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Insect Excretory Mechanisms

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1 Introduction

Excretion refers to the processes which remove from the metabolic pool substances which interfere with metabolism. In this chapter I will consider three separate aspects of excretion, as first proposed by Maddrell (1971): (1) removal of molecules which are undesirable or perhaps even poisonous at all except very low concentrations; (2) excretion of molecules which are not toxic but merely useless, and which would become obstructive if allowed to accumulate and (3) excretion of molecules which are useful or essential but are present to excess. This would include excretion of water and physiological ions in some circumstances.

In writing this chapter I have attempted to link recent advances in our understanding of the physiology, biochemistry, genetics, molecular biology and chemical ecology of insect excretory mechanisms to the fundamental principles of design established more than 30 years ago. For example, the primary ATP-dependent transporter in insect epithelia was referred to in the literature until the early 1990s as the ‘electrogenic alkali cation pump’. This pump is now known to reflect the integrated actions of two separate transporters, an electrogenic vacuolar-type H⁺-ATPase and an exchanger of Na⁺ or K⁺ and H⁺ (Maddrell and O’Donnell, 1992; Wieczorek, 1992; Rheault et al., 2007). Similarly, the high osmotic permeability of insect epithelial cells noted in early studies (O’Donnell et al., 1982) is now seen to be based on aquaporins (Echavarria et al., 2001; Kaufmann et al., 2005). A third example is the multialkaloid transporter responsible for elimination of alkaloids such as nicotine, morphine and atropine by insect excretory organs (Maddrell and Gardiner, 1976). Recent studies suggest that this transporter is a P-glycoprotein (P-gp)-like transporter (Gaertner et al., 1998; Leader and O’Donnell, 2005) produced by expression of multidrug resistance (MDR) genes (Tapadia and Lakhotia, 2005).

A recurrent theme of this review is that studies of insect excretory mechanisms frequently provide illustrations of the Krogh principle: “For a large number of problems there will be some animal of choice or a few such animals on which it can be most conveniently studied.” In many instances, these examples of the Krogh principle are a consequence of unusual diets, such as blood, or life in extreme environments, as is the case for the tenebrionid beetles of the Namibian desert or mosquito larvae found in saline lakes containing very high levels of magnesium and sulfate.
Adaptations to extreme environments, coupled with the large number of unusual food sources exploited by insects and their extraordinary diversity, provides a fertile ground for research into excretory mechanisms.

Several reviews in the last decade have analyzed literature relevant to this work. Comprehensive studies of insect diuretic and antidiuretic hormones are provided by Gade et al. (1997) and Coast et al. (2002). A briefer analysis of recent work on endocrine control of salt balance in insects has been written by Coast (2007), and Orchard (2006) has reviewed the orchestration of diuresis and other physiological events in the hemipteran Rhodnius by the amine serotonin. Much of what we know about the functioning of the insect hindgut is due to the extensive work of Phillips and co-workers (Phillips et al., 1996). The dominant role of the vacuolar-type H\(^+\)-ATPase in driving solute transport across plasma membranes of insect epithelial cells has been reviewed frequently (e.g. Wieczorek et al., 2000; Beyenbach and Wieczorek, 2006). The mechanism and regulation of secretory transport in Malpighian tubules of the yellow fever mosquito Aedes aegypti has been reviewed by Beyenbach (2003). The application of genetic, molecular biological and physiological tools has lead to tremendous advances in our understanding of the mechanisms and control of solute and water transport by the Malpighian tubules of the fruit fly Drosophila melanogaster. This area has been reviewed frequently, primarily by Julian Dow and Shireen Davies (Dow and Davies, 2003, 2006; Davies and Day, 2006).

\section{Design principles underlying the insect excretory system}

The insect excretory system is comprised of the Malpighian tubules and the gut, especially the hindgut (Fig. 1). Each Malpighian tubule consists of a single-layer of squamous epithelial cells which form a blind-ended tube. Tubules range from 2 to 70 mm in length, 2–250 in number and up to 100\(\mu\)m in diameter (Phillips, 1981). Transport of ions (primarily K\(^+\), Na\(^+\) and Cl\(^-\)) and osmotically obliged water into the tubule lumen produces a near isosmotic secreted fluid. Fluid secretion is accompanied by passive diffusion of small solutes into the lumen, as well as selective secretion of solutes, including toxins. Some KCl and water may be reabsorbed downstream in a proximal segment of the Malpighian tubule or the anterior hindgut. The bulk of water, ion and metabolite reabsorption occurs downstream in the posterior hindgut, in particular the rectum, resulting in strongly hyperosmotic or hypoosmotic excreta. In many terrestrial species of insect, the hindgut can recover virtually all the water from the gut contents and fluid secreted into the gut by the Malpighian tubules. So effective are these water recovery systems that in the mealworm Tenebrio (Ramsay, 1971) and in the firebrat Thermobia (Noble-Nesbitt, 1970), for
example, the fecal pellets are powder dry. In addition, water vapour can be taken up from moist air drawn into the hindgut in both these species (O'Donnell and Machin, 1988). An essential feature of the insect hindgut for such extreme water reabsorption is the presence of a cuticular lining. The hindgut cells are thus protected by this cuticular lining from high concentrations of solutes, wastes and toxins in the lumen during water recovery (Phillips and Dockrill, 1968).

Adjustments of hemolymph fluid and solute levels are required in response to processes such as desiccation, feeding or osmotic influx across the external surfaces of aquatic species. However, several features of insect physiology and morphology mean that there has been less evolutionary pressure for rapid control of hemolymph composition by the excretory system. Firstly, respiratory gas exchange is accomplished by the ramifying tubes of the tracheal system whose termini extend into cells and tissues. Insects are not reliant, therefore, upon a blood-borne respiratory pigment whose functions would be compromised by changes in the volume and composition of the extracellular fluids. Secondly, sensitive tissues such as the eyes, the testis and the central nervous system are protected from the extracellular fluids by the actions of a series of epithelia, characterized by

![Schematic diagram of the insect excretory system.](image-url)
the presence of simple tight junctions. Such epithelia are not found in other invertebrates, but they play a crucial homeostatic role for the underlying tissues in insects. The regulatory properties of the perineurium, which envelopes the insect central nervous system, have been particularly well described (e.g. Treherne et al., 1984). Thirdly, one of the pre-eminent adaptations of insects for terrestrial life, is the possession of a waxy cuticle. A consequence of the small size of insects is that they have a very large ratio of surface area to volume and they are thus faced with the potential for relatively rapid and large scale changes in the composition of their extracellular fluids. Insects are commonly found in osmotically and ionically stressful environments and life on land, in particular, poses the threat of desiccation. The waxy cuticle is thus a feature which limits water loss.

A key role of an excretory system is to eliminate automatically toxins not encountered before. Given the large number of plant-derived toxins which have evolved to minimize herbivory, this requirement is particularly important in insects. Arthur Ramsay was among the first to appreciate that the Malpighian tubule provides a means for removal from the hemolymph of all soluble substances of low molecular weight, and that in this respect it has analogies with both the glomerulus and the tubule of the vertebrate nephron. His classic paper (Ramsay, 1958) examined the diffusion of organic solutes such as sugars, urea and amino acids across the Malpighian tubule wall of the stick insect *Carausius morosus*.

The view that emerged from this and subsequent work is that the absolute permeability of the Malpighian tubule epithelium of insects is quite low when compared to the excretory epithelia of other invertebrates and the vertebrate kidney. This arrangement seems paradoxical (O'Donnell et al., 1984; Skaer et al., 1987), given the need to clear toxic compounds from the hemolymph. The solution to this paradox lies in the restriction of the area for passive permeation, namely the paracellular clefts, rather than the restriction of the permeability of these sites. The adjacent cells in insect epithelia such as the Malpighian tubule are separated by septate junctions rather than the occluding, or tight, junctions of transporting epithelia in vertebrates (Lane and Skaer, 1980). An intercellular space which is about 17 nm in width separates the large flattened cells of the Malpighian tubules of *Rhodnius*, for example. The ratio of the areas of the basal side of the cells (i.e. the transcellular route) and the intercellular cleft is 3000:1. However, when the 40-fold amplification of basal surface area due to membrane folding and convolution is accounted for, the ratio of transcellular to paracellular area increases to 120 000:1 (Skaer et al., 1987). Compounds which are not transported actively diffuse through the intercellular clefts and are excreted; useful components also move slowly, and can be recovered from the excretory fluid by active transport.

Molecules cross the tubule epithelium by passive diffusion at rates which vary inversely with molecular mass (Maddrell and Gardiner, 1974;
Dalton and Windmill, 1981). Molecular mass also determines the route of transport for solutes which are not actively transported. Uncharged solutes whose molecular mass is less than that of sucrose cross into the lumen of the upper Malpighian tubule of *Rhodnius* by a cellular route (O'Donnell et al., 1984). Surprisingly, mannitol, a relatively large molecule which has often been used as an extracellular space marker, readily permeates through the cells and enters the lumen about 10 times more rapidly than sucrose, even though this route entails crossing at least two cell membranes in series. Rapid permeation by the transcellular route in *Rhodnius* presumably reflects the large area of cell membrane available, as noted above. Uncharged solutes which have a molecular mass greater than that of sucrose as well as charged solutes are excluded from the tubule cells and can cross only via the intercellular junctions.

It is important to emphasize the relatively non-selective nature of the filtration process by any excretory system. The vertebrate glomerulus passes compounds as large as inulin, whose molecular weight is \( \sim 5000 \), and the coelomosaic of the crustacean antennal gland is even more permeable, passing molecules as large as 50 000 Da. The pill millipede, *Glomeris marginata* is not an insect but it does have Malpighian tubules which rapidly secrete fluid very similar in composition to the bathing fluid (Farquharson, 1974). Even large solutes as inulin and dextrans of molecular weight up to 16 000 appear in the secreted fluid at concentrations only slightly below those of the bathing fluid (Farquharson, 1974).

In sharp contrast, the Malpighian tubules of insects secrete fluid which contains even such small molecules as sugars and amino acids at concentrations much below those in the hemolymph. For example, during diuresis the Malpighian tubules of the blood-feeder *Rhodnius prolixus* secrete fluids which contain glycine and inulin at concentrations only 1% of those in the bathing fluids. This low overall permeability of the tubule wall is a consequence of the limited area of the permeable sites rather than by restriction at the sites of permeation. As pointed out by Maddrell (1981) this arrangement means that toxic molecules, even those of moderate size, will still be removed passively, albeit slowly, from the insect’s hemolymph. By contrast, the tubules can secrete fluid at extraordinary rates in the presence diuretic factors, as described below.

In sum, the excretory system of insects operates relatively slowly and the passive rate of movement of hemolymph solutes into a slowly secreted primary excretory fluid is restricted by a reduction in the area available for passive transfer. Maddrell (1981) has pointed out several advantages of this arrangement. Firstly, less energy is required both to produce the primary excretory fluid and to reabsorb useful substances from it. Secondly, relatively high concentrations of molecules such as amino acids, trehalose and lipids can be maintained in circulation. Flying insects use the disaccharide trehalose and/or diglyceride lipid in the hemolymph as fuels.
for flight, and these compounds may be present at concentrations as high as 100 mM (Weis-Fogh, 1967). Many species in the orders Lepidoptera, Hymenoptera and Coleoptera contain levels of amino acids in the hemolymph as high as 100–200 mM (Maddrell, 1971). Hormones that are released into circulation will also be removed slowly by the excretory system and the amounts which must be released may therefore be reduced.

During diuresis, insects can eliminate excess fluid at very high rates and yet lose only trace amounts of hemolymph solutes. Comparisons presented in Maddrell (1981) show that most vertebrates filter their extracellular fluid 10–20 times more rapidly than do most insects. Given that many hemolymph solutes of even quite small molecular size only appear in the filtered fluid at concentrations in the range 10–50% of those in the insect’s hemolymph, reabsorption of such solutes need only occur in insects at about 1–2% of the rate required in many vertebrates, and there is thus a large saving in the energetic costs associated with reabsorption.

This energy saving may be a driving force behind the evolution of very rapid rates of fluid secretion by Malpighian tubules stimulated with diuretic factors. When fluid is secreted at high rates, dilution of solutes in the tubule lumen is associated with reduced rates of diffusion back into the hemolymph. However, once the concentration of a solute in the lumen reaches a very low level, further increases in the rate of fluid secretion have virtually no more effect on solute entry. If excess fluid is very quickly removed, and contains only low levels of hemolymph solutes, no reabsorption of the solutes is needed. This may explain in part why many blood-sucking insects can excrete fluid at such extraordinarily high rates; after feeding the hemipteran Rhodnius can eliminate fluid equivalent in volume to its total hemolymph in as little as 30 min.

Passive diffusion of solutes from hemolymph to tubule lumen is backed up by active mechanisms that can rapidly transport a very wide range of substances that insects repeatedly need to reabsorb, such as glucose (Knowles, 1975a), or excrete, such as organic anions, urate, phosphate, magnesium, sulfate, sulfonates, acylamides, alkaloids and glycosides (Maddrell, 1977; O’Donnell et al., 1983; Linton and O’Donnell, 2000; Torrie et al., 2004). The importance of specific transcellular transporters for many classes of potential toxins has been emphasized in a DNA microarray study of Malpighian tubule-specific gene expression by Wang et al. (2004). There is high enrichment (9- to 29-fold) of 3 of the 5 organic anion transporter genes in the tubule-specific microarray, relative to whole flies. Similarly, the genes for 11 of 21 organic cation transporters are enriched 4–37-fold in the tubules relative to whole flies. The prime significance of this study is that it suggests a re-interpretation of the major functions of the Drosophila tubule. The epithelium is now viewed less as one primarily concerned with the secretion of inorganic salts and water and more as an epithelium richly endowed with solute transporters, including
transporters for many potential toxins. An additional significance of the microarray study is that enrichment of genes for transporters for so many organic and inorganic solutes implies that a non-specific paracellular pathway plays only a minor role for many toxins (Wang et al., 2004; Dow and Davies, 2006).

3 Secretion of physiological ions by insect Malpighian tubules

The hemipteran R. prolixus provides a useful model species for studying excretion of physiological ions and water that are present in excess of the insect’s needs. Rhodnius ingests blood meals which may exceed the animal’s unfed weight by a factor of 10 and during the subsequent diuresis it voids urine at a rate equal to its original body weight every 20–30 min. The prodigious rate of urine production means that each cell in the secretory segment of the Malpighian tubule secretes a volume of fluid equal to its own volume every 15 s (Maddrell, 1991) and cellular Cl\(^{-}\) content is exchanged every 5 s (O’Donnell et al., 2003). These extraordinary rates of ion and water transport, coupled with the large size of the cells, ~100 \(\mu\)m in length, make the Rhodnius Malpighian tubule highly suitable for studies of epithelial transport.

The major transporters and pathways for movements of Na\(^{+}\), K\(^{+}\) and Cl\(^{-}\) across the upper (secretory) segment of the Malpighian tubule of the hemipteran R. prolixus are shown in Fig. 2. This model of fluid secretion is based on data from electrophysiological recordings using intracellular and ion-selective microelectrodes, the effects of ion substitution and transport inhibitors and also on immunofluorescent localization of transporters. Tubules of Rhodnius and the ant Formica polyctena are composed of a single cell type in the secretory segment. In the following sections, each of the inorganic ion transporters in the Rhodnius tubule as well as the paracellular pathway are discussed separately. Tubules of dipteran insects such as mosquitoes and fruit flies, as well as those of orthopterans such as the locusts Locusta migratoria and Schistocerca gregaria, contain more than one cell type in the secretory segment. The arrangements of ion transporters and ion permeation pathways for these tubules are discussed in Section 2.2. The control of the Malpighian tubules and hindgut by diuretic and antidiuretic hormones has been extensively reviewed by Coast et al. (2002), and is therefore considered only briefly in the sections below.

3.1 Ion transport across Malpighian tubules composed of a single cell type

The blood upon which Rhodnius feeds has an osmolality of 320 mOsm kg\(^{-1}\), approximately 14% hypoosmotic to the insects hemolymph osmolality of
370 mOsm kg$^{-1}$ (Maddrell and Phillips, 1975b). Homeostasis therefore requires production of a hypoosmotic urine through a two-stage process involving secretion of Na$^+$, K$^+$, Cl$^-$ and water across the upper Malpighian tubule followed by reabsorption of K$^+$ and Cl$^-$ but not water downstream in the lower Malpighian tubule, as described in Section 4.1. The fluid voided from the insect during diuresis contains $\sim 120$ mM NaCl, matching the composition of the plasma fraction of the blood.

The fluid secreted by the upper segment of Malpighian tubules of *R. prolixus* during diuresis consists of approximately 100 mM NaCl and 80 mM KCl (Maddrell and Phillips, 1975b). Secretion of ions and osmotically obliged water by tubules of *Rhodnius* and other species is driven primarily by an apical vacuolar-type H$^+$-ATPase (Wieczorek et al., 1989; Maddrell and O’Donnell, 1992). As a general rule, the H$^+$-ATPase plays a dominant role in energizing transport across insect epithelia, in contrast to the role of the Na$^+$/K$^+$-ATPase in vertebrate epithelia. This difference may reflect the co-evolution of insects with plants, which provide a K$^+$-rich and Na$^+$-poor food source. It is suggested that electrogenic transport of H$^+$ from
the cell to the lumen energizes amiloride-sensitive exchange of cytoplasmic K$^+$ and/or Na$^+$ for luminal H$^+$ (Fig. 2). Entry of Na$^+$, K$^+$ and Cl$^-$ through a basolateral Na$^+$–K$^+$–2Cl$^-$ cotransporter has been proposed for tubules of *R. prolixus* on the basis of the effects of bumetanide, Na$^+$-free saline and Cl$^-$-free saline on fluid secretion and transepithelial potential (O’Donnell and Maddrell, 1984; Ianowski and O’Donnell, 2001). Na$^+$:K$^+$:2Cl$^-$ cotransport has also been implicated in basolateral entry of ions into Malpighian tubules of other species, including *Ae. aegypti* (Hegarty *et al*., 1991), *F. polyctena* (Leyssens *et al*., 1994), *Manduca sexta* (Audsley *et al*., 1993; Reagan, 1995), *Teleogryllus oceanicus* (Xu and Marshall, 1999b) and *L. migratoria* (Al-Fifi *et al*., 1998).

The Na$^+$/K$^+$-ATPase and K$^+$ channels, although present in the basolateral membrane of the upper tubule, play a minor role in net transtubular transport during diuresis in *Rhodnius*. In tubules of the ant *F. polyctena*, however, K$^+$ channels play a significant role in transepithelial transport when hemolymph K$^+$ levels are high, as described below in Section 3.1.5.

Measurements of transepithelial, basolateral and apical membrane potential have been used to identify the sequential activation of ion transporters when *Rhodnius* Malpighian tubules are stimulated with 5-hydroxytryptamine. A characteristic triphasic change in transepithelial electrical potential (TEP) has been reported when fluid secretion by isolated Malpighian tubules of *R. prolixus* is stimulated by serotonin (O’Donnell and Maddrell, 1984). From an initially negative value in unstimulated tubules (−2.5 mV, lumen-negative), TEP shifts to −33 mV in phase 1, +30 mV in phase 2 and −32 mV in phase 3 (Fig. 3). TEP was measured in the study of O’Donnell and Maddrell (1984) using the Ramsay technique, in which electrodes are positioned in bathing and secreted fluid droplets for tubules isolated under paraffin oil. The validity of this method of TEP measurement has been questioned on the grounds that, in tubules of small diameter, it may permit shunting of current from lumen to bath through the cells or through the thin layer of fluid adherent to the surface of that portion of the tubule in the oil (Aneshansley *et al*., 1988; Isaacson and Nicolson, 1989). TEP measurements using the Ramsay technique were therefore compared with two other methods in which the voltage-sensing electrode is positioned in the tubule lumen by impalement of the Malpighian tubule with a microelectrode or by means of a cannula advanced into the tubule (Ianowski and O’Donnell, 2001). TEP measured during each phase of the stimulation with 5-hydroxytryptamine was similar irrespective of the measurement technique, validating the original measurements of O’Donnell and Maddrell (1984). The large diameter of the *Rhodnius* tubule (∼100 μm) presumably minimizes the shunting of current described in the smaller diameter tubules of species such as *Ae. aegypti*. 

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Ion substitution experiments and the effects of specific pharmacological reagents were used to support the proposal that the three phases of the response of TEP to serotonin correspond to sequential activation of an apical Cl\(^{-}\) channel, an apical V-type H\(^{+}\)-ATPase and a basolateral Na\(^{+}\):K\(^{+}\):2Cl\(^{-}\) cotransporter (Fig. 3). Pre-incubation of tubules in Cl\(^{-}\)-free saline abolishes phase 1, whereas pre-incubation in the presence of the H\(^{+}\)-ATPase inhibitor bafilomycin abolishes phase 2. Pre-incubation of the tubules in saline containing bumetanide, an inhibitor of Na\(^{+}\):K\(^{+}\):2Cl\(^{-}\) cotransporters, abolishes phase 3 but does not alter phases 1 and 2. Negative going shifts in TEP during the first and third phases reflect enhanced movement of Cl\(^{-}\) across the apical membrane from cell to lumen. An increase in apical Cl\(^{-}\) conductance during phase 1 could account for this change in TEP. The transepithelial potential will thus reflect contributions of the V-type H\(^{+}\)-ATPase, tending to drive the lumen to more positive values, and compensating movement of Cl\(^{-}\) from cell to lumen, tending to drive the lumen to more negative values. Once the apical permeability to Cl\(^{-}\) has increased in phase 1, the apical membrane potential will be influenced by changes in intracellular Cl\(^{-}\) activity following enhanced entry of Cl\(^{-}\) into the cell by activation of a basolateral Na\(^{+}\):K\(^{+}\):2Cl\(^{-}\) cotransporter in phase 3.

**FIG. 3** Changes in transepithelial potential of the *Rhodnius* Malpighian tubule in response to stimulation with serotonin (5-HT). The current model suggests that the triphasic response corresponds to successive activation of (1) an apical Cl\(^{-}\) conductance, (2) the apical V-type H\(^{+}\)-ATPase and (3) the basolateral Na\(^{+}\):K\(^{+}\):2Cl\(^{-}\) cotransporter. Based on data in Ianowski and O'Donnell (2001).
3.1.1 The apical vacuolar $H^+\text{-ATPase}$

The structure, function and regulation of insect and other V-ATPases has been reviewed extensively in recent years, primarily by Wieczorek, Harvey and co-workers (Harvey and Wieczorek, 1997; Wieczorek et al., 2000). Beyenbach and Wieczorek (2006) provide an historical overview in a recent review. V-ATPases are multisubunit, heteromeric proteins which consist of two structural domains, a peripheral, catalytic $V_1$ domain ($\sim$ 500 kDa) and a membrane-spanning $V_o$ domain (100–250 kDa). Regulation of V-ATPase activity may be accomplished by multiple mechanisms, including disassembly/reassembly of $V_1$ and $V_o$ domains, oxidation of SH groups, complexation with activator or inhibitor proteins or small signalling molecules, and sorting of the holoenzyme or its subunits to target membranes (Merzendorfer et al., 1997). The first gene knockout of a V-ATPase in any animal was accomplished in *Drosophila* (Davies et al., 1996). This study reported that epithelial dysfunction may be detectable at an early stage in those tissues where V-ATPases play a plasma membrane role and that the genes are essential for normal embryonic development (ibid.). An experimental and in silico survey of the V-ATPase gene family in *Drosophila* has revealed that 13 of the more than 32 genes are specialized for epithelial roles. When these genes are disrupted, they display a clear tubule phenotype that is conserved across mutant alleles for almost all the V-ATPase subunits; the transparent tubules result from a loss of birefringence in the tubule lumen and is a consequence of a failure to excrete uric acid (Allan et al., 2005).

V-ATPases transduce the energy from ATP hydrolysis into a proton current across a membrane. The resultant electrochemical gradient $\Delta\mu_H$ is termed the proton motive force (PMF) and it is used to drive the movements of many other ions through carriers and channels. V-ATPases are inherently electrogenic because their activity pumps protons without an associated anion. In the absence of other ion carriers, the V-ATPase will generate a large voltage across a membrane and produce a modest change in $H^+$ concentration. In the presence of anion channels (e.g. Cl$^-$) a large acid flux is generated. By contrast, an alkalizing flux is generated in the presence of a cation: $nH^+$ antiporter or symporter and appropriate counterions (anions).

In contrast to P-type-ATPases, such as the $Na^+/K^+$-ATPase and the mammalian gastric $H^+\text{-ATPase}$, V-ATPases do not form a phosphorylated intermediate and are relatively insensitive to vanadate. They are sensitive to the potent and specific inhibitor bafilomycin A1, a macrolide antibiotic isolated from the fermentation of the Gram-positive actinobacteria *Streptomyces* spp. (Drose and Altendorf, 1997). Fluid secretion by the tubules of *Rhodnius* is inhibited 49% by 1 μM bafilomycin (Gutierrez et al., 2004), for example, and secretion by
Formica tubules is inhibited 50% by 10 μM bafilomycin (Dijkstra et al., 1994).

Immunohistochemical techniques have been used to determine the cellular location of V-ATPases. Fluorescently labelled antibodies to the vacuolar-ATPases of Ma. sexta midgut and bovine kidney bind to the apical membrane of the Malpighian tubules of Formica (Garayoa et al., 1995) and Ma. sexta (Klein, 1992). In Malpighian tubules of Formica, specific labelling of the brush border of the epithelium, extending along the entire length of the tubules, is consistent with an apical location for the V-type ATPase.

3.1.2 Apical K⁺/H⁺ and Na⁺/H⁺ exchange

The V-ATPase provides the electrical and/or chemical gradients necessary to drive exchange of cellular K⁺ (or Na⁺) for luminal H⁺ across the apical membrane of insect epithelia. In the midgut of Ma. sexta, 2H⁺ are exchanged for each K⁺ ion. The exchanger is thus electrogenic and is driven by the large lumen-positive electrical gradient established across the apical membrane by the V-ATPase (Wieczorek et al., 1991). Exchange of 2H⁺ for each K⁺ or Na⁺ also seems to be required in Malpighian tubules of Ae. aegypti (Beyenbach and Wieczorek, 2006). By contrast, measurements of membrane potential, intracellular and lumenal pH in Malpighian tubules of F. polyctena (Zhang et al., 1994) and R. prolixus (Ianowski and O'Donnell, 2006) show that electroneutral exchange (1H⁺:1K⁺ or 1H⁺:Na⁺) is sufficient to drive alkali cations from cell to lumen (Fig. 4A, B). When Rhodnius tubules are stimulated with the diuretic factor 5-HT there is a small acid shift in intracellular pH from 6.97 to 6.92 and an accompanying alkaline shift in lumen pH from 6.08 to 6.32 (Ianowski and O'Donnell, 2006). These changes suggest that the apical Na⁺/H⁺ and K⁺/H⁺ exchange activity is stimulated to a greater extent by 5-HT than is the V-ATPase, thus tending to drive the lumen slightly alkaline and the cells slightly acid, relative to unstimulated tubules. When stimulated Rhodnius tubules are exposed to the drug amiloride, which blocks Na⁺/H⁺ exchangers, the lumen pH shifts to a value approximately 0.5 pH units more acidic and fluid secretion is inhibited, consistent with continued operation of the apical V-ATPase when the apical Na⁺/H⁺ exchanger is blocked. The molecular identity of the apical Na⁺/H⁺ exchanger in the tubules has been the subject of several studies using dipteran insects and is therefore discussed in Section 3.2.1.

3.1.3 Na⁺:K⁺:2Cl⁻ cotransport

Basolateral cotransport of Na⁺, K⁺ and Cl⁻ plays an important role in uptake of these ions from the hemolymph in Malpighian tubules of the ant
FIG. 4 Electrochemical gradients and ion activities in (A) unstimulated and (B) 5-HT-stimulated Malpighian tubules of *Rhodnius prolixus*. Values for pH\textsubscript{bath}, pH\textsubscript{lumen}, and luminal a\textsubscript{K}, a\textsubscript{Cl} and a\textsubscript{Na} are presented as means \pm S.E.M. for (N) tubules. Values in (B) were recorded 30 min after stimulation with 10\textsuperscript{-6} M 5-HT. Intracellular and bath activities for Na\textsuperscript{+}, K\textsuperscript{+} and Cl\textsuperscript{−} (italicized) and electrochemical gradients (Δμ/F, in mV) for Na\textsuperscript{+}, K\textsuperscript{+} and Cl\textsuperscript{−} across the basolateral membrane are taken from Ianowski et al. (2002). Figure from Ianowski and O’Donnell (2006, Fig. 1). Reprinted with permission from the Company of Biologists.
F. polyctena (Leyssens et al., 1994), the cricket Te. oceanicus (Xu and Marshall, 1999b) and the hemipteran R. prolixus (Ianowski et al., 2002). Measurements with ion-selective microelectrodes for Na+, K+ and Cl− show that Na+:K+:2Cl− cotransport is thermodynamically feasible in both unstimulated and 5HT-stimulated Rhodnius tubules (Ianowski et al., 2002). Chloride activity in the cell is higher than expected on the basis of passive distribution of Cl− across the basolateral membrane in both unstimulated and 5HT-stimulated cells (Fig. 5). Cl− must therefore be actively transported across the basolateral membrane into the cell. K+:Cl− involvement in net transepithelial ion transport can be ruled out on the basis of these measurements, because the net electrochemical potential for such a transporter is positive, indicating that it would mediate net movement of K+ and Cl− from inside the cell to the bath, opposite to the direction of transepithelial transport during fluid secretion.

Operation of the Na+:K+:2Cl− cotransporter at a low rate in unstimulated tubules was suggested by Maddrell and Overton (1988) and is consistent with measured net electrochemical potentials. In unstimulated tubules, K+ may also enter cells from the bath both through an ouabain-sensitive Na+/K+-ATPase (Maddrell and Overton, 1988). Addition of ouabain to unstimulated tubules reduces one path for K+ entry and blocks the transport of Na+ from cell to bath, with a resulting increase in Na+ transport from cell to lumen (Maddrell and Overton, 1988).

Transepithelial ion transport increases nearly 1000-fold in response to serotonin. Stimulation increases both the apical ion transporters as well as

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**FIG. 5** Schematic diagram showing net electrochemical potentials (Δμ_{net}/F) for three cation:Cl− cotransporters in serotonin-stimulated tubules of *Rhodnius prolixus*. Corresponding values for unstimulated tubules are given in parentheses. (A) Na+:K+:2Cl−, (B) Na+:Cl− and (C) K+:Cl− (Ianowski et al., 2002, Fig. 2). Reprinted with permission from the Company of Biologists.
the basolateral Na\(^+\):K\(^+\):2Cl\(^-\) cotransporter (Maddrell and Overton, 1988; Maddrell, 1991). Under these conditions ion flux through the basolateral cotransporter is so much greater than that through the Na\(^+\)/K\(^+\)-ATPase that fluid secretion in stimulated tubules is insensitive to ouabain.

An intriguing property of the basolateral cation:Cl\(^-\) cotransporter in *Rhodnius* Malpighian tubules is the ability for K\(^+\) to be replaced by Na\(^+\). The study of Ianowski et al. (2004) is the first description of competitive inhibition of K\(^+\) transport by Na\(^+\) in a bumetanide-sensitive mechanism. The likely function of K\(^+\) replacement by Na\(^+\) is to permit homeostatic regulation of hemolymph K\(^+\) levels even when tubules are secreting fluid at near maximal rates. Although tubules bathed in saline containing 6 or 10 mM K\(^+\) and 137 mM Na\(^+\) secrete fluid at equal rates, the fluxes of Na\(^+\) and K\(^+\) differ greatly. For tubules in saline containing 6 mM K\(^+\), Na\(^+\) flux is 3-threefold that of K\(^+\), whereas the fluxes of Na\(^+\) and K\(^+\) are equal in saline containing 10 mM K\(^+\). Total cation (Na\(^+\) and K\(^+\)) flux is the same in the two salines. Moreover, addition of bumetanide reduces fluid secretion rate to the same extent in both salines. These findings are consistent with replacement of K\(^+\) by Na\(^+\) in a bumetanide-sensitive cotransport mechanism during serotonin-stimulated fluid secretion. In the extreme case of complete removal of K\(^+\) from the bathing saline, Malpighian tubules secrete fluid at 45% of the rate in the control saline and remain sensitive to bumetanide. However, transport is unaltered by the drug hydrochlorothiazide, an inhibitor of Na\(^+\):Cl\(^-\) cotransporters in vertebrate tissues. Dose–response curves relating percentage inhibition of fluid secretion to bumetanide concentration are identical for tubules bathed in control and K\(^+\)-free saline, suggesting that fluid secretion involves the same bumetanide-sensitive cotransport system in the presence or absence of K\(^+\). Malpighian tubules bathed in K\(^+\)-free saline also show the characteristic triphasic change of apical membrane potential in response to stimulation with serotonin. Addition of bumetanide to such tubules produces a lumen-positive shift in apical membrane potential, as described above for tubules in control saline. Taken together, measurements of fluid secretion rates, cation fluxes and apical membrane potentials indicate that Na\(^+\) can replace K\(^+\) in a single bumetanide-sensitive cotransport mechanism in the basolateral membrane.

Kinetic analysis also supports the idea that Na\(^+\) competes with K\(^+\) for transport by a basolateral bumetanide-sensitive cation/Cl\(^-\) cotransporter during fluid secretion. Increasing Na\(^+\) concentration in the bath increases the Michaelis–Menten constant (K\(_m\)) for K\(^+\) transport while the maximum transepithelial K\(^+\) flux (J\(_{\text{max}}\)) remains constant, consistent with competitive inhibition of K\(^+\) transport by Na\(^+\). This is in stark contrast to the common finding that increasing bathing saline Na\(^+\) concentration
produces an increase in K\(^+\) flux through bumetanide-sensitive Na\(^+\)/K\(^+\)/2Cl\(^-\) cotransporters in vertebrate epithelia (for reviews, see Russell, 2000).

During diuresis, *Rhodnius* produces copious amounts of NaCl-rich urine, effectively eliminating the nutrient-poor plasma fraction of the blood meal and retaining the nutrient-rich blood cells. Studies by Maddrell *et al.* (1993) showed that the upper tubule responds to reductions in hemolymph K\(^+\) concentration by reducing the K\(^+\) concentration in the secreted fluid. This reduction enhances reabsorption of K\(^+\) by the lower tubule, thereby contributing to homeostatic regulation of hemolymph K\(^+\). Competition between Na\(^+\) and K\(^+\) for transport by a single bumetanide-sensitive cotransporter in the upper tubule provides a mechanism for reducing secreted fluid K\(^+\) concentration, thereby reducing the need for K\(^+\) reabsorption by the lower Malpighian tubule. Importantly, the increase in Na\(^+\) flux when saline K\(^+\) concentration is reduced not only minimizes the loss of K\(^+\) but at the same time permits the rate of urine production to remain very high.

