the organ excised and processed into methacrylate. Two micrometer sections were stained with Lee's methylene blue basic-fuschin (Luna, 1968). Slice were examined and the total number of cells and mitotic figures counted. Muclei which appeared between late prophase and telophase Were counted as mitotic figures. The number of nuclei counted per animal (n=8) ranged from 2000 to 3000. % MI = number of mitotic figures counted multiplied by 100, divided by the total number of nuclei counted. 39

For light microscopy autoradiography, animals (n=44) from sampling intervals of 1, 2, 10 and 20 weeks were ased. Prior to killing of animals, they were pulse injection intra ~peritoneally with 1.0 microcurie of tritiated thymidine/g body weight (specific activity 59ci/mmole ICN Canada Ltd) at two one-hour intervals, killed and processed as above for glycol methacrylate. Two micrometer methacrylate sections were dipped into undiluted Ilford K.5D emulsion. The slides were drained and dried in a vertical position for one hour at room temperature, placed in a black box containing drierite, tape sealed and stored in the dark at 4°C for 3 weeks. Slides were developed in Dektol (Eastman Kodak, Rochester, NY) for two minutes, washed in water for 30 seconds, fixed in Kodak fixer for 5 minutes, washed in Water for 15 minutes and counterstained with either methylene blue basic-fuschin or Gill's hematoxylin and Cosin for two minutes (Budd, 1971) (Appendix V). To establish a labelling index (LI), the amount of background labelling determined the minimum grain count necessary to score a cell as labelled. In this study, the background

labelling was moderate and nuclei that had 15 grains or more overlying the nuclei were considered labelled. LI number of labelled cells counted multiplied by 100, divided by the total number of cells counted. Analysis of mitotic and labelling indices was performed using the one-tailed unpaired Student's t test.

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Eight micrometer thick paraffin sections of control (n=4) and one week treated animals (n=4) were deparaffinized and stained with high iron diamine/alcian blue pH 2.5 (Spicer 1965), and potassium hydroxide/ alcian blue pH 1.0/ PAPS (Reid <u>et al.</u>, 1985). The results of these stains are:

high iron diamine/AB pH 2.5

- Sulphomucins: grey-purple-black

- Sialic acids: blue (aqua)

KOH/AB1.0/PAPS:

- Sialic acids with side chain substitutions: red

- tissue vicinal diols: red

- O-sulphate esters: blue (Appendix VI)

For lipid nistochemistry, control (n=4) and one week experimental (n=4) animals were killed and gallBladder tissue was fixed in Baker's formol solution for 24 hours,

54. 1 frozen in isopentane immersed in liquid nitrogen, sectioned in a cryostat and stained with oil red O "(Luna, 1968).

For localization of cholesterol at the electron microscopic level, digitonin-complexing experiments were conducted using control (n=4) and one week experimental (n=2) animals. Tissue was fixed in 2.5% gluteraldehyde for one hour at room temperature. The gallbladder was then split along its longitudinal axis, rinsed of its luminal contents and placed in fresh fixative for an additional 2 hours. The tissue was then washed with phosphate buffer for one hour, changing the solution every 15 minutes, and placed in a 2% digitonin solution dissolved in 0.1M Millonig's phosphate buffer for 4 hours at room temperature (Okros, 1967; Williamson, 1968; Koga <u>et al</u>., 1975). The specimen was post-fixed in 1% osmium tetroxide and then processed routinely for electron microscopy (Appendix IV).

For electron microscopic mucin autoradiography, control (n=4) and one week experimental (n=4) animals were matched by weight, injected with H<sup>3</sup>galactose, 1.0 microcurie/g body wt. (specific activity 5-20ci/mmole, ICM Canada, Ltd.) intraperitoneally, and killed by cervical dislocation at 25 and 40 minutes interval thereafter. The gallbladder was removed and processed as above for electron? microscopy. Ultrathin sections of 600-900A<sup>0</sup> were cut on a Reichert Ultracut microtome and mounted on 200 mesh copper grids attached to glass slides by 0.5% parlodion. Slides were dipped in Ilford L4 emulsion diluted 1:6 in water, allowed to dry for one hour, placed in a black box, taped sealed, and stored at 4<sup>0</sup>C for 4 weeks. Slides were developed in Kodak D-19 for 3 minutes, fixed in 25% sodium thiosulphate for 6 minutes, washed in distibled water for 6 minutes and stained in uran soctate for minutes and lead citrate for 5 minutes prendix IV, Grids were, then examined with a Philips 41 sectron microscope at an accelerating voltage of 801%. Micrographs were taken on Kodak fine grain positive am film. RESULTS

Control

Electron microscopic examination of the gallbladder. wall of the Richardson's ground squirrel revealed that it was composed of three Tayers: mucosa, muscularis and adventitia or serosa. The mucosa was composed of a layer of simple columnar epithelial cells and lamina propria. The lamina propria was a well defined layer under the basal lamina. It was composed of collagenous fibrils, capillaries, arterioles, venules and fibroblasts.

Capillaries were observed in close proximity to the lining epithelium especially in mucosal folds (fig. 1). The endothelium was fenestrated, each fenestration closed by a diaphragm (fig. 2). Pinocytotic vesicles were observed in the endothelium. A continuous basal lamina surrounded the endothelium which was distinct from that of the overlying  $\int$ epithelium.

Beneath the lamina propria the muscularis was composed of smooth muscle bundles arranged in thin sheets of varying extent. The spaces between layers were occupied by characteristic connective tissue fibers and cells as well as blood vessels. In the lamina propria, unmyelinated nerve fibers were observed in close proximity to larger

blood vessels (fig. 3). Piriform swellings, or varicosities of postganglionic neurons were seen intramuscularly and in the lamina propria beneath the basal lamina (figs. 4, 5). Membrane specialization of these varicosities was not observed. They contained granules of various electron density, each of which exhibited a prominent surrounding membrane, a peripheral electron lucent halo and a dense central core similar to those observed in non-direct adrenergic nerves. External to the muscularis the adventitia, or serosa, consisted of lobules of fat, collagenous and elastic fibers, blood vessels, fibroblasts and lymphoid cells.

Three types of cells were observed in the lining epithelium: light, dark or pencil and edematous . Cell classification was based on the shape of the cell and nucleus as well as density of the cytoplasmic matrix.

## Light cells

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The majority of the cellular population in the lining enthelium were light cells (fig. 1). The cytoplasmic density of these cells varied depending on the functional activity and the organelles present. Non-active, or resting cells, contained few organelles and hence the cytoplasm appeared less dense than active cells. Active

cells had a heterochromatic, serrated nucleus that contained a prominent nucleolus. These cells ranged from 19 to 21 micrometers in height and 4.5 to 5.5 micrometers in diameter. A distinct apical zone devoid of cellular organelles, 0.8 to 1.5 micrometers in depth, was present in the cell. The shape of this zone varied from a slight convexity to an inverted U, both exhibiting intact microvilli (figs. 1, 6).

Electron dense mitochondria with transverse cristae were seen in the basal and subapical region of the cell, more numerous in the latter. Other organelles in the subapical cytoplasm were rough endoplasmic reticultam, lysosomes, residual bodies and centrioles (figs. 1, 6, 7). The residual bodies were membrane-bound, of different sizes and contained a variable amount of lipid. A feu-lipid droplets were occasionally observed in the basal region of the cell (fig. 8). No lipid droplets were seen in the lamina propria. Microfilaments were scattered throughout the cytoplasm of the cell but more concentrated near the centrioles (fig. 7). The Golgi apparatus was a prominent supranuclear feature in most cells. It generally consisted of 3 to 5 layered saccules with vesicles of various sizes seen near the maturing face (fig. 9).

#### Mucus secretion

Light microscopic histochemistry revealed that the gallbladder epithelium of control dnimals secreted both sialylated and sulphated mucin at a basal level (figs. 10, 11). The scanning electron microscope provided evidence of "mucus secretion as thin viscoid strands issuing from the luminal aspects of these cells (fig. 12). Secretory activity was not simultaneous through wide areas of the epithelial sheet but appeared restricted to groups of cells scattered over the luminal surface.

Autoradiographic studies at the ultrastructural level of control and seven day experimental animals, using a labelled glycoprotein precursor H<sup>3</sup>galactose, demonstrated the mucin granules to be membrane bound. These granules contained a flocculent material of moderate electron density (fig. 13). Twenty five minutes after intraperitoneal injection of the precursor, those mucus granules which contained the label were found predominantly in the supranuclear and subapical regions of the cell (fig. 14). Forty minutes after injection, labelled mucus was commonly observed in the lumen of the gallbladder (fig. 15).

· Examination of epithelial cells of control.

gallbladder at the ultrastructural level showed mucuscontaining granules of varying diameter in the supranuclear region of cells that contained prominent Golgi complexes. These granules were a characteristic feature of the subapical zone where they appeared to coalesce into larger granules prior to extrusion of their contents into the gallbladder lumen (figs. 16, 17).

The apical plasma membrane exhibited prominent microvilli that ranged in length from 0.8-1.6 micrometers with a diameter of 0.15 micrometers. The terminal web was poorly developed below the microvill<sup>\*</sup>. In a few instances, cytoplasmic bullae were observed to arise between microvilli and contained particulate material but were devoid of cellular organelles (fig. 18).

Mitotic activity was rarely observed in the epithelial sheet but figure 19 shows a cell in the prometaphase stage. Mitochondria in the basal and apice Le regions, microvilli, mucus granules in the apical zone, desmosomes between adjacent cells, a centricle and microfilaments were observed in the dividing cell. Cells undergoing mitoses were larger than those adjacent, the chromosomes were at the same or higher level than the interphase nuclei.

### Dark cells

The dark rod-shaped, or pencil cells, had a higher electro. lensity than the principal light cells (fig. 1). They ranged in height from 19 to 21 micrometers but had a narrower diameter of 3 to 3.5 micrometers at the nuclear level that became wider at the apical surface. The dark cells made contact with the basal lamina and lumen of the gallbladder (figs. 20, 21). The apical surface was populated with microvilli of normal density and dimension. There was a distinct apical zone devoid of cellular organelles but containing granules (fig. 20). Residual bodies and numerous mitochondria with transverse cristae were observed in the subapical and basal region of these cells but a Golgi apparatus was not seen. Dark cells appeared to occur singly, their lateral interdigitating processes standing out strongly against the less electron dense cytoplasm of the light cells. Desmosomes remained intact between light and dark cells (figs. 20, 21).

<u>Edemàtous cells</u>

Light barrel-shaped edematous cells were infrequently observed in the lining epithelium (figs. 22, 23). These cells were wider than the surrounding light cells and displayed a pyknotic or karyolytic nucleus. The

apical membrane was generally ruptured with cytoplasmic organelles extruded into the lumen (fig. 22). Numerous hydrated and vacuolated mitochondria with disrupted and distorted cristae and a lower electron density than normal were observed. Venicular profiles of degenerated rough endoplasmic reticulum and lysosomes were also observed in the edematous cells (fig. 23). No other cellular organelle was observed.

