EXCRETORY SYSTEM

- Rectal papillae
- Malpighian tubules
Generalized insect alimentary tract, including excretory system
# Excretory System in Humans and Insects

**Humans**

1. Liquid system tied in with the circulatory system. Includes kidneys and a urinary bladder.
2. Main excretory product is urine (all ages)

**Insects**

1. System tied in with the digestive tract but includes the circulatory system.
2. Main excretory product is uric acid (adults). Main product depends on habitat.
FUNCTIONS OF THE EXCRETORY SYSTEM IN INSECTS

Problems insects face in their environments
1. Losing water because of the size/volume ratio of being small
2. Controlling the ionic balance of the body fluids
   a. Freshwater insects tend to lose ions to the environment
   b. Insects in salt water tend to gain ions

THESE PROCESSES BASED ON OSMOSIS AND DIFFUSION

Maintain a nearly constant internal (HOMEOSTASIS), osmotic environment of the hemolymph tissues, and cell environment by:
1. Elimination of excretory products
2. Reabsorption of water from the feces
3. Reabsorption and/or elimination of various ions
4. Absorption of materials produced by the symbionts in the hindgut of those insects housing them
What is one of the major problems facing insects?

What kinds of excretory products would one expect to find in insects and why would one expect these to be the kind of products they would produce?

**WATER LOSS** - INSECTS, BECAUSE OF THEIR SIZE MUST CONSERVE WATER

Cuticle and excretory system maintain proper water and ion balance

\[
\text{Uric acid} \rightarrow \text{allantoine} \rightarrow \text{allantoic acid} \rightarrow \text{urea} + \rightarrow \text{ammonia} + \text{CO}_2
\]

\[
\text{uricase} \quad \text{allantoinase} \quad \text{allantoicase} \quad \text{urease} \quad \text{glyoxylic acid}
\]
The excretory product in insects is usually colorless, it may be yellow or greenish in color depending on the food. Malpighian tubules may be whitish in color (Uric acid) or contain a yellow pigment, thus they appear yellow.

Amino acids are derived from proteins in foods. They are used by cells for synthesis of new body protein or other nitrogen-containing molecules. The amino acids not used for synthesis are oxidized to generate energy or are converted to fats or carbohydrates that can be stored. In either case, the amino groups (-NH$_2$) must be removed because they are not needed for any of these purposes. Once the amino groups have been removed from the amino acids, they may be excreted from the body in the form of ammonia, urea, or uric acid, depending on the species.
Notice in the diagram at the right that uric acid is not very soluble in water, whereas urea and ammonia are. **What does this mean to the Insect?**

In insects the waste product is usually 80% uric acid but this varies on their life style.

**It is NOT know how uric acid is transported to the Malpighian tubules.**
The synthesis of uric acid occurs primarily in the fat body. Since ammonia has 3 Hydrogens for every nitrogen, compared to uric acid having 1 to 1, the hydrogen for the ammonia must come from somewhere. It may come from water. Thus, it takes more water to get rid of ammonia.
This chart shows the type of excreta used by different insects. One should be able to correlate the life style of the insect with that of the main component of the excreta.

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This slide shows several things:

1. The hemolymph is a major storage area for amino acids.

2. Potassium is actively pumped into the Malpighian tubule, as is proline.
The great disparity between the ionic concentration of the animal’s hemolymph/tissues may be another reason why insects did not invade the oceans.
Structure of the Malpighian tubules

The Malpighian tubules are surrounded by muscles. They actually are moving in the hemolymph and can carry out peristaltic movements to move material from the terminal end to the opening in the hindgut. The Malpighian tubules produce the primary urine while the hindgut produces the secondary or final urine. They are absent in aphids and Collembola. In Diplura and Protura they are represented only by papillae. No. varies from 2 in coccids to 250 in desert locust.
Diagram of structures in *Rhodnius* Malpighian tubules. Note brush border made of microvilli. 