The contribution of K\(^+\) channels to transport of K\(^+\) from bath to lumen during fluid secretion by *R. prolixus* tubules can be ruled out on the basis of two types of evidence. First, treatment with Ba\(^{2+}\) does not inhibit fluid secretion, suggesting that K\(^+\) channels are not a significant component of transepithelial K\(^+\) transport at physiological concentrations of extracellular K\(^+\). Second, basolateral membrane potential depolarises in response to addition of Ba\(^{2+}\) (Ianowski *et al.*, 2002). Depolarization is consistent with blockage of K\(^+\) leakage into the bath, since the basolateral membrane potential appears to be determined primarily by the K\(^+\) conductance (i.e. basolateral membrane potential is very similar to the Nernst equilibrium potential for potassium, \(E_K\)). The depolarisation of the basolateral membrane potential after addition of Ba\(^{2+}\) is consistent with a small but positive value for \(\Delta m_K/F\) that favours movement of K\(^+\) from cell to bath through ion channels.

### 3.1.4 Transport of Cl\(^-\) across the apical membrane

Intracellular chloride activity in *Rhodnius* tubules declines when the apical membrane potential increases (i.e. as the transepithelial potential becomes more lumen-positive) in Na\(^+\)- or K\(^+\)-free saline or in the presence of bumetanide (Ianowski *et al.*, 2002). This finding is consistent with the presence of apical Cl\(^-\) channels which would mediate transport of Cl\(^-\) from cell to lumen when entry of Cl\(^-\) into the cell across the basolateral membrane is blocked. In Na\(^+\)-replete saline, intracellular Cl\(^-\) activity is near equilibrium across the apical membrane both in unstimulated and in serotonin-stimulated tubules (Fig. 4A, B), as would be expected if Cl\(^-\) channels mediate vectorial ion transport across the apical membrane. Cl\(^-\) would be expected to be very close to electrochemical equilibrium.
across the apical membrane if Cl\(^{-}\) channels account for most of the apical membrane conductance. Although 4,4'-diisothiocyanato-2,2'-disulfonic acid stilbene (DIDS) inhibits fluid secretion by *Rhodnius* tubules, it has no effect on transepithelial potential or intracellular pH, as would be expected if DIDS blocked apical Cl\(^{-}\) channels or Cl\(^{-}\)/HCO\(_3\)\(^{-}\) exchangers, respectively. This suggests that Cl\(^{-}\) channels, if present, are DIDS-insensitive and that the effects of fluid DIDS on fluid secretion may reflect indirect effects of DIDS, such as inhibition of mitochondrial functioning. It is worth pointing out that the possibility of a DIDS-sensitive channel mediating Cl\(^{-}\) entry into the cells across the basolateral membrane can be ruled out on the basis of the measured electrochemical gradients described above. Chloride activity is above electrochemical equilibrium, and the opening of basolateral Cl\(^{-}\) channel would result in an efflux rather than an uptake of Cl\(^{-}\).

The contribution of other pathways for Cl\(^{-}\) movement, such as apical K\(^{+}\):Cl\(^{-}\) cotransport and/or paracellular Cl\(^{-}\) movement, can be ruled out because they are thermodynamically unfeasible. Although the transepithelial potential across tubules of most insects is lumen-positive, the TEP in fully stimulated *Rhodnius* tubules is lumen-negative and a passive paracellular pathway for Cl\(^{-}\) movement is thus precluded. The predicted Nernst equilibrium activity of Cl\(^{-}\) in the lumen for a bathing saline Cl\(^{-}\) activity of 93 m\(\text{M}\) (Ianowski *et al.*, 2002) and a lumen negative transepithelial potential of –28 mV is 26 m\(\text{M}\), which is ~3.5-fold less than the measured activity of 109 m\(\text{M}\) (Ianowski and O’Donnell, 2006). The calculated electrochemical gradients also show that if a K\(^{+}\):Cl\(^{-}\) cotransporter were active in the apical membrane of *Rhodnius* Malpighian tubules, it would mediate net movement of these ions from the lumen into the cell, in the opposite direction to that required for fluid secretion. A proposal of K\(^{+}\):Cl\(^{-}\) cotransport from cell to lumen was based on the inhibition of fluid secretion by dihydroindenyl)oxy]alkanoic acid (Gutierrez *et al.*, 2004). However this drug is also known to cause mitochondrial damage (Pond *et al.*, 2004). In addition, the experiments involved perfusion with fluid containing 8 mM K\(^{+}\) (Gutierrez *et al.*, 2004). The potassium activity of this solution is ~ 6 m\(\text{M}\), more than 8 times less than the physiological level of 49 m\(\text{M}\) in the lumen of 5HT-stimulated tubules.

### 3.1.5 K\(^{+}\) channels

Although K\(^{+}\) channels are not implicated in net transepithelial K\(^{+}\) and fluid secretion by *Rhodnius* Malpighian tubules, they do appear to be important in tubules of the ant *F. polyctena* under some conditions. Multiple mechanisms of K\(^{+}\) transport by tubules of this species may be necessary because hemolymph K\(^{+}\) levels are quite variable. A study by
Van Kerkhove et al. (1989) showed that the hemolymph $K^+$ concentration in ants of one nest ranged from 6 to 20 mM, whereas in the hemolymph of ants from another nest the potassium concentration ranged from 12 to 70 mM. The Malpighian tubules may therefore be exposed to a wide range of $K^+$ concentrations in vivo, and this may explain the findings that a cascade of different mechanisms of $K^+$ uptake across the basolateral membrane operate as bathing saline $K^+$ concentration increases. The model of Leyssens et al. (1994) proposes that a $Na^+:K^+:2Cl^-/C0$ cotransporter which is sensitive to $10^{-5}$ M bumetanide is the primary mechanism of uptake at the lowest bathing saline $K^+$ concentrations of 5–10 mM. At the intermediate concentration of 51 mM $K^+$, a $K^+:Cl^-/C0$ cotransporter sensitive to high (10$^{-4}$ M) concentrations of bumetanide becomes active. At very high bathing saline concentrations (113 mM $K^+$) $K^+$ enters through basolateral $Ba^{2+}$-sensitive $K^+$ channels (Leyssens et al., 1994).

A role for basolateral $K^+$ channels has also been proposed for the main segment of the tubules of the black field cricket *Te. oceanicus* (Xu and Marshall, 1999b). In this species as well as in the ant *F. polyctena*, hemolymph $K^+$ and $Na^+$ probably vary as a result of omnivory. Under some conditions, $K^+$ may enter the Malpighian tubule cells through $Ba^{2+}$-sensitive $K^+$ channels. At lower hemolymph $K^+$ concentrations, bumetanide inhibits fluid secretion by 80% and inhibits $Na^+$, $K^+$ and $Cl^-$ secretion by 60%, 70% and 50%, respectively. This result suggests that a large proportion of net transepithelial ion transport is mediated by cotransport of these three ions (Xu and Marshall, 1999b).

3.2 ION TRANSPORT ACROSS MALPIGHIAN TUBULES COMPOSED OF PRINCIPAL CELLS AND STELLATE CELLS

The Malpighian tubules of dipterans such as the fruit fly *D. melanogaster* and the mosquito *Ae. aegypti* contain two cell types in the fluid secreting segment (Fig. 6). The structure and functioning of the Malpighian tubules of these species has been reviewed by Dow and Davies (2003) and by Beyenbach (2003). There is general agreement that cations and possibly some of the chloride moves through the larger principal cells which make up 80% of the cells. For the tubule secreting at the basal rate chloride is transported through the smaller stellate cells which comprise 20% of the cells in the secretory segment. When *D. melanogaster* tubules are stimulated with the amine tyramine or the peptides Drosokinin (Terhzaz et al., 1999) or leucokininin, a pathway through chloride channels has been proposed. An alternative proposal based primarily on the analysis of the *Ae. aegypti* Malpighian tubule indicates a paracellular pathway that is stimulated by leucokinin. Roles for potassium channels, $Na^+:K^+:2Cl^-/C0$ cotransporters and the $Na^+/K^+$-ATPase have been proposed for entry of
ions across the basolateral membrane of the principal cells. Ion movement from cell to lumen appears to involve much the same mechanisms as proposed for tubules of *Rhodnius* and *Formica*. An apical V-type H\(^+\)-ATPase drives protons from cell to lumen, and protons are then recycled back into the cell in exchange for potassium or sodium. The V-type proton pump has been extensively characterized in other insect epithelia such as the midgut of *Ma. sexta* (Beyenbach and Wieczorek, 2006) and the molecular characterization of the *D. melanogaster* Malpighian tubule V-type H\(^+\)-ATPase has been reviewed by Dow (1999) and Dow et al. (1997). The study of Allan et al. (2005) shows that the plasma membrane V-ATPase holoenzyme is a single isozyme and is composed of polypeptides derived from expression of 14 genes, not just in the Malpighian tubule but in all the major transporting epithelia (Allan et al., 2005).

3.2.1 *The apical and basolateral cation: proton exchangers*

The nature of the putative amiloride-sensitive putative Na\(^+\)/H\(^+\) and K\(^+\)/H\(^+\) exchangers has been the subject of much recent attention. The sodium
proton exchanger (NHE) gene family has been characterized in the *Drosophila* Malpighian tubules by Giannakou and Dow (2001). The order of inhibition of fluid secretion by amiloride derivatives (ethylisopropylamiloride ≫ 2,4-dichlorobenzamil > N,N-dimethylamiloride > amiloride ≅ benzamil) is consistent with the presence of NHE rather than epithelial Na⁺ channels (ENaC) in Malpighian tubules of *D. melanogaster* and *Ae. aegypti* (Petzel, 2000). The presence of Na⁺/H⁺ exchange is also consistent with RT-PCR data showing that NHE genes are expressed in Malpighian tubules of *Drosophila* whereas there is no evidence for expression of ENaC genes (Giannakou and Dow, 2001). The latter finding is consistent with the finding that amiloride inhibits transepithelial sodium secretion in *Ae. aegypti* tubules but does not alter membrane resistance, as would be expected for channel inhibition. Evidence presented by Petzel (2000) is most consistent with a basolateral location of NHE and a role in pH regulation. Malpighian tubules respond rapidly to changes in bathing saline pH and intracellular pH acidifies when sodium is removed from the bathing saline, consistent with the basolateral location of NHE. If NHE is located on the apical membrane and its function is to pump sodium into the lumen in exchange for protons, then exposure to sodium free bathing saline would be predicted to result in an alkalinization of the cell if the apical proton-ATPase continued to pump protons from cell to lumen.

The sodium/proton exchanger isoform 3 (NHE3) is localized to the basolateral membrane in tubules of *Ae. aegypti* and co-localizes with the Na⁺/K⁺-ATPase. Na⁺ uptake in transgenic cells expressing this isoform is insensitive to amiloride (Pullikuth et al., 2003, 2006). The authors propose that *AeNHE3* is not the exchanger operating in concert with the apical V-ATPase. However, *AeNHE3* may, by virtue of its basolateral localization, play a role in regulating intracellular pH that indirectly impacts V-ATPase and apical exchanger functions.

Kang’ethe et al. (2007) suggest that the apical Na⁺/H⁺ exchanger which transports cations into the lumen, driven secondarily by the proton gradient created by the V-type H⁺-ATPase, is sodium/proton exchanger isoform 8 (NHE8). Immunolocalization studies show *AeNHE8* expression in the apical membranes of Malpighian tubules, gastric caecae, and rectum. Heterologous expression of *AeNHE8* in NHE-deficient fibroblast cells results in an amiloride-sensitive sodium uptake. When membranes from yeast cells expressing the protein are reconstituted into lipid proteoliposomes, fluorometric assays reveal saturable exchange of Na⁺ and K⁺ for H⁺. *AeNHE8* may be coupled to the inward H⁺ gradient across the Malpighian tubules and may play a role both in the extrusion of excess sodium and potassium and in maintenance of intracellular pH (Kang’ethe et al., 2007).

An alternative explanation is that an electrophoretic antiporter (Na⁺/2H⁺) in the apical membrane of tubules and gut is driven by the
electrical potential difference across the membrane created by the V-type H\(^+\)-ATPase (Rheault et al., 2007). This would contrast the role of NHEs in catalyzing electroneutral Na\(^+\)/H\(^+\) exchange using the Na\(^+\) chemical gradient generated by the Na\(^+\)/K\(^+\) P-ATPase. A cDNA encoding an electrophoretic antiporter has been cloned from the alimentary canal of larval African malaria mosquito, Anopheles gambiae, and is termed AgNHA1. Immunolabeling of whole mounts and longitudinal sections of isolated alimentary canal show that AgNHA1 is expressed in proximal Malpighian tubules and also in the cardia, gastric caeca, anterior midgut, posterior midgut and rectum, as well as in the suboesophageal and abdominal ganglia (Rheault et al., 2007). AgNHA1 is a likely candidate for the mosquito homolog of the caterpillar K\(^+\)/2H\(^+\) antiporter identified by Wieczorek et al. (1991).

3.2.2 Ancillary role of the Na\(^+\)/K\(^+\)-ATPase

The role of the Na\(^+\)/K\(^+\)-ATPase in the basolateral membrane has long been controversial. In Drosophila, the Malpighian (renal) tubule contains large amounts of Na\(^+\)/K\(^+\)-ATPase that is known biochemically to be highly sensitive to ouabain, whereas the effects of the drug on the intact tissue appear to be variable and depend whether the tubules are secreting fluid at a basal rate or have been stimulated to secrete at higher rates by diuretic factors or cyclic AMP (cAMP). Ouabain, a specific inhibitor of the Na\(^+\)/K\(^+\)-ATPase, at concentrations of 10 \(\mu\)M depolarizes the electrical potential across the basolateral membrane (\(V_{bl}\)) and also causes a slight increase in the rate of fluid secretion by unstimulated tubules and an increase in the concentration of Na\(^+\) in the secreted fluid (Fig. 7). The rapid depolarization of \(V_{bl}\) by ouabain is consistent both with inhibition of the contribution of an electrogenic pump (3Na\(^+\):2K\(^+\)) and with a small decline in intracellular K\(^+\) levels. An increase in the level of Na\(^+\) within the cell in the presence of ouabain, in conjunction with the high Na\(^+\) affinity of the apical cation/H\(^+\) exchanger (Maddrell and O'Donnell, 1993), means that more Na\(^+\) is transported into the lumen of the tubule, resulting in a higher Na\(^+\) concentration in the secreted fluid and a slightly higher secretion rate (Fig. 7B, C). Similar effects were observed when ouabain was applied to the unstimulated Malpighian tubules of R. prolixus (Maddrell and Overton, 1988). In contrast, fluid secretion by unstimulated Malpighian tubules of Ae. aegypti (Hegarty et al., 1991) and L. migratoria (Anstee et al., 1979) is inhibited by 0.1–1 mM ouabain, as is secretion by cAMP-stimulated tubules of Drosophila (Linton and O'Donnell, 1999).

The Na\(^+\)/K\(^+\)-ATPase provides a minor route of K\(^+\) entry into the tubule cells of Drosophila given that there is only a slight increase in the secretion rate and the [Na\(^+\)]/[K\(^+\)] ratio of the secreted fluid when ouabain is applied. These data suggest that the functions of the Na\(^+\)/K\(^+\)-ATPase
are to maintain the potential across the basolateral membrane by maintaining differential ion concentrations. Moreover, transport of Na\(^+\) from the cell to the bath by the Na\(^+\)/K\(^+\)-ATPase may permit high levels of solutes to be accumulated by Na\(^+\)-coupled entry mechanisms with little loss of Na\(^+\) by secretion into the lumen. Maintenance of low intracellular Na\(^+\) concentrations by the Na\(^+\)/K\(^+\)-ATPase may be necessary in part because of the high affinity of apical cation/H\(^+\) exchangers for Na\(^+\) (Maddrell and O'Donnell, 1993). The concentration of ouabain required to inhibit fluid secretion by isolated Malpighian tubules in standard bathing medium (SBM). The sample recording in (A) shows the change in V\(_{\text{bl}}\) in response to the addition of ouabain for the period indicated by the horizontal bar. Downward and upward arrows indicate cell impalement and withdrawal of the microelectrode into the bathing saline, respectively. In (B) and (C) the values are means±S.E.M. (n = 9–16) before and 30 min after the addition of either ouabain (experimental group, open columns) or ethanol (0.1%; control group, filled columns). Asterisks indicate significant differences between the experimental and control groups (P<0.05). Redrawn from Linton and O'Donnell (1999, Figs. 4 and 5). Reprinted with permission from the Company of Biologists.
active sites in intact cells and that this effect can be minimized through the addition of competing substrates such as taurocholate for the ouabain transporter (Torrine et al., 2004). However, it must be noted that measurements of the concentrations of organic anions and organic cations near the cell surface typically indicate concentrations only a few percent below those in the bulk medium (e.g. O'Donnell and Rheault, 2005). One possibility is that co-localization of the ouabain transporter organic anion transporting polypeptide (OATP) with the Na\(^+/K^+\)-ATPase prevents ouabain from reaching inhibitory concentrations within the basolateral infoldings of principal cells, although direct evidence for such an effect is lacking. An alternative explanation is that the competing substrates may indirectly inhibit fluid secretion, as occurs in response to high concentrations (> 0.5 mM) of organic anions such as salicylate (O'Donnell and Rheault, 2005). For example, 0.06 mM taurocholate inhibits fluid secretion by Drosophila tubules by ~25%, whereas in combination with 1 mM ouabain the effect is increased to 50% (Torrine et al., 2004).

An additional complication is the finding of high levels of Na\(^+/K^+\)-ATPase in the stellate cells of the Malpighian tubules of the mosquito Ae. aegypti (Patrick et al., 2006). The role of the stellate cells in transcellular Cl\(^-\)/C\(_\text{O}\) transport is discussed below in Section 3.2.4, but it is uncertain as to whether the cells are also involved in transcellular cation transport, or if the Na\(^+/K^+\)-ATPase plays a role in energizing ion transport for homeostatic functions such as pH regulation or cell volume regulation.

3.2.3 Na\(^+\):K\(^+\):2Cl\(^-\) cotransporter

Fluid secretion by the Malpighian tubules of D. melanogaster is inhibited approximately 25% by bumetanide (Linton and O'Donnell, 1999). When bumetanide alone is added to the saline bathing Drosophila tubules, K\(^+\) flux decreases but Na\(^+\) flux is unchanged. Under these conditions, however, much of the Na\(^+\) that enters the cells is recycled back to the bathing saline by the Na\(^+/K^+\)-ATPase. Blockade of the Na\(^+/K^+\)-ATPase with ouabain increases secreted fluid Na\(^+\) concentration from 23 to 45 mM, and subsequent addition of bumetanide reduces transepithelial fluxes of both Na\(^+\) and K\(^+\) by nearly equal amounts, consistent with inhibition of an Na\(^+\):K\(^+\):2Cl\(^-\) cotransporter (Ianowski and O'Donnell, 2004). Bumetanide also inhibits Na\(^+\), K\(^+\) and Cl\(^-\) secretion by Aedes tubules which have been stimulated with cAMP, consistent with the presence of a cAMP-stimulated Na\(^+\):K\(^+\):2Cl\(^-\) cotransporter (Hegarty et al., 1991).

In tubules of both species, the Na\(^+\) gradient across the basolateral membrane provides the driving force for secondary active Cl\(^-\) transport through a bumetanide-sensitive Na\(^+\):K\(^+\):2Cl\(^-\) cotransporter. The Na\(^+\) gradient is maintained by the actions of two other transporters; the
basolateral Na\(^+\)/K\(^+\)-ATPase and the apical Na\(^+\)/H\(^+\) exchanger, which in turn is driven by the proton gradients established by the V-type H\(^+\)-ATPase. The Na\(^+\)/K\(^+\)-ATPase is thus poised to play a pivotal role in setting the Na\(^+\)/K\(^+\) ratio of the secreted fluid. In particular, in unstimulated Malpighian tubules the basolateral Na\(^+\)/K\(^+\)-ATPase may return much of the Na\(^+\) that enters the cell through Na\(^+\)-linked exchangers or cotransporters back to the hemolymph. The result will be high levels of K\(^+\) in the secreted fluid. When the Na\(^+\)/K\(^+\)/2Cl\(^-\)/C\(_0\) cotransporter is stimulated along with the apical vacuolar H\(^+\)-ATPase in response to increases in cellular levels of cAMP, there is an increase in the entry of both K\(^+\) and Na\(^+\) and in fluid secretion rate. As a result, the Na\(^+\) content of the secreted fluid increases. This is particularly the case if the cotransporter is able to accept alternative stoichiometries. As noted above in Section 3.1.3, there is evidence that the alkali cation chloride cotransport or in the R. prolixus Malpighian tubule is able to accept Na\(^+\) in place of K\(^+\). As a result, the Na\(^+\) content of the secreted fluid can increase above that of K\(^+\). In addition, some of the K\(^+\) which enters through the Na\(^+\)/K\(^+\)-ATPase or through the alkali cation:chloride cotransporter may leak into the bathing saline through K\(^+\) selective channels in the basolateral membrane.

3.2.4 Na\(^+\) conductance in the basolateral membrane of Aedes tubules

In Malpighian tubules of Aedes, the natriuretic peptide or its second messenger cAMP activates both basolateral Na\(^+\):K\(^+\):2Cl\(^-\) cotransport (Hegarty et al., 1991) as well as a basolateral Na\(^+\) conductance (Sawyer and Beyenbach, 1985). The depolarization of the basolateral membrane voltage together and the concomitant decline in transepithelial resistance and the fractional resistance of the basolateral membrane are consistent with an increase in the Na\(^+\) conductance of the basolateral membrane. Enhanced entry of Na\(^+\) into the principal cell presumably increases cytoplasmic [Na\(^+\)], thereby increasing its availability for extrusion across the apical membrane. The net effect is selective stimulation of transepithelial NaCl and water secretion.

3.2.5 Cl\(^-\) channels and paracellular pathways

Patch clamp studies have revealed two types of Cl\(^-\) channel in excised, inside-out apical membrane patches of the stellate cells of Ae. aegypti. Type I channels have a conductance of 24 pS, an open probability of 0.82 and a mean open time of 867 ms (O’Connor and Beyenbach, 2001). Type II Cl\(^-\) channels have a conductance of 8 pS, an open probability of 0.07 and a mean open time of 7.5 ms. The high density and halide selectivity sequence of the type I Cl\(^-\) channel is consistent with a role in transepithelial Cl\(^-\)
secretion under control conditions. The permeability of the tubule wall and type I channels share Eisenman halide selectivity sequence I (in order of decreasing permeability): $I^- > Br^- > Cl^- > F^- >$ isethionate. Type I channels, therefore, appear to be part of the transepithelial pathway taken by $Cl^-$ under control conditions. In contrast, when stimulated by leucokinin, the tubule wall exhibits Eisenman selectivity sequence III: $Br^- > Cl^- > I^- > F^- >$ isethionate (Yu and Beyenbach, 2001), suggesting that type I channels are not part of the leucokinin-stimulated pathway. Excised patches show no effects of removal and/or chelation of $Ca^{2+}$ upon type I currents, as might be expected if type I channels respond to the elevation of intracellular $Ca^{2+}$ evoked by leucokinin. However, this effect may be due to loss of essential cell components (e.g. protein kinases) required to complete signal transduction in response to leucokinin. An essential future experiment will be to determine whether current through stellate cell ion channels increases in response to leucokinin when current is recorded using the cell-attached configuration.

DIDS-sensitive ‘maxi’ $Cl^-$ channels with a conductance of 256 pS are found in areas of the apical membrane consistent with their presence in stellate cells of the *Drosophila* Malpighian tubule (O’Donnell et al., 1998). Both fluid secretion and the calcium-mediated increase in anion conductance are inhibited by $Cl^-$ channel blockers such as niflumic acid. Vibrating probe analysis has revealed a small number of current density hot spots, coincident with stellate cells, and currents associated with the stellate cells are reduced by low-chloride saline or chloride channel blockers. Targeted expression of an aequorin transgene has been used to demonstrate that the neurohormone leucokinin elicits a rapid increase in intracellular calcium levels in stellate cells before physiological effects are evident. In sum, these data show that leucokinins act on *Drosophila* tubule stellate cells to elevate intracellular calcium and thereby increase transcellular chloride conductance through channels.

For tubules of both *Drosophila* and *Aedes* secreting at the basal rate there is agreement that transepithelial $Cl^-$ flux is mediated by stellate cell $Cl^-$ channels (O’Donnell et al., 1998; O’Connor and Beyenbach, 2001). In *Aedes* tubules, however, a paracellular pathway has been proposed in response to stimulation with leucokinin. The transepithelial resistance of *Aedes* tubules falls 4.3-fold in response to leucokinin, whereas the input resistance of the principal cells falls only 1.7-fold (Masia et al., 2000). Coupled with the finding that leucokinin reduces transepithelial resistance in specific regions of tubules apparently lacking stellate cells (Yu and Beyenbach, 2004), this suggests that leucokinin acts on a site separate from both cell types, i.e. the paracellular pathway. In support of this finding, the large transepithelial $Cl^-$ diffusion potentials induced by leucokinin are similar for lumen-to-bath and bath-to-lumen transepithelial gradients.
The latter finding is most easily explained on the basis of a single diffusion barrier such as a septate junction.

3.3 AQUAPORINS AND OSMOTIC PERMEABILITY OF MALPIGHIAN TUBULES

The osmotic permeability of the *Rhodnius* Malpighian tubule wall has been measured by simultaneous perfusion of internal and external surfaces. The measured permeability of the upper tubule, $4.3 \times 10^{-3} \text{ cm s}^{-1} \text{ osmol}^{-1}$, can be increased 35%, to $5.8 \times 10^{-3} \text{ cm s}^{-1} \text{ osmole}$, when corrections are made for unstirred layer effects (O’Donnell et al., 1982). The high osmotic permeability of the tubule wall is one factor which indicates that the route for transepithelial water movement during fluid secretion is transcellular. A transcellular route, rather than a paracellular one, is also more likely given the very small frontal area of the intercellular clefts. The paracellular clefts between *Rhodnius* tubule cells are about 17 nm in width and occupy about 0.039% of the frontal area of the tubules. If all the fluid moved through the clefts (120 nl min$^{-1}$) it would pass through at 660 $\mu$m s$^{-1}$ and the fluid would take only 22 ms, therefore, to traverse the 15 $\mu$m length of the cleft. Small ions and solutes such as urea take 2–3 times as long to diffuse 15 $\mu$m, and larger solutes such as sucrose would take longer still. One would therefore expect a strong correlation between the rate of fluid transport and passive transepithelial movements of extracellular markers. In particular, entrainment of solutes into a rapidly flowing fluid will accelerate transepithelial solute movement. However, neither sucrose nor inulin crosses the wall of the tubule any faster during rapid fluid secretion (Maddrell, 1981). This absence of significant solute/solvent coupling in the clefts, together with the small number and area of the clefts, militates against significant paracellular water flow during secretion by the tubules.

Measurements by Whittembury et al. (1986) raised the possibility of some paracellular fluid flow, based on entrainment of some solutes (urea, mannitol, sucrose) but not others (inulin, raffinose, dextran) in fluid flow. Interpretation of these findings is complicated by the finding that changes in intraluminal hydrostatic pressure can greatly alter the rates of permeation of molecules such as sucrose (O’Donnell et al., 1983).

The most convincing argument against paracellular flow and in support of transcellular flow is the discovery of aquaporins (water channels) in Malpighian tubules and the finding that tubule osmotic permeability is reduced by mercurial agents which block aquaporins (Hernandez et al., 2001; Echevarria et al., 2001). Expression of the mRNA for the *Rhodnius* aquaporin is increased after a blood meal and in tubules treated with 5-hydroxytryptamine or cAMP (Martini et al., 2004). In the tubules of the cricket *Acheta domesticus*, aquaporin expression appears to be hormonally regulated in both apical and basolateral membranes (Spring et al., 2007). It thus appears that transporters used for the excretion of excess water and
excess ions can be regulated in insect Malpighian tubules. Aquaporins are also expressed in both the principal cells and stellate cells of the *Drosophila* Malpighian tubule (Kaufmann *et al*., 2005). One of the aquaporins, *Drosophila* integral protein (DrIP) is similar to human AQP4, a channel that exhibits the highest rates of water transport, and it is localized to the stellate cells. DrIP contains several residues consistent with its actions as a water-specific channel.

4 Secretion and reabsorption by downstream segments of the Malpighian tubule and the hindgut

A common feature of the insect excretory system is the modification of the fluid secreted by Malpighian tubules in a downstream segment of the tubules themselves or in the hindgut. Although reabsorption of useful ions and water is accomplished by the hindgut in most species, in some cases a proximal (lower) segment of the Malpighian tubule is involved.

4.1 REABSORPTION BY THE MALPIGHIAN TUBULES

For the blood feeder *R. prolixus*, the high surface area offered by the lower Malpighian tubule relative to the volume of fluid in its lumen permits rapid reabsorption of KCl during the post-prandial diuresis. In *Drosophila*, reabsorption of both K\(^+\) and water in the lower Malpighian tubule probably serves a preparatory role, prior to additional reabsorption in the hindgut. The fluid secreted by the upper Malpighian tubule of *Rhodnius* (approximately 100 mM NaCl, 80 mM KCl) is modified by reabsorption of almost all the KCl but not water as the secreted fluid passes through the 30% of the lower tubule’s length closest to the hindgut. Osmotic permeability of the lower tubule is much lower in this region relative to the upper tubule or the upper region of the lower tubule (O’Donnell *et al*., 1982). Water is thus not reabsorbed as a consequence of KCl reabsorption, so what passes into the hindgut is both NaCl-rich and hypoosmotic. *Rhodnius* feeds on blood which is hypoosmotic to its own hemolymph, so this process of isosmotic secretion by the upper tubule followed by KCl reabsorption by the lower tubule preserves osmotic and ionic homeostasis.

The transporters implicated in reabsorption of K\(^+\) and Cl\(^-\) by the lower Malpighian tubule differ fundamentally from those which secrete these ions across the upper Malpighian tubule (Fig. 8). The current model proposes that K\(^+\) is moved thermodynamically uphill from lumen to cell by an ATP-dependent pump, resembling the H\(^+\)/K\(^+\)-ATPase of the gastric mucosa, and that K\(^+\) moves passively from cell to bathing saline (hemolymph) through an electrodiffusive pathway, i.e. K\(^+\) channels. K\(^+\) reabsorption is unaffected by amiloride (which blocks a putative K\(^+\)/H\(^+\)
exchanger in the upper tubule) and by the V-type-ATPase inhibitor bafilomycin A₁, applied to the basolateral or apical services of the lower tubule (Haley and O’Donnell, 1997). By contrast, K⁺ reabsorption is inhibited by drugs such as omeprazole, SCH 28080, SKF 96067 and SKF 96356. These drugs are either activated by low pH or are weak bases which tend to accumulate in acidic compartments. The pH of fluid in the lower tubule is 8.48, approximately 1.2 units more alkaline than the fluid secreted by the upper tubule. The concentration of omeprazole and the other drugs which inhibit lower tubule KCl reabsorption are higher, therefore, than those required to inhibit gastric H⁺ secretion in vertebrates, but comparable to the levels which inhibit H⁺/K⁺-ATPases in other tissues (Haley and O’Donnell, 1997). K⁺ reabsorption by the lower tubule is also blocked by the K⁺ channel blocker Ba²⁺. Changes in basolateral

FIG. 8 Schematic diagram summarizing the working hypothesis for the mechanisms of KCl reabsorption by the lower Malpighian tubule of Rhodnius prolixus. On the apical membrane, an H⁺/K⁺-ATPase inhibitable by drugs such as omeprazole and SCH 28080 exchanges lumenal K⁺ for cellular H⁺ (Haley and O’Donnell, 1997). Lumenal acidification is prevented by a compensatory flux of base through a putative SITS-insensitive Cl⁻/HCO₃⁻ exchanger. Cl⁻ exits the cell through channels which can be blocked by diphenylamine-2-carboxylate (DPC) and 5-nitro-2-(3-phenylpropylamino) benzoic acid (NPPB), 4-acetamido-4'-isothiocyanatostilbene-2,2'-disulfonic acid (SITS) and diisothiocyanatostilbene-2,2'-disulfonic acid (DIDS). K⁺ exits the cell through Ba²⁺-sensitive channels (Haley and O’Donnell, 1997). Acetazolamide (ACZ) inhibits carbonic anhydrase (CA), thereby inhibiting KCl reabsorption by preventing formation of HCO₃⁻ and H⁺ for the apical exchangers. Redrawn from Haley et al. (1997, Fig. 5). Reprinted with permission of Elsevier Science.
membrane potential in response to changes in bathing saline K⁺ concentration and blockade of these changes by Ba²⁺ also indicate the presence of basolateral K⁺ channels.

The current model for the mechanism of chloride reabsorption by the lower tubule proposes that chloride moves from lumen to cell through a stilbene-insensitive Cl⁻/HCO₃⁻ exchanger and then exits the cell through basolateral chloride channels (Fig. 8; Haley et al., 1997). Changes in basolateral membrane potential and transepithelial potential in response to bathing saline Cl⁻ concentration indicate the presence of Cl⁻ channels in the basolateral membrane. The blockade of Cl⁻-dependent changes in potential and KCl reabsorption by Cl⁻ channel blockers but not by compounds that inhibit K⁺:Cl⁻ and Na⁺:K⁺:2Cl⁻ cotransporters are also indicative of a role for chloride channels. Inhibition of KCl reabsorption by carbonic anhydrase inhibitors such as acetazolamide suggests that cellular CO₂ is a source of both H⁺ and HCO₃⁻ for exchange of K⁺ and Cl⁻, respectively. Some H⁺ and HCO₃⁻ may combine in the tubule lumen to form CO₂, which may diffuse back across the tubule ball into the bathing saline. The pH of the secreted fluid will thus be set by the rates of both H⁺ and HCO₃⁻ influx and CO₂ loss.

The lower tubule of D. melanogaster reabsorbs fluid and transports ions into or out of the lumen (O'Donnell and Maddrell, 1995). Secretion of Ca²⁺ by the lower tubule and acidification of the lumen are discussed in Section 6.1. The K⁺-rich fluid secreted by the upper tubule is modified by reabsorption of K⁺ and water. Modification of the urine by reabsorption is also accomplished by the lower tubule or ampulla or both in the house cricket Ac. domesticus (Spring and Hazelton, 1987), and possibly by reabsorptive cells scattered throughout the epithelia in tubules of the cricket Te. oceanicus (Marshall et al., 1993). Reabsorption by specific cells or regions of the Malpighian tubule thus reduces the amount water and ions that must be reabsorbed by the hindgut. A similar role is performed by the ileum of the locust, S. gregaria (Phillips et al., 1986). Reabsorption of water and ions in the ileum allows the rectum, downstream of the ileum, to process more thoroughly a reduced volume of fluid delivered to it.

4.2 THE INSECT HINDGUT

The hindgut, and the rectum, in particular, is a major site of water conservation in insects. The hindgut modifies the fluid secreted by the Malpighian tubules by reabsorbing useful solutes and water and eliminating those molecules which are present in excess or are toxic. Up to 80–95% of the ions and water secreted by the Malpighian tubules is reabsorbed in the hindgut of stick insects (Ramsay, 1955a,b), water-stressed locusts and cockroaches (Phillips 1964a,b; Wall, 1971), and about 65% in blowflies. In insects such as the mealworm larvae of
Tenebrio molitor water reabsorption in the rectum is so efficient that the fecal pellets which are expelled are almost completely dry. Water reabsorption in the insect hindgut is tightly regulated. For example, less than 10% of water is reabsorbed from the hindgut when locusts are fed on a succulent diet (Loveridge, 1975), and blood-feeders such as Rhodnius recover virtually no water during diuresis.