A prominent basal lamina lay under all three cell types. It was continuous and followed the contour of the basal plasma membranes. The attachment of the plasma membrane to the basal lamina was by numerous peg and socket interdigitations rather than hemi-desmosomes (fig. 24). These interdigitations were more prominent in organs not actively involved in water resorption at the time of fixation.

The lateral cell membranes showed finger-like interdigitations, or microfolds, between adjacent cells. In the apico-lateral region the plasma membranes formed a prominent junctional complex between neighbouring cells. Extending down the lateral membrane to the basal plasma membrane, prominent desmosomes were observed. The lateral membranes between cells often exhibited various degrees of separation with dilation of the intercellular space. The

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apical regions were held closely apposed by the junctional complexes while the basolataral regions displayed a superation of the lateral interdigitations to become widely distended (fig. 25). The desmosomes in these regions remained intact anchoring the finger-like projections across the dilated intercellular space (fig. 26). The basal plasma membrane remained apposed to the basal lamina although the peg and socket interdigitations common to this region were no longer observed (fig. 27).

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Small round infiltrating leucocytes or observed migrating through the basal lamina into the interce lular spaces (fig. 28). Leucocytes were generally observed near the basal lamina and occasionally at Higher levels approximating the junctional complex (fig. 29). The most freq: 'y encountered cell type was the lymphocyte.

The scanning electron microscope demonstrated mucosal fords, or rugae, on the luminal surface of the gallbladder. The number and height of these folds was dependent on elative volume of fluid in the organ attime of fixat. figs. 30, 31). Semi-filled organs exhibited pocket-like depressions between mucosal fords? These pockets disappeared when the gallbladder was full of fluid, the surface showing only widely spaced, relatively small bumps. Sections through these projections failed to demonstrate any underlying structure responsible for their formation.

Higher resolution scanning microscopy revealed the cobble-stone appearance of the epithelial sheet. The apical surface of these cells were dome-like and protruded to variable degrees into the lu en (fig. 12). Examination of fractured edge revealed the extensive lateral membrane interdigitations (fig. 32). A deeper from the showed the large oval nuclei located near the base of the cells (fig. 33). Surface morphology of the basal lamina showed projections that may interdigitate in a peg and socket arrangement with the overlying epithelial cell. Small defects in the basal lamina were also noted and may be the result of the thixotropic nature of this layer (fig. 34).

Experimental animals

#### Mucus secretion

The 2% cholesterol-enriched diet stimulated an increase in mucus secretion by the gallbladder epithelium as early as 18 hours following ingestion, clearly demonstrated by scanning electron microscopy (fig. 35). This hypersynthesis and secretion of mucus was confirmed a quantitatively by morphometry on the ultrastructural level

(table I, page 64). Mucus hypersecretion occurred when the lithogenic index was 0.742 (table II, page 65), ificantly higher than control animals. In one and the day treated animals, SEM studies revealed a thick sludge-like layer covering the epithelial surface (figs. 36, 37, 38). TEM studies showed numerous mucus granules in the apical and subapical regions of the epithelial cells. The secretory granules maintained their discrete shape but the luminal surface of the cell changed from a slight convexity to an inverted U (fig. 39). Scanning EM studies continues to demonstrate a thick sludge-like layer during the one and two week intervals which was observed throughout the experimental period (figs: 41, 42). The continued hypersecretion of epithelial glycoprotein, coupled with other bile constituents such as bilirubin, formed the sludge layer. The extent of this layer, both in thicknessand area increased throughout the experimental periody This intraluminal environment appeared to be suitable for the continued growth of these crystals into platy units, microliths and stones.

In experimental animals and occasionally in controls, cytoplasmic bullae were observed on the luminal surface of epithelial cells. These bullae were devoid of microvilli (figs. 40, 43). Goblet cells were not observed either in control or experimental animals.

Morphometric studies confirmed mucus hypersecretion throughout the experimental period (table I, page 64), although at a lower level in the 10 and 20 week treated animals. Animals fed a lithogenic diet for 3 weeks and then control diet for 3 weeks, indicated that the rate of mucus secretion returned to control values (fig. 44).

# Crystal precipitation and stone formation

Scanning electron and polarizing more copy failed to demonstrate the presence of cholesterol monohydrate crystals on the luminal surface or in the bile of control animals. Biochemical analysis of the bile components from control gallbladders indicated the lithogenic index (L.I.) averaged 0.356 while that of animals on the experimental diet, with evidence of ingestion, for 6 hours was 0.337, thus exhibiting no significant increase (table II, page 65). By 12 hours however, the lithogenic index had risen significantly to 0.568. Neither group exhibited alterations in epithelial morphology or physiological activity detectable by SEM. The first evidence of altered mucosal physiology occurred in animals after 18 hours on the lithogenic diet.

One day after ingestion of the diet the lithogenic index of the bile had risen to 0.974. Scanning electron microscopic examination of the mucosal surface showed large numbers of rhomboidal cholesterol monohydrate crystals precipitating from the bile, many of which had the notched appearance noted in humans by Juniper and Burson (1957). The crystals were observed stuck between the microvilli on the luminal surface of the epithelial cells (fig. 45). Appearance of the precipitated crystals was indicative of cholesterol saturation the bile, a physiologic feature supported by biochemical analysis. Animals on the diet for 3 days had a lithogenic index of 1.133 and continued to exhibit cholesterol monohydrate crystal precipitation. Animals on the diet for 5' days, with a lithogenic index of 0.989, continued to demonstrate crystal precipitation by SEM. The occurrence of platy units was observed in the luminal surface in these animals (fig. 38).

Table II showed the variation in lithogenic indices of bile during the remainder of the experimental period. It indicated the significant increase of the lithogenic index during the early experimental period which leveried off after the first seven days to a value in the 0.651-0.842 range through the first 10 weeks. The lithogenic index by 20 weeks averaged 2.615, still significantly higher than controls. Animals in the 20 week group that had their cystic ducts filled with stones had lower lithogenic indices than those whose ducts were not

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occluded. The difference in values however, was not significant at this point.

One week animals continued to Memonstrate cholesterol

monohydrate crystal precipitation observable by SEM (fig. 46). After two weeks on the diet, the number of aggregated crystals seen on the mucosal surface was diminished. Examination of luminal contents by direct light microscopy revealed the presence of concrements. Luminal rinsings filtered and examined by SEM demonstrated continued aggregation, and growth of crystals through parallel lamination into larger platy units (figs. 47; 48). After three weeks, luminal rinsings contained refractive microliths, pure white in color and observable by light microscopy. The SEM showed these concrements to be clusters of platy band units (fig. 49). These platy bands are regarded as the basic unit required for the formation of gallstones in this model. These 3 week microliths had an irregular external surface formed predominantly by a random subparallel aggregation of the smaller platy subunits. This configuration gave the appearance of being arranged radially:

At ten weeks, concrements removed from the luminal washings were macroscopically visible and appeared to be of varying size (figs. 50, 51). This array of varying configuration was illustrative of the aggregative, growing
process of stone formation. The largest stones observed at 10 weeks exhibited a yellowish color and measured roughly 1mm. in diameter by direct light microscopy. Examination of these stones by SEM showed them to be composed of an aggregation of a number of cholesterol microliths. The surface was not smooth and the aggregate-like appearance gave these stones a typical mulberry shape (fig. 52). Higher resolution scans revealed that often these clusters were free of any matrix-like material in the grooves between them, adhering to one another through interdigitation of the crystals and platy units near each surface (fig. 53). In other areas of the stone however, component subunits were observed embedded in a thick layer of sludge on the surface of the stone (fig. 54).

Visual examination of the cystic duct and gallbladders of 20 week animals revealed many of the dystic ducts to be severely congested with a yellowish material while the bile was watery and turbid. The lumon of many organs was filled with these concrements. Light microscopic examination of the material rinsed from the organs and ducts demonstrated concrements that ranged from a fine sandy material to macroscopically visible stones. The largest of these stones was approximately 2mm. in diameter and those over 1mm. were generally brownish-yellow in. color. Examination of these stones with the SEM showed a smoother surface with occasional areas still exhibiting the parallel array of crystals and platy units comprising each lobe of the mulberry shaped stone (figs. 55, 56). Fracture of the stones revealed the formation of a much smoother, homogeneous, capsule-like outer layer which at higher resolution was observed to be composed of cholesterol plates that extended down into the central region of the stone in a radial fashion (fig. 57). These stones were not solid enough to allow a smooth fracture and the existence of a central cavity was not observed. The contral region of the stone was composed of aggregation of leaf-like cholesterol crystals (fig. 58).

Biochemical analysis of the concrement aggregations revealed them to have a cholesterol content ranging from 61 to 74%, the amount increasing with prolonged exposure to the diet.

#### Mucosal changes

#### Cell proliferation and damage

Mitoses and labelled cells were observed throughout the epithelial sheet of control and also experimental, animals and did not show any preference for either the valleys or crests of the mucosal folds (figs. 59, 60).

Light, electron microscopic and autoradiographic studies revealed a low mitotic activity with a mean mitotic index (MI) of 0.027% and a labelling index (LI) of 0.160% (table III, page 66).

In animals exposed to the lithogenic dict for one week, the MI and LI were significantly higher than control values with a mean MI of 0.250% (p<0.05) and LI of 0.394 (p<0.01)<sup>8</sup> (table III, page 66). The highest frequency of mitoses and labelled cells occurred in the two week-treated group which exhibited a mean MI of 0.384% (p=0.01) and LI of 1.312% (p<0.001). Transmission EM examination of tipsue from the latter group revealed prominent mitoses on the crests of mucosal folds (fig. 59). Light microscope autoradiography demonstrated labelled cells on both the sides and valleys of the folds (fig. 60). Crystal and microlith formation was observed during these early stages of the experimental period but no macroscopic stones were seen.

Groups of light, barrel-shaped edematous cells with pyknotic or karyolytic nuclei were frequently observed in the 10 and 20 week experimental animals by TEM (fig. 62). The apical membranes of these cells were often ruptured with cytoplasmic organelles extruded into the lumen. SEM observations revealed gaps in the epithelial sheet (fig. 61) while TEM demonstrated that adjacent epithelial cells slid under the degenerated cells to protect the basal lamina (fig. 62). During these later stages of the experimental period macroscopically visible stones were commonly observed.

In 20 week cholesterol-fed animals areas of the epithelial sheet revealed hyperplasia and hypertrophic cells on the crests of mucosal folds examined by light and scanning electron microscopy (figs. 63, 64). The normal hexagonal configuration of the epithelial cells was lost and their boundaries were indistinct. Frominent mitoses were often observed near hyperplastic lesions (fig. 64). The nuclei of the columnar cells were no longer basal, the cells themselves exhibited a variety of shapes. This arrangement gave the appearance of being pseudostratified.

The muscle layer of the gallbladder in 20 week animals appeared thicker, especially where prominent diverticula configurationally similar to Rokitansky-Aschoff sinuses were seen extending down to the adventitia (fig. 65). Electron microscopic studies revealed the presence of plasma cells and other loukcrytes in the lamina propria and between the basal lamina and epithelial cells of these animals (figs. 66, 67).