- **a** = basement matrix,
- **b** = invagination of plasma Membrane with mitochondria (e),
- **c** = endoplasmic reticulum,
- **f** = mineralized granule,
- **g** = microvilli with mitochondria entering the microvilli.

Enlarged view of microvilli showing droplets released into hemolymph.
TEM of larval Malpighian tubules of *A. taeniorhynchus*. L=lumen; SC= stellate cell; BL=basal lamella or matrix. PC=primary cells. H=hemolymph. SC have wider extracellular spaces or infoldings than does the primary cell.
Remember – $K^+$ in the hemolymph is at least 10 times lower than it is in the Malpighian tubule. This is due to the proton pump.
The point to note from the table to the right is that basically, the osmolarity of the hemolymph and that of the primary urine in the Malpighian tubule is nearly equal or isomotic. The exception, however, is for the ions like Na and K that are actively transported across the tubule against concentration gradients.

Potassium and proline are higher in the urine because they are actively excreted.
Hemolymph and fluid in the Malpighian tubule are isosmotic.

To carry out all of the active transport needed, mitochondria in the microvilli provide the energy for the proton pump.
If the hindgut or rectum area of the insect is involved in water uptake, ion movement, and amino acid uptake, what might be the characteristics that have to be met for this to take place?

1. Change in cuticle permeability
2. Active uptake mechanisms must be in place

What physical characteristic of the hindgut would make it difficult for water and ion movement?

**HINDGUT IS LINED WITH CHITIN**
Structure of anal papillae, anal organs and rectal papillae

The permeability of the cuticle of the hindgut is highly permeable compared to that of the foregut (i.e. crop here) (see below) and it is usually much thinner.
TEM of papillar cell within the rectum of *Calliphora*.

This cell is involved in uptake of water and ions from the rectal lumen (Lu).

Note that the rectum is lined by cuticle (Cu).

The apical cell membrane is thrown into a regular border of leaflets involved in the transport of water and ions.
Rectal papillae of flies and rectum

Various types of papillae in the rectum of insects are involved in reabsorption of water and the movement of ions for osmoregulation.
Notice the cuticle of the anal gland in the photo on the right is about half as thick as that of the adjacent cuticle. Also, note that it is delineated from the surrounding cuticle, thus preventing materials from moving laterally instead of just in and out/or vice versa of the gland. In the photo above, notice how the anal organ in fig. 6 is delineated from the rest of the cuticle. Remember, its cuticle is produced by epidermal cells that produce it while the adjacent cells produce the normal cuticle.
Terrestrial insects lose water. How do they recoup it?

How do they lose water?

Sonoran desert cicada. Pores 7X size of pore canals located on dorsal mesonotum + connected to special dermal glands via cuticular ducts are involved in water transport to the surface. Cooling of 2-5°C below ambient of 42-45°C.
Terrestrial insects lose water. How do they recoup it?

How do they lose water?
1. Through cuticle
2. Water loss from the respiratory surfaces
3. Water loss in excretion

How do they gain water?
1. Drinking
2. Uptake through cuticle
3. Metabolic water (grain beetles)

Other ways to prevent water loss
1. Cryptonephridial tubes
CRYPTONEPHRIDIAL TUBES

1. Found in larvae of Lepidoptera, many Coleoptera and antlion immatures

2. FUNCTION(S)
   A. Reabsorb water from rectum
   B. Absorb atmospheric water
Water vapor is taken up by the hygroscopic fluid on the hypopharynx and enters a duct that then takes the liquid water into the pharynx via the action of the cibarial pump.
Some dipterous larvae span a broad range of salinity tolerances.
Freshwater insects tend to lose salts to the environment because of their highly permeable cuticle. K, Na, and chloride are reabsorbed in the rectum but water is not.