Four distinct physiological mechanisms of water conservation are seen in the hindgut of different species (Bradley, 1985; Phillips et al., 1986):

1. absorption of hypoosmotic fluid by rectal pads or papillae in cockroaches, locusts and adult blowflies. The classic paper by Wall (1971) demonstrated the importance of local osmosis and solute recycling during fluid absorption. Active transport of Na\(^+\), K\(^+\) and Cl\(^-\) creates the osmotic gradients for absorption of water from the rectal lumen. Fluid and ions move from the lumen into the rectal epithelial cells and then into the intercellular spaces between the epithelial cells. As the solutes and water move along the intercellular channels, the ions are reabsorbed into the epithelial cells for subsequent return to the intercellular spaces so as to permit further osmotic reabsorption from the lumen. Microsampling of fluid from the rectal tissues provides direct support for this scheme (Wall, 1971). Fluid in the lateral intercellular spaces near the lumen is hyperosmotic, and the total ion concentration in the intercellular channel declines toward the hemocoel.

The elegant work of Phillips and co-workers suggests that in the locust rectum a primary mechanism of Cl\(^-\) transport is responsible for absorption of ions and water (Phillips et al., 1986, 1996). This Cl\(^-\) transport is not coupled to or driven secondarily by movements of Na\(^+\), K\(^+\), HCO\(_3^-\), Ca\(^{2+}\) or Mg\(^{2+}\). Although an apical V-type H\(^+\)-ATPase acidifies the hindgut lumen, it does so at a rate that is only 10–15% of Cl-dependent short-circuit current across the rectum. Chloride transport is stimulated by chloride transport stimulating hormone (CTSH) and ion transport peptide (ITP), both of which act through intracellular cAMP.

S. gregaria ITP (SchgrITP) is the first reported insect member of a large family of crustacean neuropeptide hormones that control blood sugar levels, molting and reproduction. A partial amino acid sequence of ITP purified from locust nervous corpus cardiacum was used to derive degenerate primers that were used to amplify a cDNA from brain RNA using reverse transcription and the polymerase chain reaction (RT-PCR). The cDNA sequence was extended using anchored PCR to yield a partial, 517bp cDNA clone which encodes a putative ITP prohormone (Meredith et al., 1996). The prohormone can be cleaved at two dibasic amino acid
sites to yield a 72 residue active amidated peptide. The deduced amino acid sequence from the cDNA agrees completely with the amino acid sequence and molecular mass (8564 Da) derived from analysis of purified ITP. Native ITP has the same actions as crude CC extracts in stimulating Cl\(^-\), Na\(^+\), K\(^+\) and fluid absorption. *Drosophila* Kc1 cells transfected with preproITP secrete a peptide (KcITP) that is very active in the locust ileal bioassay (Wang et al., 2000; Zhao et al., 2005). Site-directed mutagenesis and expression of KcITP mutants has shown that that amidated leucine (residue 72) at the C-terminus is essential for receptor stimulation.

(2) secretion of hyperosmotic salt solutions by salt glands. Salt water mosquito larvae which are capable of hypoosmotic regulation produce a concentrated urine by the secretion of a hyperosmotic fluid across the posterior rectum. Reabsorption of ions and useful metabolites may occur across the anterior rectum (Strange and Phillips, 1985; Bradley, 1987). Na\(^+\), K\(^+\), Mg\(^{2+}\) and Cl\(^-\) are all actively transported by the posterior segment to the lumen side (Bradley and Phillips, 1977).

(3) secretion of hyperosmotic salt solutions within the cryptonephridial complexes of beetle larvae. The distal portions of the Malpighian tubules of *Te. molitor* are held closely to the surface of the rectum by an enveloping perinephric membrane, so that the rectum and the tubules act in concert. The tubules secrete K\(^+\), Cl\(^-\) and to a lesser extent Na\(^+\) into the lumen. The concentration of KCl in the tubule lumen in some tenebrionid larva approaches and may even transiently exceed saturation. The resultant ‘osmotic sink’ (Machin, 1983) is used for fecal dehydration or atmospheric water vapour absorption. The ultimate source of ions for transport into the tubule lumen is the hemolymph. The perinephric space is a functional compartment beneath the perinephric membrane and surrounding the tubules. Potassium activities in the perinephric space are close to electrochemical equilibrium with the hemolymph, whereas the activities of Na\(^+\) and H\(^+\) are reduced fivefold and threefold, respectively, below the corresponding Nernst equilibrium values. It appears that cations move from hemolymph across the perinephric membrane into the perinephric space, which is the immediate source for cations transported into the tubule cells and then the lumen (O’Donnell and Machin, 1991). The main function of the perinephric membrane is to provide an osmotic barrier so water does not move from the hemolymph into the complex in response to the osmotic sink within the Malpighian tubule lumen; instead, water is extracted from the rectal lumen.

(4) the use of anal sacs in thysanurans (silverfish and firebrats) and flea larvae. In this case, there is no evidence for high concentrations of...
5 Nitrogenous waste excretion

Ammonia resulting from catabolism of nitrogen-rich compounds such as amino acids is toxic and must be either eliminated from the body or converted into less toxic forms. Whereas aquatic species typically excrete ammonia as the predominant nitrogenous waste, many terrestrial insects expend metabolic energy to synthesize uric acid. Some larger species, or those with access to water, excrete ammonia, either alone or in combination with uric acid.

5.1 Uric Acid

5.1.1 Synthesis and storage of uric acid

Two properties suit uric acid for the purpose of nitrogen excretion: it contains 33% nitrogen, and its solubility is particularly low (1.1 mM at pH 5.5) under the acid conditions found in the terminal section of the excretory system (Phillips, 1977). Elimination of uric acid or urate salts thus leads to the excretion of large amounts of nitrogen without incurring much loss of water. Uric acid is synthesized primarily in the fat body. During larval growth, uric acid is released into the hemolymph shortly after synthesis and is subsequently excreted by the Malpighian tubules.
Xanthine oxidase, which catalyses uric acid formation from its immediate precursor xanthine, is present in fat body, Malpighian tubules and intestine in the silkworm *Bombyx mori* (Hayashi, 1960; Komoto *et al.*, 1999). In *Drosophila*, a search for xanthine oxidase in FlyAtlas: the *Drosophila* adult gene expression atlas (Chintapalli *et al.*, 2007), shows that the genes are much more enriched in the tubules than in the gut or fat body.

The excretory system is non-functional during metamorphosis of holometabolous insects. In lepidopterans, therefore, uric acid is sequestered in the fat body during larval-pupal metamorphosis (Buckner, 1982). In fly pupae, uric acid is accumulated not only in the fat body, but also in the intestine and the Malpighian tubules in response to the shut down of the excretory organs during this phase of the life cycle (Schwantes, 1990). The switch from excretion to storage is under hormonal control in larval *Ma. sexta* and it occurs during the transition from the feeding to the wandering stage (Buckner, 1982). There is an increase in fat body urate levels and a corresponding decline in hemolymp urate concentration at this time. An increase in the titre of 20-hydroxyecdysone and a decrease in the titre of juvenile hormone I control the switch from uric acid excretion to storage in the fat body. Fat body cells respond to 20-hydroxyecdysone and switch from secretion to storage of urate only in the absence of juvenile hormone. When exogenous juvenile hormone I or the analog methoprene are injected, storage is prevented or reversed and there is no decline in hemolymph uric acid levels.

During the subsequent transformation from pupa to adult, the fat body releases stored urate, and the timing of release corresponds to declining levels of 20-hydroxyecdysone in the hemolymph (Ehresmann *et al.*, 1990). Release of uric acid can be delayed by exogenous 20-hydroxyecdysone, but not during the final stages of transformation. A second ancillary hormone may be involved, therefore.

Urate is stored in the fat body of *Ma. sexta* pupae as discrete membrane bound vacuoles, termed uric acid storage vacuoles, which can be stained with reduced silver to reveal stored uric acid. Electron microscopy shows the vacuoles to be spherical or biconcave, and 0.5–2 µm in diameter (Buckner *et al.*, 1985). Biochemical and elemental analyses indicate that the storage vacuoles contain 75% uric acid, 5% protein, 0.5% carbohydrate, 6–7% water, 6.7% potassium and 0.5% sodium. At high magnification, the granules appear in micrographs as concentric whorls of fibrous or filamentous material. Granules appear to be constructed from tightly coiled fibres of crystalline uric acid or urate salts, and each fibre is enveloped with protein (Buckner *et al.*, 1990).

Uric acid is also stored in wings or larval cuticle in Lepidoptera, presumably to provide pigmentation. Levels of uric acid in the hemolymph increase during development of *Ma. sexta* pupae, possibly as a consequence of the release of uric acid from epidermal stores.
(Buckner and Newman, 1990). The uric acid is subsequently either excreted or stored in the fat body. Uric acid is also stored in the eyes in *Drosophila*; this storage excretion allows the crystals to act as reflectors to help light penetrate more deeply into the eye. In flies with the mutation *rosy* the eyes are dull red, because they lack uric acid crystals around their ommatidia (Reaume *et al.*, 1991).

### 5.1.2 Active transport of urate by Malpighian tubules

Although insects fed a nitrogen-poor diet may retain uric acid internally, those with a protein-rich diet require mechanisms for elimination of urates or uric acid. In *Rhodnius* urate is excreted at a rate of up to 1.2 mg day\(^{-1}\), more than 15% of the basic body weight per day (O’Donnell *et al.*, 1983). The tsetse fly, *Glossina morsitans* Westw., must excrete nearly 50% of the dry weight of the ingested blood in the form of nitrogen-containing compounds (mainly uric acid, but also arginine and histidine) merely to dispose of surplus nitrogen (Bursell, 1965). It is important to point out that the conclusion of active transport is based on measured concentrations of urate and in hemolymph and tubule lumen and on the transepithelial potential. However, if urate is immediately converted to uric acid and precipitated, a gradient favouring passive movement of urate into the lumen may be created.

Although uric acid is now known to be transported actively by the lower Malpighian tubule of *Rhodnius* (O’Donnell *et al.*, 1983), a ‘secretion-reabsorption’ model for passive uric acid transport in the upper Malpighian tubules of *Rhodnius* was first proposed by Wigglesworth (1931), and is reproduced in many current textbooks of entomology or comparative physiology. Wigglesworth proposed that urate moved passively across the cells and into the lumen of the upper secretory portion of the tubules, and was then carried by entrainment in the flow of fluid into the lower tubule, where precipitation of uric acid occurred as a result of acidification and water reabsorption. The model proposed that acidification resulted from reabsorption of bicarbonate. Wigglesworth’s model was subsequently modified by Miles (1966), who suggested that acidification may be a direct result from anisotropy of oxidation-reduction during electron transport in the cells of the excretory system.

A requirement of the models of both Wigglesworth and Miles is a significant flux of urate across the upper tubule. However, measurements of the rate of passive movement of urate across the upper fluid secreting segment of the tubule indicate that this segment of the tubule accounts for less than 2.3% of the uric acid eliminated by the insect. Measurements with isolated tubules demonstrate that urate is actively transported against an electrochemical gradient into the lumen of the lower Malpighian tubule (O’Donnell *et al.*, 1983). The lower 1/3 of the lower tubule is also...
responsible for reabsorption of $K^+$ and $Cl^-$ but not water. Subsequent histochemical studies by Wigglesworth confirmed the exclusive role of the lower tubule in urate transport (Wigglesworth, 1987a,b). Wigglesworth notes that he had favoured the ‘secretion/reabsorption’ hypothesis in order to accommodate the low solubility of uric acid combined with the massive excretion of it. However, his histochemical studies some 46 years after his initial study revealed that urates or uric acid pass through the basal lamina and the cytoplasm of the cells of the lower segment in high concentration (presumably in association with protein).

Urate transport has been identified in the isolated Malpighian tubules of locusts, butterfly larvae and mantids, as well as Rhodnius, suggesting that the mechanism is widespread (O’Donnell et al., 1983). Although most studies have dealt with nymphs or adult insects, uric acid deposition in the tubule lumen has been demonstrated in the embryo in Drosophila, prior to emergence of the first instar larva (Ainsworth et al., 2000). In Calliphora, urate oxidase activity in the Malpighian tubules converts uric acid to a related nitrogenous compound, allantoin, which is then excreted. Urate oxidase activity is not detectable during embryonic development of Drosophila, in first and second instar larvae, or in pupae, but is present in third instar larvae and adults. Through control by a developmental clock within the Malpighian tubules, urate oxidase activity in the tubules reappears at the time of emergence of the imago from the puparium (Friedman and Johnson, 1977). Dow and Davies (2003) have included a section on purine metabolism in Drosophila in their comprehensive review. They note that the presence of allantoinase in third instar larvae and adults implies that urea could be produced. However, the flies lack a functional urease, so production of ammonia from any urea via this route seems unlikely.

In contrast to the mechanism of urate transport in vertebrates, transport by isolated tubules is ouabain-insensitive and results in precipitation of free uric acid rather than urate salts (O’Donnell et al., 1983). In vivo, higher urate transport rates are induced in Rhodnius by increases in hemolymph urate concentration as the blood meal is digested in the days following feeding (Fig. 9). A similar induction of transport of organic anions such as para-aminohippurate after the blood meal has been reported by Maddrell and Gardiner (1975). In vitro experiments with isolated tubules of the tsetse fly suggest that induction of urate transport may be an autonomous response of the tubule cells to increases in bathing fluid urate concentration, and neurohormonal mediation of the response may not be not required (O’Donnell et al., 1983).

In addition to entrainment in fluid movement down the lower Malpighian tubule of Rhodnius, uric acid crystals are also moved as a consequence of the beating of axopods (Bradley and Satir, 1979), structures more commonly found in protozoans. The axopods arise from
the luminal cell membrane and are 0.2–0.8 μm in diameter and 10 μm or more in length. Each contains up to 46 microtubules. The axopods of the Rhodnius lower Malpighian tubule differ from those in protozoa in that they branch as they extend outward from the luminal cell surface. The beating of the axopods moves urate crystals, as observed in short lengths of lower tubules, isolated in vitro and open at both ends. Tubules isolated in this way are unaffected by fluid movements produced by peristalsis in the rectum or fluid secretion by the upper tubule. The lower tubule itself is reabsorptive rather than secretory, and time lapse photography studies by Bradley and Satir (1979) show an absence of peristalsis during movement of urate crystals in the lumen. Crystal movement has been shown to be restricted to the open lumen, near the tips of the axopods, and does not occur at the periphery of the lumen, near microvilli.

The need for movement of urate crystals by axopods is apparent when one considers the other functions of the lower tubule of Rhodnius. This segment reabsorbs K⁺ and Cl⁻, but not water. It has thus taken over the function of the hindgut in many other insects. However, the same characteristics which suit the lower tubule for KCl reabsorption, such as a narrow lumen, extensive microvilli and a high-surface area to volume ratio, create a potential for blockage by urate crystals, particularly given the low flow rates of water when the blood meal is digested 3–4 days after diuresis and the high rates of urate transport at this time. Movement of uric acid crystals down the lower tubule will thus tend to minimize the tendency for

FIG. 9 Induction of (A) urate secretion and (B) para-aminohippuric acid (PAH) in Rhodnius Malpighian tubes in response to feeding. (A) Urate secretion rates (mean ± S.E.M) for the indicated number of 5th instar tubules were measured in vitro in saline containing 0.5 mM urate. Animals were fed a blood meal on day 0. Redrawn from O’Donnell et al. (1983, Fig. 6). (B) PAH transport in Malpighian tubules isolated from 4th stage larvae of Rhodnius starved until day 10 then fed and allowed to moult to the 5th instar on day 21. Tubules were bathed in saline containing 0.05 mM tritiated PAH. Redrawn from Maddrell and Gardiner (1975, Fig. 1). Reprinted with permission from the Company of Biologists.
the tubule to become blocked. Such translocation is normally accomplished by ciliated epithelia in other animal groups. Cilia are not found in the somatic cells of arthropods, so it has been suggested that the axopods of the *Rhodnius* lower tubule may represent an example of convergent evolution (Bradley and Satir, 1979).

Although the Malpighian tubules provide a potent means for elimination of excess urate, some species retain high levels of this molecule because it may serve other purposes in the insect’s biology, in addition to its role as a nitrogenous waste. In *Rhodnius*, high levels of urate in the hemolymph acts as an antioxidant, as described below in Section 7.3. Cockroaches utilize stored urate as part of an osmoregulatory/ionoregulatory strategy. Urate levels in the fat body increase when cockroaches are dehydrated or starved. Urate appears to be stored primarily as salts of Na\(^+\), but also K\(^+\) and possibly NH\(_4\)\(^+\) when concentrations of these ions increase, during dehydration. These ions can then be released during rehydration. Urate appears to be present in fat body ‘spherules’, formed as concentric laminations and typically about 9–11 \(\mu\)m in diameter (Tucker, 1977a). Urate and K\(^+\) are mobilized from the spherules during dehydration, whereas sodium urate deposits increase at the same time (Tucker 1977a,b). Sodium urate may be present as amorphous granules as well as spherules. It is suggested that uptake of Na\(^+\) by the fat body enables the animal to maintain a fairly constant level of Na\(^+\) in the hemolymph during dehydration, without having to excrete large amounts of Na\(^+\).

Cochran (1985) has championed the idea that uric acid plays a central role in cockroach physiology. In addition to the ionoregulatory role for urates in maintenance of hemolymph Na\(^+\) levels, urate excretion plays a regulatory role in the nitrogen economy of cockroaches, with excretion in *Parcoblatta fulvescens* occurring only when excess nitrogen is available in the diet. In most cockroaches, urates are stored in the fat body when the animals are fed nitrogen-rich diets, and are mobilized when the insects are fed carbohydrate-rich diets (Mullins and Cochran, 1975). Animals fed 14C-labelled hypoxanthine incorporate the label into uric acid, and animals on low nitrogen diets release 14CO\(_2\) to the atmosphere, consistent with degradation of uric acid. Degradation appears to involve microbial symbionts in the fat body cells. Although the majority of cockroaches excrete ammonia rather than uric acid, as described below, wood cockroaches and a few other species in sub-families Blatellinae and Plecopterinae void pellets which are high in urates, and urate spherules are present in the lumen of the Malpighian tubules of these species (Lembke and Cochran, 1988). The spherules form an opaque white slurry in the tubule lumen. It is suggested that tubules of all cockroaches can transport urate, but that it is bound to hemolymph protein (Hopkins and Lofgren, 1968) in those species which do not void it. Birefringent vacuoles within the tubule cells may indicate
vesicular transport of urate, although other materials could be the source of the birefringence (Lembke and Cochran, 1988).

5.2 AMMONIA TRANSPORT IN INSECTS

5.2.1 Aquatic species

Although most terrestrial insects are uricotelic, classic work by Staddon (1955, 1959) showed that aquatic species have reverted to ammoniotelism. This finding is in keeping with the original proposal of Delaunay (1931) that synthesis of more complex molecules from ammonia by terrestrial invertebrates is essentially a detoxification mechanism necessitated by a restricted water supply. Aquatic larval stages of insects such as lacewings (Neuroptera; Staddon, 1955) and dragonflies (Staddon, 1959) excrete ammonia as the primary nitrogenous waste. Lacewing larvae excrete 90% of their total nitrogenous waste as ammonia, and secretion of ammonium by the Malpighian tubules is suggested (Staddon, 1955). The latter proposal is in contrast with measurements in locusts (see below), although it is well known that Malpighian tubules of some terrestrial insects can secrete NH$_4^+$ in place of K$^+$ (Maddrell, 1969). In dragonfly larvae there is a transient increase in ammonia excretion following a protein-rich meal, whereas there is no change in the output of uric acid (Staddon, 1959). At least 60% of the nitrogen absorbed by the larvae within 24–48 h of feeding is excreted as ammonia during the same period.

The aquatic larvae of the mosquito Ae. aegypti also excrete NH$_4^+$, and in this species an important site of excretion is the anal papilla (Donini and O’Donnell, 2005; Donini et al., 2006). The four papillae arise from an extension of the terminal segment and project into the external medium. Studies using the scanning ion-selective electrode technique have shown that the papillae take up Na$^+$, K$^+$ and Cl$^-$ from the water and that they also excrete H$^+$ and NH$_4^+$. Although the mechanism of NH$_4^+$ transport is unknown, the data are consistent with a role for diffusion trapping by protons. Earlier studies of whole larvae using radioisotopes did not show an effect of NH$_4^+$ on Na$^+$ influx, suggesting that Na$^+$/NH$_4^+$ exchange is not involved (Stobbart, 1967). NH$_4^+$ flux across the papilla in fresh water at pH 7.4 is 360 nmol cm$^{-2}$ h$^{-1}$, similar in magnitude to fluxes across the isolated locust hindgut (580 nmol cm$^{-2}$ h$^{-1}$; Thomson et al., 1988) and fluxes of 220 nmol cm$^{-2}$ h$^{-1}$ calculated by the latter authors using whole animal data of Mullins (1974) for Periplaneta americana. Possible involvement of a two ATPases in driving ion transport by the papillae is suggested by the work of Patrick et al. (2006). Immunohistochemical labelling shows that P-type Na$^+$/K$^+$-ATPase and V-type H$^+$-ATPase are localized to the basal and apical surfaces, respectively, of the anal papillae.
5.2.2 Terrestrial species

There also appear to be numerous exceptions to the generalization that terrestrial insects are primarily uricotelic. Studies of cockroaches *(Mullins and Cochran, 1972, 1973a)* and locusts *(Thomson et al., 1988)* indicate an important role for ammonia excretion. Up to 90% of the nitrogenous material in the excreta of cockroaches such as *Periplaneta* is in the form of ammonia and little or no uric acid is detectable. Amino acids are also found in the excreta, possibly as a result of microbial activities in the gut or incomplete absorption from food. Ammonia is also a major excretory product in some lepidopteran larvae as well *(Kuzhivelil and Mohamed, 1987)*, although the possible contribution of gut microbial activity was not ruled out in the latter study. Urea can also be a major excretory product in some lepidopteran larvae *(Lazar and Mohamed, 1979)*. Detailed studies of desert locusts (*S. gregaria*) have revealed excretion of ammonium at three times the rate of total urate and ammonium concentrations in the excreta as high as 400 mM *(Harrison and Phillips, 1992)*. Ammonium excretion is compatible with water conservation in this species because ammonium is present in precipitated form as ammonium urate. Harrison and Phillips *(1992)* point out that many early studies of nitrogenous excretion in insects may have underestimated the importance of ammonium excretion because pellets were either oven-dried, resulting in volatilization and loss of NH₃, and/or dilution levels during preparation of pellets for analysis were insufficient to release ammonium from precipitated form.

The majority (> 50%) of NH₄⁺ is excreted across the ileum in locusts; less than 10% of NH₄⁺ excretion is accounted for by the Malpighian tubules *(Stagg et al., 1991)*. The ileum performs several important and interlinked homeostatic functions in locusts *(Peach and Phillips, 1991)*. Amino acids such as proline are reabsorbed from the gut lumen and metabolized to produce ammonia. Cellular NH₄⁺ is then exchanged for Na⁺ in the lumen, thus driving reabsorption of Na⁺, which is in short supply in the herbivorous diet. This exchange is suggested by the finding that the rate of transport of ammonia, but not H⁺, is directly affected by Na⁺ substitution or amiloride. The ileum is also a major site of acid excretion, in the form of H⁺ trapped by NH₃ as NH₄⁺. However, ammonia transport is independent of transmembrane pH gradients and luminal proton secretion. These findings thus rule out the ‘diffusion trapping’ mechanism of ammonia excretion typical of mammalian transporting epithelia *(e.g. Knepper et al., 1984)*. Instead, it appears that ammonia crosses the apical membrane of the locust ileum via an amiloride-sensitive Na⁺/NH₄⁺ exchange mechanism *(Thomson et al., 1988)*. Whereas proline is absorbed from either the lumen or the hemolymph in the ileum, it must be supplied to the luminal side in the rectum. In the latter tissue, proline transport exceeds the metabolic requirements of chloride transport more
than 10-fold, and appears to function not just as an energy source but also as an important osmolyte in fluid reabsorption (Chamberlin and Phillips, 1982), as described in Section 5.3. Proline also functions as an important osmolyte in mosquito larvae (Section 5.3).

Larvae of blowflies and flesh flies are commonly faced with high concentrations of ammonia as they burrow into and feed upon rotting flesh or living tissue. Larvae of the flesh fly Sarcophaga bullata ingest ammonia, reabsorb it from the midgut, and then excrete it across the hindgut epithelium. In addition, they are exposed to excess ammonia resulting from protein metabolism. Deamination of amino acids and active secretion of \( \text{NH}_4^+ \) have been studied in isolated hindgut preparations of blowflies by Prusch (1972, 1976). Ammonia can be secreted against a pH gradient (lumen alkaline) which would not be favourable for \( \text{NH}_3 \) diffusion. Both \( \text{K}^+ \) and \( \text{Cl}^- \) are actively secreted into the lumen, and the mechanism of \( \text{NH}_4^+ \) secretion is independent of that for \( \text{K}^+ \). The mechanism of \( \text{NH}_4^+ \) secretion differs from the process of \( \text{Na}^+/\text{NH}_4^+ \) exchange in the locust ileum. \( \text{Na}^+ \) is passively distributed between the hindgut lumen and hemolymph, accumulating in the lumen in response to the lumen negative transepithelial potential of about \(-15\) mV. \( \text{NH}_4^+ \), \( \text{K}^+ \) and \( \text{Na}^+ \) can be concentrated in the lumen simultaneously, so exchange of \( \text{NH}_4^+ \) with \( \text{K}^+ \) or \( \text{Na}^+ \) seems unlikely. However, ammonium secretion is reduced by about 60% when either \( \text{K}^+ \) or \( \text{Na}^+ \) is removed, suggesting that both alkali cations are implicated in some way in \( \text{NH}_4^+ \) secretion.

The larvae of sheep blowflies are also exposed to high concentrations of ammonia. The larvae live in urine-soaked wool, proteinaceous sera and blood from the lesions produced by the larvae themselves. The larvae are exposed to excess nitrogen resulting not only from metabolism of proteins, but from the urine soaking the wool around the lesions. The larval environment fluctuates widely in ionic and osmotic composition, since sheep urine varies from 800 to 2500 mOsm kg\(^{-1}\) and osmotic pressures of fluids in the lesions will rise above the plasma value of 290 mOsm kg\(^{-1}\) as wounds dry out. Studies of ionic and osmotic regulation by larvae of the sheep blowfly Lucilia cuprina show that \( \text{NH}_4^+ \) concentrations in the hemolymph are not altered significantly from the very high control value (16 mM) when the concentration of \( \text{NH}_4^+ \) in the rearing medium is varied between 3 and 560 mM (Marshall and Wood, 1990). Hemolymph \( \text{Na}^+ \) concentration is also well regulated, increasing less than 20% above the control value of 115 mM when the \( \text{Na}^+ \) concentration of rearing medium is increased from 15 to 510 mM. Potassium levels in the hemolymph vary from 13 to 36 mM when rearing medium \( \text{K}^+ \) concentration is increased from 31 to 460 mM. Marshall and Wood (1990) also report the somewhat surprising finding that the optimal pH for growth of Lucilia larvae is 8–9, even though much of the ammonia in the medium is then present in the non-ionic and presumably, more permeant form (\( \text{NH}_3 \)). The hindgut has
been proposed as the site of NH$_4^+$ excretion (Marshall and Wood, 1990), based on early reports of high levels of NH$_4^+$ in the hindgut tissues of L. cuprina larvae and also on the demonstration of active secretion of NH$_4^+$ by larvae of Sa. bullata (Prusch, 1974, 1976).

Another group of insects which are faced with excess ammonia are adult mosquitoes. More than 60% of the amino acids derived from the digestion of blood meal proteins by Ae. aegypti females is oxidized to CO$_2$ to provide energy and only 3% is used for egg protein production. Mosquitoes tolerate potentially toxic loads of ammonia resulting from amino acid catabolism by high rates of ammonia excretion in the feces coupled with the synthesis of glutamine and proline (Goldstrohm et al., 2003; Scaraffia et al., 2005). Ammonia subsequently recovered from proline can be excreted and the carbon skeleton used for synthesis of different compounds or for energy production. Ammonia excretion from mosquitoes fed 80 mM NH$_4$Cl in 3% sucrose increases markedly for the first day and reaches a peak of $\sim$ 120 nmol animal$^{-1}$ by 2 days, approximately sixfold greater than the level of uric acid excretion. Hemolymph glutamine and proline concentrations increase markedly within an hour when females are fed solutions containing NH$_4$Cl or a blood meal, whereas concentrations of aspartate, asparagine, glutamate and alanine do not change after NH$_4$Cl feeding. Glutamine synthetase is an important enzyme in the process of ammonia storage, since addition of an inhibitor of the enzyme added to the meal reduces hemolymph glutamine concentration and causes a corresponding increase in proline concentration. Glutamate synthase contributes to the production of glutamate for proline synthesis since azaserine, an inhibitor of glutamate synthase, results in an increase in glutamine concentration and a corresponding decrease in proline concentration after feeding. The mRNA expression and activity of several enzymes implicated in ammonia metabolism increases in the fat body after a blood meal, suggesting that it is the main tissue involved in ammonia detoxification.

5.2.3 Uptake of ammonia

Ammonia transport has also been studied in the caterpillar Ma. sexta (Weihrauch, 2006). All three sections of Ma. sexta midgut (anterior, median, posterior) actively absorb from the gut lumen toward the hemolymph (Weihrauch, 2006). Larvae feed on a diet which is relatively low in protein, suggesting that active ammonia absorption might be necessary for larval protein anabolism. Active ammonia uptake, which is de facto a net acid absorption, may also enhance midgut alkalinization. A model of ammonia uptake across the midgut epithelium suggests that NH$_4^+$ may enter the columnar cells via apical H$^+$/cation exchange, possibly by an amiloride-sensitive member of the NHE family. In addition, ammonia may enter as NH$_3$ across the plasma membrane, exploiting the high pH (10–11) of the
lumen contents relative to that of the cells (\(\sim\) pH 7). Dissociation of cytosolic \(\text{NH}_4^+\) to \(\text{H}^+\) and \(\text{NH}_3\) is accompanied by diffusion of \(\text{NH}_3\) into vesicles acidified by a bafilomycin-sensitive V-type \(\text{H}^+\)-ATPase. It is further proposed that the ammonia-loaded vesicles then move via colchicine-sensitive microtubules to the basal membrane, where they fuse with the membrane, releasing \(\text{NH}_4^+\) into the hemolymph space (Weihrauch, 2006). In addition, a small amount of ammonia may exit from cell to hemolymph through \(\text{Ba}^{2+}\)-sensitive \(\text{K}^+\) channels.

One consequence of ammonia uptake by the gut is that hemolymph ammonia concentration of the fifth instar larvae of *Ma. sexta* is 0.87 mM, more than twice as high as blood concentrations found in the human portal vein and about eight times as high as ammonia concentration levels in body fluids of aquatic crustaceans. Caterpillars may tolerate high ammonia levels in the hemolymph in part because high levels of a Rhesus-like ammonia transporter in the ganglia may protect neuronal tissues from ammonia toxicity. In addition, excess hemolymph ammonia may be secreted by the Malpighian tubules. Presumably ammonia is then released for excretion into the hindgut where, in contrast to midgut sections containing \(\sim\) 1 mM ammonia, very high ammonia concentrations (ca. 60 mM) are found. High levels of ammonia in the lumen of the hindgut suggest that ammonia is a major nitrogenous excretion product in *Manduca* larvae. In contrast to the midgut, the hindgut of the larvae exhibits no transepithelial active net transport, and high concentrations of ammonia in the hindgut lumen reflect entry via the Malpighian tubules and release of metabolic ammonia across the apical membrane.

Evidence for a Rhesus-like ammonia transporter (RhMS) was found with high expression levels in the hindgut, Malpighian tubules and ganglia of *Manduca* larvae but low RNA expression in midgut tissues. A full length cDNA (RhMS) has been transcribed from midgut mRNA and translation of the open reading frame indicates a 425 amino acid protein with 10 predicted transmembrane domains and more than 40% identity in amino acid sequence to known mammalian Rhesus-like ammonia transporters (RhAG, RhBG and RhCG). Low levels of mRNA expression in all midgut sections, and the fat body suggest a minor, probably regulatory role for RhMS in ‘ammonia management’ in these tissues (Weihrauch, 2006).

### 5.3 FREE AMINO ACIDS

The blood volume of insects varies greatly during changes in hydration state or in association with food intake, suggesting that the hemolymph may act as a reservoir for water. This capacity is probably a consequence of respiratory gas exchange via the tracheal system, so that changes in hemolymph volume do not impede transfer of oxygen to the tissues. In spite of changes in hemolymph volume, levels of amino acids in the
hemolymph remain elevated. Whereas vertebrates have about 0.05 g% free amino acids in the blood, the amount ranges from 0.29–2.43 g% in the hemolymph of insects (Wigglesworth, 1972).

One role for hemolymph amino acids is to act as metabolic substrates for energy consuming tissues such as the flight muscles and the hindgut. Transport of amino acids has been examined in detail in the desert locust Schistocerca americana. Hemolymph proline and glutamate are actively secreted by the Malpighian tubules, and are subsequently reabsorbed in the rectum. The primary function of this translocation of amino acids appears to be the provision of metabolic substrates for the highly active rectal epithelial cells. However, only a small fraction of proline, alanine, serine or threonine that are actively transported across the epithelium are subsequently metabolized. Glutamate, though metabolized, is not transported across the epithelium. Surprisingly, the amount of proline transported across the rectum exceeds by a factor of 10 the amount necessary for metabolism of the rectal tissue. This extra capacity is associated with rectal water transport (Phillips et al., 1986). In other words, proline may function as an osmolyte driving water reabsorption from the rectal lumen against the large osmotic differences which develop as the rectal contents are dehydrated. Neutral amino acids are also reabsorbed, probably by a common carrier system because addition of one substrate such as L-serine competitively inhibits uptake of a second, such as glycine (Balshin and Phillips, 1971).

In larvae of osmoconforming mosquitoes, proline functions as both an extracellular and intracellular osmolyte. Larvae of Culex tarsalis increase the hemolymph levels of both proline and the disaccharide trehalose in response to increases in environmental salinity (Patrick and Bradley, 2000a,b). In 64% sea water, hemolymph proline and trehalose are accumulated approximately 50-fold and twofold, respectively, relative to freshwater values. Proline accumulation is cued by increases in salinity, and the proximal cue for an increase in proline synthesis appears to be a rise in hemolymph Na\(^+\). In contrast, trehalose increases in response to changes in osmolality produced either by addition of either electrolytes (sea salt) non-electrolytes such as sorbitol to the rearing medium (Patrick and Bradley, 2000b).

The amino acid tryptophan is of special significance in insects because of its role as a precursor of eye pigments. Excretion of tryptophan metabolites is a consequence of the ommochrome pathway used for synthesis of eye pigments. Eye-color mutants and their corresponding genes can be used as scorable marker systems to follow the results of genetic crossing experiments in Drosophila and other insects. The initial step in ommochrome synthesis involves the conversion of tryptophan to N-formylkynurenine by tryptophan oxygenase. In the lepidopteran, Plodia interpunctella, silencing of tryptophan oxygenase through the use of RNA interference (RNAi) during embryonic development results in loss of eye-color pigmentation.
(Fabrick et al., 2004). In the stick insect, end products of tryptophan metabolism are the ommochromes ommin and xanthommatin, which accumulate in the epidermis, and kynurenic acid, which accumulates in the faeces (Stratakis, 1979). Excretion of tryptophan metabolites accounts for up to 2% of nitrogen excretion in cockroaches, and probably represents a means of dealing with excess dietary tryptophan rather than an important component of nitrogen excretion (Mullins and Cochran, 1973a).