Examination of sections from 10 week cholesterol-fod animal continued to demonstrate a significant increase in the number of mitoses (p<0.005) with a mean MI of 0.329, while in the 20 week group, animals exhibited a mean MI of 0.289% (p<0.005). Table III indicated that the fI values for both groups followed this general trend. The incidence of mitoses and labelled cells at this point were lower than the peak values of the two week group but remained significantly higher than in control animals. Labelled fibroblasts were repeatedly observed in the lamina propria and adventitia as early as two weeks in animals fed the lithogenic diet, but were infrequent in the controls (figs. 68, 69).

Cholesterolosis-like flesion

In the twelve hour treated animals, the intercellular spaces were distended and contained electron lucent material with dense particulate material and osmiophilic bands among them, (fig. 70). Similar material was also observed intracellularly, especially in the apical and supranuclear regions of the cells. The apico-lateral membranes remained firmly intact.

One, three, five and seven day treated animals exhibited numerous residual bodies within the epithelial cells (fig. 71). The cells continued to accumulate lipid in massive amounts in the supranuclear region while basal regions contained discrete lipid droplets of varying electron density (figs. 72, 73, 76).

Frozen sections of control animals stained with oil red O failed to demonstrate lipid material in epithelial cells while in a five day-treated animal showed neutral lipid in the supranuclear region. (figs. 74, 75).

Electron microscopic observations on these tissues revealed that the supranuclear accumulations were nonosmiophilic and predominantly electron lucent exhibiting some flocculent material. At higher magnification discrete lipid masses were observed around the periphery of the accumulations. Centrally however, these droplets impinged upon one another, coalesced and became an ill-defined mass. Scattered among the lipid masses were lysosomes which appeared to be ingesting portions of the lipid mass to form residual bodies. Endoplasmic reticulum, mitochondria and Golgi of normal appearance were seen around the periphery of these accumulations (fig. 73).

Tissue from seven day treated animals, complexed with digitonin, revealed that the supranuclear accumulations contained free cholesterol incorporated into the overall

hass (fig. 77). Discrete lipoid masses were observed in dilated smooth endoplasmic reticulum appeared to the membrane bound. Numerous ribosomes and a few lysosomes were often observed near these lipid accumulations. Examination of the residual bodies at higher magnification revealed digitonin precipitation indicating the presence of free cholesterol (fig. 78), Digitonin-complexed cholesterol crystals were also observed on the luminal surface of the repithelial cells (fig. 79). The density and staining property of this precipitation was similar to that seen in the supranuclear accumulations and residual bodies.

One, two and three week treated animals also revealed supranuclear accumulations and distinct lipid droplets in the basal regions of epithelial cells. These droplets had distinct osmiophilic rings and central regions of varying electron density (fig. 76). Digitonin-complexed tissue failed to demonstrate the presence of any free cholesterol in the basal area of these cells.

The epithelial cells of ten and twenty, week treated animals continued to demonstrate accumulation of lipits in the supranuclear and basal regions, but now also in the lamina propria. The electron density of the lipid in the lamina propria ranged from electron lucent to highly osmiophilic. Many of these lipid droplets had a myelihoid

http://mode (fig. 00). Acrophages exhibiting dense bundophilic boules v le also observed in the lamina propria fill. 30, Foam coll however, were not observed.

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TAB	LE	Ι	:	
	2.4			

Intervals	Mean ± SEM
Cetrol	1.395 ± 0.180
6 hours	$1.716 \pm 0.234^{\#}$
12 hours	1.759 ± 0.281
,18 hours	2.367 ± 0.260
21 hours	3.680 ± 0.360
3 days	3.486 ± 0.450
5 days	4.090 ± 0.31()
a~. 7 days	4.569 ± 0.513
2 weeks	4.357 ± 0.610
3 weeks	3.949 ±,0.592
10 weeks	2.495 ± 0.480
20 weeks	2.335 ± 0.330.
3W 3R*	1.593 ± 0.335 <sup>#</sup>
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Morphometric analysis of muchs secretion in the galgebladder opithelium · 64

<sup>1</sup> all values were compared to controls and unless indicated by '#' were statistically significant to p<0.05 as assessed by the unpaired Student's t test.

\* animals on the cholesterol-enriched diet for 3 weeks followed by 3 weeks on the control diet.

SEM refers to standard error of the mean

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TABLE 41. Analysis of biliary lipids in gallbladder bile. 1, 2, 3. 65

	a Cholestergi, ششما/11	Phospholi id (mmol/1)	Bile acils (mmol/l)	Lithogenic Index
Control	19.15±12.41	20.9 <u>5</u> 2,3	237.0±11.0	0.356±.086
, 6) hours	6.03±0.71	00.50±6.51	_175.1±19.2	0.337±.031
1.1 mary	11.11±0.46	36.17±4.32	211.2±13.9	0.562±.079
le hear.	1 11. + ±1.2	,46.17±11.66	217.0±22.0	.6.742±.000
1 day	1.09±1.47	2. <b>*.</b> 49±4.21	136.9±36.2	0:974±:161
3 days	.5.33£0.50'	35.55±1.64	181.6±3.20	1.133±.086
the tays	21.54±1.42	32.73±1.68	104.4±21.1	0.980±.100
n 2 day∋	14.74±1.02	31.78±2.29	19516±27.5	0.707±.058
D weeks	15.85±1.61	41.80±9.50	249.5±76.3	0:665±64
B weekg	14.40±2.20	39.09±54.77	173.3±29.0	0.651±.016
10 weeks	<ul> <li>145÷10±1.50</li> </ul>	37.2.13.35	190.0259.0	-0:#43±.207
191 - Weiger Halls	3 17.38±1.67	44.5024.05		0.015±.146

I nede in control group, ned in all other groups Values are expressed as the mean ± standarl error

unless indicated by '\*', all lithedenic index values were statistically significant to p<0.05 as assessed by the unpaired Student's to test



Mitotic and labelling index values (0/00) during the various experimental intervals. Experimental values are compared to control and the level of significance is in parentheses below each.\*

Condition.	Mitotic Index(MI)	Labelling Index(LL)		
Control	0.037 ± .013	0.169 ± .016		
1 week-treated	0.250 ± .097 (p<0.05)	0.394 ± .084 (p<0.05)		
2 week-treated .	0.384 ± .093 (p <sup>s</sup> 0.01)	1.312 ± .209 (p<0.001)		
10 week-treated	0.329 ± .009 (p<0.005)*	0.516 ± (19) (p<0.05)		
20 week-treated	0.289 ± .083 (p<0.05)	0.317 ± .020 (p<0.05)		

. \* values represent mean  $\pm$  standard error n = 4 to 8 animals per interval





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Summary of crystal precipitation and stone formation

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	L.1.	Mulaus husur-		Constal	Platy units a		
	4. e. 4. e. X	secretion	oruq,e	ULISUALS	microliths	o concs	
in the second sec		. :::		•••• •• •• •• •.			
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12 h <del>r</del> s		<u> </u>		- -			
18 hrs #	$= \frac{1}{2} \sum_{i=1}^{n} $	+	• <b>*</b>		-	- ,	
24 hrs	<b>†</b>	+		+	-		
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5 days			+ .	+	·		
7 days		+.	+	+		. –	
2 wks		+ ,	+	+	÷.	-	
3 wks		+	ہے۔ اب ہے ا	+	.+		
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20 wks	↓ · · · · · · · · · · · · · · · · · · ·	+	,	+	+	+	
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- denote	s abso	nt		, ,			

+ denotes present a

# L.I. denotes lithogenic index

÷	Mitosis & L.I.	Hypertroph hyperplasi		nd RA-sinus n plasma ce	& Cholesterolo lls like lesion	osis- n .
Control		-	-	- -		••••
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12 hrs	- 		-	-	$+ \frac{\sqrt{2}}{f}$	
18 hrs	-	-	-	 	· _ + :/	·
24 lirs 3 days	•	-	_	-	+	; •
5 days	-	- ,	. <u>.</u>	-	+	, ,
7 days	*	۰ 	• –	- 		
2 wks.	*	- * *.		C 2	+	
3 Wks.	#		• •	-	+ •	
10 wks.	* ^ *	+ .			+	• -
20 wks				,	,	

denotes absent

denotes present 🐒 à +

denotes higher values than Control?  $^{\star}$ 

denotes not done įį.

L.I. denotes labelling index

# DISCUSSION

# Control animals

The gallbladder of the Richardson's ground squirrel was thrown into numerous mucosal folds or rugae, the height of which reflected the fullness of the organ at the time of fixation. Mueller <u>et al</u>. (1972) examined guinea pig gallbladder in empty and filled states and found that the folds or rugae disappeared when the gallbladder was fixed in the filled state. These authors suggested that the instimulated filled gallbladder should be considered as the normal resting state of the organ. In addition they found that the residual elevations in a completely filled gallbladder were caused by large underlying vessels. The ground squirrel gallbladder exhibited these small mucosal humps when filled as well but further examination failed to reveal any underlying structures.

Fenestrated capillaries with diaphragma were observed in close proximity to the epithelium and were present in these folds. Hayward (1968) described fenestrated capillaries in the lamina propria in cata. Faye <u>et al.</u> (1966) showed fenestrated capillaries in the gallbladder of rabbits with different degrees of distention depending on the functional activity of the lining

epithelium. The capillaries below non-transporting cells were collapsed while those of active cells were distended.

Unmyelinated nerves were seen between muscle layers and in the lamina propria in close proximity to blood vessels. Cai and Gabella (1983, 1984) described ganglionated nerve plexuses in the serosa, near the cystic artery and its branches and in the lamina propria of the guinea pig gallbladder. Neurohistochemical observations by these authors (1983) revealed both acetylcholine-positive and catecholamine-containing fibers in the gallbladder of the guinea pig. In the present study, electron microscopic observations revealed prominent adrenergic varicosities intramuscularly and beneath the basal lamina. Cai and Gabella (1983) and Baumgarten and Lange (1969) did not. observe any adrenergic varicosities intramuscularly in the guinea pig, rheses monkey or cat, but saw a rich supply in the periods and lamina propria where they form part of the mucosal plexus. Kyosola and Penttila (1977) observed adrenergic fibers in the human gallbladder, especially in the fibromuscular layer, and a few axons beneath the basal lamina. Bjorck et al. (1982) have shown that adrenergic nerves enhance the concentrating function of the feline gallbladder by stimulating water absorption from the lumen. In this study, the adrenergic varicosities reflected a non-direct system of merves to the musculature and

epithelium of the ground squirrel gallbladder. However, neurohistochemical studies are required to identify the distribution of fiber types.

The lateral cell membranes exhibited microfolds that appeared to interd gitate with one another in TEM sections. Similar observations were made by Mueller et al. (1972) in the guinea pig and MacPherson et al. (1983) in the dog. They may also be similar to the microvilliform projections described by Hayward (1968), finger-like evaginations of Tormey and Diamond (1967), and elaborate digital processes of Wheeler (1971), all seen in cross section in a number of species with the transmission electron microscope. The microfolds of adjacent cells separated from each other and allow extensive distention of the intercellular space during active transport of solutes to the lamina propria. Spring (1983) postulated that some sodium was actively  $\lesssim$  pumped into the lateral intercellular space while the rest was exocytosed through basal plasma membrane by the Na-F pump. The ATP for these pumps was supplied by mitochondria which were found concentrated in the basal region of the cell. Tormey and Diamon (1967) calculated that the basal region of the cells compared about 60-80% of the mitochondria present in the subapical region. Rough estimates indicated a similar ratio in the light cells of the ground squirrel gallbladder.