Ways to recoup salts in freshwater larvae
1. Special chloride cells in some aquatic larvae
2. Rectal gills in dragonfly naiads
Chloride cells in the gills of the mayfly naiad for retrieval of salts
Chloride cells in the rectal chamber of dragonfly naiad for retrieval of salt ions
Chloride cells in the rectal chamber of dragonfly naiad for retrieval of salt ions
Saltwater insects gain salts and water with their food, thus losing water osmotically. Some insects like *Aedes campestris* or *Aedes sollitans* (common along salt marshes of Massachusetts). Also, *Ephydra cinerea* lives in Utah’s Salt Lake, which is 20% NaCl.
Mosquito larvae respire by using a respiratory siphon that breaks the water thus providing for gaseous exchange. They also have anal papillae (see white arrows) that are involved in osmoregulation and function in removing chloride, sodium and potassium ions from the water and putting them back into the hemolymph.

Structure compliments function at the ultrastructural level.
Click on the website below to go there:
Drosophila malpighian model
Diuresis—rapid flow of urine elimination from the body

Many insect species decrease urine output and increase blood volume prior to the molt. WHY? Following the molt, they increase urine output and decrease blood volume after cuticular expansion.

**Primary urine production** in Malpighian tubules

**Regulation by different mechanisms**

**Resorption of salts** in the hindgut

Rates of each are
Insect diuretic and antidiuretic hormones

**Diuretic hormones** generally act on the *Malpighian tubules* to stimulate urine production

**Antidiuretic hormones** generally increase fluid reabsorption by act on the *hindgut*

*Malpighian tubules* are not innervated, thus they must be regulated by hormones released in the blood

Muscles of the tubules can be modulated by diuretic hormones and Myotrophic peptides. Increase writhing movement in hemolymph

This made Ramsay’s assay a useful bioassay

Using his assay, it was shown that an extract from the fused mesothoracic ganglion mass in *Rhodnius* increased urine production up to 1,000-fold. *A. aegypti* diuretic hormone-loss of 40% of water in the blood meal with 2 hrs of feeding.
Bioassay technique developed by Ramsay for determining what factors influence excretory rates and secretions by the Malpighian tubules. Air bubble provides the tubule cells with oxygen to respire. Urine is insoluble in the liquid paraffin so it remains as a droplet at the proximal end of the tubule that would lead into the hindgut for excretion.

**Can add substances to the droplet to see their effect**

Since then there have been major technological advances to further the identification, isolation, and purification of peptides (2 or more amino acids linked together).

1. HPLC=high performance liquid chromatography
2. Automated peptide sequencing
3. MS=mass spectrometry
4. Development of routine molecular protocols for
   a. mRNA isolation
   b. amplification and sequencing of genes
   c. gene expression
5. Sequencing the entire genome of *Drosophila melanogaster* (2000) and *Anopheles gambiae*
6. Genomic databases for searching for genes encoding neuropeptides and their receptors
Click on the website below to go there:
The Interactive Fly

http://www.sdbonline.org/fly/aimorph/maltubls.htm
With the exception of serotonin, all of the factors that have been identified as having diuretic or antidiuretic activity are all **NEUROPEPTIDES**

Some insect neuropeptides are similar to those of vertebrates, thus indicating a long evolutionary history as the two groups diverged about 6000 million years ago.