The most detailed studies of the role of the Malpighian tubules in excretion of eye pigments have been done on Drosophila. Tearle (1991) has examined tissue-specific effects of ommochrome pathway mutations in Drosophila, including those that alter excretion of pigment by the tubules. Dow and Davies (2003) note that the first Drosophila mutation described (white) was for eye color and that now over a hundred distinct loci are known that produce an eye-color phenotype. The white gene encodes an ABC transporter found in the eye and in the Malpighian tubule. Many of the eye color mutants have Malpighian tubule color phenotypes, possibly reflecting an original excretory function of the pathways that generate and transport the major classes of metabolite (ommochromes and pteridines). These pathways may have subsequently taken on specialized roles in pigment granule formation (Dow and Davies, 2003).

5.3.1 Phloem-feeders

Phloem-feeders such as silverleaf whiteflies (Bemisia tabaci Grennadius) ingest a diet that is rich in soluble carbohydrates and also contains relatively high levels of free amino acids (Buchanan et al., 2000). Although plant phloem sap is generally considered to be limiting in nitrogen (Byrne and Miller, 1990), whiteflies and other phloem-feeding insects such as aphids excrete amino acids, especially non-essential amino acids such as glutamine, asparagine, glutamate and aspartate (Douglas, 1993). These four amino acids comprise the majority of the free amino acids commonly found in phloem sap (Douglas, 1993) of cotton and plants in general (Buchanan et al., 2000).

The amount of amino nitrogen excreted per day exceeds by nearly twofold the total amino acid pool in the body of whiteflies reared on well-fertilized cotton plants. Phloem sap of these plants thus contains supra-optimal amounts of nitrogen with regard to whitefly nutrition. However, after just a few days of feeding on low nitrogen plants, excretion of amino nitrogen essentially stops and there is an adjustment of pools of individual amino acids within the insect bodies (Crafts-Brandner, 2002). Total nitrogen content of whiteflies reared on plants with reduced nitrogen content does not decline significantly relative to whiteflies reared on plants with higher levels of nitrogen. However, the free amino acid content of whiteflies feeding on low-nitrogen plants declines >90% relative to controls. The predominant
amino acids in whiteflies are glutamine (26% of the total), alanine (19%), proline (13%) and glutamate (10%), whereas the predominant amino acids in phloem sap from high nitrogen cotton plants are aspartate, glutamate and arginine, with relatively large amounts of glutamine and asparagine. The predominant free amino acids in the honeydew are asparagine (46% of total amino acid content) and glutamine (12%).

Given that the amino acid pools in whiteflies feeding on high nitrogen cotton plants are not closely related to the amino acid composition of the phloem sap, dietary amino nitrogen must be rapidly assimilated into metabolites or protein. Synthesis of at least some of the essential amino acids in aphids is dependent on endosymbiotic bacteria (Sasaki and Ishikawa, 1995; Wilkinson and Douglas, 1996; Douglas, 1998). High levels of free proline and alanine in whiteflies indicate an important role for proline metabolism in energy balance in the whitefly. The hemolymph and flight muscle tissue of several insect species contains high levels of proline that is used for fuel, via oxidation to alanine, for flight (Gäde, 1992).

It also appears that glutamine metabolism plays an important role when whiteflies adjust to changes in dietary nitrogen levels. Presumably a large and alterable pool of glutamine permits maintenance of metabolite synthesis via transamination reactions and/or formation of tricarboxylic acid cycle intermediates. Changes in the glutamine pool may be necessary to maintain levels of proline and alanine that are likely to be important metabolic fuels during flight (Crafts-Brandner, 2002).

6 Transport and excretion of divalent ions and bicarbonate

6.1 Calcium

Vertebrates and insects appear to use fundamentally different mechanisms to regulate the level of Ca$^{2+}$ in their extracellular fluids. Whereas calcium homeostasis in mammals relies upon precise control of transcellular calcium absorption by the duodenum, control of excretion and not absorption is the major means of hemolymph calcium regulation in flies (Calliphora vicina; Taylor, 1985). When fed 12.5 mM CaCl$_2$ solution in place of water the calcium content of Cas vicina rapidly increases to more than double the original content but then remains stable. However, midguts isolated from flies fed either 12.5 mM CaCl$_2$ solution or water absorb calcium at the same rate, and this rate is unaltered by treatment with the putative transport stimulants dopamine, octopamine, proctolin or 5-hydroxytryptamine. Rapid calcium absorption from the blowfly midgut appears to be an adaptation not for regulation of hemolymph calcium concentration but for removal of calcium from the lumen. This removal may facilitate other midgut functions such as phosphate absorption
that would be impeded by high Ca\(^{2+}\) levels in the gut lumen as the gut contents are concentrated by water absorption. The calcium content of the Malpighian tubules is higher than that of other tissues, and rates of turnover of \(^{45}\)Ca when flies are fed excess calcium are highest in the Malpighian tubules, indicating the tubules as the means by which hemolymph calcium concentration is regulated (Taylor, 1985).

Subsequent studies of fruit flies have shown that calcium homeostasis is accomplished through the combined effects of elimination of calcium in fluid secreted by the Malpighian tubules and the sequestration of calcium in granules, especially within the distal segment of the anterior pair of Malpighian tubules (Dube et al., 2000a,b). Adult flies ingest 30 times more calcium than they retain, yet they regulate hemolymph calcium concentration and whole animal calcium content quite precisely. In response to a 6.2-fold increase in dietary calcium level, for example, calcium content of whole flies increases only 10%. Hemolymph calcium concentration (~0.5 mM) is similar in flies raised on diets differing more than sixfold in calcium content (Dube et al., 2000a).

Studies with isolated tubules of \(D.\) melanogaster indicate that all segments of both the anterior and posterior Malpighian tubules transport Ca\(^{2+}\) at high rates and are thus capable of regulating whole animal and hemolymph calcium levels (Dube et al., 2000b). Approximately 85% of calcium which enters the tubule is sequestered, and ~15% is secreted in soluble form into the tubule lumen. Tubules secreting fluid at maximal rates can remove an amount of calcium equal to the whole animal calcium content in ~ 9 h. The distal segments of the anterior pair of Malpighian tubules can sequester the same amount of calcium in <2 h. The processes of secretion and sequestration are controlled independently. For example, although cAMP increases both basolateral uptake and transepithelial secretion of Ca\(^{2+}\), the calcium channel blockers diltiazem and verapamil inhibit only basolateral uptake. Increases in basolateral membrane potential in response to reduction of bathing saline K\(^+\) concentration enhance uptake of Ca\(^{2+}\) across the basolateral membrane, further supporting a role for Ca\(^{2+}\) uptake through channels, whereas transport from cell to lumen requires some form of Ca\(^{2+}\) pump, either an-ATPase or an electrogenic exchanger (e.g. 3Na\(^+\)::Ca\(^{2+}\)). Ca\(^{2+}\) sequestered within the tubule is released into the lumen even when basolateral uptake is blocked. Transepithelial Ca\(^{2+}\) secretion is increased by treatments which depolarize the transepithelial potential by increasing Cl\(^-\) conductance, or which acidify the secreted fluids (bicarbonate-free saline). The latter treatment may favour dissolution of intracellular granules through changes in intracellular pH.

Mineralized granules enriched in calcium, magnesium and phosphorous are also found in the lumen of the Malpighian tubules of larvae of the face fly \(M.\) autumnalis (Krueger et al., 1987). These mineral stores are
subsequently mobilized and used to harden the cuticle of the puparium (Krueger et al., 1988). The spherical granules range from 0.2 to 10 μm in diameter and are composed of many concentric layers (Grodowitz et al., 1987). Layers appear to be covered by an organic-rich material which may bind one layer to the next. Calcium, phosphorous and magnesium make up 98% of the inorganic content, which is about 85% of granule weight.

The decrease in pH from 8.08 in the lumen of the distal region of the larval tubules to 7.35 in the proximal region appears to be a major effector of granule dissolution in *Mu. autumnalis*. Release of minerals from the granules increases exponentially as the bathing medium pH decreases. The calcium and magnesium salts which are released are transported from the proximal tubule lumen to the hemolymph, and this transport may sustain or augment the rate of granule dissolution. Calcium is transported directly from tubules to the cuticle of the puparium via the hemolymph. Hemolymph calcium levels then decline as calcium is deposited in the puparial cuticle. Granule deposition, in common with granule dissolution, involves a change in pH. Cuticular pH measured with flat-surface pH electrodes increases from 7 to 8.4. This shift favours precipitation of calcium and magnesium phosphates and carbonates, which have low solubility products at alkaline pH.

Calcium is also mobilized from intracellular CaPO4 concretions (‘spherites’) in Malpighian tubules of the house cricket *Ac. domesticus* during the early phases of diuresis (Hazelton et al., 2001). When tubules are treated with the diuretic hormone second messenger cAMP, there are dramatic changes in tubule ultrastructure within 15–420 s. The cytoplasm becomes extensively vacuolated and the mitochondria move into areas adjacent the transport membranes. In addition, the CaPO4 spherites become associated with both mitochondria and vesicles during this period. Although some of the spherites dissolve, many appear to be exocytosed intact into the tubule lumen (Spring and Felgenhauer, 1996). Presumably, release of spherites during diuresis in effect flushes them into the hindgut. In this way, the tubule lumen does not become obstructed, as might occur if the spherites were released into the lumen during periods of slow fluid secretion.

Blood-feeders such as *Rhodnius* are faced with a considerable excess of dietary calcium when they ingest up to 10 times their own body weight in blood, which may contain 1.5 mM calcium in the case of avian blood. Calcium accumulation by Malpighian tubule cells is correlated with the appearance of membrane bound spherical concretion bodies in the upper Malpighian tubule cells (Maddrell et al., 1991). The net result of calcium deposition in the tubule is that almost none is excreted from the body. By contrast, Ca distribution between hemolymph and Malpighian tubule lumen is passive in the ant *Formica*, and the concentration of calcium in the luminal fluid is 0.9 mM (Van Kerkhove et al., 1989). Calcium sequestration
in *Rhodnius* tubules may be necessary to prevent elevation of calcium concentrations in the lower Malpighian tubule or hindgut during water reabsorption. High concentrations of metal ions might interfere with rectal ion and water transport. Alternatively, calcium deposits might subsequently be mobilized for production of eggs or spermataphores after ecdysis of adults.

The concretion bodies within the tubule cells of *Rhodnius* appear as concentric layered structures in electron micrographs, suggesting that they are deposited in distinct phases, perhaps after each meal. Comparison of X-ray diffraction patterns from tubular hydroxyapatite and dried tubules indicate that the concretions are amorphous, in form, not crystalline. Concentrations of Ca in the tubules exceed 21 mM, 400 mM and 1 mM when the insects are fed on low-K⁺ sheep blood, rabbits and chickens, respectively. For all three diets, the hemolymph calcium concentration is 8 mM, and its activity, as determined by an ion-selective electrode, is about 2.5 mM. ⁴⁵Ca recently incorporated in the previous 48 h into the tubules is released at the rate of about 25% per hour when the tubules are transferred to ⁴⁵Ca-free saline, indicating that the process is a dynamic one. This rate declines to 13% per hour for animals allowed to incorporate ⁴⁵Ca for 9 days following injection of the isotope into the hemolymph. Calcium sequestration within the bodies is a continuous and metabolically dependent process, but calcium becomes less accessible to the hemolymph over time, presumably since the ‘oldest’ calcium is near the center of the concretions. The post-prandial increase in calcium uptake by the tubules *(Maddrell et al., 1991)* is similar to that seen for para-aminohippuric acid and uric acid *(Maddrell and Gardiner, 1975; O'Donnell et al., 1983)*. For the latter compounds it has been suggested that an inducible transport system, stimulated by some correlate of feeding, is involved. In each case, uptake rates peak 2–4 days after a blood meal and remain elevated for about 2 weeks.

### 6.2 Magnesium

Larvae of the salt marsh mosquito *Aedes campestris* are able to grow, moult and pupate in hyperosmotic lakes found in central British Columbia. These lakes are unusual in that they contain high levels of magnesium and that the anionic component consists almost entirely of sulfate ions *(Scudder, 1969)*. These mosquitoes cope with osmotic water loss by drinking and absorbing the water in which they live and then excreting the excess salts *(Kiceniuk and Phillips, 1974)*, an osmotic strategy analogous to that of marine fishes. The larvae absorb into the hemolymph virtually all the water that they ingest and thus face excess loads of sodium or magnesium and considerably larger loads of sulfate *(Kiceniuk and Phillips, 1974)*. Malpighian tubules of the larvae secrete fluid containing 30–40 mM magnesium when bathed in
solutions containing 4–20 mM Mg$^{2+}$ (Phillips and Maddrell, 1974). Magnesium is secreted against an opposing transepithelial potential difference of 15 mV, and against concentration gradients as high as 10-fold. Transport is saturable, and the Michaelis–Menten constant for transport ($K_t$) is about 2.5 mM. Secretion of Mg$^{2+}$ by the tubules permits regulation of hemolymph concentrations of 1.5–4 mM for *Ae. campestris* larvae living in and ingesting large amounts of media containing as much as 100 mM magnesium.

The highest known rates of epithelial magnesium secretion have been observed in isolated tubules of black field cricket *Te. oceanicus* (Xu and Marshall, 1999a). The small distal segment of the Malpighian tubules of the *Te. oceanicus* and the house cricket *Ac. domesticus* are characterized by very high rates of fluid secretion. Fluid secreted by the distal segment in *Te. oceanicus* is hyperosmotic and contains primarily Mg$^{2+}$ (125 mM), Cl$^-$ (242 mM) and Na$^+$ (43 mM). The concentration of magnesium in the fluid is 15 times greater than that of the bathing fluid. Surprisingly, the total volume of fluid secreted per unit time by the distal segment is constant, possibly because short distal segments are of larger diameter than long distal segments. In contrast, as the main segment increases in length the volume of fluid secreted in unit time also increases. The rate of magnesium secretion by the distal segment, 75 pmol min$^{-1}$ mm$^{-1}$ tubule length, is the highest known for any epithelium and is 6 times higher than the average rate of Mg$^{2+}$ secretion by the Malpighian tubules of *Ae. campestris* (Phillips and Maddrell, 1974) and 47 times higher than the rate measured for flounder renal tubules (Beyenbach, 1990).

High rates of magnesium secretion by the distal segments of cricket tubules may play a homeostatic role by eliminating excess magnesium present in the diet. Feeding on lettuce leads to ingestion of as much as 1790 nmol of magnesium per day. All of the tubules can together secrete Mg$^{2+}$ at the rate of 417 nmol h$^{-1}$, so the entire daily intake can be eliminated from the hemolymph, which contains about 431 nmol of Mg$^{2+}$, in about 5 h$^{-1}$. Some of the magnesium secreted by the distal segment appears to be reabsorbed by the main segment, raising the possibility of an interaction of the two segments in regulating hemolymph and whole animal levels of magnesium.

6.3 SULFATE

As noted above, high concentrations of sulfate are also present in the magnesium-rich lakes inhabited by larvae of the mosquito *Ae. campestris*. However, the larvae maintain sulfate concentration in the hemolymph well below that of the water. When the external concentration of sulfate is increased 30-fold from 2.5 to 73 mM, the hemolymph concentration increases only fourfold, from 1.5 to 6.6 mM. Experiments using
radiolabelled sulfate have shown that the regulation of sulfate is not accomplished by precipitating the anion in the tissues. However, isolated Malpighian tubules set up in Ramsay fluid secretion assays actively transport sulfate from bath to lumen against a threefold concentration gradient. The $K_t$ for transport is 10 mM and the maximum transport rate is 50 pmol min$^{-1}$ tubule (Maddrell and Phillips, 1975a).

It also appears that transport of sulfate by the Malpighian tubules occurs against an opposing electrical gradient of as much as 20 mV. The latter conclusion requires confirmation because the method used for measurement of the transepithelial potential has been shown in subsequent studies to be a possible source of artefacts. Although the transepithelial potential across the large diameter tubules of $R.\ prolixus$ can be measured by simply placing one electrode in contact with the secreted fluid droplet and the reference electrode in the saline bathing the outer surface of an isolated tubule set up in a Ramsay assay (Ianowski and O’Donnell, 2001), this method is prone to errors in tubules whose diameter is less than ~50 \mu m. The error results partly from shunting of current along the surface of the tubule between the bathing droplet and the secreted fluid droplet (Aneshansley et al., 1988). As a result, the measurements of transepithelial potential reported in Maddrell and Phillips (1975a) may be incorrect. However, by varying the level of potassium in the bathing saline, Maddrell and Phillips (1975a) demonstrated that there is no strong relationship between the resulting changes in transepithelial potential and the secreted fluid to bath concentration ratio for sulfate. These data indicate that sulfate ions are not transported into the secreted fluid merely as a response to an electrical potential gradient favouring their entry.

Calculations based on the rate of sulfate transport by isolated tubules indicate that secretion by the Malpighian tubules is sufficient to remove all of the sulfate ingested by larvae living in waters containing less than 100 mM sulfate. At higher concentrations, sulfate ions are probably also excreted elsewhere, perhaps by the rectum or anal papillae. Larvae of another mosquito, $Aedes taeniorhynchus$, which can also live under hyperosmotic conditions, can secrete into the rectal lumen a markedly hyperosmotic fluid (Bradley, 1987). A similar mechanism in $Ae.\ campestris$ may contribute an increasing proportion of sulfate excretion as the osmotic concentration of the external medium rises. The rectum of $Ae.\ campestris$ has an extra posterior region which is lacking in its freshwater relative $Ae.\ aegypti$ (Meredith and Phillips, 1973). Given the important role of the posterior rectum in secretion of hyperosmotic fluids by larvae of $Ae.\ taeniorhynchus$, it is quite plausible that this segment of the rectum excretes sulfate ions under hyperosmotic conditions in $Ae.\ campestris$.

Larvae of $Ae.\ taeniorhynchus$ are also exposed to high levels of sulfate in salt water pools. They absorb most of the water and nearly all of the sulfate (and probably the other ions as well) from the fluid that they ingest.
Much of the water ingested by unfed larvae is to replace that lost by osmosis to the hyperosmotic medium. Sulfate absorption by the midgut does not proceed against even a small concentration difference. Since most epithelia are relatively impermeable to ions as large a sulfate, these findings indicate that its uptake across the midgut epithelium probably involves some form of facilitated diffusion.

The sulfate transport system in tubules from *Ae. taeniorhynchus*, as in tubules of *Ae. campestris*, is characterized by high capacity and low affinity, transport being half saturated in the range 5–10 mM. However, tubules from insects reared in sulfate-free water have only a weak ability to transport sulfate ions; indeed the fluid they secrete is scarcely if ever found to contain sulfate ions at a higher concentration than in the bathing medium. In contrast, Malpighian tubules from *Ae. taeniorhynchus* reared in sulfate-containing waters develop an ability to transport sulfate at a rate which varies directly with the sulfate content of the environment (Maddrell and Phillips, 1978). Malpighian tubules from larvae reared in sulfate-enriched sea water containing 89 mM sulfate transport sulfate ions at rates twice those from insects reared in normal sea water and the fluid secreted contains sulfate ions at concentrations several times higher than in the bathing medium. For example, tubules bathed in fluid containing 3.3 mM sulfate secreted fluid containing an average of 18.5 mM sulfate and tubules bathed in fluid containing 30 mM sulfate secreted fluid containing more than 130 mM sulfate. The transport ability is not retained in adults which developed from larvae reared in sulfate-rich water. The induction of transport appears to involve a change in capacity with no change in affinity and the half-time for induction of this transport ability is approximately 16 h. The sulfate-transporting ability induced in the Malpighian tubules is more than sufficient to match the rate at which sulfate ions are taken up across the gut. Induction may involve one or more of the following mechanisms: synthesis of more pump molecules, the incorporation of more pumps into the cell membranes, or the activation of inactive pumps already in the membranes (Maddrell and Phillips, 1978).

The rate of sulfate transport by the isolated Malpighian tubules depends on the rate of fluid secretion. Fluid secretion rates by the Malpighian tubules can be increased up to sixfold above the basal level by addition of extracts of the brain and of the thoracic ganglia. Sulfate ions are removed more slowly from the bathing medium at the lower rates of fluid secretion. This relationship is very similar to that seen in the transport of organic anions and cations by Malpighian tubules of other insects (Maddrell *et al*., 1974; Maddrell and Gardiner, 1976). The passive permeability of the tubule wall reduces the effectiveness of transport at the lower rates of fluid secretion through passive diffusive backflux of the transported molecules from the lumen to the bath, as described in Section 8.3.
Malpighian tubules of the blowfly *Calliphora erythrocephala* also secrete fluid containing levels of sulfate above those in the bathing medium over the range of bathing medium sulfate concentrations from 1 to 50 mM (Knowles, 1975b). Earlier studies (cited by Knowles, 1975b) show that tubules *Calliphora* as well as those of the hemipteran *Rhodnius* and the stick insect *Ca. morosus* all cease fluid secretion when sulfate is the only anion present in the bathing fluids. However, lower concentrations of sulfate do not suppress fluid secretion by tubules of *Calliphora* or *Carausius*. For *Calliphora* tubules, the highest ratio of sulfate in the secreted fluid to that in the bathing medium (SF/M) is 4.8 and is found when the bath contains 6 mM sulfate. Intact flies also excrete sulfate in the urine.

The mechanism of sulfate transport by *Calliphora* tubules is unclear. The measured transepithelial potential is not sufficiently positive to account for the observed concentrations of sulfate in all experiments, and the 40-fold increase in sulfate transport between 0.5 and 5 mM sulfate in the bathing saline is not associated with a sufficiently large change in transepithelial potential to explain the increase. However, the method of transepithelial measurement used in this and other early studies has subsequently been shown to result in erroneous values in small diameter tubules such as those of blowflies and mosquitoes, as discussed above.

The function of sulfate transport in insects not normally exposed to high levels of this anion is unknown. However it has been pointed out by Knowles (1975b) that sulfate secretion may be linked to the metabolism of hormones. Malpighian tubules of *Ca. erythrocephala* conjugate β-ecdysone with sulfate and metabolites of insect juvenile hormone can combine with sulfate to form conjugates which are then excreted.

### 6.4 BICARBONATE

#### 6.4.1 Transport of bicarbonate by Malpighian tubules

Bicarbonate transport by insect epithelia has been examined primarily in species adapted to alkaline lakes. For the freshwater species of water boatmen, *Cenocorixa blaisdelli* (Corixidae, Hemiptera) it appears that extrarenal pathways contribute to hemolymph ion and CO₂ regulation when the animals are acclimated to high pH (NaHCO₃, Na₂CO₃) water typical of athalassohaline lakes (i.e. those not of marine origin but from evaporation of fresh water) found in western Canada (Cooper et al., 1987). Although hemolymph ion concentrations in alkaline-acclimated insects do not differ from those of insects reared in fresh water, urinary loss of ions and CO₂ cannot fully compensate for increased intake of sodium and CO₂. Chloride cells are present on the bodies (Komnick and Schmitz, 1977) of corixids and on the labium of the alkaline lake species *Cenocorixa bifida* (Jarial et al., 1969), raising the possibility that increases in hemolymph
chloride concentration under alkaline conditions reflect extrusion of basic equivalents by Cl⁻/HCO₃⁻ exchange (Cooper et al., 1987). Regulation of hemolymph Na⁺, however, is more problematic. The possibility of extrarenal Na⁺/H⁺ exchange seems unlikely since, the elevated pH and high sodium content of the surrounding water would make Na⁺ uptake more likely than Na⁺ loss. It is worth noting that changes in osmotic water influx may also be important in ion-regulation by corixids. The water permeability of Ce. bifida increases when ambient salinity increases, so corixids may be able to obtain some water without an ion load if the cuticle acts as an ion filter during osmotic influx of water (Cannings, 1981).

A role for the Malpighian tubules in secretion of bicarbonate has been suggested by studies of fresh water and alkaline lake species of corixids. Four distinct segments are present in Malpighian tubules of these species. Segment 1 is connected to the hindgut and appears to act as a passive conduit. In response to cAMP, segment II increases the pH of the luminal fluid, from 7.0 to 8.1 in the fresh water species Ce. blaisdelli (Cooper et al., 1989), and from 7.0 to 8.6 in the alkaline lake species Ce. bifida (Szibbo and Scudder, 1979). The alkalinity increase is correlated with increased total CO₂ content of luminal fluid, suggesting that bicarbonate is secreted by segment II. The carbonic anhydrase inhibitor acetazolamide blocks the pH change, and does so in the absence of exogenous bicarbonate, suggesting that bicarbonate in the lumen is derived from metabolically produced CO₂. cAMP may activate carbonic anhydrase, therefore, or a bicarbonate conductance. The positive transepithelial potential would favour anion movements into the lumen, although a cell-to-lumen concentration gradient of bicarbonate is presumably required as well to explain an elevation of pH by one unit or more.

Transport and storage of bicarbonate and carbonate play important roles in the biology of shore flies (Ephydridae, Diptera). These insects are well known for their tolerance of extremes in temperature, salinity and pH which occur in the larval habitats such as hot springs, hypersaline desert lakes and tidal splash pools. Ephydra hyans inhabits alkaline salt lakes in western North America, and is capable of hyperosmotic and hypooosmotic regulation of hemolymph osmolality (at around 300 mOsm kg⁻¹) in alkaline lake water, but less capable of regulation in non-alkaline sea water/sodium chloride solutions (Herbst et al., 1988). Although hemolymph osmolytes consist primarily of sodium and chloride, carbonate and bicarbonate are major components of the species habitat. The Malpighian tubules on one side of the gut appear to excrete and store an insoluble salt of CO₃²⁻ and HCO₃⁻. Although the morphology of the anterior pair of Malpighian tubules is typical of most insects, the posterior pair of tubules is modified into an enlarged gland containing spherical concretions of nearly pure calcium carbonate (Herbst and Bradley, 1989). These lime gland tubules accumulate calcium more rapidly than normal tubules.
It appears that carbonate and bicarbonate are regulated by means of chemical precipitation of these compounds with calcium in the lumen of the lime gland tubules Herbst et al. (1988). This storage excretion role of the Malpighian tubules contrasts with the structural modification of ileum and rectum that appear to be involved in ionoregulation by other ephydrids (Marshall and Wright, 1974).

Lime gland mass increases with the proportions of $\text{HCO}_3^-/\text{CO}_3^{2-}$ in the surrounding waters, and also with increased osmolality due to NaCl. It has been suggested that the lime gland serves not only to regulate carbonates, but to provide negative buoyancy (Herbst and Bradley, 1989). Larvae will tend to float and be exposed to higher mortality in wave-washed shallows of their lakeshore habitat in the absence of a lime gland ‘weight belt’.

6.4.2 Transport of bicarbonate by the rectal salt gland of mosquito larvae

Larvae of the mosquito *Aedes dorsalis* inhabit alkaline (pH 10.5) lakes rich in bicarbonate (250 mM) and carbonate (100 mM). Although larvae ingest the lake water at rates up to 130% body weight per day, they maintain hemolymph pH between 7.55 and 7.70 and hemolymph bicarbonate concentration between 8 and 18.5 mM (Strange et al., 1982). Regulation is achieved through active secretion of $\text{HCO}_3^-$ and $\text{CO}_3^{2-}$ by the rectal salt gland. The rectum is divided into two structurally distinct regions termed the anterior and posterior segments. Hyperosmotic secretion occurs in the posterior segment, whereas chloride uptake by exchange with $\text{HCO}_3^-$, along with organic solute reabsorption, occurs in the anterior segment (Bradley, 1987). Bicarbonate and carbonate are excreted against both chemical and electrical gradients. The lumen:hemolymph chemical gradient is 21:1 for $\text{HCO}_3^-$ and 241:1 for $\text{CO}_3^{2-}$, and the rectal transepithelial potential is 14–25 mV, lumen negative. The net electrochemical potential is 69 mV for $\text{HCO}_3^-$ and 76 mV for $\text{CO}_3^{2-}$, indicating active transport of both ions into the lumen.

Salt glands isolated and microperfused *in vitro* maintain transepithelial potentials as high as $-$50 mV, lumen negative, and secrete CO$_2$ in the absence of Na$^+$ or K$^+$ in lumen or serosal bathing salines. Bicarbonate secretion is inhibited by removal of chloride or addition of acetazolamide or DIDS, and net chloride reabsorption equals the rate of total CO$_2$ secretion, suggesting a 1:1 exchange of luminal Cl$^-$ for serosal $\text{HCO}_3^-$ (Strange and Phillips, 1984).

Both CO$_2$ secretion and Cl$^-$ reabsorption occur primarily in the anterior rectal gland, whereas the posterior segment is responsible for 75% of total hyperosmotic NaCl secretion, with the anterior segment contributing the remainder (Strange et al., 1984). It seems likely that both segments secrete NaCl rich fluids *in vivo*, and that the composition of the luminal fluid is subsequently modified by ion exchange and reabsorptive processes.
Fluid movements *in vivo* are not unidirectional or continuous, since the salt gland empties only after becoming filled with secretion and faecal material. Fluid secreted by the posterior segment, therefore, can come in contact with, and be modified by, the anterior segment.

A tentative model of ion transport by the gland was developed based on data from recordings with intracellular voltage-, pH- and Cl\(^{-}\)-selective microelectrodes, coupled with calculations of bicarbonate activity by the Henderson–Hasselbalch equation (Strange and Phillips, 1985). Bicarbonate crosses the basolateral membrane into the anterior rectal cell by an active mechanism against an electrochemical gradient of 77 mV and exits the cell at the apical membrane down a favourable electrochemical gradient of 28 mV (Fig. 10). The model proposes that chloride crosses the apical membrane into the cell by passive, electrodiffusive movement through a chloride-selective channel, and bicarbonate moves from cell to lumen by a passive or active electrogenic mechanism. Changes in serosal Cl\(^{-}\) concentration have no effect on basolateral membrane potential, indicating an electrically silent mechanism for Cl\(^{-}\) movement from cell to hemolymph. This step is DIDS-sensitive, raising the possibility of direct coupling of chloride and bicarbonate movements through Cl\(^{-}\)/HCO\(_3\)\(^{-}\) exchange across the basolateral membrane (Strange and Phillips, 1985).

**FIG. 10** Tentative cellular model of HCO\(_3\)\(^{-}\) and Cl\(^{-}\) entry and exit steps in anterior rectal salt gland cells of the larva of the mosquito *Ae. dorsalis* (c.a. = carbonic anhydrase). Note that the actual species (i.e. HCO\(_3\)\(^{-}\), H\(^{+}\) or OH\(^{-}\)) involved in transepithelial HCO\(_3\)\(^{-}\) secretion are not known with certainty. Redrawn from Strange and Phillips (1985, Fig. 12). Reprinted with permission from the American Physiological Society.
7 Excretion and sequestration of toxic metals

7.1 Sites and mechanisms of cadmium accumulation and transport

High levels of metals are accumulated by the midgut epithelium in many insects (e.g. Postma et al., 1996), in part in metal-containing mineral concretions. In ants, for example, measurements of metal levels (Pb, Cd, Cu, Zn, Fe, Mn) in different tissues of workers of *Formica pratensis* Retzius, *F. polyctena* (Forster) and *Camponotus ligniperda* (Latreille) collected from a metal-polluted site reveal that highest levels are found in the midgut, followed by the Malpighian tubules and the hindgut (Rabitsch, 1997). Levels of the metals in the tissues are correlated with the extent of metal pollution of the collection sites. The midgut appears to act as an effective barrier for many of the minerals, thereby limiting the extent to which toxic metals cross into the hemolymph, although the Malpighian tubule cells also contain concretions which may act as a form of storage excretion. A switch from accumulation to elimination presumably occurs in long-lived species when the storage capacity is exceeded.

Microautoradiography of whole larvae of *Chironomus riparius* exposed to cadmium in water or in water and sediment has shown that the alimentary canal is the primary site of cadmium accumulation (Craig et al., 1998). The digestive tract of *C. riparius* larvae has four structural divisions: the oesophagus (stomadeum), the anterior midgut, posterior midgut and the hindgut (proctadeum). Cadmium accumulates only in the anterior portion of the posterior midgut, a section which is rich in mitochondria, particularly within the long apical microvilli and alkaline phosphatase. These characteristics are consistent with a role in absorption. Accumulation of cadmium by the posterior midgut is related presumably to the function of the midgut as the site of absorption of calcium, and cadmium uptake probably occurs via calcium channels (Craig et al., 1998). Uptake of Cd$^{2+}$ via Ca$^{2+}$ channels is to be expected on the basis of similarity in ionic radii (0.097 and 0.099 nm for Cd$^{2+}$ and Ca$^{2+}$, respectively). Consistent with a role for Ca$^{2+}$ channels, Cd$^{2+}$ accumulation can be reduced as much as 88% by increasing the Ca$^{2+}$ concentrations of the water from 0.1 to 10 mM (Craig et al., 1999). There is also a concentration-dependent decrease in Cd$^{2+}$ accumulation in response to the Ca$^{2+}$ channel blockers lanthanum (73% at 10 μM and 92% at 100 μM) and verapamil (59% at 100 μM and 85% at 300 μM; Craig et al., 1999). A role for Ca$^{2+}$ channels in Cd$^{2+}$ uptake has also been proposed from studies of the *Aedes albopictus* C6/36 cell line. Uptake is saturable and the value of the Michaelis–Menten constant ($K_t$) is 1.7 μM. Uptake is unaffected by pre-treatment with dinitrophenol but is reduced by verapamil, suggesting that the metal is taken up into the cells by mediated transport not requiring metabolic energy (Braeckman et al., 1999).
The latter study also indicated that production of proteins which are likely members of the heat shock protein (HSP) family is induced by Cd\textsuperscript{2+} exposure. HSPs, metallothioneins and a family of proteins in invertebrates known as cadmium-binding proteins (CBPs) are all known to bind cadmium.

The Malpighian tubules were included with the gut in the analyses of the larvae of *Ch. riparius* by Postma *et al.* (1996), and separate values for the levels of cadmium in each tissue, therefore, are not available for this species. Seidman *et al.* (1986b) note little accumulation of radiolabelled cadmium in the anterior midgut, exoskeleton or Malpighian tubules, although the loss of more than 30% of the isotope during fixation of the tissue reduced the sensitivity of their technique. However, in the silkworm (*B. mori*) reared on an artificial diet containing cadmium at concentrations of 5 and 80 \( \mu \text{g g}^{-1} \) wet diet, cadmium accumulated in the alimentary canal and the Malpighian tubules at concentrations of 1100 and 470 \( \mu \text{g g}^{-1} \) wet weight, respectively. The finding of high levels of cadmium in the Malpighian tubules indicates that cadmium crosses the gut epithelium into the hemolymph, and that it is then transported into the Malpighian tubules (Suzuki *et al.*, 1984).