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Prominent desmosomes were seen scattered along the lateral cell membranes of adjacent cells as far down as the basal plasma membrane. In a number of studies, (Kaye et al. 1966; Tormey and Diamond, 1967; Dietschy, 1966), desmosomes were not described between apposing cell membranes except in the apical zone. Kaye et al. (1966)& considered desmosomes a rare occurrence in rabbit gallbladder. Mueller et al. (1972) have shown prominent desmosomes in the guinea pig gallbladder and postulated that the absence of desmosomes in the above studies may be the result of artifactual disruption due to manipulation and interruption of blood supply to the organ while the tissue was being obtained. The random distribution of desmosomes appeared to ensure a labyrinth of interconnecting channels between adjacent cells for the movement of solutes as postulated by Diamond and Bossert (1967) for the standing-gradient osmotic flow model. In the present study, basolateral desmosomes were not disrupted even when the intercellular space was extensively distended.

Dark, or pencil; cells were seen singly among the light cells. The dark cells exhibited the same features as the light cells only the cytoplasm was more electron dense and the organelles closely packed together. - Dark cells were described in humans (Evett <u>et al</u>. 1964), and rabbits (Hayward <u>et al</u>. 1968). Togari and Okada (1953) regarded

these cells as being effete while Yamada (1962) showed an increase in enzyme activity. Hopwood et al. (1980) suggested loss of cytosol might explain this increase in enzyme activity, prevented the free movement of organelles and impaired normal function of these cells. In the gallbladder of man, Bader (1965) proposed that the dark cell was an intermediate cell type in the course of development and differentation from the undifferentiated basal cell to normal epithelial cell. The presence of mitosis in the lining epithelium showed that replacement epithelial cells are not derived from basal cells. Studies on the gallbladder epithelium of sheep (Hayward, 1965) and the dog (Johnson et al. 1962) suggested a declining functional activity for these cells. The dark cells observed in the present study were normal and lend support to the suggestion of Hopwood et al. (1980) that loss of cytosol impaired organelle movement and function. The functional significance of the dark cells needs further study.

A light edematous barrel-shaped cell was observed infrequently in the lining epithelium. The ultrastructural features of these cells suggested that they were aged and degenerative with a high degree of hydration. They appeared to be in the process of being extruded from the lining epithelium into the gallbladder lumen during normal

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epithelial turnover. These cells exhibited the classical features of cell death as described by Trump and Arstila (1976). Apoptotic bodies were not observed in the gallbladder lumen or in neighbouring cells. Several authors (Kerr et al., 1972; Wyllie et al. 1980; Harmon et a., 1984) described apoptotic bodies as specific for cell death in which membrane bound organelles are extruded in the lumen of an organ or the degenerated cell is phagocytosed by neighbouring cells. Yamada (1968) observed light barrelshaped cells with degenerated organelles in the mouse gallbladder and suggested that they were aged cells with declining activity. Dyban (1973) has shown that the normal turnover rate of the gallbladder epithelium of guinea pigs was 32 days. In the present study, the marked infrequency of mitotic figures and edematous cells, may also suggest a slow turn over rate for the ground squirrel gallbladder.

Leucocytes were seen migrating through the basal lamina and were found between epithelial cells near the basal lamina and occasionally in intercellular spaces at the level between the nucleus and junctional complex. Leucocytes were first described by Virchow (1857) and have been subsequently described in the literature as cask cells (Togari and Okada, 1953), and basal cells (Ferner, 1949). Kilburn <u>et al</u>. (1973) have shown random migration of leucocytes between epithelial cells in the trachea of

hamsters and guinea pigs exposed to toxic substances. In the present study a random transepithelial migration of leucocytes was observed but the functional implication remains to be elucidated.

Mucus

The gallbladder epithelium of control ground squirrels exhibited a basal level of both sialylated and sulphated glycoprotein secretion, a phenomenon similar to that described in the gallbladders of other models (Hayward, 1968; Lee, 1980; Lee and Scott, 1982; Yamada, 1962b; Lee, 1981; Wahlin, 1979; Esterly and Spicer, 1968; Smith and LaMont, 1985b).

Intracellular mucin granules have had their ultrastructural characteristics defined as membrane bound electron lucent granules with a dense core (Hayward, 1963); Lee, 1980; Koga, 1973) which led to contradictions. Hora is and Schultz (1970) described lipid droplets in the supranuclear regions of epithelial cells of human gallbladder while Koga (1973) suggested these droplets were mucin granules. Light microscopy autoradiography failed to conclusively identify mucus granules but verified the incorporation of tritiated glycoprotein precursors as well as the pathway of mucus secretion (Wahlin, 1977; El<sup>6</sup>y et a) ., 1971). In the present study, autoradiography at the electron microscope level demonstrated the occurrence of silver halide over these cellular inclusions following administration of radiolabelled galactose, a mucin precursor. This positive identification of mucin granules, rather than sole reliance on morphology, was integral to the interpretation and verification of observations made in this study.

Mucus, because of its visco-elastic properties, is extruded from the epithelial cells into the lumen in the form of strands. Scanning EM confirmed the transmission observations that not every cell was in a similar stage of the secretory cycle. Active secretion was observed to occur in groups of cells. Transmission EM showed these cells contained prominent rough endoplasmic reticulum, a Golgi complex, and microtubules, all features indicative of secretory potential and/or activity.

The mechanism of mucus release is still a matter of controversy. Hayward <u>et al</u>. (1968) proposed a holocrine mode for rabbit, others Koga, 1973; <sup>7</sup>Lee, 1980; Yamada, 1962b) described a merocrine secretion in the mouse and guinea pig, while Laito and Nevalainen (1972) advocated apocrine secretion in the human gallbladder. In the present study, mucin granules were observed in close proximity to

the apical surface, in some instances fusing with the plasma membrane and discharging their contents into the lumen of the gallbladder. This process was similar to that demonstrated electron microscopically in the rabbit (Lee, 1980), mouse (Wahlin <u>et al</u>. 1974, 1976; Wahlin, 1977), and guinea pig (Wahlin and Schieber, 1975), indicating a merocrine secretory mode.

The stimulus for and mechanism of mucus release and expansion is not fully understood. Lee <u>et al</u>. (1981b) demonstrated that ligation of the cystic duct before feeding prairie dogs a lithogenic diet prevented mucus hypersecretion and concluded the stimulus for hypersecretion was in the lithogenic bile.

The mechanism of mucin granule release is subjected to speculation. S.P. Lee proposed that there is a change in the ionic content of the granule which resulted in water absorption and an explosive extrusion of the granule content over the epithelial surface. This allowed the mucin to spread over non-secreting cells and formed a continuous layer (Personal communication).

The increased frequency in observation of apical cytoplasmic bullae by scanning elgetron microscopy in, experimental groups, suggested an apocrine secretory

process similar to that observed in humans. Correlative transmission electron microscopy however, did not support this hypothesis. Sections through these apical bullae revealed that they contained few, if any, mucus granules. Apical regions of cells exhibiting various degrees of convexity contained masses of mucus secretory granules but the plasma membrane retained their microvilli, a feature not consistent with that of true apical bullae. Goblet cells were not observed in the mucosa of the ground squirrel. Apical bul'ae of similar morphology were observed in the gallbladder epithelium of newborn rabbits (Hayward, 1968) and dogs (MacPherson et al., 1983). None of these authors was able to attribute any definite function to bullae. Hopwood et al. (1980) claimed that the terminal web might be responsible for keeping organelles from entering the bullae. In the ground squirrel, the terminal web was poorly developed and as such would be unable to effectively ensure the absence of organelles in bullae. A poorly developed terminal web was also described in the mouse and rabbit (Yamada, 1955; Hayward, 1968). Trump and Arstila (1976) however, considered these structures to represent part of the cellular reaction to injury as they were common features in uschemic bowel syndromes. The absence of cellular organelles in the bullae and their function requires elucidation.

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## Experimental animals

#### Mucus secretion

Hypersecretion of epithelial mucins was observed in gallbladder epithelium of experimental animals as early us 18 hours after the ingestion of the cholesterol-enriched diet. Precipitation of cholesterol monohydrate crystals from the lithogenic bile was not observed until 24 hours in treated animals. Previous studies have demonstrated mucin to be a nucleating agent for crystal precipitation and its hypersecretion occurred prior to crystal precipitation (LaMont <u>et al.</u>, 1984; Whiting and Watts, 1985; Hayward et <u>al.</u>, 1968; Lee and Scott, 1979; Ice and Scott, 1982; Smith and LaMont, 1985b). In the present study, the ground squirrel-cholesterol model confirmed this pattern and has allowed quantitation of the phenomenon prior to and during cholekithiasis.

Previous studies have suggested that the stimuli for mucus hypersecretion was in the lithogenic bile (LaMont et al., 1984; Leevet al., 1981b). The latter autnors demonstrated that lightion of the systic duct in the prairie dog before cholesterol feeding prevented mucus hypersecretion. In the present study, bile lithogenicity increased significantly in the 13 hour treated animals and

responsible in whole or in part, for the stimulation of cus hypersecretion. Florey (1970) postulated that mucous membranes in response to a noxious stimuli will increase mucus secretion and cell proliferation as a protective mechanism. Increased cellular proliferation was also observed in ground squirrels fed a lithogenic diet.

Quantitation of epithelial mucins demonstrated their continued increase in synthesis and secretion throughout the 20 week experimental period. These values peaked at the one-two week interval, an ideal time for the mucus to act as a nidus in the aggregation of crystals into platy units, microliths and stones. Continued hypersecretion facilitated the progressive growth of these smaller concrements into macroscopic stones.

Crystal precipitation and stone formation

Cholesterol gallstone formation was not merely a matter of producing supersaturated bile and was far morcomplex than the situation initially suggested by Admirand and Small (1968). Supersaturation of the bile with cholesterol was essential for stone formation but not everyone with supersaturated bile forms stones. The difference between stone forming and non-stone forming subjects appeared to be their ability to form cholesterol crystals (Whiting and Watts, 1984). This was thought to be f, related to the presence of nucleation factors, their

ence, or the presence of inhibitors in the bile. Precipitation of cholesterol crystals from mildly supersaturated bile required the addition of particulate material, a process referred to as heterogenous nucleation (Bouchier, 1984). Whiting and Watts (1985) concluded the difference between stone and non-stone forming bile lay in the nucleation stage of crystal formation rather than in the presence of inhibitors. Initially many factors were thought to be able to initiate nucleation including bacteria, desquamated cells, epithelial.glycoprotein mucina and mucus. Particular attenti has focused on the role of epithelial glycoproteins in their many forms as mucins, mucus, and sludge (LaMont <u>et al.</u>, 1984; Smith and LaMont, ~ 1985b).