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**TABLE 1** List of identified diuretic and antidiuretic peptides, detailing the species from which they were first identified, abbreviated name, whether they have been fully sequenced (Seq), primary site of action, second messenger and activity.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Species</th>
<th>Name</th>
<th>Seq.</th>
<th>Site</th>
<th>Messenger</th>
<th>Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRF-related</td>
<td>M. sexta</td>
<td>Manse-DH</td>
<td>Yes</td>
<td>MT</td>
<td>cAMP/Ca^{2+}?</td>
<td>Diuretic/antidiuretic</td>
</tr>
<tr>
<td>Kinin</td>
<td>L. maderae</td>
<td>Leuma-K</td>
<td>Yes</td>
<td>MT</td>
<td>Ca^{2+}</td>
<td>Diuretic</td>
</tr>
<tr>
<td>Cardio-acceleratory</td>
<td>M. sexta</td>
<td>Manse-CAP_{28}</td>
<td>Yes</td>
<td>MT</td>
<td>cGMP</td>
<td>Diuretic/antidiuretic</td>
</tr>
<tr>
<td>Calcitonin-like</td>
<td>D. punctata</td>
<td>Dippu-DH_{31}</td>
<td>Yes</td>
<td>MT</td>
<td>cAMP</td>
<td>Diuretic</td>
</tr>
<tr>
<td>AVP-like</td>
<td>L. migratoria</td>
<td>AVP-IDH</td>
<td>Yes</td>
<td>MT</td>
<td>cAMP</td>
<td>Diuretic</td>
</tr>
<tr>
<td>Antidiuretic factor</td>
<td>T. molitor</td>
<td>Tenmo-ADF</td>
<td>Yes</td>
<td>MT</td>
<td>cAMP</td>
<td>Diuretic</td>
</tr>
<tr>
<td>Antidiuretic factor</td>
<td>A. aegypti</td>
<td>MNP</td>
<td>No</td>
<td>MT</td>
<td>?</td>
<td>Antidiuretic</td>
</tr>
<tr>
<td>Antidiuretic factor</td>
<td>F. polyctena</td>
<td>FopADF</td>
<td>No</td>
<td>MT</td>
<td>cAMP</td>
<td>Antidiuretic</td>
</tr>
<tr>
<td>Antidiuretic factor</td>
<td>M. sexta</td>
<td>Manse-ADF-B</td>
<td>No</td>
<td>CNC</td>
<td>cAMP</td>
<td>Antidiuretic</td>
</tr>
<tr>
<td>Ion transport peptide</td>
<td>S. gregoria</td>
<td>Schgr-ITP</td>
<td>Yes</td>
<td>Ileum</td>
<td>cAMP/++?</td>
<td>Antidiuretic</td>
</tr>
<tr>
<td>Cl^- transport stimulating hormone</td>
<td>S. gregoria</td>
<td>CTSH</td>
<td>No</td>
<td>Rectum</td>
<td>Ca^{2+}</td>
<td>Antidiuretic</td>
</tr>
<tr>
<td>Neuroparsins</td>
<td>L. migratoria</td>
<td>Nps</td>
<td>Yes</td>
<td>Rectum</td>
<td>Ca^{2+}</td>
<td>Antidiuretic</td>
</tr>
</tbody>
</table>

MT, Malpighian tubules; CNC, cryptonephric complex.
Insect brain or nervous tissue (ganglia)
  a. Neuropeptides with diuretic activity

Corpus cardiacum
  a. Storage and release of the peptides into the

HEMOLYMPH

Malpighian tubules
  a. Diuretic peptide hormone released into the blood and increases in titre in the hemolymph. Goes to the Malpighian tubule and activates diuresis
Cuticular plasticization in blood feeder and rapid excretion of water

*Rhodnius prolixus*-kissing bug and vector of trypanosome that is causative agent of Chaga’s

Occurs as a result of the action of hormones or neurohormones.
The rapid acquisition of a blood meal by hematophagous insects could produce a major osmotic problem if all of the water in the bloodmeal were to get into the hemolymph and stay there. Also, such a load greatly hinders the movement, especially flying, of these insects. They have solved this problem by using diuretic hormones that are released by stimulation of stretch in the abdomen in *Rhodnius*. These hormones cause rapid movement of water from the bloodmeal into the hemolymph where it rapidly moves into the Malpighian tubules for elimination as a droplet of urine (see photos).
Malpighian tubule of 60 hr old house fly larva. Note the lumen, the irregular waviness of the overall structure and the fuzziness of the tissue surface due to the microvilli. Note reticulate fat body.
Excess water from the blood meal enters the hemolymph and then into the Malpighian tubules.
Rhodnius prolixus as a model