Metal-adapted and non-adapted populations of *Ch. riparius* differ in rates of storage and excretion of cadmium (Postma *et al.*, 1996). The adapted populations both accumulate more cadmium from the water and show higher rates of cadmium elimination when transferred to cadmium-free water. Cadmium may be initially sequestered by metal-binding proteins but eventually it is stored in the form of membrane bound concretions that have been observed in the posterior midgut of *Chironomus* (Seidman *et al.*, 1986a; Craig *et al.*, 1998). Cadmium-containing granules in the midgut may be expelled into the lumen by exocytosis or degeneration of whole cells. Degeneration of posterior midgut cells is seen in larvae exposed to high concentrations of cadmium (Seidman *et al.*, 1986a). The effects of crossing adapted and non-adapted chironomid populations on larval growth and EC\textsubscript{50} values during cadmium exposure has been examined by Groenendijk *et al.* (2002). A rapid loss of metal adaptation in the first generation hybrid offspring is evident in crossings of the adapted and non-adapted populations. Reciprocal crossings rule out any maternal effects and show a major genetic component for metal tolerance in exposed chironomid populations.

It is important to point out that the high tolerance of *Chironomus* larvae to cadmium exposure is not explicable solely by induction of cadmium-binding proteins. Reports that exposure to metals such as copper and cadmium induces metal-binding proteins in species such as the caterpillar *Galleria mellonella*, the housefly *Musca domestica*, the fruit fly *D. melanogaster* and several species of chironomids have been discussed by Postma *et al.* (1996). Duplication of the metallothionein gene may be an
adaptation to heavy metals in *D. melanogaster*. In Malpighian tubules of the latter species, the importance of metallothioneins can be deduced from the finding that metallothionein A is the most abundant transcript of any gene in the Malpighian tubule (Wang *et al.*, 2004). In *Chironomus*, however, cadmium uptake is very rapid, and appears to be non-specifically absorbed onto high molecular weight proteins from which it is rapidly released when larvae are transferred to cadmium-free water. Although a low molecular weight cadmium-binding protein is induced by exposure to cadmium in the water, the induction is very slow relative to the rate of cadmium uptake (Yamamura *et al.*, 1983). The cadmium-binding protein contains little zinc or copper, in contrast to vertebrate metallothioneins. Displacement of cadmium by copper, the prevalence of cysteinyl residues and absence of aromatic amino acids in the protein are common to metallothioneins in other species. The high tolerance of *Chironomus* larvae cannot be explained by the induction of these cadmium-binding proteins because cadmium is mostly not bound to them during the period of rapid cadmium uptake (Yamamura *et al.*, 1983). Nor is complexation with proteins the sole mechanism of cadmium detoxification in *Sarcophaga peregrine*. In this insect most of the cadmium accumulated from the diet is retained through the pupal stage but is excreted by the adult during the first day after emergence. Levels of a cadmium-binding protein gradually decrease after pupation, so that although most of the cadmium accumulated in the larvae is retained in the pupae, only limited amounts of it are bound to protein. Protection of pupal tissues and those of the newly emerged adult from the toxic effects of cadmium therefore requires a mechanism other than complexation with proteins such as metallothioneins (Yamamura *et al.*, 1983).

Results of a study on predatory beetles which had been fed mealworm larvae containing Cd, Zn, Hg or methyl mercury (Me-Hg) also indicate that complexation in the midgut cannot be the sole mechanism of detoxification (Lindqvist *et al.*, 1995). It appears that loss of the metals reflects faecal excretion of metals that were not absorbed from the food plus the discharge of midgut epithelial cells which had taken up each metal. Levels of cadmium and mercury are reduced to less than 1% of the initial quantity present within 2 weeks. By contrast, 60% of Me-Hg and 20% of Zn are still present in the beetle tissues 30 d after ingestion of metal-loaded food. Slower rates of discharge of Me-Hg reflect the more hydrophobic nature of the compound and consequent bioaccumulation. Autoradiography of slices of the beetles 2–3 weeks after ingestion of cadmium show that the metal is located primarily in the midgut but also in the integument. Some cadmium must therefore cross the gut epithelium into the hemolymph for subsequent deposition in the integument. In *Drosophila* larvae which have ingested cadmium, most of the metal is found in the digestive tract, with smaller quantities present in the Malpighian tubules,
hemolymph, fat body and integument (Maroni and Watson, 1985). The latter findings indicate passage of cadmium across the midgut epithelium in *Drosophila* as well. The significance of this passage is that cadmium detoxification cannot be explained solely through processes within the midgut, and a role for sequestration and or secretion within the Malpighian tubules and/or hindgut seems plausible.

Whereas terrestrial insects may face metal loading through the diet, aquatic species may also take up metals across the external organs used for ionoregulation. The number of ionoregulatory cells (chloride cells) is related to uptake rates for dissolved Cd\(^{2+}\) and Zn\(^{2+}\) (Buchwalter and Luoma, 2005). Dissolved Cd\(^{2+}\) uptake rates are determined by the numbers of Cd\(^{2+}\) transporters rather than by the affinities of those transporters for Cd\(^{2+}\), and Cd\(^{2+}\) uptake is not related to body size or gill size.

### 7.2 REGULATED STORAGE OF ZINC AND COPPER IN *DROSOPHILA*

The presence of zinc, copper and iron in storage vesicles within the Malpighian tubules (Sohal *et al*., 1977) lead to the hypothesis that accumulation of these metals was a detoxification mechanism. The gut may also be implicated in detoxification. For example, copper cells found in the *Drosophila* midgut are named for their capacity to accumulate the metal and are also of interest because they show structural similarities to the acid-producing gastric parietal cells of the mammalian stomach (Dubreuil, 2004). Maddrell (1977) suggested that metals would be reabsorbed into the hemolymph if they were excreted from the Malpighian tubule into the gut. Metal ions are small relative to organic toxins and would therefore diffuse more readily through the cells of the hindgut into the hemolymph and would thus simply be returned to the Malpighian tubules. This futile cycling would be particularly problematic if a substantial proportion of the metals passing through the gut were absorbed, and the storage excretion of metals would thus minimize energy expenditure and protect sensitive tissues from exposure to toxic levels of divalent cations.

The detoxification hypothesis has been challenged by Schofield *et al*. (1997), who propose that accumulation of Zn and Cu in tissues of the fruit fly *Drosophila* represents regulated storage rather than deposit excretion. Several pieces of evidence support this view. Measurement of elemental contents of whole flies and organs by a penetrative ion microprobe technique shows that zinc is accumulated primarily in the main segments of both the anterior and posterior Malpighian tubules at concentrations up to 2.8% of tissue dry weight (Schofield *et al*., 1997). By contrast, iron and copper are distributed throughout the body. Accumulation of a specific divalent metal is considered to indicate regulation of biological availability rather than deposit excretion. The most critical challenges to the
detoxification hypothesis were the findings that less than 1% of copper and zinc are absorbed from the food and that there is no proportionality between the quantities of zinc and copper consumed and the quantities accumulated.

Regulated storage of zinc rather than deposit excretion is consistent with the requirement of zinc in several biological processes. In contrast to copper and cadmium, zinc does not bind to metallothioneins nor does it induce their synthesis (Maroni and Watson, 1985). Although high dosages of zinc are toxic, zinc is a component of at least 300 enzymes and is also chelated by zinc finger proteins which are implicated in gene regulation. Zinc is also incorporated into the arthropod cuticle during moult and is accumulated at concentrations as high as 25% in the mouthparts and claws as a means of increasing hardness.

7.3 DETOXIFICATION AND EXCRETION OF IRON AND HEME

Multiple tissues are involved in maintaining appropriate levels of iron within insects (Locke and Nichol, 1992). The midgut plays roles in uptake of iron from the diet, storage and also excretion. The hemolymph plays a role in transport and storage, and the Malpighian tubules are also implicated in uptake from the hemolymph, storage and excretion. Lastly, the pericardial cells are important for uptake and heme recycling and the fat body plays a role in intermittent storage of iron. Within cells, diverticula of the rough endoplasmic reticulum contain concretions of calcium salts along with other metals such as zinc and iron.

Blood-feeders, such as mosquitoes and the hemipteran R. prolixus, face a massive release of heme during hydrolysis of host hemoglobin in the digestive tract. Subsequent degradation of heme by heme oxygenase (HO) leads to production of iron, carbon monoxide and biliverdin. Both iron and heme lead to formation of reactive oxygen species that can damage a variety of biological molecules. Although biliverdin does have antioxidant properties, it must be excreted to avoid cytotoxicity presumably related to the effects of high levels of hydrophobic molecules on membrane structure and function as well as the effects of bile pigments on DNA and protein synthesis.

Studies by Paiva-Silva et al. (2006) have identified a unique and complex heme-degradation pathway, with four distinct intermediates, in R. prolixus. In mammals the heme ring is cleaved by HO at the alpha methene bridge to form biliverdin which is subsequently converted to bilirubin by biliverdin reductase. In Rhodnius, heme is first modified by addition of two cysteinylglycine residues before cleavage of the porphyrin ring, followed by trimming of the dipeptides. The end product of heme detoxification in Rhodnius is a dicysteinyl-biliverdin IXβ. The addition of hydrophilic residues to heme molecules may be a way to facilitate the excretion of large
amounts of biliverdin, a relatively hydrophobic molecule, produced after blood digestion (Paiva-Silva et al., 2006). Similarly, in mammals, bilirubin is excreted only after conjugation to glucuronic acid, and transport by members of the multidrug resistance-associated protein (MRP) family and the OATP family of anion transporters has been implicated.

Although both heme and iron cause oxidative stress, they promote lipid peroxidation by different mechanisms. Iron-induced oxidative stress results from generation of hydroxyl radicals (\(\text{OH}^*\)) by the Fenton reaction:

\[
\text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH}^- + \text{OH}^*
\]

Hydroxyl radicals can then initiate lipid peroxidation chains by removing electrons from other molecules such as unsaturated fatty acids, thereby generating an alkyl radical (\(\text{R}^*\)). In contrast, heme-induced formation of radical species results from conversion of less reactive organic hydroperoxides (ROOH) into highly reactive alkoxyl (RO\(^*\)) and peroxyl (ROO\(^*\)) radicals (Graca-Souza et al., 2006).

An important antioxidant role is played by urate in the hemolymph of the blood feeder \textit{R. prolixus}. Although the Malpighian tubules of \textit{Rhodnius} secrete urate at high rates, the rates of urate synthesis and excretion are balanced so that the hemolymph contains urate concentrations as high as 5 mM (Souza et al., 1997). Urate synthesis by the fat body \textit{in vitro} is stimulated by the presence of hemin (the Fe\(^{3+}\) oxidation product of heme) through a mechanism involving protein kinase C activation. Hemolymph urate concentration increases \textit{in vivo} in response to blood feeding or injection of hemin into the hemocoel and also when animals are exposed to hyperoxic conditions (70% \(\text{O}_2\)). Preventing urate synthesis \textit{in vivo} by addition of the xanthine oxidase inhibitor allopurinol results in increased levels of free radicals in the hemolymph. Urate synthesis thus appears to be an important adaptation to the oxidative stress produced by hemin degradation during digestion of the blood meal. Additional protection is provided by the \textit{Rhodnius} heme binding protein (RHBP).

In mosquitoes, protection against the oxidative challenge presented by excess iron derived from digestion of the blood meal in mosquitoes is provided by the iron-binding protein ferritin (Geiser et al., 2003) and also be excretion (Zhou et al., 2007). The primary site of expression of the ferritin in the yellow fever mosquito, \textit{Ae. aegypti}, is the midgut and expression of both the ferritin heavy-chain homolog (HCH) and the light-chain homolog (LCH) are induced by blood-feeding. Messenger RNA levels for both LCH- and HCH-genes increase with iron, hydrogen peroxide and hemin treatment, and the temporal pattern of gene expression of the genes is very similar. Ferritin is also present in the peripheral tissues of blood-feeders to provide further protection against the effects of iron-induced oxidative stress. It appears that iron required for short term
nutrient needs is retained in cytoplasmic ferritin, whereas excess iron is stored inside ferritin that is subsequently secreted from the cells into hemolymph. Secreted ferritin may thus be a primary iron storage protein in mosquitoes and the hemolymph may be the primary tissue involved in iron storage. Ferritin secretion may thus provide a mechanism by which mosquito cells protect against iron overload and reduce the potential for iron-mediated oxidative stress inside cells (Geiser et al., 2006).

It also appears that excretion plays a major role in preventing iron toxicity after the blood meal. The blood meal provides high levels of iron primarily as hemoglobin in erythrocytes, and to a small extent as ferric-transferrin. A quantitative evaluation of the fate of blood-meal iron during the first gonotrophic cycle in *Ae. aegypti* has been provided by Zhou et al. (2007). Most of the heme iron provided by the blood meal is excreted, whereas iron provided as ferric-transferrin is highly absorbed and transferred to the ovaries and eggs. Because iron provided from heme in blood is in much greater quantity than that provided by transferrin, the majority of iron absorbed from the blood meal comes from heme. Of the iron supplied in the form of heme, 87% of the ingested total is excreted, 7% accumulates in the eggs and 6% is stored in other tissues. Of the iron supplied in a blood meal, approximately 7% appears in the eggs and of this iron 98% is from hemoglobin and 2% from ferric-transferrin.

A role for the Malpighian tubules in iron metabolism has been suggested on the basis of studies of the mosquito *Anopheles albimanus*. Martinez-Barnetche et al. (2007) cloned a homolog (AnaNRAMP) of the divalent metal transporter 1/natural resistance-associated macrophage protein (NRAMP) family. This group of proton-coupled divalent cation transporters is widely distributed in prokaryotes and in eukaryotes and is involved in Fe and Mn transport. AnaNRAMP mRNA induces iron incorporation when injected into *Xenopus* oocytes and Western blots show expression of AnaNRAMP in head, midgut and at high levels in Malpighian tubules of unfed female mosquito. AnaNRAMP levels in the midgut decline after blood-feeding, but increase in the Malpighian tubules. AnaNRAMP is localized in the membrane of intra-cellular concretions or spherites in the principal cells of the Malpighian tubule. Localization of AnaNRAMP in concretions suggests that divalent cations such as Fe$^{2+}$ and Mn$^{2+}$ may also be deposited there for storage. AnaNRAMP may thus function as a divalent cation efflux pump transporting iron from the vesicle to the cytosol, where it can be stored by cytosolic ferritin. At present it is not known if AnaNRAMP-mediated cation transport is involved in iron retention/storage or iron excretion (Martinez-Barnetche et al., 2007).

In honeybees iron is obtained from pollen in the diet, and the rate of iron accumulation is directly related to the levels of iron in the diet. Honey bees store iron both as granules in the midgut and in the trophocytes of the fat body. However, the iron content of the fat body reaches a maximum level
regardless of the amount of iron available for ingestion, and is reached at
the commencement of foraging behavior, suggesting a possible role in
orientation. The honey bee iron granules are larger than other forms of
iron granules (0.32 μm diameter), contain significant amounts of calcium
and phosphorous, and have an amorphous electron structure when
examined by X-ray diffraction. Given that magnetite is absent or only a
minor component (<0.33%) and that the trophocytes are not innervated,
1
a direct role for the granules in magnetic field detection seems unlikely.
However, the granules may be paramagnetic and affect local intracellular
enzymatic activity in response to fluctuations in external magnetic fields
(Kuterbach and Walcott, 1986a,b; Kuterbach et al., 1982). Alternatively,
the granules may be involved in iron homeostasis. Many of the enzymes
required for energy production during flight, for example, contain iron
(e.g. cytochrome c oxidase, NADPH oxidase).

8 Transport of organic cations and organic anions

Renal tissues of vertebrates and invertebrates transport a wide range of
organic cations (OCs), most with a quaternary nitrogen, a hydrophobic
region and a positive charge at physiological pH. They include numerous
exogenous compounds and metabolites, many of which are toxic.
Transport and elimination of OCs is therefore essential for the
maintenance of homeostasis. Multiple transporters are involved, in part
reflecting the need to transport molecules from two broad structural
classes. Type I organic cations are relatively small (< 400 Da) hydrophilic
monovalent compounds and include both endogenous molecules such as
choline or N-methyl nicotinamide (NMN) and drugs such as tetraethyl-
1
lammonium (TEA). Type II organic cations are larger amphiphilic
compounds that contain a positively charged group situated close to or
within a large aromatic ring structure (Wright and Dantzler, 2004).
Examples of type II organic cations are nicotine and quinidine. Organic
cation transport by renal organs commonly involves carrier-mediated
potential-driven uptake at the basolateral membrane, intracellular
sequestration that reduces the free concentration of the cation, and
luminal exit by P-gps or through organic cation-proton exchange
(Pritchard and Miller, 1993). Studies of organic cation transport often
utilize radiolabelled compounds such as the prototypical organic cations
TEA or NMN. Organic cation transport can also be quantified through the
use of fluorescent compounds such as [2-(4-nitro-2,1,3-benzoxadiazol-7-yl)
aminoethyl]-trimethylammonium (NBD-TMA) (Bednarczyk et al., 2000)
or quinidine (Leader and O’Donnell, 2005).
Early qualitative studies indicated the accumulation of basic (cationic) dyes such as methyl green and methylene blue by the proximal Malpighian tubules of *Ma. sexta* (Nijhout, 1975). The first quantitative estimates of excretion of type I organic cations such as TEA or NMN in insects were presented for *D. melanogaster* (Rheault and O'Donnell, 2004). Electrophysiological techniques for assessing the fluxes of tetraalkylammonium compounds, especially the prototypical organic cation TEA, were applied to analysis of organic cation transport by isolated guts and Malpighian tubules. The techniques exploit the high selectivity of the cation exchanger potassium tetra(*p*-chlorophenyl)borate (Corning 477317) for quaternary ammonium compounds such as TEA and tetramethylammonium (TMA). Although originally used for measurement of $K^+$ activity, the selectivity of microelectrodes based on this exchanger for TEA and TMA exceeds that for $K^+$ by factors of $10^7$ and $10^{2.7}$, respectively.

TEA-selective microelectrodes are used to measure TEA transport in two ways, and related microelectrode methods can be applied to measuring the transport of other ions ($Na^+$, $K^+$) and also organic compounds such as salicylate (Section 8.3) by Malpighian tubules and other tissues. Figs. 11–14 are therefore presented both with a view to explaining specific features of TEA transport and with a view to showing the methodologies for quantifying ion fluxes through the use of ion-selective microelectrodes. Firstly, the concentration of TEA is measured in fluid droplets secreted by isolated Malpighian tubules of *Drosophila* set up in a Ramsay secretion assay (Fig. 11). Flux across the entire tubule is calculated from the product of secretion rate and secreted droplet TEA concentration. Secondly, TEA-selective microelectrodes based on Corning 477317 ion exchanger are also used with the scanning ion electrode technique (SIET) to assess spatial and temporal variations in TEA flux in different regions of the Malpighian tubules and gut. A TEA-selective microelectrode is moved between two positions within the unstirred layer next to the surface of a cell, and the measured difference in TEA concentration between the two positions is converted into a corresponding TEA flux using the Fick equation (Fig. 12).

Measurements based on the Ramsay assay and the SIET measurements have revealed TEA transport by the Malpighian tubules, ureter and posterior midgut (Figs. 13, 14). TEA transport by the posterior midgut may serve to minimize absorption of potentially toxic organic cations from the gut lumen into the hemolymph, whereas transport by the tubules presumably serves to clear the hemolymph of toxins. Given that the transepithelial potential across the main (secretory) segment of the tubule is 30–80 mV lumen-positive (O'Donnell *et al.*, 1996) and that the lumen concentration of TEA is ~12-fold above that in the bath for main
segments bathed in 5 μM TEA, TEA must be actively transported into the secreted fluid against an opposing electrochemical gradient.

Active secretion of TEA by the lower tubule (Figs. 13, 14) is accompanied by acidification of the urine, active secretion of Ca^{2+} and reabsorption of K^{+} and Cl⁻ by this segment (O'Donnell and Maddrell, 1995). A possible mechanism for TEA secretion across the apical membrane of the lower tubule is exchange of organic cations for protons (Bijelic and O'Donnell, 2005), as proposed for organic cation transport across the apical membrane of vertebrate renal tissues (Pritchard and Miller, 1991).

The mechanism of TEA transport across the basolateral membrane of Drosophila tubules was studied using measurements of basolateral...
membrane potential ($V_{bl}$) and uptake of [14C]-labelled TEA (Rheault et al., 2005). In common with vertebrate renal tissues, uptake of [14C]-labelled TEA is dependent on metabolism and saturable. TEA uptake in Drosophila tubules is inhibited by a number of other type I organic cations (e.g. other quaternary ammonium compounds, cimetidine, quinine) and is dependent on basolateral membrane potential. Conditions that depolarize basolateral membrane potential (e.g. high K+ saline) reduce TEA uptake, whereas hyperpolarizing conditions (low K+ saline) increase uptake. Addition of TEA to the saline bathing Malpighian tubules rapidly depolarizes the basolateral membrane potential in lower Malpighian tubules of Drosophila, indicating that TEA uptake is electrogenic. However, the effects of TEA on basolateral membrane potential are not altered when K+ channels are first blocked with Ba2+, indicating that TEA uptake does not occur through K+ channels in the lower tubule (Fig. 15).
FIG. 13  (A) Representative scan of tetraethylammonium (TEA) flux at locations along the secretory segment of the Malpighian tubule (MT) and the lower MT (LMT) of *Drosophila melanogaster*. Tubules were bathed in saline containing 100 μM TEA. The common ureter and part of the other LMT of the pair are shown. The tip of the TEA-SeR microelectrode is located just above the asterisk. The image is a collage formed from two images. At each site, indicated by arrowheads, ASET software calculated the TEA-specific signal differences (ΔV; μV) between the two limits of microelectrode excursion by subtracting the voltage at the outer limit of the excursion from that measured at the inner limit. The length of each arrow corresponds to the magnitude of TEA influx. (B) TEA influx as a function of distance from the ureter along lower, main and distal segments of the MT. An influx of TEA reduces TEA concentration in the unstirred layer adjacent to the surface of the tissue, and the corresponding voltage difference is therefore negative. Distance 0 on the abscissa corresponds to the junction of the ureter and the LMT. Both the differential signal recorded by the TEA-SeR microelectrode (right ordinate) and the calculated TEA influx (left ordinate) are shown (N = 4). Redrawn from Rheault and O’Donnell (2004, Fig. 1). Reprinted with permission from the Company of Biologists.
FIG. 14  Concentration–response curves for tetraethylammonium (TEA) influx in the (A) lower Malpighian tubule (LMT) of *Drosophila melanogaster*, (B) Malpighian tubule main segment and (C) midgut. Each point is the mean ± S.E.M. of \( N = 4–7 \) preparations. Values for \( J_{\text{max}} \) and \( K_i \) were determined by non-linear regression analysis. Redrawn from Rheault and O’Donnell (2004, Fig. 4). Reprinted with permission from the Company of Biologists.
Addition of verapamil, a type II organic cation, also causes a concentration-dependent inhibition of TEA uptake. Previous studies on snake renal proximal tubules have demonstrated that verapamil competes with TEA for the basolateral organic cation uptake pathway (Kim and Dantzler, 1995). Moreover, the plant alkaloid, nicotine, a demonstrated P-gp substrate in insect Malpighian tubules, inhibits basolateral TEA uptake. Taken together, these results suggest a broad overlap in specificity of the basolateral organic cation transport pathway for type inorganic cations such as TEA and type II organic cations such as nicotine and other alkaloids. Studies in mammals also indicate that type II OC’s inhibit transport of type I OC’s, but not vice versa (Wright and Dantzler, 2004).

Secretion of TEA by Malpighian tubules appears to be common but not universal among insects. TEA is also transported from bath to lumen in Malpighian tubules of seven species of insect from six orders. Based on the ratio of secreted [TEA] to that in the bathing medium, as well as the value of the transepithelial potential recorded in earlier studies by several groups,
Rheault et al. (2006) concluded that TEA is actively secreted by tubules of the orthopterans *L. migratoria*, and *Ac. domesticus*, the dictyopteran *P. americana*, the coleopteran *Te. molitor*, the lepidopteran *Trichoplusia ni* and the hemipteran *Oncopeltus fasciatus*. By contrast, Malpighian tubules of two blood-feeders, the hemipteran *R. prolixus* and the dipteran *Ae. aegypti* do not actively transport TEA under normal physiological conditions. Transport of TEA but not nicotine increases during the moult in the Malpighian tubules of *Rhodnius*, but the concentrations of TEA in the secreted fluid are still consistent with passive accumulation in response to the lumen-negative transepithelial potential. Both of the latter species actively transported the type II organic cation nicotine. The absence of active transport of the prototypical type I organic cation TEA by the Malpighian tubules of these species is puzzling and suggests that toxic levels of type I organic cations in the hemolymph may be avoided by excretion across other epithelia such as the midgut, as observed in *Drosophila* (Rheault and O'Donnell, 2004). Alternatively, type I OCs in *Rhodnius* and *Aedes* may be metabolized into other compounds, which are either non-toxic or are readily excreted by the tubules or other epithelia.

Several types of evidence suggest that there are separate pathways for transport of the type I organic cation TEA and the type II organic cation nicotine in insect Malpighian tubules. Verapamil reduces TEA transport by the lower tubule but not main segment of *Drosophila* Malpighian tubules (Rheault and O'Donnell, 2004). Inhibition by verapamil suggests the additional involvement of a P-gp-like transport mechanism for TEA in the lower tubule that is not involved in the secretion of TEA by the main segment.

The possibility that excretory mechanisms are altered by exposure to dietary toxins was examined by feeding *Drosophila* larvae a diet enriched in TEA (Bijelic et al., 2005). More than 75% of third instar *Drosophila* larvae survive on a diet containing 100 mM TEA. There is a synergistic, rather than additive, increase in larval mortality when the organic cations quinidine or cimetidine are added to a diet containing 100 mM TEA; the combined response was significantly greater than the sum of the mortalities associated with TEA and either cimetidine or quinidine alone. The synergism may reflect the competitive inhibition of TEA transport by either quinidine or cimetidine, resulting in higher hemolymph levels of TEA. The rate of decline in hemolymph TEA after transfer to TEA-free diet is reduced by 58% when larvae are maintained on diet that contains 10 mM cimetidine and 100 mM TEA as compared to the rate of decline when the diet contains 100 mM TEA alone. Cimetidine presumably slows the rate of decline of the TEA concentration in the hemolymph because it reduces the rate of TEA secretion by transporters in the Malpighian tubules and posterior midgut. When larvae are fed a diet that contains <20 mM TEA and are subsequently transferred to TEA-free diet, the
concentration of TEA in the hemolymph declines at a rate that can be
accounted for by active and saturable transport of TEA across the
Malpighian tubules and the midgut. However, when larvae are transferred
from a diet containing 100 mm TEA to a TEA-free diet, hemolymph TEA
concentration declines at least 10 times more rapidly than predicted on the
basis of active, saturable transport across these tissues. Passive transport
into the gut may provide a non-saturable means for rapid clearance of
hemolymph TEA when animals are fed TEA-free diets (Bijelic et al., 2005).
The gut contents may thus act as a passive ‘sink’ for toxins in the
hemolymph, providing that the insect can feed upon toxin-free foods after
ingestion and absorption of high levels of toxins.

Transepithelial TEA secretion and secreted fluid TEA concentration
increase significantly in tubules isolated from third instar larvae that are
maintained for 24 h on media containing 50 or 100 mM TEA (Bijelic et al.,
2005). This increase suggests an adaptive response, whereby ingestion of a
TEA is associated with up regulation of TEA excretion by the Malpighian
tubules. The effects of long-term exposure (~ 10 d) to another toxin, the
organic anion salicylate, are discussed in Section 8.3.

8.2 TRANSPORT OF TYPE II ORGANIC CATIONS BY P-GLYCOPROTEIN-LIKE
MECHANISMS

P-gps function in MDR in humans by acting as ATP-dependent drug efflux
pumps to maintain the intracellular concentrations of anticancer drugs
below the cytotoxic levels. Expression of P-gp on the secretory surfaces in a
number of tissues including adrenal gland, kidney, liver, intestinal tract
and uterine epithelium also suggests a role either in transporting substances
away from tissues and out of the body or in decreasing absorption from the
surroundings. P-gps in aquatic species aid survival in environments which
contain high levels of anthropogenic pollutants or toxins produced by
other organisms. The term multixenobiotic resistance (MXR) is used to
distinguish this phenomenon from MDR in humans (Bard, 2000).

Substrates of P-gps are usually moderately hydrophobic planar
molecules, including type II organic cations (Wright and Dantzler, 2004).
In addition, P-gps transport relatively hydrophobic compounds that are
uncharged or neutral, including digoxin, colchicine and some corticoster-
oids. It is generally thought that P-gps function as ‘flippases’ to flip
substrates from the inner to the outer leaflet of the plasma membrane
against an intramembrane concentration gradient. The substrate then
diffuses out of the outer leaflet into the extracellular medium (Higgins and
Gottesman, 1992). In the sections below, the possible roles of P-gps in
transporting insecticides are discussed first, followed by a section dealing
with evidence for P-gps in specific insect tissues such as the Malpighian
tubule. The subsequent sections deal with the chemical ecology of P-gp substrates such as nicotine and other alkaloids.

8.2.1 \textit{P-glycoproteins and insecticide effects}

The P-gp inhibitor verapamil (100 \textmu M) increases by 1.5 to 2-fold the toxicity of examples of three classes of insecticides (cypermethrin, endosulfan and ivermectin) in three strains of \textit{Culex} mosquitoes but has no effect on the toxicity of the organophosphate chlorpyrifos (Buss et al., 2002). Although verapamil is best known as a potent blocker of L-type calcium channels, it is also a well known example of agents known as chemosensitizers, which have been shown to reverse multidrug resistance. Although relatively non-toxic themselves, chemosensitizers interact with P-gp so as to increase the toxicity of compounds such as insecticides. Two of the insecticides also strongly inhibit the efflux of a fluorescent P-gp substrate out of CH1 cells, a standard test of P-gp inhibition. The organophosphate insecticides fenithion and chlorpyrifos and the organochlorine DDT have no effect on R123 efflux, whereas ivermectin and the cyclodiene organochlorine endosulfan reduce efflux. Ivermectin is the most potent P-gp inhibitor in these assays. Widely used as an antihelminthic and as a commercial insecticide, ivermectin is a semi-synthetic analogue of avermectin, which is produced by the actinomycete \textit{Streptomyces avermitilus}. Ivermectin is well known as a P-gp inhibitor in studies of mammalian cells and tissues as well as clinically.

The interactions of insecticides with P-gps have also been studied in cultured murine melanoma cells transfected with a human MDR gene (Bain and LeBlanc, 1996). Six out of eight organophosphate insecticides, including chlorpyrifos, inhibit P-gp, as measured by the inhibition of efflux of the P-gp substrate doxorubicin. Although carbamate and pyrethroid insecticides do not interact with P-gp, many organophosphorus and organochlorine pesticides do bind to the transporter. However, many pesticides that are capable of binding to P-gp are not transported by it. Based on structure-activity analyses, pesticides that are good candidates for P-gp inhibitors have lipophilicity (measured as the log of the octanol: water partition coefficient, \( K_{ow} \)) of between 2.5 and 4.5 and a molecular mass of 300 or more (Bain and LeBlanc, 1996). Using these criteria, pesticides likely to inhibit P-gp are ranked in the order endosulfan > chlorpyrifos > DDT > fenithion. Although chlorpyrifos is not an inhibitor of P-gp based on reduction in R123 efflux from CH1 cells (Buss et al., 2002), chlorpyrifos increases P-gp expression in rats and the active metabolite chlorpyrifos oxon inhibits P-gp substrate binding. Differences in the effectiveness of chlorpyrifos as a P-gp inhibitor may relate to differences in substrate specificities in P-gps derived from expression of different MDR genes (Buss et al., 2002).
The significance of P-gp interaction with commercial pesticides is twofold. Firstly, P-gp may be a component of insecticide resistance, especially given its potential to confer resistance against a wide range of pesticides. This adds a complication to control programs for mosquitoes and possibly other disease vectors and agricultural pests. However, the finding that insecticide toxicity is increased by chemosensitizers such as verapamil opens up a new avenue for development of novel, environmentally benign insecticide synergists. In the study of Buss et al. (2002) for example, verapamil was itself non-toxic but increased the toxicity to examples of three insecticide classes. The idea is that compounds of low toxicity themselves may permit reduced insecticide application rates because they block elimination of insecticides from pest species.

8.2.2 P-glycoproteins in the gut, Malpighian tubules and anal papillae

Both the gut and the Malpighian tubules are important sites of P-gp expression in insects. In *D. melanogaster*, P-gps are the products of expression of three genes, MDR49, MDR50 and MDR65. Larvae of *D. melanogaster* fed on diets containing a non-toxic level of the P-gp substrate colchicine (10 μM) show increased expression of MDR genes in brain and gut relative to control larvae raised on colchicine-free diet (Tapadia and Lakhotia, 2005). Increased MDR expression is evident in all sections of the gut (foregut, midgut, hindgut and Malpighian tubules).

Prompted by evidence that promoters of human multidrug resistance MDR genes harbour stress-responsive elements, which respond to heat shock, heavy metals and cytostatic drugs, Tapadia and Lakhotia (2005) also studied expression of *Drosophila* MDR genes under heat stress. After heat shock at 37°C for 40 min, third instar larvae were either immediately dissected and different tissues were fixed or were allowed to recover from heat shock for different time periods at room temperature before tissue fixation. A 40-min heat shock significantly increases the expression of MDR genes in different larval tissues such as the brain and salivary glands. Levels of the MDR49 transcripts remain elevated above control levels for more than 60 min after heat shock, and expression returns to basal non-heat shock levels only after 90 min recovery. By contrast, the levels of MDR65 transcripts are unaffected by heat shock.

These differences in the effects of heat shock on expression of MDR49 versus MDR65 are explicable on the basis of differences in the actions of the heat shock transcription factor (HSF), which has been widely implicated in the induction of the heat shock genes in eukaryotes by heat and other forms of stress. The involvement of HSF in the up regulation of MDR genes in *Drosophila* larvae has been examined after heat shock by immunostaining polytene chromosomes with anti-HSF antibody. Although HSF is found at the MDR49 gene locus on polytene
chromosomes from heat shocked salivary glands, it is not present at the 65A region, where the MDR65 gene is located, either in control or in heat-shocked salivary glands. These findings are consistent with a role for HSF in mediating the increased expression of MDR49 relative to MDR65 in response to heat shock. MDR genes use alternative promoters in response to different stimuli in human cell lines and tissues and there are distinct nuclear protein-binding sites in the promoter of the murine MDR gene. It will be of interest to analyze promoter regions of the MDR genes of Drosophila to further examine differences in the regulation of these genes.

Several lines of evidence indicate the presence of a P-gp-like pump in the anal papillae of the larvae of the non-biting midge Ch. riparius (Podsiadlowski et al., 1998). The anal papillae are stained in vivo by the fluorescent P-gp substrate rhodamine 123 in the presence of the P-gp inhibitor verapamil. There is no staining in the absence of the inhibitor, consistent with the role of a P-gp-like pump in transporting rhodamine 123 out of the cells. Immunohistochemical analysis also reveals P-gp-like molecules in the anal papillae. Positive staining is evident in the anal papillae when a monoclonal antibody against human P-gp is used in conjunction with secondary antibodies conjugated to peroxidase. There is no staining when non-immune mouse serum replaces the primary antibody, or when antibodies against human MRP are used (Podsiadlowski et al., 1998). Hydrolysis of ATP in homogenate fractions from the posterior segments (including the anal papillae) of Chironomus larvae is enhanced when P-gp substrates such as vinblastine and ivermectin are present. By contrast, addition of the MRP substrates arsenate and glutathione does not enhance ATP consumption. Hydrolysis of ATP by other ATPases was blocked in these experiments by addition of a cocktail containing appropriate inhibitors such as ouabain for the Na\(^+\)/K\(^+\)-ATPase. Lastly, synergism between the toxicity of the P-gp substrate ivermectin and P-gp inhibitors is consistent with a role for P-gp in protection of the larvae against P-gp substrates such as ivermectin. The LD\(_{50}\) of ivermectin was reduced nearly threefold by sublethal concentrations of either of the P-gp inhibitors verapamil or cyclosporin.