Most gallstones exhibited a core of mucus and these glycoproteins contributed to the lattice structure of the stone (Lee et al., 1979). Mucin can act as an area of epistatic contact in crystal growth (Bouchier, 1983), bound bilirubin (Smith and LaMont, 1983), and enhanced <u>in vitro</u> ( nucleation of cholesterol crystals (Levy <u>et al.</u>, 1983). Strong indicators of the role of mucin in nucleation have been provided by <u>in vitro</u> studies (Levy <u>et al.</u>, 1984) but particularly by <u>in vivo</u> animal models of cholelithiasis where excessive mucus secretion was the underlying common

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observation (Doty <u>et al</u>., 1983a; Freston <u>et al</u>., 1969; Lee, 1981; Lee <u>et al</u>., 1981b). Hypersecretion of mucus has been noted to either accompany or precede stone formation in these models. Lee and Nicholls (1986) found a striking ... increase in the amount of mucus glycoprotein in galibladder bile of patients with biliary sludge, even greater than those with gallstones. This high molecular weight glycoprotein is now considered a pronucleating agent in experimental and human gallstone disease (LaMont <u>et al</u>., 1984).

The ground squirrel model has demonstrated, through SEM observations, that evel cholesterol crystal precipitation was preceded by increased mucus secretory activity. An increase in both the amount, and number of actively secreting cells was observed as early as 18 hours on the diet. The lithogenic indices revealed that the morphological documentation of increased mucus activity occurred just before, or in concert with, the supersaturating of the bile with cholesterol. Holan et al. (1979) was the first to demonstrate that accelerated nucleation of cholesterol month crystals was a distinguishing feature of lithogenic bile. LaMont et al. (1984) considered the stimulus for gallbladder mucus hypersecretion to be a component of the lithogenic bile. Prostaglandin regulation of mucin release from the

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gallbladder has been demonstrated in the prairie dog using aspirin to block this process and thereby inhibited gallstone formation (Lee <u>et al.</u>, 1981a).

The lithogenic index of the ground squirrel bile rose quickly to exceed 1.0 but then fell as low as 0.615 at 20 weeks. Gallstone formation however, continued during this period and analysis of biochemical components required to calculate the lithogenic index indicated the bile was not desaturated and contained a nigh proportion of cholesterol.

Biochemical analysis of the bile showed that the phospholipid level increased rapidly during the first 18 thours and dropped to normal level at 24 hours at which time cholesterol crystal precipitation occurred. Holzbach (1936) proposed that phospholipid vesicles maintained cholesterol in solution in the metastable zone and above this saturation level, cholesterol crystallized out of solution. During the same experimental period, bile acids level dropped below normal while cholesterol level increased thereby facilitating the saturation of bile. The increased lithogenic index may be due to decrease secretion of bile, dilution of secreted bile or randure of the gallbladder to absorb water adequately.

Scanning electron microscopic observations on the

mucosal surface of the ground squirrel gallbladder during the pathogenesis of cholelithiasis, revealed accumulation of a thick sludge-like layer. This became evident following the initial stages of cholesterol monohydrate crystal precipitation and continued throughout the resperimental. period. Gallbladder sludge, observed by sonography in humans; has been defined as thick bile that may either contain, or consist of, a fine suspension of pigment granules, mucin, calcium bilirubinate, cholesterol crystals and small stones of less than 3mm. diameter (Allen etcal., 1981; Smith and LaMont, 1985a). Juniper and Burson (1957) described the presence of cholesterol crystals in a microscopic examination of gallbladder sediment. crystals appeared colorless, bransparent, and thin, with / parallel edges often having a notched corner resembling pieces of broken window glass. Scanning electron microscopy of the cholesterol monohydrate crystals precipitated from ground squirrel bile confirmed these This indicated a similarity in earlier observations. configuration between these and human crystals.

Sludge occurred in gallbladders where stasis was evident and often disappeared after return of normal contractility. Doty <u>et al</u> (1983b) concluded that gallbladder stasis was an important link between hepatic secretion of cholesterol saturated bile and listed a number

of factors that might impair gallbladder emptying. Several worker's have shown that sludge is an integral part of both cholesterol and Ligment stone formation (Allen <u>et al.</u>, 1981; Been <u>et al.</u>, 1979; Bernhoft <u>et al.</u>, 1983; Soloway <u>et</u> <u>al.</u>, 1977). The thick sludge-like layer observed in the ground squirrel by SEM occurred at an ideal time to become involved as an accessory nucleation agent and aiding in the aggregation of small platy band subunits into larger concrements. Lee and Nicholls (1986) determined this sludge to be a sediment composed of cholesterol monohydrate crystals and bilirubin granules embedded in a matrix of mucus gel.

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Smith and LaMont (1985a) identified the chemical nature of a brown-black material that composed the matrix of human cholesterol gallstones. This material was a non-lipid component identified as a mucin-bilirubin complex and had a composition similar to that of biliary sludge. Ground squirrels on the lithogenic diet for 10 weeks exhibited stones where the sludge-like material was embedding cholesterol crystals into its growing surface. After 20 weeks the larger cholesterol gallstones began to take on a brownish-yellow coloration when examined macroscopically. The SEM showed this matrix-like material on the surface of these stones. It helped to fill in the crevices between larger subunits in the stone, smooth out

the surface and continued to embed cholesterol crystals. Morphological observations on this model leave little doubt that mucus hypersecretion played a role in the initial stages of nucleation followed by the formation of biliary sludge which then aided in the aggregation of smaller subunits into concrements of increasingly largensize.

Osuga and his colleagues (1974) used the scanning electron microscope to investigate experimentally-induced gallstone development in the squirrel monkey and compare it with observations made on luminal contents of human gallbladders resected for tholecystitis (Osuga <u>et al.</u>, 1975). Summarizing their findings they claimed that although the sequence of stone formation was similar, then were significant differences between the concrements of man and squirrel monkeys (Osuga <u>et al.</u>, 1975).

When the pattern of stone development observed in the ground squirrel gallbladder are compared to the schematic' diagrams provided by Osuga <u>et al</u>. for monkey (1974 and human (1975), the observations made in this study most closely fit those for the human. Neither the "wagon wheel" concrements nor the central cavity in simple stones of the monkey were common to either the ground squirrel or human. Similar confiduration of ground squirrel microliths to those of human and the sand-like stones of the monkey, indicated an underlying common growth pattern in the early phases of stone formation. The larger mulberry shaped stones, and the initiation of condensation, in the outer layers of stones from 20 week ground squirrels, when the ... organs were filled with concrements, indicated a pattern that will ultimately form mature stones with a configuration similar to those of the human.

The similarity in the pattern of stone formation and, morphology of the resultant concrements to human stones is encouraging. When this is coupled with the similarity of bile chemistry, the reproducibility and frequency of incidence, the non-toxicity of the diet, and the relatively short time factor volved from introduction of the diet to gross stone formation, the ground squirrel is an excellent animal model for induction of experimental cholelithiasis.

# <u>Mucosal changes</u>

### Cell proliferation and damage

The data from this study demonstrated an increase in the rate of cell proliferation and progressive mucosal damage, both of which were intimately related to time on the diet and the presence of macroscopic stones. Some of

the changes observed were similar to those noted in previous studies and tt, 1978; Putz and Willems, 1981; Hubigler and Palme, 1982; Lee and Scott, Scott, 1976 : No. 1982; Haya (et al. Fel; Fel; 1972). A number of those observed spectacity in enremology, were significantly different. The occurrence of m toses throughout the epithe ial chest of the add experimental animals, without any processes or welleys or crests, was similar to studies in the human of at and Willems, 1978; Putz and Willems, 1979) a. 1977 (sea pig (Scott, 1974; Jacoby, 1958), but dissimilar to the rabbit (Kaye et al., 1966). The latter author suggested that proliferative compartments existed in the gallbladder of rabbits, the valleys more active in mitoses. Mueller et al. (1972) however, have shown that valleys and crests of mucosal folds were transient structures dependent upon the state of the gallbladder when fixed. The present study supported this finding.

Mitotic figures observed in the epithelial sheet of control animals were relatively rare. Previous studies in human (Putz and Willems, 1978; Putz and Willems, 1979), guinea pig (Scott, 1978; Jacoby, 1958; D. C., 1973), meuse (Scott, 1978; Lee and Scott, 1982) and rapid (Scott, 1978; Kaye<u>set al.</u>, 1966) gallbladder have demonstrated that the normal cell turnover rate in this organ was very slow.

. go Bargmann (1959) failed to observe mitosis in the human gallbladder and suggested this tissue was amitotic. A significant difference between this and previous studies (Putz and Willems, 1978; Evett <u>et al.</u>, 1964) centered on which cell was actually undergoing mitosis. Electron microscopic observations in the ground squirrel showed that columnar epithelial cells, and not the basal cells seen near the basal lamina, were involved in mitosis. Light microscopic studies of human gallbladder had earlier suggested the basal cells as the progenitors for this organ (Putz and Willems, 1978; Evett <u>et al</u>., 1964). The findings of this study clearly do not support this hypothesis.

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In the one and two week experimental groups, the mitotic and labelling indices were significantly highers than those of control animals. These indices were considerably lower than those of other experimental models (Scott, 1978) but closer to the values obtained in human gallbladder epithelium (Putz and Willems, 1978; 1979). This increase in pro' erative activity was observed before the occurrence of macroscopic stones, but during the period of crystal and microlith formation. Studies on the mouse (Scott, 1978, Put) Willems 1981; Marsch-Zeigler and Palme, 1982; Lee (Scott, 1982) and other models (Scott, 1978) have also shown this rapid increase in the number of mitotic and labelled cells long before the presence of

stones. These authors suggested that abnormal bite might have been pesponsible for this increased proliferative activity and that mechanical irritation to the mucosa was Putz and Willens (1978) examined human secondary. lithiasic gallbladder and suggested that chronic mechanical irritation by gallstones might cause increased cell proliferation. This phenomenon was thought to be one form of response to irritation or injury (Florey, 1970). Cel'l proliferation may help increase the surface area to compensate for supersaturated bile (Marsch-Zeigler and Palme, 1982 . In the present study, cholesterol-fed ground squirrels demonstrated modified gallbladder bile as early as eighteen hours, crystal precipitation within 24 hours, and microliths at 2 weeks.

Previous studies in guinea pig (Scott, 1974; Jacoby, 1958), mouse (Putz and Willems, 1981), and man (Putz and Willems, 1979) have shown that distended gallbladders also exhibited an increase in proliferative activity. The presence of stones in human (Putz and Willems, 1978) and mouse (Lee and Scott, 1982) gallbladder on the other hand, did not cause distention and therefore did not stimulate the increase proliferative activity. The crystals and microliths present early in the cholesterol-fed ground squirrel did not block the cystic duct, nor were the organs distended. The increase in mitotic activity in this model
was also clearly not related to distention. The gallbladders of animals from the early experimental period also failed to show any inflammatory response, another mitotic stimulant.