1. Extremely rapid loss of water from the bloodmeal in blood feeders
2. The rate of water movement across the midgut must somehow match that entering the Malpighian tubules otherwise their will be a drastic change in the osmotic balance of the insects hemolymph.
3. Maddrell removed some of the Malpighian tubules from Rhodnius and did the measurements. Somehow, water leaving the bloodmeal across the midgut slowed down to match what was coming in.
4. Evidence suggests that hormonal control over the midgut is the same as that over the Malpighian tubule takeup.
5. Human blood contains a lot of calcium, which Rhodnius stores in crystaline form in the Malpighian tubules.
6. It is the stretch of the abdomen (monitored by stretch receptors in the abdomen) by the bloodmeal that triggers the release of serotonin from the abdominal nerves in the hemocoel.
7. At the same time, a diuretic hormone is released. Serotonin and the diuretic hormone act synergistically to regulate primary urine production by the Malpighian tubules.
Drawing of histology of Malpighian tubule of *Rhodnius* and an SEM of the same. Note spheres of calcium.

\[ V = \text{contains spheres of calcium} \]
SEM of Malpighian tubule of *Aedes taeniorhynchus*. H = hemolymph; V = spherical vacuoles in the cells of the tubule containing concentric crystals of Ca. L = lumen of the tubule. M = mitochondria in microvilli of the cells.
TEM of principal cell of the Malpighian tubule of *Calpodes ethlius* larva. Note the presence of spherocrystals that are produced from materials taken up from the hemolymph and packaged into these spherules that can contain uric acid, Ca, Mg, and/or Phosphates.
FAT BODY revisited-

Fat body cells are involved in:

1. **Intermediary metabolism** (glycogen to glucose; glycerol production; synthesis of trehalose from glucose)
2. Contain MFOs or cytochrome P450 enzymes (similar to vertebrate liver)
3. Fat body as a protein factory. It takes precursors from the hemolymph and produces the female specific protein or vitellogenin (Vg) and puts it into the hemolymph
4. Takes wastes out of hemolymph and produces uric acid
5. Production of antibacterial proteins known as Cecropins
6. Serves as a storage organ for lipids, etc.
7. Hormonal modulation of fat body (JH makes the fat body competent to make Vg)
8. Can house mycetocytes
9. Can house uric acid in special cells called urocytes, which are found amongst the fat body cells
Fat body can be categorized on where it is found

1. Subcuticular or peripheral fat body
2. Perivisceral fat body
1. **Fat body cells**

Fat body cells are involved in:
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3. Take precursors from the hemolymph and produce the female specific protein or vitellogenin and put it into the hemolymph
4. Takes wastes out of hemolymph and produces uric acid
5. Production of antibacterial proteins known as Cecropins and defensins
Fat body, at one time was considered to be composed of only one cell type. Now we know that this is not true. The principal cell type of the fat body is the trophocyte. The fat body may also contain bacteriocytes (=mycetocytes), urate cells and hemoglobin.
Trophocytes-the principle cell of the fat body

Trophocytes are held together by desmosomes and are surrounded by a basal lamina, thus the appearance that they are one mass of tissue.
Bacteriocytes (=Mycetocytes in fat body)
Many insects contain specific microorganisms, which are present in every individual, and are transmitted from one generation to the next by elaborate mechanisms. Associated with gut, gonads or fat body. Key role in the nitrogen economy of the host. Provide host with amino acids and vitamins.

Mycetocytes: Located in various tissues
1. **Midgut**
   a. *Glossina* and *Acheta* in wall of midgut
2. **Gonads**
   a. *Cimex, Formica*
3. **Malpighian tubules**
4. **Fat body**
   a. Aphids, Coccids, Aleurodidids, Cicadids (Most Homoptera)
Bacteriocytes, or cells containing micro-organisms, are found in various parts of the body in a number of insects. Bacteria and/or yeast seem to be the major endosymbionts. These organisms, shown below in a TEM reveal bacterioids in bacteriocytes of the fat body in the American cockroach. Presumably they are able to use uric acid produced by the fat body.
Most endosymbionts are passed from one generation to the other via transovarial transmission.