A P-gp-like mechanism in the Malpighian tubules of the tobacco hornworm is responsible for active excretion of nicotine and other alkaloids (Gaertner et al., 1998). Malpighian tubules were cannulated and P-gp substrates were applied to the bathing fluids or to both the bathing fluids and the lumen. The luminal contents were flushed out at the end of the experiment and lumen to bath ratios of the substrates were measured. Lumen to bath ratios of 4.0, 10.1 and 3.2 were measured at bathing saline nicotine concentrations of 0.05, 0.5 and 5 mM, respectively. Application of the P-gp inhibitor verapamil (0.025 mM) reduced the lumen to bath ratio from 10 to 1.2 for tubules in saline containing 0.5 mM nicotine. Stirring the bathing saline to minimize unstirred layer gradients (i.e. depletion of
nicotine in the unstirred layer at the basolateral surface) increased the lumen to bath ratio from 4 to 12.3. Body movements tend to minimize unstirred layer gradients in vivo (cf. Collier and O’Donnell, 1997), so the highest lumen to bath ratios are probably indicative of the maximal capacity of the tubules to secrete nicotine. Malpighian tubules of the tobacco hornworm are immunopositive for P-gp, as is the blood–brain barrier (Murray et al., 1994). P-gp in the latter tissue thus protects the nervous system from the effects of nicotine until the toxin can be secreted into the lumen of the Malpighian tubule and eliminated.

It is difficult to estimate the contribution of P-gp to elimination of toxins by the Malpighian tubules using only measurements of lumen to bath ratios. The problem is that changes in the rates of fluid secretion rate influence the concentration of the solute in the lumen, particularly if the permeability of the tubule wall to the solute is high. A stimulant of fluid secretion will thus appear to inhibit transport of a P-gp substrate if only lumen to bath concentration ratios are measured, since the concentration of the solute in the lumen will be reduced. However, the transepithelial flux of the solute may actually increase, since the reduction in luminal concentration will reduce the extent passive lumen to bath diffusion of the solute. It is thus essential to measure both the concentration of the transported solute in the lumen and the fluid secretion rate, so that the transepithelial flux may be calculated. For tubules of most species, fluid secretion rates are usually expressed in units of nanoliters per minute and concentrations of solutes in the lumen as μM. Transepithelial flux is calculated as the product of fluid secretion rate (nl min⁻¹) and luminal solute concentration (μM) and is usually expressed in units of fmol min⁻¹ for Drosophila tubules.

Measurement of fluid secretion rate permits non-specific toxicity of P-gp inhibitors or of high concentrations of P-gp substrates themselves to be distinguished from specific inhibition of transepithelial flux through direct effects on P-gps or other types of transporter. A study of transepithelial flux of several fluorochromes revealed inhibition of transepithelial flux of the P-gp substrates rhodamine 123 and daunorubicin by verapamil (Leader and O’Donnell, 2005). The results of the latter study also indicate that rhodamine 123 is not an appropriate substrate for analysis of P-gp function in the tubules of species such as Drosophila because it dramatically inhibits fluid secretion. Measurements of only lumen-to-bath ratios of rhodamine 123 would have led to profound errors in estimates of the kinetic parameters (Jₘₐₓ, Kₜ) for rhodamine 123 transport. Rhodamine 123 is also used as a mitochondrial dye, so its effects on cell metabolism and hence on fluid secretion may be particularly apparent in the highly active cells of Malpighian tubules. Rhodamine 123 inhibits fluid secretion by tubules of crickets (Teleogryllus commodus) and Drosophila at concentrations of 10 and 0.1 μM, respectively. The latter value is nearly
40-fold less than the concentration at which rhodamine 123 transport is half maximal ($K_t$) in cricket tubules. Analysis of dye transport solely by image analysis of the cellular and luminal compartments would not reveal this additional information about the effects of the dye on cell function. The use of rhodamine 123 is also complicated by the finding that principal cells of dipteran tubules retain rhodamine 123 for a short period and the stellate cells reabsorb the dye from the lumen. The dye accumulates in stellate cell mitochondria and intensely fluorescent vesicles which are probably lysosomes (Meulemans and De Loof, 1992).

A novel fluorometric method for the analysis of transepithelial fluxes of fluorescent dyes, including some P-gp substrates, has been developed by Leader and O’Donnell (2005). The method is also suitable for use with fluorescent substrates of other transporters for organic cations or organic anions. Isolated tubules are setup in Ramsay secretion assays and nanoliter samples of secreted fluid are collected in optically flat rectangular glass capillaries. The measurement of fluorochrome concentration in the secreted droplets circumvents many of the problems associated with confocal laser scanning microscopy (CLSM) of the isolated tubules. The lumen and the cells are often difficult to distinguish by CLSM because of the presence of numerous opaque concretions and extensive apical microvilli. Problems with confocal laser scanning microscopy of fluorochrome transport of isolated tissues were first noted by Gaertner and Morris (1999). The fluorescent P-gp substrates rhodamine 123 and daunorubicin did not appear to accumulate in the lumen in confocal laser scanning micrographs, in contrast to the results of an earlier study demonstrating lumen to bath ratios $>1$ for the radiolabelled P-gp substrates nicotine and vinblastine (Gaertner et al., 1998). Gaertner and Morris (1999) suggested that the apparent lack of transport of the fluorescent P-gp substrates may have been the result of quenching, reflection or absorption of the exciting and/or emitted light by the tissues and by opaque uric acid crystals present in the lumen. Similarly, it was not possible to clearly visualize fluorochromes and the lumen of many of the cricket Malpighian tubules and all Drosophila tubules examined because of opaque concretions in the cells and/or lumen (Leader and O’Donnell, 2005). Additional concerns are that concretions move in the lumen during fluid secretion, thereby altering the amount of laser light passing through the lumen, and that concretions may be transferred from the cell to the lumen, particularly when fluid secretion is stimulated (Hazelton et al., 2001). By contrast, collection of secreted fluid samples in optically flat glass capillaries and subsequent analysis by CLSM avoids the errors associated with the presence of concretions, which sink to the bottom of the secreted fluid droplet and are not collected in the capillary. Estimates of dye concentration are less affected by variations in cytoplasmic pH and the calibration solutions can be adjusted to resemble closely the pH and ionic
strength of the secreted fluid. Measurement of dye transport by the method of Leader and O'Donnell (2005) also provides a faster, cheaper and more sensitive method than the use of radiolabelled substrates.

The evidence for P-gps in insect Malpighian tubules and the finding that nicotine is transported by a P-gp-like mechanism suggests that P-gps may be involved in excretion of other alkaloids as well. Alkaloids are nitrogenous bases which protect the plants that produce them from attack by insects and other herbivores. In turn, insects have evolved means to reduce alkaloid toxicity. In many species of insect which feed upon alkaloid-rich plants, the Malpighian tubules remove alkaloids such as nicotine, morphine and atropine at high rates (Maddrell and Gardiner, 1976). Nicotine is transported unchanged in *Rhodnius* and *Pieris*, whereas it is transformed into an unidentified substance in *Musca* and *Calliphora*. Transport occurs at high rates, and large lumen-hemolymph concentration differences are established. Transport is saturable \( J_{\text{max}} = 700 \text{ pmol min}^{-1} \text{ tubule, } K_t = 2–3 \text{ mM in } Rhodnius \) and is not coupled to ion transport. Competition studies suggest that a common transport mechanism is used for nicotine, morphine and atropine. Nicotine transport by *Rhodnius* tubules is unaffected by organic anions, such as amaranth and benzyl penicillin, or by the organic anion transport inhibitor, probenecid. In vertebrates, atropine and morphine are also transported by the same renal transport system which handles organic bases such as \( N \)-methylnicotinamide, choline and histamine. By contrast, nicotine and TEA appear to be substrates of different transporters in *Rhodnius* and other insects (Rheault *et al.*, 2005).

8.2.3 Roles of metabolism and excretion in adaptation of Lepidoptera to nicotine-rich diets

Nicotine tolerance is well known for *Ma. sexta* and for several other sphingids of the subfamilies Macroglossinae and Sphinginae, whereas members of the subfamily Smerinthinae are sensitive to poisoning by nicotine (Wink and Theile, 2002). A simple target-site modification as a basis for nicotine tolerance has been ruled out based on the absence of apparent amino acid substitution in the putative nicotine binding site of the f-subunits of nicotinic acetylcholine receptors from nicotine-tolerant and nicotine-sensitive sphingids. This conclusion is supported by the results of feeding experiments which show that larvae of *Ma. sexta* and other sphingids of the Macroglossinae and Sphinginae tolerate not only nicotine, but also many other alkaloids that affect neuroreceptors other than acetylcholine receptors.

When nicotine is injected into larvae of nicotine-tolerant taxa, 80–90% is apparently converted to polar conjugates or degradation products such as nicotine \( N \)-oxide or cotinine and only 10–20% is recovered as free nicotine
(Wink and Theile, 2002). Usually more than 98% of the recoverable alkaloids are found in the faeces and excretion peaks <6 h after injection in tolerant taxa. Rearing larvae of *M. sexta* on a nicotine-rich diet, results in higher nicotine degradation (Snyder et al., 1994) and faster nicotine elimination relative to naïve larvae (Fig. 16). Application of the cytochrome P450 inhibitor SKF 525A (proadifen) reduces the formation of nicotine N-oxide and the rate of alkaloid degradation. Sphingidae are thus able to live on host plants rich in otherwise toxic secondary

![Graph](image)

**FIG. 16** Levels of nicotine and its metabolites in tissues of fifth instar larvae of *Manduca sexta*. The larvae were fed diet containing 0.75% nicotine diet for 24 h (0–24 h) and were then returned to a nicotine-free diet (24–72 h; Snyder et al., 1994, Fig. 1). Reprinted with permission of Elsevier Science.
metabolites by the combined actions of an inducible detoxification mechanism and rapid and inducible excretion.

The high capacity for excretion of alkaloids is clearly evidenced by the results of Self et al. (1964), Maddrell and Gardiner (1976) and Snyder et al. (1994). Competition between morphine, atropine and nicotine indicates a broad specificity of the alkaloid transporter in Malpighian tubules (Maddrell and Gardiner, 1976). It seems likely that the same mechanism would also actively transport the nicotine metabolites nicotine-N-oxide and cotinine-N-oxide, thus resulting in a rapid clearance from the insect (Snyder et al., 1994).

It has been suggested by Snyder et al. (1994) that active transport of alkaloids or more generally high excretion rates are not sufficient, and perhaps not necessary, to explain the adaptation to nicotine in larvae. Malpighian tubules of Pieris brassicae larvae also actively transport nicotine at very high rates (Maddrell and Gardiner, 1976), yet this insect is not tolerant of dietary nicotine (David and Gardiner, 1953). However, the study of Murray et al. (1994) suggests a possible explanation for differences of nicotine toxicity to Manduca and Pieris. P-gp-like transporters in the blood brain barrier of Manduca protect the central nervous system from the effects of nicotine (Murray et al., 1994). These authors suggest that detoxifying enzymes as well as the nicotine pump are likely to account for the metabolic blood–brain barrier to nicotine. In this scenario, then, the sensitivity of Pieris to nicotine may reflect the absence of an efficient blood–brain barrier toward nicotine and its metabolites. The fate of nicotine in Ma. sexta appears to resemble a general mechanism for handling allelochemicals in the Lepidoptera. In other insects that are unaffected by chemicals such as xanthotoxin, precocene II, or α-terthienyl, tolerance also involves rapid metabolism followed by excretion (Haunerland and Bowers, 1985; lyengar et al., 1990; Nitao, 1990).

8.2.4 What other compounds are transported by P-glycoprotein-like mechanisms in insects?

Insects which feed on conifer needles are also exposed to high levels of piperidine alkaloids that may be suitable substrates of P-gps. Spruce and pine needles contain ~0.2% wet weight of several disubstituted piperidine alkaloids, some of which have insecticidal or teratogenic activity. Although some beetles sequester such compounds as a defence against predation by other insects and vertebrates, they are accumulated in the tissues of western spruce budworm larvae (Choristoneura occidentalis) feeding on the needles (Kamm et al., 1998). The frass of the larvae contain the same alkaloids found in the needles and also some more polar metabolites, indicating processing of these compounds and excretion of metabolized and unaltered forms. The absence of sequestration of deterrent or toxic alkaloids is
consistent with the cryptic coloration of spruce budworms, in contrast to aposematic coloring of beetles that synthesize some of the same alkaloids de novo (Kamm et al., 1998).

P-gps have also been implicated in resistance to the toxin α-amanitin, a compound found in mushrooms which are fed on by members of the Drosophila quinaria species group (Begun and Whitley, 2000). Toxicity of α-amanitin is due to interference with RNA polymerase activity. Although D. melanogaster feed on fruit and are unlikely to be exposed to α-amanitin, some strains are resistant to the toxin and genetic studies indicate that polymorphism in a MDR gene (MDR65A) is implicated in resistance. However, DNA sequence analysis does not show any differences in the amino acids coded for by the MDR65A genes of α-amanitin sensitive and resistant strains (Begun and Whitley, 2000). Given that MDR in mammalian cells is often the result of amplification of MDR genes, the latter authors suggest that a mutation in gene regulation may be the basis for α-amanitin resistance in D. melanogaster. The mutation might result in either increased expression of a P-gp or its modification by other molecules such as protein kinases.

P-gps may also be involved in excretion of some types of terpenoids in insects, although direct evidence is lacking. The neem tree is grown in most tropical and sub-tropical areas of the world for shade, reforestation and as a source of natural insecticides and medicines. Toxic effects in insects are produced primarily by azadirachtin, a complex tetranortriterpenoid limonoid from the neem seeds. Terpenoids are a large and structurally diverse class of naturally occurring organic compounds formed from 5-carbon isoprene units which are often arranged to form multicyclic structures. In addition to azadirachtin, terpenoids include menthol, camphor and the cannabinoids and they contribute to the flavours of cinnamon and cloves and the scent of eucalyptus. Terpenoids inhibit transport of digoxin by P-gps in a dose-dependent manner and the IC\textsubscript{50} values in cultured mammalian cells are near or below the value for the commonly used inhibitor verapamil (Yoshida et al., 2006).

Azadirachtin acts as an antifeedant and repellent in insects and it also interferes with normal development by inhibiting the release of prothoracicotropic hormones and allatotropins (Mordue and Blackwell, 1993; Williams and Mansingh, 1996). Tritiated dihydroazadirachtin A, fed in a blood meal to R. prolixus, is absorbed, transported into the hemolymph and then excreted unmetabolized by the organism (Garcia et al., 1989). The level of dihydroazadirachtin A in the hemolymph rises steeply to a peak 24 h after feeding, then declines gradually as it is excreted. The highest amount per milligram tissue is found in the Malpighian tubules and midgut, the lowest in the head and the remaining body. Unchanged labelled compound appears in the urine, consistent with excretion of unmetabolized dihydroazadirachtin A through the Malpighian tubules.
It will be of interest in the future to determine if other terpenoids are transported by insect Malpighian tubules and which transporters are involved.

8.2.5 Transport of cardiac glycosides and miserotoxin

The milkweed bug *O. fasciatus* feeds exclusively on milkweeds, which contain cardiac glycosides as a defence against vertebrate and invertebrate herbivores. Toxicity of cardiac glycosides results from inhibition of the Na\(^+\)/K\(^+\)-ATPase. Cardiac glycosides are composed of a sugar (glycoside) conjugated to an aglycone (steroid) moiety and are divided into two classes: cardenolides, which have an unsaturated butyrolactone ring, and the bufadienolides, which have an \(\alpha\)-pyrone ring. The toad venom bufalin is a well-known bufadienolide and examples of cardenolides are ouabain, digoxin and digitoxin. Milkweed bugs tolerate diets rich in cardiac glycosides by removing these compounds from the hemolymph and sequestering them in the dorsolateral space (Duffey and Scudder, 1974). About 60–95% of the cardenolide is stored in the dorsolateral space, with the remainder found in ventral metathoracic gland secretions of adults or middorsal abdominal gland fluid of larvae. Concentrations of cardenolides in the urine–faeces mixture or hemolymph are low. Sequestration not only permits the bugs to exploit milkweeds as a food source, but provides the warningly colored insect with a chemical defence against predators. For defensive purposes, droplets of dorsolateral space fluid are released from an orifice in each segment.

Physical sequestration of glycosides in special spaces presumably avoids toxicosis that would result from elevated hemolymph levels of these compounds. Sequestration is energy independent (i.e. insensitive to the mitochondrial uncoupler dinitrophenol) and follows first order kinetics (Duffey *et al.*, 1978). The sensitivity of sequestration is quite dramatic, and begins at hemolymph concentrations as low as \(3.5 \times 10^{-7}\) M of ouabain, for example. Moreover, sequestration of one cardenolide is independent of other cardenolides. Phase contrast microscopy reveals the presence of a distinct emulsion phase, and physical sequestration in emulsion particles is consistent with the observed energy independence, first order kinetics and absence of inhibition by presence of related compounds.

Polar cardenolides, such as ouabain, but not non-polar ones such as digitoxin, are sequestered. It appears that both preferential uptake and metabolism may be important in this regard. Digitoxin is converted to at least two more polar cardenolides, for example. Both polar and non-polar cardenolides cross the gut, and a protein carrier may transport them via the hemolymph to the special organs or glands of the dorsolateral space or other body regions.
The sequestration process is enhanced by a reduction in loss of cardiac glycosides through the excretory system (Meredith et al., 1984). Malpighian tubules of *O. fasciatus* consist of two morphologically and physiologically distinct segments, both of which metabolize ouabain. In segment II, the distal segment, ouabain transport into the tubule lumen appears to be a passive phenomenon, since secretion does not occur against a concentration gradient. By contrast, segment I, the proximal segment, reabsorbs fluid and ouabain but not its metabolites. Ouabain reabsorption is independent of fluid secretion, and can occur against concentration gradients as large as 23-fold. Sodium concentration in the reabsorbed fluid (53 mM) is also elevated relative to the bathing saline (28 mM) or whole tubule secretion (9 mM), suggesting that ouabain reabsorption may be linked to sodium cotransport. More than 50% of ouabain is reabsorbed in slowly secreting tubules and as much as 93% is reabsorbed in rapidly secreting tubules. Reabsorption of ouabain by the tubules thus contributes to accumulation of glycosides in the dorsolateral space.

The grasshopper *Zonocerus variegatus* also feeds on plants containing cardiac glycosides (Rafaeli-Bernstein and Mordue, 1978), and in this case excretion rather than sequestration of the ingested toxins is important for survival. The tubules are passively permeable to ouabain, as are those of *Locusta*. However, exposure of *Zonocerus* to dietary ouabain for more than 12 days induces more effective excretion of the glycosides, resulting in lumen/hemolymph concentration gradients of radiolabelled ouabain (or its metabolites) exceeding 3. Although *Locusta* tubules are not poisoned by ouabain, other tissues account for the toxicity of ouabain given to whole animals.

The glycoside miserotoxin (3-nitro-1-propyl-β-D-glucopyranoside) is found in the diet (4.5–5.2% by dry weight) of the melanopline grasshoppers *Melanoplus bivittatus* and *Melanoplus sanguinipes* (Orthoptera: Acrididae). They are able to tolerate high levels of the glycoside, in part by excretion of the intact molecule. Miserotoxin is a causative agent in cattle poisoning on rangelands in the western United States (Johnson et al., 2001), and was originally isolated from *Astragalus miser* var. *oblongifolius* (Leguminosae). It is also found in *As. miser* var. *serotinus* (timber milkvetch), which occurs in the southern interior of British Columbia. Other glycosides of 3-nitropropanol (NPOH) are also present in timber milkvetch. Miserotoxin is rapidly hydrolyzed by microbial enzymes of the rumen, and the aglycone is absorbed and converted to 3-nitropropionic acid (NPA) a potent inhibitor of mitochondrial enzymes essential to respiration. When the aglycone is administered to melanopline grasshoppers, detoxification is achieved by two routes: by oxidation of the aglycone to NPA which is then conjugated with glycine, and by glucosylation of the aglycone to miserotoxin, in each case followed by excretion (Johnson et al., 2001). Karakin, a glucose trimester of NPA is
also found in *Astragalus*, and is detoxified after hydrolysis *in vivo*, conjugation of the NPA to the glycine, serine or glutamate, and excretion of the resulting amides (Majak et al., 1998).

Very little is known of the molecular nature of cardiac glycoside transport by insect cells. In vertebrates, glycosides are transported by P-gps (e.g. Koren et al., 1998). However, a member of the OATP family is responsible, at least in part, for transport of compounds such as ouabain by *Drosophila* Malpighian tubules (Torrie et al., 2004). There are eight OATP genes in *Drosophila*, of which six are abundantly expressed in tubule, and of which three actively accumulate ouabain when expressed in S2 cells. *In vivo*, RNAi against just one of these genes (oatp58Db) reduces ouabain transport in *Drosophila* tubules by 50%. Colocalization of OATP and the Na\(^+\)/K\(^+\)-ATPase in the basolateral membrane of the principal cells of the tubule may explain the relative refractoriness of insect Malpighian tubules toward ouabain. In addition, transepithelial transport in Malpighian tubules is dependent upon the actions of the V-ATPase, as described in Section 2, with the Na\(^+\)/K\(^+\)-ATPase playing an ancillary or subordinate role.

### 8.3 TRANSPORT OF TYPE I ORGANIC ANIONS

Organic anions can be separated into two structural groups, based on molecular weight, net charge and hydrophobicity (Wright and Dantzler, 2004). Type I OAs are small (generally <400 Da) and typically monovalent compounds, such as p-aminohippurate (PAH), probenecid and fluorescein. Type II OAs are bulkier (generally >500 Da), frequently polyvalent and include, calcein, methotrexate and many glutathione, glucuronide and sulfate conjugates. Type I and type II organic anions are often preferentially transported by different transporters in vertebrate renal tissues (Wright and Dantzler, 2004), and this appears to be the case in insect Malpighian tubules as well.

The Malpighian tubules have long been known to accumulate acidic dyes that have been injected into the hemolymph (Lison, 1937). Acidic dyes include carboxylates and sulfonates, and are derivatives of more toxic compounds resulting from metabolism or xenobiotic compounds in plants which form part of the insect’s diet. Insect Malpighian tubules actively excrete both acylamides, such as PAH acid, and sulfonates, such as indigo carmine and amaranth (Maddrell et al., 1974). Transport of acidic dyes such as amaranth and indigo carmine into secreted fluid can be measured spectrophotometrically. For compounds such as PAH and benzyl penicillin K, C\(^{14}\)-radiolabelled compounds are used and the concentrations of the compounds in samples of fluid secreted by Malpighian tubules are measured by liquid scintillation spectrometry. Chromatographic analysis established that neither of these compounds is metabolized by the
Malpighian tubules. Acidic dyes such as indigo carmine are secreted more rapidly at higher bathing saline pH, suggesting that it is the concentration of the anionic form of the dye which determines the rate at which it is transported (Maddrell et al., 1974). Although competition studies using isolated tubules of R. prolixus suggest separate transport systems for acylamides (such as PAH acid) and sulfonates (Maddrell et al., 1974), analysis of fluorescein transport by cockroach tubules suggest one common carrier for organic acids (Bresler et al., 1990).

Transport of type I organic anions such as fluorescein and PAH by insect Malpighian tubules requires the presence of sodium in the bathing saline. Sodium-dependence is also a characteristic of organic acid transport by proximal tubules of the vertebrate kidney. Although concentration gradients exceeding 1000 are accomplished for organic acid transport by the vertebrate proximal tubules, more modest gradients of 30-fold are established by the relatively permeable Malpighian tubules of Calliphora and 300-fold by the less permeable tubules of Rhodnius (Maddrell et al., 1974). Comparisons can also be made on the basis of the $J_{max}/K_t$ ratio, where $J_{max}$ is the maximum rate of transport and $K_t$ the Michaelis–Menten constant, corresponding to the substrate concentration at which transport is half maximal. $J_{max}/K_t$ is the effective rate constant when the transport rate is proportional to the substrate concentration. This proportionality occurs when substrate concentrations are less than $K_t$, as occurs in vivo (Bresler et al., 1990). For transport of the organic anion fluorescein, the value of $J_{max}/K_t$ is 10.6 in locusts, higher than that in rats (2.72), fish (0.9) and turtles (0.3).

The extent to which organic anions are concentrated in the fluid secreted by tubules depends on several factors such as the passive permeability of the tubule wall and the rate of fluid secretion. An important aspect of the paper by Maddrell et al. (1974) is the explanation for differences in the relationship between fluid secretion rate and dye transport by tubules isolated from different species. In the blood-feeding hemipteran R. prolixus and the stick insect Ca. morosus dye transport is independent of fluid secretion rate. By contrast, dye secretion by the tubules of the blowfly Calliphora is very greatly affected by changes in the rate of fluid secretion, being approximately linearly related.

A classic method for estimating the passive loss of dye from lumen to bath over a wide range of fluid secretion rates was developed by Maddrell et al. (1974). In order to vary fluid secretion rate, the tubule lumen is cannulated and perfused. The isolated Malpighian tubule is placed in a droplet of saline under liquid paraffin and a pair of fine forceps mounted on a micromanipulator is brought close to the bathing drop. The distal end of the tubule is then looped round the nearer point of the forceps. The forceps are arranged so that their points open and close in the horizontal plane and the tips of the forceps are held together with a small clamp.
A fluid filled cannula is connected to a motor-driven syringe through thick-walled plastic tubing. The cannula is mounted on a second micromanipulator and positioned so that it is coaxial with the lumen of the tubule. The cannula is then pushed through the wall of the tubule and on down the lumen until the taper of the cannula fills the lumen. Using this technique, the rate of perfusion of the tubule lumen can easily be varied > 40-fold.

Net dye transport by cannulated Malpighian tubules of *Calliphora* is low at low rates of fluid perfusion, increases with the rate of luminal perfusion below about 10 nl min⁻¹ but is not much affected by further increases in the rate of perfusion (Maddrell *et al.*, 1974). When the rate of perfusion is low, measurements of samples of fluid emerging from the open end of the tubule reveal that the concentration of dye in the secreted fluid is many times higher than both that of the bathing fluid and that of the fluid perfused at high rates. By assuming that passive dye efflux is negligible at very high rates of fluid transport, the maximum rate of dye transport can be estimated. By subtracting the net dye efflux measured at lower rates of fluid perfusion from this maximum value, the passive dye efflux at each fluid perfusion rate can be calculated (Fig. 17).

In *Calliphora*, then, active dye secretion into the lumen proceeds at a high rate so that a high concentration of dye is reached in the lumen at all but high rates of fluid secretion. However, because the tubule wall is quite permeable to dye, a large proportion of transported dye diffuses passively back out of the lumen into the bathing fluid. As a result, the net amount of dye transported into the lumen depends very much on the rate of fluid secretion. In *Rhodnius*, by contrast, the tubule walls are much less permeable (Maddrell and Gardiner, 1974). As a result, much higher dye concentration gradients can be maintained across the tubule wall. This has the result that far less of the transported dye leaks back into the bathing fluid and so the net amount transported into the lumen is very little affected by the rate of fluid secretion.

Transport of the organic anion salicylate by the Malpighian tubules and gut of insects has been studied using two ion-selective microelectrode methods described above for analysis of transport of the organic cation TEA (Section 7.1; Figs. 11, 12). Salicylate is of particular interest because of its role as a signal which activates defence genes in plants in response to herbivory or pathogens. Salicylate-selective microelectrodes based on the anion exchanger tridodecylmethylammonium (TDMA) are typically >2000 times more selective toward salicylate than to Cl⁻. In the first method, the concentration of salicylate in fluid droplets secreted by isolated insect Malpighian tubules set up in a Ramsay secretion assay is measured with a salicylate-selective microelectrode and flux across the entire tubule is calculated as the product of secretion rate and secreted droplet salicylate concentration (Fig. 18). In the second method, TDMA-based microelectrodes for salicylate are used in conjunction with the
scanning ion electrode technique (SIET, also known as the self-referencing ion-selective microelectrode technique) for non-invasive spatial and temporal analysis of ion flux (Smith et al., 1994). A salicylate-selective microelectrode is moved between two positions within the unstirred layer adjacent to the surface of a tissue. Salicylate transport by the tissue perturbs the salicylate concentration in the unstirred layer, and the difference in salicylate concentrations measured between the two microelectrode positions can be converted into a corresponding salicylate flux using the Fick equation.

Dose–response curves relating fluid secretion rate for *D. melanogaster* Malpighian tubules, secreted fluid salicylate concentration and salicylate flux to bathing saline salicylate concentration are shown in Fig. 18. The maximum concentration of salicylate in the secreted fluid was \( \sim 7 \text{ mM} \).
FIG. 18 The effects of bathing saline salicylate concentration on (A) fluid secretion rate, (B) secreted fluid salicylate concentration and (C) salicylate flux. Each point is the mean ± S.E.M. for 6–12 tubules. The inset in (B) shows the relationship between bathing saline salicylate concentration (mmol l⁻¹) and S/M, the ratio of salicylate concentration in the secreted fluid to that in the bathing medium. Curves in (B) and (C) were fitted to the Michaelis–Menten equation using non-linear regression analysis. $J_{\text{max}}$ refers to the maximum rate of transport and $K_t$ to the concentration of salicylate required to produce 50% of the maximum flux (O'Donnell and Rheault, 2005, Fig. 7). Reprinted with permission from the Company of Biologists.
At lower bathing saline concentrations (0.0025–0.125 mM) the concentration of salicylate in the secreted fluid was elevated 50- to 100-fold (Fig. 18B, inset). Flux data were fit to the Michaelis–Menten equation to yield the values for $J_{\text{max}}$ (2.72 pmol min$^{-1}$) and $K_t$ (0.046 mM; Fig. 18C). There is good agreement between salicylate flux determined using salicylate-selective microelectrodes and that determined using liquid scintillation counting of radiolabelled salicylate (O’Donnell and Rheault, 2005). However, the microelectrode technique is faster, cheaper and provides greater resolution for small (< 1 nl) samples. However, because some competitive inhibitors of salicylate transport interfere with TDMA-based electrodes, radiolabelled salicylate is used for pharmacological characterization of transport.

Salicylate concentrations in hemolymph samples can also be measured with TDMA-based microelectrodes after injection of salicylate into the hemocoel or after insects are fed salicylate-rich diets. The rate of salicylate secretion by *Drosophila* Malpighian tubules *in vitro* is sufficient to account for the measured rate of decline of salicylate concentration in the hemolymph *in vivo*, suggesting that the Malpighian tubules play a dominant role in elimination of organic anions (O’Donnell and Rheault, 2005).

Measurements with TDMA-based microelectrodes in conjunction with the scanning ion electrode technique show that salicylate is also secreted across the midgut, ileum and rectum when the saline contains 0.5 mM salicylate (Fig. 19). When insects are fed high concentrations of salicylate in the diet there is an efflux of salicylate from gut lumen to hemolymph. SIET measurements confirmed transport of salicylate across the main and distal segments of the Malpighian tubule, but also demonstrated that there is a small efflux of salicylate (from tubule lumen to bath) across the lower Malpighian tubule. The significance of the latter finding is unclear. Perhaps organic anion levels in the hemolymph are regulated through the combined actions of the secretory and reabsorptive segments of the tubule. Taken together, the results suggest that the gut plays a role in limiting absorption of salicylate from the food, and that salicylate which does reach the hemolymph is cleared through active transport by the Malpighian tubules.

The mechanism of transepithelial salicylate transport by *Drosophila* tubules appears similar to that previously described for other organic anions (Bresler et al., 1990; Linton and O’Donnell, 2000). $^{14}$C-salicylate secretion by isolated tubules is competitively inhibited by other carboxylates such as probenecid, unlabelled salicylate, fluorescein and PAH acid. In common with transport of PAH and fluorescein, transepithelial transport of salicylate is strongly sodium-dependent. Transport is nearly abolished in Na$^+$-free saline, and is inhibited also by ouabain or K$^+$-free bathing saline, treatments that will reduce the Na$^+$ gradient across the basolateral membrane through inhibition of the Na$^+$/K$^+$-ATPase (Ruiz-Sanchez and O’Donnell, 2007a). Transepithelial secretion of
Salicylate is accomplished by Malpighian tubules in at least 9 species of insect from 4 orders (Ruiz-Sanchez et al., 2007).

When fluid secretion rate is altered by variations in bathing saline osmolality, 64% of the changes in salicylate transport can be explained on the basis of changes in fluid secretion rate (Ruiz-Sanchez and O’Donnell, 2007a). The increases in transepithelial salicylate transport produced by cAMP, cGMP or leucokinin are most easily explained as a consequence of the effects of these compounds on fluid secretion rate. Increases in fluid secretion rate presumably minimize passive diffusive backflux of salicylate from lumen to bath, and this effect is most apparent when the concentration of salicylate in the bathing saline is high. When the bathing saline concentration of salicylate is below the value of the Michaelis–Menten constant ($K_t$) the concentration in the lumen is also lower. Under these conditions the gradient for diffusive backflux is reduced and there is therefore no relationship between dye transport and fluid secretion rate (Ruiz-Sanchez and O’Donnell, 2007a).

Although renal transport of type I organic anions is $Na^+$-dependent in both vertebrates and insects, the mechanism of uptake of the organic anion...
salicylate across the basolateral membrane of the Malpighian tubules of *D. melanogaster* nonetheless differs fundamentally from mechanisms proposed for vertebrate renal tubules (Ruiz-Sanchez and O’Donnell, 2006). In vertebrate kidneys the uptake process is composed of three steps arranged in parallel. The Na⁺/K⁺-ATPase maintains an inwardly directed (blood to cell) Na⁺ gradient which drives a Na⁺-dicarboxylate cotransporter (Na⁺-DC), thereby establishing an outwardly directed (cell to blood) dicarboxylate (α-keto acid) gradient. In turn, the dicarboxylate gradient is utilized by a dicarboxylate/organic anion exchanger to move the organic anion into the cell. This cascade of events indirectly links organic anion transport to metabolic energy and Na⁺ transport. The activity of the organic anion uptake mechanisms in the vertebrate kidney is characteristically influenced by α-keto acids; preloading cells with these compounds (trans-configuration) stimulates uptake of organic anions, whereas simultaneous addition of high concentration of α-keto acids to the bathing medium (cis-configuration) inhibits uptake of organic anions. By contrast, preloading Malpighian tubule cells with the α-keto acid glutarate does not trans-stimulate salicylate or fluorescein uptake, nor is salicylate uptake cis-inhibited by high concentrations of various α-keto acids (α-ketoglutaric acid, glutaric acid, succinic acid and citric acid) in the saline (Linton and O’Donnell, 2000; Ruiz-Sanchez and O’Donnell, 2006). Together, these results suggest that Na⁺-dependent organic anion transport by the Malpighian tubules of *D. melanogaster* is not mediated by an OAT1-like transporter similar to that of the vertebrate kidney. The absence of stimulation of salicylate uptake by acidic saline also militates against a role for a pH-dependent monocarboxylate transporter (MCT), similar to the H⁺:carboxylate cotransporter of the vertebrate gut. Current models propose a Na⁺:monocarboxylate transporter for salicylate uptake across the basolateral membrane of *D. melanogaster* Malpighian tubules (Fig. 6). The transport system is strongly inhibited by monocarboxylic acids with a ring structure and it is not dependent on basolateral membrane potential (Ruiz-Sanchez and O’Donnell, 2006). Inhibition of salicylate uptake by the ‘classical’ MCT inhibitor, α-cyano-4-hydroxycinnamic acid suggests that salicylate uptake across the basolateral membrane of the *D. melanogaster* Malpighian tubules may resemble a Na⁺-dependent active monocarboxylate transport system previously reported in the brush-border membrane of rat enterocytes (Storelli et al., 1980).