In the 10 and 20 week experimental groups, cellular damage and extrusion, hypertrophy, hyperplaska, Rokitansky-Aschoff sinuses, inflammatory cells and continued presence of significantly higher mitotic and labelling indices were phenomena similar those observed in the mouse (Marsch-Zeigler mid Palme, 1982; Lee and Scott, 1982) and guinea pig (Scott, 1976). The presence of inflammatory cells in the lamina propria, thickening of the muscular layer and occurrence of Rokitansky-Aschoff sinuses were standard criteria used for microscopic diagnosis of chronic ' cholecystitis (Halpert, 1961; Edlund and Olsson, 1961). These earlier studies in the cholesterol-cholic acid mouse model showed ce ular damage and extrusion as early as two days on the diet, hypertrophy and hyperplasia by the sixth day, the changes occurring long before the presence of macroscopically visible stones (Marsch-Zeigler and Palme, 1982; Lee and Scott, 1982). In the ground squirrel however, a significantly different till frame was involved, the changes were only observed in the presence of stones. This suggested that the increase in cellular damage might be due to either the continued toxicity of supersaturated bile or

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a direct mechanical effect. This discrepancy between model: may be either species or diet-related. The gallbladder of the ground squirrel appeared to have reacted slower to the cholesterol-enriched diet and may reflect more accurately the changes occurring in the human gallbladder:

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A previous study using the mouse model suggested that dividing epithelial cells were extruded from the epitmelial A sheet (Marsch-Zeigler and Palme, 1932). In the ground squirrel, electron microscopic observations revealed that epithelial cells in preparation for division, on the crests of mucosal folds were larger than adjacent cells. They were distinctly different in morphology however, from the edematous cells that were actually extruded. Epithelial cell debris has been implicated as a nucleating factor for . stone format fon (Small, 1980; Sedaghat and Grundy, 1980; Lee and Scoty, 1982). The extrusion of edematous cells during normal epithelial turnover in the ground squirrel may also act as a nucleating factor in the presence of saturated bile. Recent studies have implicated mucus hypersecretion as an important phenomenon contributing to stone formation (Lee et al., 1981b; LaMont et al., 1984).

Transmission EM revealed that no actual defect in the epithelial sheet existed when one, or even two, cells, were extruded. Correlative scanning and transmission EM showed that neighbouring epithelial cells slid under those being extruded to protect the underlying layers. This may not be possible however, when large areas of the epithelium are denuded as in acute and chronic cholecystitis of human gallbladders (William and Smith, 1978). Other studies on human gallbladder (Hopwood <u>et al.</u>, 1980; Myllarniemie and Nickels, 1977) also showed that damaged cells occurred singly or in groups. Scanning EM revealed actual defects in the epithelial sheet and the authors suggested that the holes represented empty doblet cells (Myllarniemie and Nickels, 1977). In the ground squirrel, goblet cells were not seen in either control or experimental animals. Cellular damage may also be attributed to ischaemia (Trump and Arstila, 1976), but no any evidence of thrombi in mucosal blood vessels was observed in this study.

The battery of changes seen in the ground squirrel closely followed the chronological pattern described for premalignant lesions of the human gallbladder (Ojeda <u>et</u> <u>al.</u>, 1985; Albores-Saavendra, 1986). The present study indicated that the Richardson's ground squirrel model is well suited to study the chronology of cholecystitis and malignant lesions.

Cholesterolosis-like lesion

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The present study demonstrated a rapid increase in lipid accumulation in the mucosa of the ground squirrel gallbladder throughout the experimental period. For ious studies have shown the gallbladder epithelium to be capable of absorbing cholesterol and phospholipids from the bile (Neiderhiser <u>et al.</u>, 1971; 1973; 1976; Hopwood et al., 1981; Kouroumalis <u>et al.</u>, 1984; Ross <u>et al.</u>, 1986). Lipid accumulation in ground squirrel gallbladder epithelium was observed intracellularly and intercellularly as early as 1. hc... after ingestion of the lithogenic diet. Coupled with this was a significant increase in bile cholesterol levels. Neiderhiser <u>et al.</u> (1976) demonstrated a rapid increase of cholesterol absorption in the gallbladder of the guinea pig, the rate of absorption dependent on the absolute and relative cholesterol content of the bile.

Previous studies have proposed one way for these cells to rid themselves of the absorbed lipids was through release into the intercellular space. The lipids subsequently moved to the lamina propria and finally into either the venous or lymphatic system (Meiderhiser et al., 1976; Kouroumalis <u>et al.</u>, 1984; Koga, 1985; Mora and Schulz, 1970; English and Hopwood, 1985). This mechanism appeared to facilitate transport of excess lipid in the gallbladder of the ground arm irrel.

The accumulation of lipid in the supranuclear and thelial cells, as well as in dilated basal regi cisternae and the stal bodies was observed in one, two, and animals. A similar accumulation of three week t lipids in the supranuclear and basal regions of the cells, as well as in tubular or vesicular reticula, has been described previously in human gallbladders with cholesterolosis (Koga, 1985; English and Hopwood, 1985; Miettinen and Tilvis, 1985). Several studies suggested that the free cholesterol absorbed from the bile was esterified in the smooth endoplasmic reticulum to form lipid droplets which were then released into the intercellular space and on into the lamina propria , Koga 1985; Nevalainen and Laito, 1972; Tilvis et al. 1982; Subbiah and Dicke, 1977; Hora and Schulz, 1970). Support for this hypothesis was obtained in the present study by the localization of digitonin complexed free cholesterol in both dilated endoplasmic reticulum and lipid droplets in the supranuclear gions of the epithelial cells. Lipid droplets in the basal regions of these cells did not complex with the digitonin.

Residual bodies were found in the subapical and supranuclear region while lipid droplets appeared only in the basal aspect of the cells. Neiderhiser and colleagues (1971, 1973; 1976) have shown that uptake of lipids 'from bile by the gallbladder epithelium was a normal phenomenon. These authors have demonstrated uptake of C<sup>14</sup>-labelled oleic acid, cholesterol, lecithin and lyolecithin by the normal guinea gallbladder. The lipid droplet was either incorporated into residual bodies or transported into dapillaries in the lamina propria. Lipid droplets observed in the ground squirrel may be on their way to capillaries in the lamina propria. The presence of residual bodies subapically, and lipid droplets basally, in these animals suggested that the lining epithelium of the gallbladder was capable of absorbing lipids from the bile and either incorporating them into residual bodies or transporting them to capillaries in the lamina propria.

Another mechanism available to the cell to rid itself of cholesterol was through heterophagocytosis, a process which results in the formation of residual bodies. Epithelial cells from ground squirrels on the lithogenic diet exhibited numerous residual bodies in the supranuclear region of the cells, a situation similar to that described in human gallblädders with cholesterolosis (Nevalainen and Laito, 1972; Hora and Schulz, 1970). Digitonin-complexed experiments revealed free cholesterol in these organelles. The residual bodies may then be released into the intercellular space. All changes observed in the present study occurred when the Lile was lithogenic and cholesterol crystal precipitation or microlith formation were observed either on the epithelial surface or in bile washings. 98

In 10 and 20 week treated animals, there was continued presence of lipid in residual bodies, supranuclear and basal cell regions. In addition, dense osmiophilic droplets was observed in the lamina propria. Lipid appearance and density varied depending upon the degree of unsaturation of fatty acids present to form a complex with osmium tetroxide (Ghadially, 1984). In the present study, chronological examination of the ground squirrel gallbladder has shown that macroscopic stones, lithogenic bile and early morphological features of chronic cholecystitis also occurred during this time frame. This suggested that epithelial cells of the ground squirrel gallbladder continued to absorb lipid from the bile and excreted it into the lamina propria. These lipids were then engulfed by macrophages and removed by venous or lymphatic drainage in a pattern similar to that suggested for human gallbladder. Failure of these macrophages to enter the ocirculation resulted in cholesterolosis (Koga, 1985; Nevalainen and Laito, 1972; English and Hopwood, 1985, Hora and Schulz, 1970). In the present study, accumulation of foam cells in the lamina proprint was not observed and the

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/lesion was different from that commonly described in cholesterolosis of human gallbladder (Koga, 1985; Miettinen 'and Tilvis, 1985). The alterations in gallbladder mucosa of ground squirrels fed a lithogenic diet were basically similar to those described in the cholesterol-cholic acid fed dog model (Holzbach et al., 1977).

The early presence of lipid intracellularly, at the time when gallbladder bile was mildly saturated, may be an adaptive mechanism of the gallbladder to prevent cholesterol precipitation. It was not dependent upon, or related to, the presence of gallstones (Holzbach <u>et al.</u>, 1977; Salmenkivi, 1964) or cholecystitis (Womack and Haffner, 1944), and occurred much earlier than either of these phases of the cholelithiasic process.

## SUMMARY AND CONCLUSIONS

Two hundred and thirty eight conditioned Richardson's ground squirrels (<u>Spermophilius richardsonii</u>) of both sexes wieghing 350-450 grams were divided into control and experimental groups. Each group contained 4 animals. Control animals were maintained on a diet of rat chow and water <u>ad libitum</u> while experimental animals were fed a 23 cholesterol-enriched diet and water <u>ad libitum</u>. Animals were killed at intervals of 6, 12, 18, 24 hours, 75, 7 days, 2, 3, 10, 20 weeks and 3 weeks on cholesterolenriched diet followed by 3 weeks on normal diet. Ų.

Animal were killed, bile withdrawn and examined by polarizing light microscopy and assayed for phospholipids, bile acids and cholesterol to establish a lithogenic index. Gallbladders and injected with the same volume of fixative, 2.5 gluteraldehyde in Miloning's buffer pH 7.2. Tissue was processed a for transmission and scanning electron microscopy, electron microscopic autoradiography, electron microscopy digitonin experiment, electron microscopy quantitative study and light microscopic: lipid histochemistry, paraffin mucin histochemistry, and methacrylate autoradiography.

The results indicated that in control animals, folds

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or rugae were transient structur s dependent upon on the volume of bile present when fixed. The lining epithelium of the gallbladder was composed of three types of cells, light, dark and edematous cells. The edematous cells were degenerating cells that were extruded from the epithelial sheet. The dark cells were narrower than the light cells but had a similar morphology. Transmission electron microscopy revealed that normal columnar epithelial cells were undergoing mitoses that was different from basal cells. The basal lamina formed peg and socket interdigitations with the basal plasma membrane and ( desmosomes were seen throughout the lateral plasma membrane in actively transporting cells. Nor-adrenergic nerve fibers were seen beneath the basal lamina and in between muscles. bundles. A basal level of both sialylated and sulphated mucins were stored in membrane-bound granules and exocytosed by merocrine secretion.

In experimental animals, mucus hypersecretion was observed as early as 18 hours before the precipitation of cholesterol monohydrate crystal. This hypersecretion continued throughout the experimental period and formed a thick sludge-like wayer over the epithelial surface. Mucus secretion peaked in the 1-2 week sampling interval, an ideal time to act as a nidus for the aggregation of crystals into platy units, microliths and stones.