1. Symbionts are released from the bacteriocytes during egg development of the host.
2. They migrate from fat body to developing ovaries
3. Gain access to the developing oocytes
4. Are taken into the oocytes by endocytosis, the process that the female specific yolk protein, Vitellogenin, is taken up by the developing oocyte.
Among bacterial endosymbionts of insects, the best studied are the pea aphid *Acyrthosiphon pisum* and its endosymbiont *Buchnera* sp. APS, and the tsetse fly *Glossina morsitans morsitans* and its endosymbiont *Wigglesworthia glossinidia brevipalpis*. As with endosymbiosis in other insects, the symbiosis is obligate in that neither the bacteria nor the insect is viable without the other. Scientists have been unable to cultivate the bacteria in lab conditions outside of the insect. With special nutritionally-enhanced diets, the insects can survive, but are unhealthy, and at best survive only a few generations.

The endosymbionts live in specialized insect cells called *bacteriocytes* (also called *mycetocytes*), and are maternally-transmitted, i.e. the mother transmits her endosymbionts to her offspring. In some cases, the bacteria are transmitted in the egg, as in *Buchnera*; in others like *Wigglesworthia*, they are transmitted via milk to the developing insect embryo.

The bacteria are thought to help the host by either synthesizing nutrients that the host cannot make itself, or by metabolizing insect waste products into safer forms. For example, the primary role of *Buchnera* is thought to be to synthesize essential amino acids that the aphid cannot acquire from its natural diet of plant sap. The evidence is (1) when aphids' endosymbionts are killed using antibiotics, they appear healthier when their plant sap diet is supplemented with the appropriate amino acids, and (2) after the *Buchnera genome* was sequenced, analysis uncovered a large number of genes that likely code for amino acid biosynthesis genes; most bacteria that live inside other organisms do not have such genes, so their existence in *Buchnera* is noteworthy. Similarly, the primary role of *Wigglesworthia* is probably to synthesize vitamins that the tsetse fly does not get from the blood that it eats.

The benefit for the bacteria is that it is protected from the environment outside the insect cell, and presumably receives nutrients from the insect. Genome sequencing reveals that obligate bacterial endosymbionts of insects have among the smallest of known bacterial genomes and have lost many genes that are commonly found in other bacteria. Presumably these genes are not needed in the environment of the host insect cell. (A complementary theory as to why the bacteria may have lost genes, Muller's ratchet, is that since the endosymbionts are maternally transmitted and have no opportunity to exchange genes with other bacteria, it is more difficult to keep good genes in all individuals in a population of these endosymbionts.) Research in which a parallel phylogeny of bacteria and insects was inferred supports the belief that the obligate endosymbionts are transferred only vertically (i.e. from the mother), and not horizontally (i.e. by escaping the host and entering a new host). Attacking obligate bacterial endosymbionts may present a way to control their insect hosts, many of which are pests or carriers of human disease. For example aphids are crop pests and the tsetse fly carries the organism (trypanosome protozoa) that causes African sleeping sickness. Other motivations for their study is to understand symbiosis, and to understand how bacteria with severely depleted genomes are able to survive, thus improving our knowledge of genetics and molecular biology.
Developmental Origin and Evolution of Bacteriocytes in the Aphid–Buchnera Symbiosis

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Symbiotic relationships between bacteria and insect hosts are common. Although the bacterial endosymbionts have been subjected to intense investigation, little is known of the host cells in which they reside, the bacteriocytes. We have studied the development and evolution of aphid bacteriocytes, the host cells that contain the endosymbiotic bacteria Buchnera aphidicola. We show that bacteriocytes of Acyrthosiphon pisum express several gene products (or their paralogues): Distal-less, Ultrabithorax/Abdominal-A, and Engrailed. Using these markers, we find that a subpopulation of the bacteriocytes is specified prior to the transmission of maternal bacteria to the embryo. In addition, we discovered that a second population of cells is recruited to the bacteriocyte fate later in development. We experimentally demonstrate that bacteriocyte induction and proliferation occur independently of B. aphidicola. Major features of bacteriocyte development, including the two-step recruitment of bacteriocytes, have been conserved in aphids for 80–150 million years. Furthermore, we have investigated two cases of evolutionary loss of bacterial symbionts: in one case, where novel extracellular, eukaryotic symbionts replaced the bacteria, the bacteriocyte is maintained; in another case, where symbionts are absent, the bacteriocytes are initiated but not maintained. The bacteriocyte represents an evolutionarily novel cell fate, which is developmentally determined independently of the bacteria. Three of five transcription factors we examined show novel expression patterns in bacteriocytes, suggesting that bacteriocytes may have evolved to express many additional transcription factors. The evolutionary transition to a symbiosis in which bacteria and an aphid cell form a functional unit, similar to the origin of plastids, has apparently involved extensive molecular adaptations on the part of the host cell.
**Urate cells**