The monocarboxylate transporter in the basolateral membrane of the Malpighian tubule may contribute to elimination of natural or synthetic toxins from the hemolymph. For example, the plant secondary metabolite jasmonic acid, which protects plants against pathogens and phytophagous insects (Li et al., 2002), would be a suitable substrate for the transport system that handles salicylate. It is also worth noting that 3-phenoxynbenzoic acid, an active metabolite of the insecticide deltamethrin, is
similar in structure to other substrates of the salicylate transporter (Ruiz-Sanchez and O’Donnell, 2006). One source of endogenous organic anions is through metabolism of the amino acid tryptophan. Three fluorescing materials excreta from the American cockroach, _P. americana_ (L.), have been identified as kynurenic, xanthurenic and 8-hydroxyquinoidal acids (Mullins and Cochran, 1973b). Concentrations of these compounds in the excreta increase as dietary nitrogen increases, and can account for 1–3% of total nitrogen excretion. Metabolism of tryptophan through the pathway involving kynurenine and 3-hydroxykynurenine leads to the synthesis of ommochrome eye pigments. In addition, kynurenine can be converted to a number of intermediates some of which are excreted.

8.3.1 Regulation of organic anion transport by Malpighian tubules

Transport of organic anions such as PAH and amaranth by the Malpighian tubules of _R. prolixus_ varies with the insect’s physiology (Maddrell and Gardiner, 1975). PAH transport declines gradually as starved insects age, but increases dramatically by more than 20-fold within 1–2 days after a blood-meal (Fig. 9). A similar pattern is seen for urate transport, as described above (Fig. 9). Thereafter the rate declines with a half-time of about 10–15 days. Ingestion of a meal of glucose-free Ringer’s solution does not result in an increase in the rate of PAH transport, which suggests that the normal induction of transport is not controlled by a hormone released in response to the same stimuli that elicit release of the diuretic and prothoracicotropic hormones. Ingestion of blood is not the only effective stimulus for induction of PAH excretion. Meals containing protein in the form of blood plasma alone, a suspension of red blood cells in saline, serum albumin, casein or milk also result in an increase in PAH transport several days later. However, there is no increase in rates of excretion of organic anions after a single injection of 0.1 μmol of PAH into the hemolymph. Ingestion of a protein-rich meal is thus sufficient to induce an accelerated transport of organic anions and this induction is not controlled by a hormone released in response to abdominal distension. Rather, induction depends instead on the sustained presence in the hemolymph of some product of digestion of the meal. As noted by Maddrell and Gardiner (1975), it is unclear why the transport of organic anions such as PAH should be regulated in this way, whereas the transport of the alkaloids nicotine and atropine is maintained irrespective of the insect’s nutritional state.

Changes in organic anion transport in response to dietary loading with organic anions have also been documented in third instar larvae of _D. melanogaster_ (Ruiz-Sanchez and O’Donnell, 2007b). Larvae chronically exposed to dietary salicylate show lower levels of salicylate in the hemolymph and increased elimination of salicylate after feeding on
salicylate-enriched diet. Exposure to dietary salicylate also leads to a fivefold increase in salicylate excretion by isolated Malpighian tubules. This increase is accompanied by an increase of more than threefold in Malpighian tubule fluid secretion rate. The increases in fluid and salicylate secretion do not appear to reflect increases in the basal levels of the second messengers cAMP and Ca^{2+} which increase apical proton pump activity and transepithelial Cl^- permeability, respectively. Instead, increases in the unstimulated level of fluid secretion in larvae fed a salicylate-rich diet may indicate increased numbers of transporters (e.g. H^+-ATPase) in the tubule cells.

An important consequence of the increase in fluid secretion rate is that the concentration of salicylate in the tubule lumen is maintained at relatively low levels. Diffusive backflux of salicylate from the tubule lumen to the hemolymph is thus minimized. Transepithelial salicylate transport increases with fluid secretion rate irrespective of whether fluid secretion is increased by intracellular second messengers or changes in bathing saline osmolality (Ruiz-Sanchez and O’Donnell, 2007a). Taken together the results indicate that exposure to salicylate in the diet leads to alterations in renal function that enhance clearance of the toxin from the insect’s hemolymph. Moreover, the increase in fluid secretion rate will enhance the excretion of other toxins.

8.3.2 Excretion of insecticides and their metabolites

One component of resistance to the organophosphate insecticide leptophos in the Egyptian cotton leafworm Spodoptera littoralis (Boisd.) is more rapid excretion of both the unchanged insecticide and its metabolite leptophos phenol (Ben-Aziz et al., 1976; Riskallah and Esaac, 1982). The half-time (t_{1/2}) for disappearance of leptophos in larvae of a resistant strain (t_{1/2} = 2.5 h) is > 8 times shorter than in a larvae of a susceptible strain (t_{1/2} = 21.5 hr). Excretion of leptophos metabolites is also more rapid in resistant than in susceptible strains. The major degradation product is the free phenol and its amount is relatively higher in resistant (17–22%) than in susceptible (5–15%) larvae. The rate of excretion of the phenol is fourfold higher in resistant than in susceptible larvae. Esterase activity is consistently higher in resistant strains of the leafworm, enabling them to degrade leptophos at a higher rate than in susceptible strains.

Control of organophosphate-resistant strains of the diamondback moth Plutella xylostella necessitates the use of compounds such as malathion at doses several thousand fold higher than those required elimination of susceptible strains. Resistance to malathion involves three changes in the physiology of the insect. Although malathion penetrates the cuticle of insects from resistant strains more rapidly, it is also metabolized and excreted more rapidly, relative to insects from susceptible strains.
As for Sp. littoralis, metabolic degradation of a toxin is thus coupled to enhanced excretion of the degradation products.

Enhanced excretion of insecticides has also been described for other species. Resistant larvae of the Egyptian cotton leafworm larvae also excrete carbaryl (Hanna and Atallah, 1971). Similarly, resistant houseflies excrete the parathion metabolite paraoxon (Plapp et al., 1961) and resistant larvae of the tobacco budworm houseflies excrete large amounts injected endrin (Polles and Vinson, 1972). Resistant larvae in each case can tolerate higher doses of insecticidal chemicals by the simple expedient of excreting the toxic chemical intact.

Although most of the studies of excretion of insecticide metabolites have been based on in vivo experiments and collection of the excreta for analysis, recent in vitro studies have suggested a specific role for the Malpighian tubules. Analysis of the transport of fluorescein by the Malpighian tubules of the cricket Ac. domesticus suggests that organic anion transporters in the tubules play a role in excretion of some malathion metabolites (Neufeld et al., 2005). Esterase activity converts malathion into malathion monocarboxylic acid (MMA) which competitively inhibits transport of the organic anion fluorescein by the tubules. Inhibition of accumulation of MMA by the tubules by probenecid, an archetypal organic anion transport inhibitor, was verified in this study by high performance liquid chromatography (HPLC). Transport of MMA by the tubules suggests that transport processes may confer resistance to xenobiotics in insects (Neufeld et al., 2005).

8.4 TRANSPORT OF TYPE II ORGANIC ANIONS BY MULTIDRUG RESISTANCE-ASSOCIATED PROTEINS

The MRPs are another group of ATP-binding cassette (ABC) transporters implicated in excretion of toxins by insects. As for P-gps, MRPs are known for their role in conferring resistance to antineoplastic agents in humans. Whereas P-gp substrates are primarily cationic or neutral compounds, MRP substrates are more commonly anions, although some neutral compounds, cations and even heavy metals such as arsenic are also transported by members of this family (Leslie et al., 2005; Deeley et al., 2006).

MRPs in insects were first identified in the Malpighian tubules of the cockroach P. americana and the cricket Ac. domesticus by Karnaky et al. (2000, 2001, 2003). Malpighian tubules of both species transport the fluorescent MRP2 substrate Texas Red (sulforhodamine 101) through the cells and into the lumen. Analysis of isolated Malpighian tubules using confocal microscopy showed that the concentration of the fluorochrome in the lumen is dependent upon metabolism and is inhibited by the
non-fluorescent MRP2 substrate chlorodinitrobenzene, which does not inhibit lumenal accumulation of the smaller, more hydrophilic organic anion chlorophenol red. Transport is unaffected by addition of the organic anion transporter substrate PAH acid, the organic cation transporter substrate TEA chloride, or by the P-gp inhibitor verapamil (Karnaky et al., 2001). Luminal cells are immunopositive for an antibody to a sequence of rat MRP2, further suggesting that a MRP2-like transporter is present in Malpighian tubules.

Transepithelial flux of the MRP2 substrate Texas Red across the Malpighian tubules of the fruit fly, D. melanogaster and the cricket, Te. commodus has been measured using confocal laser scanning microscopy (Leader and O’Donnell, 2005). Texas Red accumulates in the cells and lumen of cricket tubules at concentrations up to 20 and 40 times, respectively, those in the lumen. Transepithelial flux of Texas Red is calculated by multiplying the dye concentration (µM), as measured by CLSM of secreted fluid samples collected in rectangular glass capillaries, by the fluid secretion rate (nl min⁻¹) measured in the Ramsay assay. Estimates based on the in vitro fluxes suggest that Drosophila tubules could clear the hemolymph of 20µM Texas Red within 6 min. Transport of Texas Red is inhibited by the MRP2 inhibitors MK-571 and probenecid. However, Texas Red transport is not reduced in Na⁺-free saline, suggesting that probenecid is not acting on the Na⁺-dependent process implicated in transepithelial transport of smaller organic anions such as fluorescein (Linton and O’Donnell, 2000) and salicylate (Ruiz-Sanchez and O’Donnell, 2006). Analysis of concentration-response curves for secretion of Texas Red by Malpighian tubules indicates that the transport mechanism is characterized by higher affinity (i.e. lower $K_t$) but lower capacity ($J_{max}$) than the transporters utilized for Na⁺-dependent-transport of fluorescein and salicylate. The value of $K_t$ for Texas Red (7.1 µM) is fivefold lower than that for salicylate (36 µM), whereas the $J_{max}$ for Texas Red (118 fmol min⁻¹ tubule⁻¹) is 23-fold lower than that for salicylate (2.72 pmol min⁻¹ tubule⁻¹).

The effects of stimulants of fluid secretion on net transepithelial transport of P-gp and MRP2 substrates by Drosophila Malpighian tubules have been examined by O’Donnell and Leader (2006). Stimulation of fluid secretion might alter the transport of an organic ion by altering the transepithelial potential. Both cAMP and cGMP produce a lumen positive potential in Drosophila tubules, thus tending to favour secretion of anions such as Texas Red. Tyramine augments fluid secretion through a Ca²⁺-dependent increase in transepithelial Cl⁻ permeability, reducing the magnitude of the lumen positive transepithelial potential. Tyramine might thus enhance the transport of type II organic cations such as daunorubicin by reducing the opposing electrical gradient. Increases in the levels of intracellular second messengers in response to diuretic factors might also
increase the levels of protein kinases (e.g. protein kinase C). Studies of vertebrate tissues have demonstrated the modulation of P-gps and MRP2 by protein kinase C, raising the possibility of some form of MRP2 modulation by protein kinases in insect Malpighian tubules.

Alternatively, stimulants of fluid secretion may alter transepithelial flux of MRP2 substrates by indirect means. An increase in fluid secretion rate will tend to dilute the MRP2 substrate in the lumen, thereby reducing the tendency for diffusive backflux of the dye from the lumen to the bathing medium. Increases in fluid secretion rate have the greatest effect on solutes that are actively transported (i.e. when the solute concentration in the lumen is much greater than that in the bathing medium) and to which the tubule is relatively permeable (Maddrell et al., 1974). For the MRP2 substrate Texas Red, dye transport is greatly affected by fluid secretion rate (O’Donnell and Leader, 2006). When the rate of fluid secretion is stimulated by cAMP or tyramine, transport of Texas Red is increased, and these effects are more pronounced when the bathing saline concentration of the dye is near or above the concentration required for half-maximal transport ($K_t$). Regression analysis indicates that 57–80% of the change in dye flux is attributable to changes in fluid secretion rate produced by the stimulants. Similarly, when the osmolality of saline bathing cAMP-stimulated tubules is increased 22% by addition of sucrose, the reduction in fluid secretion rate accounts for 88% of the reduction in Texas Red flux. Taken together, these results suggest that increases in fluid secretion rate minimize diffusive backflux, and that stimulants of fluid secretion thus increase transepithelial fluxes of Texas Red indirectly. A similar relationship between fluid secretion rate and solute flux has been described for Na$^+$-dependent transport of the type I organic anion salicylate (Ruiz-Sanchez and O’Donnell, 2007a) in Section 7.3.1.

### 8.4.1 Metabolism and excretion of acetylchromenes

Precocenes, natural 2,2-dimethyl-chromene constituents of the plant *Ageratum houstonium*, destroy the corpora allata (the source of juvenile hormone) in sensitive insect species and thereby cause precocious moulting. However, precocenes are atypical of the > 30 more widely distributed plant chromenes in which an acetyl group is linked to the aromatic ring. The insecticidal compound Encecalin (from *Encelia* spp., Asteraceae) is one of the commonest acetylchromenes. Only 20–40% of the applied dose of encecalin and demethylencecalin (DME) is recovered in the frass of treated grasshoppers (*Me. sanguinipes*) after 48 h, almost all of which is excreted within 24 h of administration. Both parent compounds and metabolites of acetylchromenes and precocene II are excreted and the metabolites are the result of phase 1 reactions (Isman et al., 1987). Other studies of precocene II metabolism in insects report that the major route of precocene
metabolism in several species of insects is via demethylation followed by rapid glucosylation. Much of the radiolabel in hemolymph of the African armyworm *Spodoptera exempta* following treatment with labelled pre-coccone II is also in the form of conjugates. Acetylchromene metabolites may occur as conjugates in the hemolymph and may be hydrolyzed subsequently with recovery of the glucose (or other conjugating group) during excretion by the Malpighian tubules (Isman et al., 1987). Given the metabolism in phase I reactions and conjugation of acetylchromene metabolites, it will be of interest to determine whether P-gps or MRPs in the Malpighian tubules are involved in excretion of these compounds. In other systems, P-gps often act on the product of phase I reactions, such as those mediated by P450 enzymes. Similarly, there is evidence that MRPs act on conjugated compounds produced through the actions of glutathione-S-transferases, as described in Section 8.5, below.

8.4.2 Excretion of sesquiterpene lactones and their metabolites

It will also be of interest in future studies to examine the role of transporters such as P-gps and MRP2 in the excretion of a number of plant-derived insecticides such as the sesquiterpene lactones. The leaves of many trees in the family Magnoliaceae, such as the tulip tree (*Liriodendron tulipifera* L.) and sweet bay (*Magnolia virginiana* L.), contain a variety of toxic constituents, including sesquiterpene lactones such as parthenolide. Sesquiterpene lactones are feeding deterrents and their toxicity is due to the presence of various electrophilic groups, such as the \( \alpha \)-methylene-\( \gamma \)-lactone. The mechanism of toxicity is relatively non-specific because these electrophilic groups react with electron-rich atoms such as sulfur and oxygen in proteins and nucleic acids. Parthenolide also causes oxidative stress in cells, leading to gut lesions and apoptosis, and this stress may be exacerbated by a concomitant depletion of the antioxidant molecule glutathione.

Caterpillars of the tiger swallowtail butterfly *Papilio glaucus* consume large amounts of parthenolide when feeding on the leaves of host plants such as tulip trees and sweet bay. They avoid the toxic effects of diets which may contain as much as 0.5% parthenolide by excreting intact sesquiterpene lactones and sesquiterpene lactone metabolites (Frankfater et al., 2005). Excretion of unaltered parthenolide from the diet indicates that either there is a barrier to absorption of allelochemicals across the gut or that there is absorption into the hemolymph followed by highly efficient excretion by the Malpighian tubules. Parthenolide is modified in the gut by reduction of an exocyclic methylene group, possibly by an (NADPH)-dependent reductase, such as an \( \alpha,\beta \)-ketoalkene double bond reductase, and also by addition of a hydroxyl group, presumably in a reaction catalyzed by a cytochrome P450 enzyme. These reactions may be
accomplished either by gut microbes or by enzymes present in the midgut or Malpighian tubules. Sesquiterpenes are transported by P-gps in mammalian cells, and the actions of P450 enzymes may render them as more suitable substrates (Barthomeuf et al., 2006). Alternatively, the increase in polarity of the metabolite as a consequence of the addition of the hydroxyl group may promote subsequent conjugation to sugars or glutathione and thereby lead to enhanced excretion by transporters such as the MRP2. It is worth noting in this context that high levels of MRP2-like gene expression are found in the gut of another lepidopteran, the cabbage looper Tr. ni (R. Labbe and D. C Donly, personal communication).

8.4.3 Secretion of cyclic nucleotides by Malpighian tubules

Organic anion transport has also been implicated in the secretion of cyclic nucleotides by insect Malpighian tubules. Cyclic AMP is secreted by isolated tubules of R. prolixus in response to stimulation with 5-HT and the compound is also excreted in the urine during diuresis in vivo (Montoreano et al., 1990). The cells of the Malpighian tubules of Drosophila also take up cAMP and cGMP from the bath (Riegel et al., 1998) and uptake is inhibited by the organic anion transport inhibitor probenecid and by other organic anions such as PAH and phenolsulfonphalein, but not by ouabain or the weak base quinacrine (Riegel et al., 1999). Cyclic GMP appears in the fluid secreted by Drosophila tubules (Day et al., 2006), indicating that there is also transport of it and presumably other cyclic nucleotides from cell to lumen. The uptake of cyclic nucleotides across the basolateral membrane may be mediated by the same Na+-dependent mechanism which transports PAH and salicylate. The mechanism of transport across the apical membrane into the lumen is unknown, although it is worth noting that members of the MRP family transport cyclic nucleotides in mammalian cells (Deeley et al., 2006).

8.5 Interaction of excretory mechanisms with phases I and II detoxification mechanisms

Detoxification through phases I and II mechanisms is commonly implicated in insecticide resistance. Elevated levels of the phase II enzyme glutathione-S-transferase are associated with resistance to all major classes of insecticides (Ranson and Hemingway, 2005). Similarly, the P450 monoxygenases are phase I enzymes which are well known for their roles in metabolism of xenobiotics, including natural and synthetic insecticides, to less toxic metabolites. However, as noted by Feyereisen (1999), “P450 enzymes will metabolize xenobiotics irrespective of the eventual fate of the metabolites, and there are many cases where P450 enzymes act as anything but detoxification enzymes.” Indeed, P450 chemistry has been exploited to
produce bioactivated insecticides. Metabolism of pesticides to more toxic compounds through the action of P450 enzymes is the basis for the efficiency of the phosphorothioate insecticides (conversion of P = S to P = O) and the cyclodiene insecticides (aldrin and heptachlor to their epoxides).

There are several reasons to suspect that phase I and phase II pathways may interact with phase III elimination pathways in insects. Several cytochrome P450 genes and glutathione-S-transferase genes are enriched in the Malpighian tubules of *Drosophila* (Dow and Davies, 2006). It is also well known that P-gps and P450 enzymes share many substrates (Bard, 2000; Abu-Qare *et al*., 2003). A moderately hydrophobic xenobiotic may therefore be transported out of cells unmodified or after hydroxylation via P450 enzymes (Fig. 20). Similarly, xenobiotic resistance mediated by GST may require removal of the glutathione conjugate via MRP2 (e.g. Smitherman *et al*., 2004). Some xenobiotics may therefore be transported through apical P-glycoprotein (P-gp) and multidrug resistance-associated protein (MRP), the Phase I detoxification enzymes, cytochromes P450 (P450) and the phase II detoxification enzyme glutathione-S-transferase (GST). It is suggested that the transporters reside in the apical membrane of the Malpighian tubule and gut. A moderately hydrophobic toxin (X) may be transported into the lumen by P-gp. Alternatively, the compound may be modified by P450 enzymes and either transported into the lumen by P-gp, or further modified by conjugation with glutathione (GSH) through the actions of GST and then transported into the lumen by MRP. Redrawn and modified from Fig. 3 of Bard (2000). Reprinted with permission of Elsevier Science.
by MRP2 either unmodified or subsequent to both hydroxylation by P450 enzymes and conjugation by GST (Fig. 20).

The esterases are another group of enzymes that play important roles in insecticide resistance (Hemingway and Ranson, 2000). Many examples of resistance to organophosphorous insecticides are based on increased detoxification of the insecticide by esterases. In the mosquito *Culex pipiens* resistance involves overproduction of esterases which have a high binding affinity for insecticides but a low rate of hydrolysis. As a result, the insecticide is sequestered and unable to reach the target enzyme acetylcholinesterase. Overproduction of the esterases involves either increased gene expression or, more commonly, gene amplification. A common feature of resistant strains of *Culex* is overproduction of esterases in the alimentary canal and the Malpighian tubules (Pasteur et al., 2001). Esterases in the Malpighian tubules are aptly located both to aid detoxification of the hemolymph and to facilitate subsequent excretion of the products of esterase activity. Esterases in the alimentary canal presumably limit transfer of active insecticide from the gut to the hemolymph, from where it can gain access to the target enzymes of the nervous system.

Given the common finding that phase I and phase II enzymes are induced by exposure to appropriate substrates, it will be of interest to see if there is an associated up regulation of phase III elimination pathways. The effects of the plant-produced signal salicylate are of interest in this context. Salicylate activates plant defence genes after herbivory or pathogenic attack. The corn earworm can ‘eavesdrop’ on this signal, so that by the time salicylate activates production of toxic allelochemicals in the corn plants, the insect’s own P450 enzymes have been activated (Li et al., 2002). However, chronic exposure to salicylate is also known to increase fluid secretion rate and salicylate secretion by isolated Malpighian tubules of *D. melanogaster* (Ruiz-Sanchez and O’Donnell, 2007b). As a result, salicylate is more rapidly cleared from the hemolymph. Given that phase I and phase II pathways modify the chemical properties of molecules to make them more suitable for elimination by transporters such as P-gps and/or MRPs, it would seem beneficial for insects to increase the capacity of their phase III pathways during exposure to xenobiotics. Exploration of the links between the detoxification (phase I and II) and elimination pathways (phase III) may thus lead to new insights into understanding insecticide resistance in pest species. In this context, it is worth noting the study by Yang et al. (2007) of insecticide metabolism in *Drosophila*. This study shows that cytochrome P450 and GST loci implicated in insecticide resistance are most abundantly expressed in the Malpighian tubule. The importance of the tubule in detoxification of insecticides is reinforced by the finding that flies become more susceptible to DDT intoxication if RNAi against just one P450, cyp6g1, is expressed in only the tubule cells of
transgenic Drosophila. This implies that the tubule can be the limiting tissue in insecticide metabolism.

9 Future directions

9.1 THE IMPACT OF GENOME SEQUENCING PROJECTS

The sequencing of the Drosophila genome has had a dramatic impact upon our understanding of ion transport mechanisms (Section 3.2), control of ion transport by extracellular and intracellular messengers and the great diversity of solute transporters in the Drosophila tubule (Wang et al., 2004). Sequencing of the genomes for other insect species will allow expression of the genes for specific ion channels, pumps, cotransporters and exchangers to be studied in the absence of other transporters. By 2007, whole-genome sequencing projects have been completed for 12 insect lineages (Zdobnov and Bork, 2007): the honeybee (Apis mellifera) representing order Hymenoptera, the red flour beetle (Tribolium castaneum) from the Coleoptera, the silkworm moth (B. mori) from the Lepidoptera and two mosquitoes (An. gambiae and Ae. aegypti) and seven fly species (D. melanogaster, Drosophila erecta, Drosophila ananassae, Drosophila pseudoobscura, Drosophila mojavensis, Drosophila virilis and Drosophila grimshawi) from the Diptera. Many important advances in our understanding of insect excretory mechanisms will be based on heterologous expression of the genes for specific transporters from the gut or Malpighian tubules in cell lines or Xenopus oocytes, so that the transporter may be characterized in isolation from other transporters present in the whole epithelium. Studies of organic ion transporters using heterologous expression may be particularly important because Malpighian tubule cells, like other renal epithelia, possess multiple transport systems with overlapping substrate specificities. An alternative approach is the use of gene-knockouts or the selective reduction of the expression of an individual gene through the use of RNAi techniques (e.g. Torrie et al., 2004). The use of tissue-specific DNA microarrays, such as that developed for the D. melanogaster Malpighian tubule (Wang et al., 2004), will be of great use in looking at the functions of the excretory system in responding to stresses such as high levels of toxins in the diet, and the interactions between membrane transporters and phase I and phase II detoxification pathways.

9.2 CHEMICAL ECOLOGY AND INSECT PHYSIOLOGY

It also appears likely that our understanding of insect excretory system will be aided by advances in chemical ecology. Improvements in separation and identification of organic compounds in the diet or excreta of insects
will aid the design of in vivo and in vitro experiments designed to reveal the sites and mechanisms of excretion. For example, compounds identified from the frass of insects feeding on particular plants may be applied to isolated Malpighian tubules in vitro with a view to determining if the compounds are secreted by the tubules, and, if so, which transporters are involved. More detailed understanding of insect excretory mechanisms through these types of experiments will aid the development of novel and environmentally-benign control measures for agricultural pests and disease vectors.

9.3 ION CHANNELS AND INSECT EXCRETORY MECHANISMS

Further application of electrophysiological techniques, including patch clamping, double-barrelled intracellular ion-selective microelectrodes and the scanning ion-selective electrode technique, will extend our understanding of the mechanisms of ion transport in insect excretory tissues. Patch clamp studies to date have examined the role of Cl⁻ channels in the tubules of dipterans (O’Donnell et al., 1998; O’Connor and Beyenbach, 2001). These studies have dealt primarily with channels in the apical membrane, which can be exposed by microdissection techniques (Section 3.2.4). Patch clamping of the basolateral membrane requires movement of the basement membrane. A clever mechanical technique for removal of the basement membrane from the tubules of Onymacris rugatipennis has been described by Nicolson et al. (1991) and Nicolson and Isaacson (1990). The basement membrane of the tubules of R. prolixus and D. melanogaster has been removed enzymatically using elastase (Levinson and Bradley, 1984). Single channel currents have been successfully recorded form tubules of both species treated with elastase (O’Donnell, unpublished observations), but the nature and characteristics of the channels have yet to be determined.

There is a paucity of information on the characteristics and regulation of K⁺ channels in the basement membrane, in spite of the importance of K⁺ channels to models of transepithelial ion secretion for Malpighian tubules of several species. An important study by Wu and Beyenbach (2003) has demonstrated the control of basolateral K⁺ channels and also the apical V-type H⁺-ATPase by ATP. When intracellular ATP levels decline to 10% of the control level, the V-ATPase is inhibited and K⁺ movement from cell to lumen is prevented. Basolateral K⁺ channels close at the same time, preventing exchange of K⁺ across the basolateral membrane. These changes facilitate intracellular K⁺ homeostasis when metabolism is inhibited, and allow the epithelium to re-initiate transepithelial K⁺ transport when ATP levels are restored.

Intracellular recording techniques and secretion assays suggest the presence of ATP-regulated potassium channels (Kₐ₅P) in the basolateral
membranes of the tubules of the mealworm *Te. molitor* and the fruit fly *D. melanogaster* (Wiehart *et al*., 2003; Evans *et al*., 2005). The K<sub>ATP</sub> channels are in addition to the Ba<sup>2+</sup>-sensitive K<sup>+</sup> channels in the membrane (Wiehart *et al*., 2003). Patch clamp studies, in particular single channel recordings, will be of great use in determining the types and of K<sup>+</sup> channels present and their contributions to parameters such as the basolateral membrane potential and net transepithelial secretion as well as homeostatic functions such as cell volume regulation. Similarly, patch clamping will be of use in examining the characteristics and regulation of the amiloride-sensitive and cAMP-induced Na<sup>+</sup> conductance in the principal cells of *Ae. aegypti* Malpighian tubule (Sawyer and Beyenbach, 1985; Beyenbach and Masia, 2002) as described in Section 3.2.3, and in determining the extent of similarity between this conductance and that associated with the epithelial Na<sup>+</sup> channel (EnaC) in vertebrate cells. Ion channels for K<sup>+</sup> and Cl<sup>-</sup> may also be involved in the reabsorption of these ions by the proximal (lower) Malpighian tubule of *R. prolifus* (Haley and O’Donnell, 1997; Haley *et al*., 1997) and *D. melanogaster* (O’Donnell and Maddrell, 1995), as described in Section 4.1. In addition to helping to understand the mechanisms of transepithelial ion secretion and reabsorption and the regulation of channels by second messengers and kinases, patch clamp studies will be of use in identification of channel blockers which may aid development of novel control measures for pest species of insect.

9.4 DIURETIC AND ANTIDIURETIC AGENTS AS CANDIDATES FOR DEVELOPMENT OF NOVEL INSECTICIDES

Control of Malpighian tubules and the insect hindgut by peptides has been discussed in Sections 3 and 4, respectively. One reason for the intense interest in studying insect peptides is the possibility of developing species-specific insecticides. Unfortunately, peptides such as insect kinins were long considered unsuitable as pest control agents because they are susceptible to both exo- and endopeptidases in the hemolymph and gut of the insect. However, when critical residues in the C-terminal pentapeptide core of insect kinins are replaced with β-amino acids or their β-homo-amino acid counterparts, potent diuretic activity is retained and the compounds are resistant to the endopeptidases which deactivate the natural insect kinins (Zubrzak *et al*., 2007). Similarly, replacement of the N-terminal lysine of a kinin with aminohexanoic acid provides enhanced resistance to aminopeptidase attack and retains biological activity (Nachman *et al*., 2002). Peptidase-resistant insect kinin analogs inhibit weight gain in larvae of the agriculturally destructive corn earworm moth (*ibid*), suggesting that peptidase resistant analogs of the insect kinins, may aid development of new, environmentally benign control measures for pest species.
The strategies for making kinins insensitive to breakdown by peptidases may in future be applied to other insect peptides, such as those controlling the hindgut (Section 4.2).

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1 Introduction

Organisms inhabit a heterogeneous world in which resources are distributed patchily, vary in size or quality over time, and trigger the interest of several competing species. In addition, the physical properties of
the environment – such as the air wind-flow for flying species or the substrate granularity for walking ones – determine the energetic costs, as well as the time investment associated with the searching and the exploitation of food resources. Hence, during its entire life, each individual forager has to choose from a range of alternatives: it has to decide not only which food type to exploit, but also which part of its home-range to explore, which foraging risk to face or which path to follow. Decision-making is even more complex for animals living in social systems (Krause and Ruxton, 2002). Indeed, the decision of each individual is closely related to the behavior of the other group members. Such social influence on individual decision-making is essential to group-living animals, for which cooperation plays a key role by enhancing fitness. It is a prerequisite for the emergence of coordinated behavior; for example, the reaching of a consensus about moving in a common direction, or the collective choice of exploiting a single food patch. Situations in which group members choose between two or more mutually exclusive actions with the specific aim of reaching a consensus are usually termed as “collective decision-making” (Deneubourg and Goss, 1989; Conradt and Roper, 2005; Couzin et al., 2005; Sumpter, 2006; Dyer et al., 2008). Examples of collective decisions were demonstrated in several different taxa and contexts such as the choice of a common feeding place in birds (see e.g. Perusse and Lefebvre, 1985) or the choice of a common moving direction in fish schools (e.g. Couzin et al., 2005) and in mammal troops (e.g. Kummer, 1968; Boinski and Garber, 2000). Put in a historical perspective, the study of collective decision-making find its roots in political sciences and several theories have been built since Condorcet’s (1785) formulation of the voting paradox. These theories have shown how individual preferences may lead to quite different social choices depending on the underlying voting procedures (Gaertner, 2001). Even though political sciences and life sciences may seek to answer similar questions – and often share the same vocabulary (List, 2004) – when they talk about decision-making, collective choices made by animals are, however, governed by procedures that are very different from voting procedures in humans.

Ants and honeybees are useful biological models for studying collective decision-making processes in group-living animals. Insect societies containing up to several thousands of individuals nevertheless succeed in behaving in a coordinated way. Both the ant nest and the beehive display striking examples of collective responses in a variety of contexts such as foraging (see references further in the text), nest moving (Franks et al., 2002; Pratt et al., 2002; Seeley and Visscher, 2004; Pratt and Sumpter, 2006; Seeley et al., 2006) or territorial defense (Lumsden and Hölldobler, 1983; Detrain and Pasteels, 1992; Millor et al., 1999). In most cases, the best food patch, the best nesting site or the safest place is selected. Amazingly, hundreds of nestmates, each with their own agenda, their own past experience, as well as
their own behavioral profile, are able to coordinate their efforts to behave in a highly cohesive way. Beehives or ant nests then appear as integrated social structures that gather a vast array of information acquired by its members, and that compare the relative value of each resource in relation to actual colony needs. These societies “wisely” deploy their workers over their home-range area. Moreover, collective decisions give rise to social-built structures that are characterized by relatively larger spatio-temporal scales than those observed at the individual level (Camazine et al., 2001). Indeed, while one ant individual can only forage over a few square meters at most, the trail network of the whole colony can extend over several acres. Likewise, while one forager, due to physiological limitations (e.g. water loss), can only stay outside the nest for, at most, a couple of hours, the colony taken as a whole can forage over food patches for relatively longer time scales – i.e. for several days or weeks.

In the past decade, life scientists have faced the exciting challenge of explaining how collective decisions may arise in societies made up of thousands of individuals. Since collective behavior may lead to a high level of cohesion, one can get the impression that “social patterns” are obeying complicated hidden blueprints or a centralized strategy imposed by one or a few leaders. This is not the case. Instead, for many social activities, the coordinated behavior of an insect colony relies on a multitude of local decisions made by individual workers. Collective structures are the dynamic by-products of interactions between colony members, and result from a decentralized decision-making process (see for example Seeley, 1995; Theraulaz and Spitz, 1997; Camazine et al., 2001; Detrain and Deneubourg, 2006; Sumpter, 2006). Any individual that makes a decision based on its local perception of the environment may contribute to the final foraging decision of the whole colony. Since each individual may influence the colony’s response and its fitness, such an individual has to constantly monitor the environment for cues and signals that indicate whether it should change its occupation, intensify its participation in current tasks and/or transfer the acquired information to nestmates. In a foraging context, the information the individual perceives about food profitability helps each worker to decide not just how, where and when to exploit a food patch, but also whether or not to recruit nestmates.