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Biochemical assay of bile showed bile lithogenicity began to rise at 18 hours while cholesterol crystal precipitation occurred at 24 hours. Platy units and microliths grew by appositional growth by two-three weeks and aggregated into mulberry stones in 10 weeks. By 20 weeks, many stones were approximately 2mm in diameter and occompanied by large numbers of various concrements. The larger stones had a cholesterol content of 61-74%.

Morphological observations of the gallbladder epithelium revealed increased cellular proliferation in one week treated animals before the occurrence of macroscopic stones but in the presence of cholesterol crystals and microliths. Electron microscopic observation of 10 and 20 week treated animals demonstrated damaged epithelial cells occurred singly or in groups and that there was no defect in the epithelial sheet. Neighbouring epithelial cells slid under the basal aspects of cells being extruded to protect the basal lamina. Hyperplasia, hypertrophy, Rokitansky-Aschoff sinuses, muscular thickening, plasma cells in the lamina propria were observed about the same time when stones were visible.

Other mucosal changes observed were the presence of lipid accumulation intercellularly and intracellularly in 12 hours tree of animals. Digtonin experiments revealed

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free cholesterol in dilated endoplasmic reticulum and in residual bodies. Neutral lipid was demonstrated by light microscopy histochemistry in the supranuce ar and basal regions of cells. In the 10 and 20 week treated animals, dense osmiophilic lipid droplets were seen in macrophages and in the lamina propria.

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The present study demonstrated that the Richardson's ground squirrel fullfilled the criteria set out by Freston and Bouchier (1968) for a model to study cholelithiasis. Potential problems in directly applying results obtained from experimental studies to the human condition will always exist. However, constant comparison of data from both animal and human sou bes will help avoid pitfalls of extrapolation and contribute to solving the problems of human pathology in cholesterol cholelithiasis Figure 1: A micrograph showing light and dark (white arrow) cells. A capillary (C) is seen in the lamina propria. Scale bar = 10µm.
Figure 2: A fenestrated capillary with diaphragm covering the fenestration (arrowhead). L- lumen. Scale bar = 0.1µm.
Figure 3: Unmyelinated nerves (N) in close proximity to a blood vessel (BV) of the lamina propria. Scale

bar =  $5\mu m$ .



Figure 4: An adrener of nerve (Ad) is seen between muscle bundles (Mu). Scale bar = 5µm.

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Figure 5: An adrenergic serve (Ad) beneath the epithelial sheet (Ep). Scale bar = 5µm.

Figure 6: Junctional complex (Jc) between epithelial cells. Residual bodies (R) are seen in the supranuclear region. N- nucleus. Scale bar = 1µm.



Figure 7: A micrograph showing microfilaments (Mf) in close

proximity to a centriole (Ce). Scale bar = 0.5µm.

Figure 8: A lipid droplet (Li) in an epithelial cell near its basal plasma membrane, BL- basal lamina. Scale bar = 0.5µm.

Figure 9: An active Golgi complex (Gc) with secretory granules (Sg) near the maturing face. Note a microtubule (Mt) near the secretory granules. Scale bar = 0.5µm.



Figure 10: A light micrograph showing both sialylated (blue-aqua) and sulphated (brown-black) mucins on the surface of epithelial cells. High iron diamine-alcian blue pH 2.5 stain. Scale bar = 20µm.

Figure 11: A light micrograph of the gallbladder epithelium showing the supranuclear areas of a group of cells secreting sialomucin with side chain substitution (red). Potassium hydroxide/alcian blue pH 1.0/phenylhydrazine-Schiff stain. Scale bar = 20µm.



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Figure 12. A scanning micrograph showing individual, and groups of cells, releasing strands of mucus into the lumen of the gallbladder ( rrows). Scale bar

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= 5µm.

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Figure 13: An electron microscopic autoradiograph of a control animal gallbladder 25 minutes after intraperitoneal H<sup>3</sup>galactose injection showing developed silver halide grains over a mucus granule: MV- microvilli. Scale bar = 0.2µm.

Figure 14: An electron microscopic autoradiograph showing

labelled material over muchs granules and endoplasmic reticulum in the supranuclear regions of cells. Seven day treated animal, 25 minutes after intraperitoneal H<sup>3</sup>galactose injection. Scale bar = 1µm.

Figure 15: An electron microscopic autoradiograph showing

labelled mucus on the luminal surface and subapical regions of epithelial cells. Mmitochondrion, Lu- lumen, MV-.microvilli, arrowhead- labelled mucus granule. Control animal 40 minutes after intraperitoneal H<sup>3</sup>galactose injection. Scale bar = 0.5µm.

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Figure 16: Mucus secretory granules (Sg) in the subapical region of the cell. Note the absence of cellular organelles in the region. M- mitochondria, Jcjunctional complex.Scale bar = 1µm.

Figure 17: A micrograph showing a secretory granule (Sg) with an irregular profile. Note the contents of one granule being extruded into the lumen (arrow). Scale bar = 1µm.

Figure 18: An apical bulla (B) containing only particulate

matter. Scale bar = 1µm,.



Figure 19: A micrograph showing mitotic activity within the epithelial sheet. M- mitochondria, Ce- centriole, N- nucleus. Scale bar = lum.

Figure 20: An underdeveloped micrograph of a dark cell to demonstrate the presence of mitochondria (M) and residual body (R) in the apical region. Note the prominent junctional complex (JC) and desmosome (D) between light and dark cells. Scale bar = lum.

Figure 21: An underdeveloped micrograph of a dark cell showing the presence of mitochondria (M) in the basal region. Scale bar. = 1µm.

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Figure 22: An edematous cell with a pyknotic nucleus (N) and cellular contents being extruded into the lumen (L). Scale bar = 5µm.

Figure 23: An edematous cell with a karyolytic nucleus (N), lysosomes (Ly) and vacuolated mitochondria (M).

Scale bar =  $5\mu m$ .

Figure 24: A micrograph showing a prominent basal lamina (BL) forming peg and socket interdigitations with the basal plasma membrane of the overlying epithelial cells. Scale bar = 1µm.



Figure 25: A micrograph showing various degrees of

distention of the basal intercellular spaces

(Is). Scale bar = 10µm.

Figure 26: A prominent desmosome (D) still intact despite the distention of the intercellular space (Is). Scale bar = 1µm.

Figure 27: The basal lamina (BL under actively

transporting epithelium demonstrates a reduction in the degree of peg and socket interdigitations with the overlying cells. D- desmosome. Scale bar = lum.



Figure 28: A'leucocyte (Lu) migrating through the basal lamina (BL). Note a pseudopod (arrow) in the intercellular space. Scale bar = 1µm.

Figure 29: A leucocyte (Lu) in the intercellular space near the junctional complex (Jc). L- lumen, arrowslateral plasma membrares. Scale bar = lum.

Figure 30: A scanning micrograph showing the degree of mucosal folding in an organ containing little or no bile. Scale bar = 0.1mm.

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Figure 31: A scanning micrograph demonstrating the

reduction in height and number of the mucosal folds when the organ is filled with bile. Scale

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bar = 0.1mm.





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Figure 32: A fractured edge of the epithelium showing the plate-like nature of the lateral plasma membrane interdigitations (arrow). Scale bar = 5µm.

Figure 33: A deeper fracture showing ovoid nuclei (N) • within epithelial cells. Scale bar = 2µm.

Figure 34: A scanning micrograph showing the basal lamina of an area close to a fracture where epithelial cells have been released. Arrows indicate small projections that were once part of the peg and . socket arrangement between the basal lamina and overlying basal plasma membrane. Scale bar = lum.



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Figure 35: Increased mucus secretion at 18 hours! Scale bar

= 10µm.

= 10µm.

. Figure 36: Increased mucus secretion at one day, Scale bar

Figure 37: Sludge-like material coalescing into a thick layer covering large areas of the epithelial surface at 3 days. Scale bar = 10µm.

Figure 38: Accumulation of mucus and initial lamination of cholesterol crystals (arrow) at day 5. Scale bar = 1µm.

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Figure 39: An electron micrograph showing numerous mucus secretory granules (Sg) in the apical region of a g 7 day-treated animal. Note the presence of microvilli on the apical convexity. Jcjunctional complex, M- mitochondria, Scale bar = lum.

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Figure 40: An apical bulla devoid of microvilli and contained cytoplasmic material. Scale bar = 10µm.



Figure 41: A thick sludge-like layer covering the epithelial surface of a two week-treated animal. Scale bar = 10µm.

Figure 42: Continued mucus hypersecretion in later sampling

animal. Scale bar = 10µm.

Figure 43: Surface morphology of a five day-treated animal showing prominent cytoplasmic bullae protruding into the gallbladder lume. Note the presence of , mucus and absence of microvilli on the surface of bullae. Scale bar = 10µm.

Figure 44: Epithelial cells of an animal fed a lithogenic diet for 3 weeks followed by normal diet for 3 weeks showing patchiness of mucus secretion similar to controls. Scale bar = 10µm.





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Figure 45: Cholesterol monohydrate crystals (arrows) stuck between microvilli at one day. Note the

hypersecretion of mucus (arrowhead). Scale bar =

Figure 46: Continued precipitation of rhomboidal cholesterol monohydrate crystals at day . Note that some appear notched (arrow) Scale bar =

1µm.

Figure 47: Crystal growth into platy subunits by parallel

lamination common at 2 weeks. Scale bar = 0.1mm.

Figure 48: Surface of a platy subunit covered with biliary

material. Scale bar = 0.1mm.

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Figure 49: A 3 week microlith composed of a number of platy subunits arranged in an initial radial configuration. Scale bar = 0.1mm<sup>\*</sup>

Figure 50: A number of concrements from a 10 week animal exhibiting various stages of growth. Scale bar =

1mm.

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Figure 51: A concrement of intermediate size from a ten week animal. Note early evidence of lobular subunit aggregation. Scale bar = 0.1mm.

Figure 52: One of the largest stones from a 10 week animal exhibiting a multilobular shape. Scale bar =



Figure 53: Higher resolution of a portion of figure 15 showing the component lobules. Scale bar = 0.1mm. 3

Figure 54: Another region of figure 15 showing biliary sludge embedding laminated cholesterol plates into the surface of the stone. Scale bar = 10µm

Figure 55: A large mulberry shaped stone from a 20 week animal. Note the relative smoothness of its surface. Scale bar = 1mm.

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Figure 56: Higher resolution of figure 20 showing polished regions where crystals are oriented parallel to the surface. Scale bar = 0.1mm.



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Figure 57: Fractured edge of a 20 week stone showing condensation of peripheral crystals into a capsule-like arrangement. The arrow indicates a septum radiating into the center of the stone. Scale bar = 0.1mm.

Figure 58: Central region of a fractured 20 week stone showing the aggregation of large numbers of

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laminated crystals. Scale bar = 0.1mm.\_\_

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Figure 59: A two week-treated animal showing columnar epithelial cells undergoing mitosis (Mc) on the crest of a mucosal fold. Scale bar = 5µm.

Figure 60: Autoradiogram of a two week-treated animal showing labelled cells in the valley and sides of mucosal folds. Scale bar = 20µm.

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Figure 61: A scanning electron micrograph of a ten weektreated animal revealing defects in the epithelial sheet (arrows). Scale bar = 10µm.