Are found in Collembola (lack Malpighian tubules), Thysanura, Blattodea and larval Apocrita (bees and wasps). Contain large spherules of uric acid. Probably for storage, which is based on some aspect of the insects life cycle or development.

Cockroaches do not excrete uric acid. Instead, they store it in the urate cells.
Waste products are taken up by the fat body from the hemolymph where they are converted to uric acid crystals (c) or uric acid in vacuoles (d).
STORAGE EXCRETION — waste products are stored in the body rather than eliminating them. Uric acid is not toxic and requires little water. It is stored in special cells (called urocytes) amongst the fat body in cockroaches and some other insects in crystalline form. Cockroaches and uric acid (Coby Schal work)

Males of several species have special storage glands called uricose glands that store uric acid and are part of their accessory reproductive glands. When males transfer the spermatophore to the female he deposits uric acid solution on it. This passes onto the female. When the sperm are released and the spermatophore is voided by the female, she eats it, thus getting the nitrogen rich uric acid from the male. Also, other species feed on a urate ‘slurry’ found in the male’s genital chamber, which increases their intake of nitrogen.

Storage of waste in the embryo—uric acid increases

Storage in the pupa also occurs

Permanent deposits in epidermal cells—Cotton stainer bug (uric acid) and pterins in white cabbage butterfly and white stripes of some caterpillars.
Insect diversity provides for wonderful examples of how a particular tissue or organ, that is used for a specific function becomes modified to serve another function. Following are examples of how the Malpighian tubules have become modified to serve various functions in different insects.

- Light producing organs
- Spittle production
- Calcified puparia
- Puparial silk
Larva of the fly *Arachnocampa luminosa*, Malpighian tubules are modified as light producing organs in Waitomo Caves of New Zealand about 600,000 tourist/yr for a total of 4 million US dollars.
Enlarged distal ends of Malpighian tubules form the luminous organ that produces the light. Tissue Cell. 1979;11(4):673-703. **Regional specialization in the Malpighian tubules of the New Zealand glowworm Arachnocampa luminosa (Diptera: mycetophilidae).** The structure and function of type I and II cells.
Various structures that normally serve a particular function, such as excretion with the Malpighian tubules, evolve in various insects to serve a totally unrelated function (i.e., here a light organ). This pattern repeats itself throughout the insect world.

Fat body cells in *Keroplatys trstaceus*, the glowworm known as the northern fungus gnat, produces its light from modified cells in the fat body.
In spittle bugs, the Malpighian tubules are involved in producing the spittle, which then comes out of the anus.

Structure and Function of the Malpighian Tubules, and Related Behaviors in Juvenile Cicadas: Evidence of Homology with Spittlebugs (Hemiptera: Cicadoidea & Cercopoidea)

Author: Rakitov R.A.¹
In the above fig. 5 one can see the puparium of the house fly on the left, which is melanized and darkened. The one of the right is of face fly, which is calcified. The calcium for puparium formation in this species comes from the transfer of calcium from the Malpighian tubules to the cuticle at the time of pupariation. Several other dipteran species have white, calcified pupal cases.
In *Chrysopa* larvae, material from the Malpighian tubules is involved with forming the silk of the pupa (see lower left). Also antlion larvae do the same thing in the Malpighian tubule modification.
What is a MECONIUM?