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Figure 62: A transmission electron micrograph of a ten week-treated animal showing two edematous cells with ruptured apical membranes and karyolytic nuclei (N). Note the adjacent cells sliding under the basal aspect of the extruding cells. Scale bar = 2µm.



Figure 63: A scanning electron micrograph of a ten weektreated animal showing hypertrophic cells (arrowheads). Note normal epithelial cells in adjacent areas (curved arrow). Scale bar = 10µm.

Figure 64: Light micrograph of a 20 week-treated animal showing a hyperplastic lesion (arrow). Note two 'epithelial cells (arrowheads) close by in the process of mitosis. Scale bar = 20µm.

Figure 65: A 20 week-treated animal showing Rokitansky-Aschoff sinuses in the hypertrophied muscle layer. Scale bar = 40µm.



Figure 66: A transmission electron micrograph of a 20 week-treated animal showing inflammatory cells in the lamina propria. Epithelial cell (Ep), plasma cell (P), fibroblast (F). Scale bar = 3μm.

Figure 67: An electron micrograph of a 20 week-treated animal showing inflammatory cell's (IC) between the basal lamina and epithelial cells Ep) and in the lamina propria. C- capillary. Scale bar =

5µm.



Figure 68: A light micrograph autoradiograph of a two week-treated animal showing humerous labelled fibroblasts in the lamina propria. Scale bar =

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<sup>2</sup>Figure 69: A light micrograph of a two week-treated animal showing numerous labelled fibroblasts in the adventitia. Scale bar = 20µm.

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Figure 70: Lipid accumulation both intercellularly and intracellularly in 12 hour treated animals. Nnucleus, Lu- lumen. Scale bar = 3µm.

Figure 71: Numerous residual bodies in the epithelial

2μm.

cells of one day treated animals. Scale bar =



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Figure 72: Lipid accumulation in the supranuclear region and lipid droplets (L) in the basal region of epithelial cells of seven day treated animals. Scale bar = 2µm.

Figure 73: Lipid accumulations (L) in the supranuclear region of an epithelial cell from a seven day treated animal. R- residual bodies, G- Golgi apparatus, rough endoplasmic reticulum (arrow), lysosome (arrowhead). Scale bar = 1µm.

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Figure 74: A frozen section of a control gallbladder

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stained with oil red O. Note the absence of any colored precipitate. Scale bar = 20µm.

Figure 75: A frozen section from a five day-treated animal stained with oil red O. Neutral lipid is demonstrated in the supranuclear regions of cells. Scale bar = 20µm.



Figure 76: An electron micrograph of a two week-treated animal showing lipid droplets in the basal regions epithelial cells. Arrow- basal lamina, - two us. Scale bar = 1µm.

Figure 77: Tissue from the gallbladder of a seven day treated animal complexed with digitonin to demonstrate free cholesterol (arrows) in dilated cisternae. Scale bar = 0.5µm.



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Figure 78: Digitonin complexing with free choresterol to demonstrate its deposition within a residual body (arrows) of a seven day treated animal. Mmitochondrion, L- lipid. Scale bar = 0.2µm

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Figure 79: Tissue from a seven day treated animal complexed with digitonin demonstrating cholesterol crystals on the surface of epithelial cells. Scale bar = 1µm.

Figure 80: Dense osmiophilic droplets in the lamina propria of a 20 week treated animal. Note the presence of a few electron lucent lipid droplets (arrows). Ep- epithelial cells. Scale bar = 1µm.

Figure 81: A macrophage in the lamina propria with numerous dense inclusions. Ep- epithelial'cells, Pl- plasma cell, C- capillary. Scale bar = 2µm.

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#### Appendix I

## 0.2M Buffer solution

Solution A: 27.6 g sodium phosphate Monobasic/l liter/DH<sub>2</sub>O Solution B: 28.4 g sodium phosphate Dibasic/l liter/DH<sub>2</sub>O

0.1M Buffer solution pH 7.4

Solution A: 190 ml/1000 ml Solution B: 810 ml/1000 ml

Dilute in 1000 ml DH<sub>2</sub>O

Adjust pH with 1% alcium chlor de solution 👘

100 ml of 2.5% gluteraldehyde in 0.1M phosphate buffer solution

1,

5 ml of 50% aqueous gluteraldehyde 95 ml of 0.1M phosphate buffer

1% osmium tetroxide in 0/1M phosphate buffer

1 g Osmium tetroxide

100 ml 0.1M phosphate buffer

<u>Epon 812</u>

Epon 812 resin

Dodecenyl succinic anhydride 11.3 g Nadic methyl anhydride 11.1 g DMP 30 0.8 g

23°. q

1.33 g

1.76 g'

0.1% Toluidine blue

0.1 g toluidine blue/100 ml  $DH_2O$ 0.1 g sodium borate in above solution

Saturated uranyl acetate.

uranyl acetate dissolved in DH20

Lead citrate

Lead nitrate

Sodium citrate

Dissolved in 50 ml<sub>i</sub>of DH<sub>2</sub>O 2 pellets of sodium hydroxide

Lee's methylene blue- basic fuchsin (MBBF).

Methylene blue stock solution

I of Salar

Methylene blue.....0.5 g Distilled water.....400 ml

Basic fuchsin stock solution Basic fuchsin.....0.5 g Distilled water.....400 ml

Solution B: Sodium phosphate dibasic......71.5 g Distilled water.....1000 ml Stock buffer pH should be 6:2-7.2. Higher pH levels produce deep blue results. Lower pH values give a pink tinge to sections.

## Working solution

-د'

Methylene blue	12	ml
Basic fuchsin solution	12	ml
Solution A (stock solution)	7	ml
Solution B (stock buffer)		
'95% ethyl alcohol	15	ml
Filter solution		

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	<i>*</i>			
<u>Methac</u>	rylate ixat and pro	ce sing	•	
•	, ,			
		`,	•	
	utral onlered form lin	•		24 h
	cchol		• • • • • • • • • • • • • •	
85% al	cohol	· · · · · · · · · ·	· · · · · · · · · · · · · · ·	1 h
	cohol (2 quanges,			
Cataly	sed sol. A 35% a cohol	(1:1)	•••••	/1 h
	•			
Soluti	on		•	
A	••••••••••••••••••••••••	••••••		ernight
Embed	tissue in mold with sol		•	
Leave	to harden at room tempe	erature		
				٠
<u>Cataly</u>	sed solution A		•	•
			·	
•			· .	
Sol. A	(JB 4)		••••••	100 ml

•.

Inpendix II

Catal/st..... 0.9 ml Mixed thoroughly for uniform solution.

## Embedding mixture

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a.

Catalysed solution A	••••••	25 parts
Solution B	V 	l parts
Stir ranidly and thoroughly to avoid	·handaning	

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Stir rapidly and thoroughly to avoid hardening

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#### Appendix III

Procedure for transmission and scanning electron microscopy Fix tissue in 2.5% gluteraldehyde for 4 hours Wash 3 x 15 minutes in Milloning's buffer Place tissue in 1% osmium for 1 hour Wash 3 x 10 minutes in distilled water Place tissue in saturated aqueous uranyl acetate Pin SEM tissue on cork with no tension 30%, 50%, 70% 90% ethanol for 15 minutes in each solution <u>TEM\_tissue</u>

100% ethanol for 1 hour

1:1 ethanol/propylene oxide for 20 minutes

1:3 ethanol/ propylene oxide for 30 minutes |

100% propylene oxide for 1 hour

3:1 propylene oxide/ epon mixture for 20 minutes
1:1 propylene oxide/ epon mixture for 30 minutes
1:3 propylene oxide/ epon mixture for 1 hour
100% epon mixture - ove hight

Embed in moulds in fresh epon Heat in oven for 48 hours at 55°C

<u>SEM tissue</u>,

100% ethanol 3x 10 Absolute acetone- 2x 15 finutes Critical-point dry

Coat with colloidal gold in sputter-coater

## MEM Digitonin Experiment

Tissue fixed in 2.5%.gluteraldehyde for 4 hours Place tissue in 2% digitonin solution dissolved in buffer for 4 hours

# Rinse in buffer

Post-fixed in 1% osmium tetroxide and process as above for TEM

# ' TEM autoradiggraphy

Dipping method

1:16 .

Grids coated with 0.5% parlodion dissolved in amyl acetate Attached to slide by parlodion

Silver sections picked in loop placed on grids

Slides dipped in Ilford L-4 emmulsion diluted with water

Dripped dried, stored in light proof pox for 3-4 weeks at 4°C and then developed:

Kodak D 19 for 3 minutes

Dipped in distilled water for 30 seconds Fixed in 25% sodium thiosulphate for 6 minutes Washed in distilled water for 6 minutes Stained in uranyl acetate and lead citrate for 10 minutes each.

### Light microscopy autoradiography

Two micrometer thick methacrylate sections were cut Slides were dipped in undiluted Ilford K.5 D emulsion Dripped dried for 1 hour

Áppendix V

Stored in light proof black box sealed with black tape at 4°C for 3-4 weeks

Developed as follows:

Dektol diluted 1:1 with water for 2 minutes

Washed in water for 2 minutes

Fixed in Kodak fixer for 5 minutes

Washed in water for 15 minutes .

#### Appendix VI

## High Iron Diamine - Alcian blue pH 2.5 Paraffin sections cleared in: Xylene for 5 minutes Xylene for 5 minutes 100% ethanol for 3 minutes 9,5% 85% **K**0% 11 ٦ Distilled water for 3 minutes. Stain for 24 hours in a fresh solution of: N,N- dimethyl-m-phenylenediamine di HCL.....1.68 g N, N-dimethyl-p-phenylenediamine HCL..... .0.28 q Dissolve these two in 750 ml of distilled water Immediately add 19.6 ml of 40% ferric chloride to activate the solution Rinse quickly in distilled water Stained in Alcian blue pH 2.5 for 20 minutes Alcian blue 8GX. . . . . . . . . . . . . ..... 1.00 g Glacial acetic acid..... 100.0 a Adjust the pH to 2.5 and fifter solution. Add a few crystal of thymol 'Rinse quickly in distilled water

70%, 85%, 95%, 100% ethanol f(r 3 minutes each Xylene (twice) for 5 minutes Mount in permount.

Results Sulphomucins grey-purple black Sialic acid Blue (aqua)

Appendix VII

Potassium hydroxide/alcian blue pH 1.0/ periodic acidphenylhydrazine- Schiff

Cleared paraffin sections as above Saponify in 0.5%.KOH in 70% ethanol for 15 minutes Wash slowly in running tap water for 5 minutes Stain in 1% alcian blue in 0.1N HCL (pH 1.0) Oxidize in 1% aqueous periodic acid for 2 hours Wash slowly in tap water for 10 minutes Treat, with 0.5% aqueous phenylhydrazine hydrochloride for 2 hours

Wash in running water for 5 minutes Stain with freshly prepared Basic Fuchsin-Schiff for 4 hours Bring to water as above

#### Results.

Sialic acids with side chain substitutions Rec
Tissue vicinal diols Rec
0-sulphate esters Blue

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