It is the waste products produced by a human, unborn baby or by an insect during the pupal stage. It contains wastes usually produced by the developing organism that is unable to pass it out because it is still inside the mother or the pupal case.

Every expectant parent hopes for an uncomplicated birth and a healthy baby. Some babies, however, do face delivery room complications. One fairly common condition that may affect your newborn's health is meconium aspiration. Meconium aspiration, also referred to as meconium aspiration syndrome (MAS), is a common cause of illness in newborns, but there is good news. Most cases are not severe, and one new treatment is saving the lives of some of the most seriously ill infants. See website below for excellent info about meconium in humans http://ak.essortment.com/whatismeconium_rhmq.htm
Ommochromes (yellow, brown and red pigments) color the meconium of Lepidoptera, especially the painted lady, which is red. Uric acid is also a large part

Is that blood coming from the newly emerged Painted Butterfly?
*Is that blood dripping from our emerging butterflies?*
No, that bright pink fluid, called “meconium,” contains the last traces of the caterpillar’s liquefied body and some other wastes that have collected during metamorphosis. The butterfly expels meconium through its anal opening.
Parasitic hiding in the Malpighian tubules, thus escaping the host response:

1. Microfilaria of pathogenic nematodes do this in Dog heartworm and Elephantiasis. When they enter the hemolymph of the mosquito, they find their way to the Malpighian tubules and enter the epithelial cells where they avoid the host response.

The effect of infection with *Dirofilaria immitis* (dog heartworm) on fluid secretion rates in the Malpighian tubules of the mosquitoes *Aedes taeniorhynchus* and *Anopheles quadrimaculatus*
Early cellular responses in the Malpighian tubules of the mosquito *Aedes taeniorhynchus* to infection with *Dirofilaria immitis* (Nematoda).

Bradley TJ, Sauerman DM Jr, Nayar JK.

Early ultrastructural changes in the Malpighian tubules of the mosquito, *Aedes taeniorhynchus*, were examined following infection with the nematode, *Dirofilaria immitis*. After ingestion by the mosquito, the microfilariae enter the cells of the Malpighian tubules, becoming intracellular. During early development, the filarial prelarvae reside in the cell cytoplasm surrounded by a clear zone without a delimiting membrane. Cells infected with prelarvae differed from uninfected cells and from cells in uninfected mosquitoes in that the volume of the apical microvilli was reduced and mitochondria were retracted from these microvilli. Morphometric analysis was used to quantify the ultrastructural consequences of infection. In infected cells, microvillar volume, the percent of microvillar volume occupied by mitochondria, and volume of mitochondria within the microvilli were significantly reduced.
Neuropeptide Analogs

• control and research applications

• peptides are not useful for control because they are:
  - susceptible to environmental degradation
  - susceptible to enzymatic degradation
  - unable to pass through insect cuticle

• nonpeptidal and pseudopeptidal agonists can overcome these obstacles
EXCRETORY SYSTEM AS A TARGET FOR PEST CONTROL

1. For terrestrial insects
   a. use diuretic agonists and/or antidiuretic antagonists

2. Problem-separate control of Malpighian tubule from that of the hindgut. If try to regulate one then the other may compensate for what has been done.

Molecular engineers
a. Design molecules that can be applied topically. Development of peptide analogues that can be applied topically and still work. Has been done with the PBAN (pheromonotropic activity) in moths and it works.
PBAN
Pheromone Biosynthesis Activating Neuropeptide

Synthesized Pseudopeptide:
• lipid moieties
• fortified peptide bonds

Results:
• cuticle penetration (esp. foregut)
• hemolymph persistence
• pheromone production

*Heliothis virescens*
SUMMARY

INSECT EXCRETORY SYSTEM INCLUDES:
1. Malpighian tubules
2. Hindgut

They regulate water and ion composition of the cells, tissues, and hemolymph and hemolymph volume.

The rates of water and ion movement are controlled by diuretic and antidiuretic hormones.

Some insects do not excrete uric acid but rather store it and use it with the aid of endosymbionts, which are housed in special cells know as bacteriocytes (=mycetocytes).