In this review, we compare collective decisions in hives and in ant nests by relating the properties of recruiting signals to the foraging strategies displayed by these two insect societies. We briefly describe the main positive and negative feedbacks that help foragers self-regulate their activities according to environmental constraints and opportunities. Even though honeybees and ants share a similar recruitment scenario, we would like to stress that the nature of their recruiting signals, as well as the ways information about food patches is conveyed to nestmates, differ greatly. In honeybees, tactile signals (i.e. the waggle dances) rule many aspects of
recruitment, whereas chemical communication (i.e. recruitment trails) is the backbone of collective foraging in ants. We emphasize the intimate link that relates the mode of locomotion of one species (i.e. walking or flying) to the media “chosen” to communicate information about food resources. In particular, we demonstrate that the distinct properties of chemical and tactile signals match the environmental constraints faced either by ant individuals on the ground or by honeybees in the air.

One goal of this chapter is to advocate that the use of either chemical or tactile signals in recruitment has provided communication constraints, which may account for different foraging patterns displayed by ant nests or by hives. Therefore, in the second part of this review, we compare how ants and honeybees make collective choices between identical sources, code information about food characteristics and select the most profitable resources. In particular, we shall argue that differences in the collective patterns displayed by ant nests or hives depend on the level of non-linearity of their recruitment process, or, in other words, on the amplifying potential of their communication. In addition, we examine how randomness may lead to diversity and flexibility of foraging patterns, instead of simply reducing the efficiency of food exploitation.

Throughout this review, collective patterns displayed by hives and ant nests are placed in a behavioral ecology perspective: we try to connect the properties of recruiting signals to environmental constraints, as well as to the efficiency of collective foraging strategies. Finally, by referring to other group-living arthropods, we address generic issues about information processing, its dissemination among conspecifics and the emergence of cooperative behavior.

2 Building up collective foraging behavior through feedbacks

The way individuals process information about current food opportunities or foraging risks, and the way they communicate with each other are crucial to the foraging success of insect societies (Seeley, 1994, 1995; Detrain et al., 1999; Sendova-Franks and Franks, 1999; Camazine et al., 2001). Insect societies not only have to communicate how far and in which direction food is located, but also have to decide how many foragers should be engaged in food exploitation. Much information can be exchanged through direct interaction between nestmates, such as antennal contacts or chemical emission of pheromones. In addition, workers can obtain information through animal-induced changes within a shared environment. Indeed, any individual that modifies its environment – and hence that of its nestmates – indirectly influences the further behavior of other group members. For instance, when an individual impoverishes an area by regularly feeding on it, it decreases the foraging success rate of
conspecifics and hence influences their further foraging decisions. This may lead to spatial specialization and related division of labor (Deneubourg et al., 1987; Beshers and Fewell, 2001; D’Ettore, 2007; Ravary et al., 2007).

Interactions between nestmates or between nestmates and their environment can be classified into two main categories: positive and negative feedback. A positive feedback is an interaction that increases or amplifies the response to an initial stimulus. One of the best-known examples of positive feedback is the emission of recruiting signals by foragers having discovered a new food source. This signal is emitted either incidentally or actively by successful scouts: it will mobilize and attract additional nestmates to the food site, which, after feeding, will in turn reinforce and amplify the recruiting signal (Hölldobler and Wilson, 1990; Seeley, 1995). However, without an antagonizing mechanism the process might become uncontrollable. Negative feedbacks, such as food exhaustion or overcrowding of food sites, act to counter or “dampen” amplifying effects that would otherwise lead to an excessive mobilization of foragers. Such mechanisms provide a stabilizing force and play a regulatory role by adjusting the number of recruited foragers to the number and size of food items. Insect societies may thus be seen as a network of coupled feedbacks. Their ecological success depends on the subtle interplay between positive and negative feedback loops that focus the colony foraging effort onto profitable sites, lead to the emergence of optimal patterns and open the way to cooperation in food exploitation and defense (see e.g. for honeybees, Seeley et al., 1991, 2000; Seeley, 1995; for ants, Hölldobler and Wilson, 1990).

2.1 RECRUITMENT: THE QUINTESSENCE OF POSITIVE FEEDBACKS

Many examples have been described and a vast body of literature has been devoted to the recruitment phenomenon, notably in social insects. Recruitment can be passive, based on inadvertent cues provided by others. For instance, the feeding activity of an animal can be perceived by conspecifics that will join it at the food source. Such passive recruitment has been reported in a variety of animal species and is associated with complex cost-benefit trade-offs, since conflict between individuals often occurs when gaining access to food sources (Thomas and Valone, 1989; Valone and Templeton, 2002; Danchin et al., 2004). Social insects are one example in which recruitment of nestmates is a truly active process. The successful scout “decides” to emit specific signals that attract and guide workers to the food. Hence, every time a forager returns to the nest, it brings home not only food but also information about the characteristics of the exploited resources, which it shares by means of recruitment signals. These signals provide a positive feedback that attracts nestmates, concentrates the foraging force around one food source, and contributes
to a social consensus about when, where and how to forage (Camazine et al., 2001).

Although both ants and honeybees are highly successful in coordinating their foraging activity and in making adaptive choices about where to forage, recruitment is achieved by using quite different communication channels.

In the case of ants, recruitment signals are essentially chemical. In some ant species, the recruiter leads one recruit (tandem recruitment) or a small group of recruits (group recruitment) from the nest to the newly discovered food source. The cohesion of the tandem or of the group is maintained by mechanical contacts between the leader’s gaster and the follower’s antennae and/or by short-range chemical signals emitted by the leader (Möglicher et al., 1974; Verhaeghe, 1982; Hölldobler and Wilson, 1990; de Biseau et al., 1991, 1994; Liefke et al., 2001; Franks and Richardson, 2006; Richardson et al., 2007). However, in ants, the most widespread recruitment procedure is to lay a chemical trail (Fig. 1A; Wilson, 1962a,b; Beckers et al., 1989; Traniello and Robson, 1995). In this case, the odor trail laid by the successful scout on its way back to the nest triggers the exit of nestmates and guides them to the food source. In turn, the recruits, having found and fed at the food source, reinforce the trail when returning to the nest. In doing so, successful foragers draw out of the nest an increasing number of nestmates and provide them with an efficient guiding trail.

In the case of honeybees, several hundred foragers bring in information about food sources and maintain a strong flow of information about food opportunities to the hive. The honeybees have a unique way of recruiting nestmates to food sources; the famous waggle dance (Fig. 1B) (von Frisch, 1967). Waggle dances, which consist of a series of waggle runs each followed by a semi-circular return, convey information about the direction and distance of the foraged site visited by the dancing bee. The angular deviation of waggle runs from the vertical corresponds to the angle of the food source relative to the current sun azimuth. The distance between the nest and the feeding site is encoded in the duration of the waggle run. Dances are thus a sophisticated means for honeybees to translate into a coded language the food location (for a review see Dyer and Seeley, 1991; Seeley, 1995; Dyer, 2002). The information encoded in this symbolic language is then read by unoccupied foragers and used to fly off and locate the foraging site. In addition to acquiring distance and direction information, bees that follow dances also learn the odor of the food and will then search for that scent in the field (Gould, 1976; Sherman and Visscher, 2002).

2.2 THE WAY YOU MOVE RULES THE WAY YOU RECRUIT

One may easily conceive that the way in which an animal guides recruits to food sources is constrained by the way the animal moves. Walking on the
ground or flying through the air has presumably imposed strong constraints on the type of recruitment signals used by ants and honeybees.

Quite obviously, the way ant scouts recruit nestmates is intimately related to their ability to lay signposts while walking between the nest and...

FIG. 1 Coupled feedbacks loops and dynamics of foraging in ants (A) and honeybees (B). Positive feedbacks such as ant trail reinforcement or honeybee dances trigger the exit of recruits. Thereafter, recruitment slows down due to the increasing impact of negative feedbacks, which take place inside the hive/the ant colony as well as around food sources. As a result, the dynamics of growth of numbers of foragers follows a logistic curve [(C) experimental curve for Lasius niger ant colonies exploiting a 1 M sucrose solution (Camazine et al., 2001)].
the food source. Successful foragers returning to the nest can easily bend their gaster downwards and hence exert a step-by-step control of the amount of recruitment signals laid over the substrate. The chemical trail, which is progressively built and reinforced by successful foragers, is then followed by ants as an invisible highway on the ground connecting the ant nest to the food site. In this case, information about the location of food, such as its spatial coordinates, is not explicitly coded by the trail or by any other signals emitted by recruiters inside the nest. In other words, recruits that exit the nest and start following the trail, have no idea where the food is located or at which distance from the nest. Foragers track the trail at a local scale through osmotropotaxis by moving their antennae over the substrate until they reach the food source (Hangartner, 1967, 1969; Detrain et al., 1988; Hölldobler and Wilson, 1990; Calenbuhr and Deneubourg, 1992; Calenbuhr et al., 1992). The recruitment trail is like an Ariadne’s thread, guiding foragers to a food source and providing them with step-by-step directional information. The use of a trail is not limited to ants, since other walking social insects, such as termites, rely on this mode of communication as well. Indeed, trail pheromones elicit recruitment and orientation for termite species in which workers forage outside the nest or move from one nesting site to another (Kaib et al., 1982; Traniello, 1982; Heidecker and Leuthold, 1984; Reinhard and Kaib, 1995, 2001; Pasteels and Bordereau, 1998; Traniello and Leuthold, 2000).

In addition to its guiding role, the following of a trail also gives an opportunity for foragers to come across many nestmates. During these head-on encounters on foraging trails, ant workers perceive several cues informing them not only about the partner’s identity but also about the type of food being exploited or about the profitability of a foraging patch (Roces, 1990; Roces and Nunez, 1993; Howard et al., 1996; Le Breton and Fourcassié, 2004; Dussutour et al., 2007a,b). For instance, contact with food residues over the recruiter’s body informs recruits about the food characteristics prior to their arrival at the feeding site and activates a food searching pattern leading to an increase in patch finding success (Le Breton and Fourcassié, 2004). Conversely, flying insects, such as honeybees, cannot exchange information about food sources along their foraging journey: they have to estimate food characteristics within the hive – namely through the perception of floral odors before flying out to the flower patch (Seeley, 1995).

Whereas the laying of chemical markers on the ground is a reliable way for walking insects to guide nestmates, it is far less convenient for flying animals, such as bees. Actually, the use of chemical trails by flying insects is limited to a few species of social stingless bees, which show a range of behavior; from odor-marking of the food alone to full trail-marking between the nest and the food source (Nieh et al., 2003). In these melipone species (Lindauer and Kerr, 1958, 1960; Michener, 1974; Jarau et al., 2003,
2004; Schmidt et al., 2006a,b; Schorkopf et al., 2007), scent-marking bees rub their proboscis over the substrate, start laying labial gland secretions and land regularly to deposit marks over twigs and leaves on their way back to the nest. The recruiting bee then triggers the exit of nestmates, which are guided by the scent trail towards the food source. The shorter foraging distances, as well as the larger sizes of stingless bee colonies compared to those of honeybees, have presumably facilitated the use of such chemical trails (Lindauer and Kerr, 1958; Beckers et al., 1989; Jarau et al., 2003, 2004; Schmidt et al., 2006a). Conversely, for honeybees, flying over long distances has set several constraints that impede the use of odor trails. First, chemical marking of the substrate requires frequent landings and takeoffs, which may become costly in time and energy for the bee scout over long distances. In addition, aerial trails emitted by a flying scout would be subjected to wind or air turbulence and too labile to be an accurate and reliable orientation cue for recruits over long distances. These flight-related constraints have prevented the building and/or the efficient following of chemical trails by honeybees. By developing their unique mode of communication about food resources through dances, honeybees have freed themselves from flight-related constraints on information transfer. The spatial information encoded within the dance allows recruits to be immediately informed about how to reach the food (Riley et al., 2005), while the floral odor inside the nest (Sherman and Visscher, 2002; Farina et al., 2005; Diaz et al., 2007) allows recruits to pinpoint the resource in the final stages of their flight as they approach the flower patch. Instead of being progressively built up through successive trail reinforcements (as in ants), information about the location of flower patches is “summarized” inside the hive by the honeybee dancers. While ant recruits walk along the trail without knowing their final destination, dance followers know the direction and the distance of flower patches before departing from the hive. In the hive, waggling runs along the comb provide an updated and accurate “flying map” to nestmates, which respond immediately with a direct flight to the specified patch.

2.3 REGULATION AND NEGATIVE FEEDBACKS

The ecological success of social insects depends not only on their ability to recruit nestmates but also to adjust the number of recruits to the available resources. To achieve such regulation of the workforce, positive feedbacks such as recruitment signals have to be coupled with negative feedbacks. The latter are often by-products of the foraging activity, such as crowding effects, food exhaustion or filling of nest reserves (Figs. 1A and 1B; de Vries and Biesmeijer, 1998, 2002; Dussutour et al., 2004; Schmidt et al., 2006b). As a result, the growth of the number of foragers at a food source can usually be defined by a logistic curve, which increases exponentially
until negative feedbacks create a plateau. These negative feedbacks include the limited number of potential recruits within the nest or limited access to the food source (Fig. 1C).

In ant populations, food exhaustion or overcrowding is known to weaken or even suppress recruitment (Fig. 1A). This adaptive response is based on a simple rule evidenced by Wilson (1962a,b); a recruited ant having no more access to food goes back to the nest without laying a trail. As the trail is no longer reinforced, the pheromone evaporates and recruitment comes to an end (Wilson, 1962a,b). Such an “all-or-nothing” response of foragers to food access is explained by a more generic rule of thumb: the “desired volume” rule. A forager having found a food droplet will lay a trail only when it is able to ingest a desired volume of its own that acts as a threshold value (Mailleux et al., 2000, 2003a, 2005, 2006). The use of such a criterion by ants is not restricted to a single source but can be generalized to multiple food sources, such as in the case of aphid honeydew droplets. This behavioral rule is a powerful and economical means for the society to adjust the number of potential food-carriers to the global quantity of available honeydew (Mailleux et al., 2003b). When an aphid colony is overexploited, aphid-tending ants can no longer ingest their desired volume, and so stop reinforcing the trail and recruitment slows down due to pheromone decay. Such negative feedback allows a constant readjustment of the foraging force to the current productivity of aphid colonies. At any time, information about the food volume still available is conveyed to the society by the number of trail-laying ants – i.e., individuals that have reached their desired volume. In some species, a volatile, non-entry pheromone may also aid in trail choice during active foraging by closing off unrewarding paths at the bifurcations of the trail network (Stickland et al., 1999; Robinson et al., 2005). The coupling between such negative feedback and amplifying recruitment signals may lead to dynamics that are more complicated than simple sigmoidal ones (Fig. 1C), such as oscillations in the number of feeding foragers (Verhaeghe and Deneubourg, 1983) or periodic activity patterns (Rissing and Wheeler, 1976; Franks and Fletcher, 1983; Goss and Deneubourg, 1989). In addition, any environmental constraint that alters the dynamics of trail recruitment can contribute to the regulation of foraging without any explicit coding of information by the worker individuals (Dussutour et al., 2006). For instance, if an ant colony is given the choice between two paths at low densities, it will forage over one trail but, in crowded conditions, both branches will be equally followed. Indeed, high rates of collision between ants leaving the nest and those returning with food force some of them to take the second path. This negative feedback, the collision rate, guarantees that an optimal rate of food return is maintained (Dussutour et al., 2004) and also generates a temporal organization of foragers’ traffic (Dussutour et al., 2005b). Negative feedbacks may also occur within the
ant nest (Fig. 1A). For example, when faced with different food sources (e.g. proteins versus sugars), workers will selectively collect the food type that best meets the colony’s requirements. In this case, we suspect that the propensity of inner nest workers to accept trophallaxis or to respond to recruiting stimuli provides a cue to the recruiter about the actual colony food demand, as well as about the level of nest reserves (Howard and Tschinkel, 1981; Sørensen et al., 1985; Camazine et al., 1998; Cassill and Tschinkel, 1999a,b; Cassill, 2003). Any difficulty experienced by a recruiter in unloading its crop content may indicate a decreased interest for such a resource and lead to its progressive neglect by foragers.

In honeybees, the mechanisms that regulate recruitment to flower patches exhibit strong similarities with those of ants (Fig. 1B). Several negative feedback loops may dampen the recruiting response triggered by dancing foragers. One should notice that, unlike in ants, most of the negative feedbacks discovered so far take place inside the hive. Indeed, a bee that returns to the nest with a load of nectar has to find a receiver bee before recruiting and going back to the flower patch. The search time experienced by this forager, before being relieved of its crop load, depends on the number of bees that are simultaneously free and is also highly correlated with receiver motivation to accept such a food resource (Seeley and Tovey, 1994). Short latencies before being unloaded increase the probability of a forager carrying out a large number of waggle dances. Conversely, long latencies result in foragers queuing up to be unloaded and performing fewer waggling runs or none at all. In some cases, the unloaded bee may start a tremble dance, which fosters additional bees to function as receiver bees (Seeley, 1992). Even though tremble dancing may somewhat prolong food retrieval, foraging progressively slows down, as no more receiver workers are available within the hive (Seeley et al., 1996). The decision of a bee scout to recruit nestmates is thus based on a simple cue: unloading time, which acts as a feedback mechanism to reduce recruitment effort towards resources of decreasing interest to the hive. This decision rule is a reliable method to assess the demand–offer balance for a given resource: it integrates information about the number of empty cells free for food storing in the comb, the food demand of larvae, as well as the number of nestmates available for food processing in the hive. Other feedback loops may also occur outside of the hive and take place at the flower patch itself. Because the patch size is limited, some bees may have to fly around and wait for a place at the source to become free or return empty (de Vries and Biesmeijer, 1998, 2002). Chemical cues, such as marking a deteriorated source with a repellent pheromone, may also be used by bees to slow down foraging recruitment and prevent further visits by foragers (Giurfa, 1993).

In insect societies, the amplifying effects of recruitment are thus counteracted by negative feedbacks, the importance of which increases
as food sources become depleted, overcrowded or of lesser interest to the colony. Even though hives and ant nests display a similar logic in foraging regulation, the nature of feedback loops involved is quite different. Besides, as discussed in the following sections, the different ways foragers are recruited lead, at the collective level, to quite different collective choices and foraging patterns that are non-intuitive and hard to predict.

3 The way you recruit rules the way you make collective choices

Under natural conditions, ant colonies or hives may be faced with several food sources and have to make efficient collective choices between these alternatives. Many factors are likely to influence such collective choices: food characteristics may differ, competitors and predators may interfere with the colony activity, temperature and other climatic conditions may influence the energy/time costs associated with foraging. Therefore, each colony has to allocate foragers to food patches in a way that maximizes energy input and minimizes foraging costs. We review simple situations that demonstrate how competition between recruitment to several food sources may lead to collective responses that are adaptive for the whole society. First, we look at laboratory experiments with ants. Although these experiments may appear somewhat artificial, they help to identify, at the individual level, the simple recruitment rules that underlie the diverse and sometimes unexpected foraging patterns displayed by the ant colonies. Moreover, they allow us to build models and to test their predictions. Such models can be extended and adapted to the recruitment logic of honeybees (Camazine and Sneyd, 1991; Seeley et al., 1991; de Vries and Biesmeijer, 1998, 2002). They are also convenient tools for comparing the impact of recruitment properties on collective decisions made by ants or by honeybees, and for discussing to what extent emergent patterns are presumably adapted to the foraging ecology of these social insects.

3.1 COLLECTIVE CHOICES AND TRAILS: EXPERIMENTAL EVIDENCE IN ANTS

The choice between identical resources is an unusual situation under natural conditions and seems of little interest from an ecological point of view. However, binary choice experiments between identical sources can be useful in highlighting how seemingly small differences in the communication process may affect collective decisions. When two identical sources are available, the most intuitive pattern would be the equipartition of foragers between the two resources. Indeed, an allocation of foragers that is strictly proportional to the availabilities of resources – i.e. an ideal free
distribution of foragers between sources (Giraldeau and Caraco, 2000) – seems the best way for a species to maximize its fitness by reducing the costs of competition between conspecifics. However, quite unexpectedly, an equipartition of foragers is seldom observed in the majority of trail-laying ant species. After a short period of equal exploitation, a bifurcation is observed, meaning that one source is selected and more strongly exploited than the other, even though the two food sources are identical (Figs. 2A and 3A). This is not the result of any asymmetry of the experimental set up since the side of the chosen source may change over successive tests (Pasteels et al., 1987; Beckers et al., 1990; Sumpter and Beekman, 2003). Similarly, when the colony has the choice between two paths of the same length, most ants will end up traveling and sharing only one foraging route (Deneubourg et al., 1990; Beckers et al., 1992b; Dussutour et al., 2004). The process of choice at the colony-level, between equivalent branches or between identical food sources, is thought to start by chance with an imbalance in the pheromone amounts between the two branches, large enough to trigger a positive feedback loop. The disproportional numbers of workers choosing and laying pheromones on the more traveled branch means that the ants’ stream to one source/path out-competes the stream to the other source/path (Goss et al., 1989).

3.2 A MODEL OF FOOD SELECTION INVOLVING CHEMICAL TRAILS

The bifurcation process – i.e. the selection of one food source among several identical ones – can be predicted by a behavioral model of food recruitment that relates the behavior of recruiting individuals to the foraging choice emerging at the society level (Deneubourg and Goss, 1989; Beckers et al., 1992a, 1993; Nicolis and Deneubourg, 1999; for a review see e.g. Detrain et al., 1999; Sumpter and Beekman, 2003; Sumpter and Pratt, 2003). This model, built upon empirical findings, describes how the amount of trail pheromone, and hence the traffic of ants on each trail, will change over time. In this model, an ant leaves the nest, chooses a path, reaches one of the two food source, ingests food and promptly returns to the nest, laying a trail pheromone. Each ant has to make two decisions: (1) to leave the nest or not, and (2) to choose one of the trails leading to equidistant food sources of identical quality (i.e. a bridge with two equal branches in a binary choice experiment). For each ant, the probability of leaving the nest increases with the total amount of trail pheromone at the nest entrance. Over time, the increasing rate of nest departure coupled with trail reinforcement by successful foragers contributes to the sigmoidal growth of the recruits’ population. One can neglect time variations in nest exits since
they do not affect the general dynamics of collective choices (Camazine et al., 2001).

An ant having left the nest has to choose one of the two paths: the probability for an ant of choosing one path $i$ depends on a general choice
function $A_i/\sum A_i$ in which $A_i$ is the attractiveness – i.e. amount of trail pheromone – of the trail $i$ leading to the source $i$ (Fig. 4). We can quantify the ant’s decision at the choice point by Eq. (1) in which the orientation toward one source and/or towards one trail compared to the others.
depends on the values of the pheromone concentration $c_i$ on each trail $i$ (Deneubourg and Goss, 1989).

$$P_i = \frac{A_i}{\sum_{l=1}^{s} A_l} = \frac{(k + c_i)^n}{\sum_{l=1}^{s} (k + c_l)^n} \quad i = 1, 2, \ldots, s. \quad (1)$$

$s = 2$ in the case of a choice between two food sources.

- $n$ determines the steepness – i.e. the level of non-linearity – of the recruitment response. A high $n$ value means that ants are highly sensitive to changes in the intensity of recruiting signals: even a slightly higher amount of pheromone over one branch will markedly increase the probability of ants to choose this branch.
- $k$ corresponds to the intrinsic degree of attraction of an unmarked branch of the bridge. For high $k$ values, high differences in trail concentration between branches are needed to get a non-random choice of one path.

On its return journey from a food source, each ant lays a quantity of pheromone $q_i$. In the case of identical food sources ($q_1 = \ldots = q_s = q$), each ant deposits the same amount of pheromone on its homeward journey.
and the pheromone concentration over one trail is directly proportional to the flow of foragers ($f$) over this trail. At each time unit, a quantity $q\phi P_i$ of pheromone is added to the trail $i$, while $vC_i$ is the decay of the trail pheromone amount. The trail is thus assumed to evaporate at a rate proportional to its concentration with $v$ being the inverse of the mean trail pheromone lifetime. The following system of $s$ differential equations describes the rate of change in pheromone concentration on trails.

$$\frac{dC_i}{dt} = \phi q P_i - vC_i \quad i = 1, \ldots, s$$

Monte-Carlo simulations based on these equations and on parameter values obtained experimentally for Lasius niger species give the time evolution of the ant population over each branch. Initially, both sources are exploited equally but a break suddenly occurs, after which one source becomes more exploited than the other. One food source is selected and exploited by more than 60% of foragers in nearly all cases (in around 90% of the simulations) (Fig. 5A). As for laboratory experiments (Fig. 2A), the asymmetrical exploitation of identical food sources arises as one trail becomes slightly stronger than the other, resulting in it being followed more often and further reinforced by trail-laying foragers in a positive feedback cycle. Due to the non-linearity ($n > 1$) of an ant’s response to the recruitment trail, the selection of one food site arises from the amplification of slight initial differences in the relative pheromone concentration on the trails. However, one can never say which trail will dominate or which source will be chosen since this selection is based on random events.

The model can be used to assess the role of each parameter in the final selection of one food source. It reveals that collective responses can be even more complex and sophisticated. One can observe that an asymmetry in food exploitation is more marked and more quickly reached when the flow of foragers or the colony size is larger, as expected from any phenomena based on amplifying processes (Figs. 3A and 5A). Below a certain number of ants or interactions, the exploitation pattern may be symmetrical, whereas above this value a collective choice is very likely to emerge, with most of the foragers being ultimately focused on one or only a subset of available resources (Nicolis and Deneubourg, 1999). Likewise, when individual trail-laying behavior is more intense, or when the rate of pheromone decay is lower, a fluctuation in the amount of trail chemical over one path is likely to be amplified and lead towards an unequal exploitation of food sources. To observe these phenomena experimentally, one may vary the “foragers’ flow” parameter by comparing colonies of different sizes, or one can increase the intensity of the trail laid by the ants ($q$) by increasing the quality of food sources (e.g. the sugar concentration). However, the influence of pheromone evaporation rate on the ants’
FIG. 5 Frequency distribution of simulations as a function of the fraction of foragers that choose food source 1 when two identical food sources are offered simultaneously (1000 simulations for each tested condition). (A) Distributions of ants at each food source. They were measured after 1 h and were based on the following parameter values: $k = 6; g_1 = g_2 = 1; v = 0.00056 \text{s}^{-1}; n = 2$ [for further explanations see Eq. (1) in text]. The bimodality of the ants’ distribution is more clear-cut for high ant flows (dark bars; probability of leaving the nest ($\phi$): 0.066 ant s$^{-1}$) than for small ant flows (grey bars; $\phi$: 0.017 ant s$^{-1}$). (B) Distributions of honeybees at each flower patch, measured after 2 h. Simulations are a simplified version of the Camazine and Sneyd (1991) model and are based on Eq. (3) (see text). Probability of remaining a dancing bee after a foraging flight $= 0.8$; flying time $= 250$ s; resting time in the dance area $= 120$ s; $k = 0; n = 1$. Honeybees tend to be equally distributed between the two food sources with a distribution that is narrower for large forager flows (dark bars; probability of leaving the hive ($\phi$): 0.05 bee s$^{-1}$) than for small foragers flows (grey bars; $\phi$: 0.0085 bee s$^{-1}$).
collective pattern is more difficult to investigate experimentally since it is determined by a multitude of parameters, such as ground temperature or the perceptive physiology of the ants. In this latter case, the model has a strong heuristic potential by allowing “numerical experiments” to be performed, which would be difficult or even impossible to carry out in the laboratory.

3.3 COLLECTIVE CHOICES AND DANCES: A MODEL FOR HONEYBEES

Similar to ant colonies, the hive has to make a collective decision about which flower patch to exploit and how many foragers should be allocated to the available resources. Under natural circumstances, the concurrent availability of several food sources is probably common as plants may undergo synchronized blooming. The model developed for trail-laying ants can be transposed to food recruitment in honeybees (Camazine and Sneyd, 1991; Seeley et al., 1991). The number of dancing bees is analogous to the trail concentration and determines the attractiveness $A_i$ of one foraging site over another. In honeybees, the probability of a recruit going to an advertised site is proportional to the number of waggle runs performed by the bees recruiting for that site (Camazine and Sneyd, 1991; Seeley et al., 1991). In addition, foragers dancing on the comb are able to attract with a vibratory signal dance-followers that are too far away to perceive tactile signals (Tautz et al., 2001). Judd (1995) observed that, although the average number of bees orienting to a dancer can be as high as five or six, at most only one or two bees are receiving the dance information. Camazine and Sneyd (1991) have incorporated this observation in their model, so that only one bee at a time can follow the dance and extract information from it. Hence, for honey bees, the attractiveness of a food source $A_i$, or in other words the number of bees recruited towards food source $i$, is directly proportional to $(D_i d_i)^n$, where $D_i$ is the number of bees in a dancing state. However, since these bees may stop for a while between successive dances, $D_i$ is multiplied by the relative amount of time ($d_i$) during which recruiting bees actually dance. When the two food sources are of identical quality, $d_1$ is equal to $d_2$.

The probability of one honeybee foraging on flower patch $i$ is then given by Eq. (3):

$$P_i = \frac{A_i}{\sum_{i=1}^{s} A_i} = \frac{(k + d_i D_i)^n}{\sum_{i=1}^{s} (k + d_i D_i)^n} \quad i = 1, 2, \ldots, s. \quad (3)$$

The parameter $n$ accounts for the non-linearity of the response and reflects the sensitivity of recruits to the relative occurrence of a dance $i$ associated with food source $i$. A value of $n > 1$ would mean that a recruit would be more likely to forage at a given flower patch when there are
several dancers coding for this source. However, in the case of honeybees, as predicted by Camazine and Sneyd (1991) and demonstrated experimentally by Seeley et al. (1991), bees do not sample dancers in an effort to compare the relative profitability of food sources. Rather, a bee recruit is influenced only by a single dancing recruiter at any one moment. Since there is no synergetic effect of concurrent dances, the value of $n$ in Eq. (3) should equal 1. Thus, the attractiveness $A_i$ of one foraging site $i$ over the others should be directly proportional to the occurrence of dances $D_i$. As a consequence, even though the number of dances is analogous to the trail concentration, the honeybee recruitment differs from that of ants in its level of non-linearity ($n = 2$ for trail-laying ants, $n = 1$ for dancing bees). This difference has only a slight impact on the probability of one individual choosing a path or a food source (Fig. 4), but it is responsible for the very different properties of collective patterns displayed by honeybees and ants. Indeed, the model predicts that the hive will occasionally select one source only and that the most frequent response of honeybee foragers will be an even distribution between identical food sources (Figs. 3B and 5B). Furthermore, unlike ants, which more frequently select one source when the number of foragers increases (Fig. 5A), the honeybees show an even distribution of foragers whatever the hive size (Fig. 5B). This prediction finds some support in the fact that small hives foraged at approximately the same number of patches as large colonies (Beekman et al., 2004). As the hive size increases, the model also predicts that the foragers will be even more equally allocated among food sites and that the presence rate of bees at one of the two food sources will shift from a flat to a mono-modal distribution with a peak at 50% (Fig. 5B). Hence, an increase in colony size has unexpected opposite effects for hives than for ant nests: larger forager flows should enhance the collective selection of one food source by ants, whereas it should draw the honeybees towards a more equal exploitation of resources (Figs. 3A and 3B). In the field, no symmetry breaking between identical sources has yet been reported for honeybees and further experimental evidence is still needed to confirm the model’s predictions of an even distribution of bee foragers between flower patches. Nevertheless, one should be aware that the selection of one food source by honeybees may occur due to some “hidden” amplifying behavioral traits associated with recruitment. For instance, once a recruiter enters the hive, it stimulates forager mates to return to a known feeding place, not only by dances, but also by nectar transfers which reactivate experienced foragers. This may have a snowball effect since experienced bees follow a significantly lower number of waggle-runs and are more easily reactivated to perform nectar collection (Gil and Farina, 2002). Another source of non-linearity could be the memorization of food location, as well as the learning of flying routes that facilitate the orientation of bees and shorten their foraging journeys. These phenomena are translated, from a
mathematical point of view, by an increase of the parameter $n$ value and may lead to symmetry breakings as predicted by de Vries and Biesmeijer (2002).

In insect societies, collective decisions arise as a result of the competition between different sources of information that are amplified by recruitment signals. Through its trail-laying, an ant colony with a large forager population will select and focus its effort on one resource even if all available sources are of identical quality. Alternatively, an ant colony may exploit all sources equally if the colony itself is small or if the number of sources is very large (Nicolis and Deneubourg, 1999). Such switching in the foraging pattern, which is not predicted by the honeybee model, matches the features of food resources exploited by ants. Unlike bees that wander among short-lived flower patches, several mass-recruiting ant species feed on honeydew and forage on rather stable food sources, such as aphid colonies. By focusing the foraging force on a limited number of food sites, ant colonies can protect and monopolize these resources through cooperative defense against competitors or predators. Such an “owner strategy” can be efficiently implemented only by large sized ant nests since the costly loss of workers may be fatal to small incipient nests (Franks and Partridge, 1993; see also Mailleux et al., 2003a).

By contrast, a hive presented with an array of identical sources will scarcely focus its activity on a single flower patch. It seems more advantageous for a honeybee colony to distribute thousands of workers among multiple flower patches rather than to be crowded at a single food source or to monopolize a flower patch of highly fluctuating profitability.

Hence, the different ways used by ants and honeybees to allocate foragers between identical resources are not primarily due to differences in the sensory or physiological capacities of the individuals, but are by-products of the properties of their recruiting signals (chemical trail versus dances), which shape the decision-making processes operating at the collective level.

4 Choosing the best food resource

The ant colony, as well as the hive, provides striking examples on how rather simple individual behavior and decision rules can lead to different but equally efficient patterns of food exploitation. The previous section considered the case study in which identical sources were simultaneously offered to a colony. However, the availability of food resources most often varies in space and time, while profitability differs according to several factors such as food quality, distance, safety and so on. Since individuals have only a local and partial knowledge of their environment, one may
wonder how insect societies adjust their foraging effort in order to exploit preferentially the most profitable resources.

In both honeybees and ants, foragers are adequately allocated to the best food sites without any of the participants having a broad knowledge of all food opportunities. An ant or a honeybee scout does not compare nor keep track of information from multiple foraging sites: its knowledge of the array of food sources is limited to its particular food site, which it assesses using functional cues (ants: Hölldobler and Wilson, 1990; Detrain et al., 1999; Detrain and Deneubourg, 2002; honeybees: Seeley et al., 1991; Seeley, 1995; Biesmeijer and Ermers, 1999; Dyer, 2002; stingless bees: Schmidt et al., 2006a). On the basis of this individual estimate of food profitability, the scout will decide how strongly to advertise the source by either laying a trail or by dancing in the hive. The best food sources are then selected by the whole colony – not by centralized decision-making, but simply by competition between recruitment signals differing in intensity and/or duration. While a tuning of recruitment allows both ants and honeybees to make adaptive choices, these insect societies are expected to somewhat differ in the robustness of their collective decisions. Indeed, while the removal of a dancing bee would lead to losing part of the information about food opportunities, the removal of one ant would hardly affect the amount of pheromone on the trail, which keeps the ants informed about food quality and food location.

The strength of amplification processes – i.e. recruitment rates towards different food sources – depends on two main categories of parameters (Detrain and Deneubourg, 2002, 2006). The first consists of any modulation of the recruiting signals that may enhance the mobilization of inactive nestmates towards a food source. For example, an individual scout bee may lengthen the dance duration or an ant may lay a higher amount of trail pheromone in relation to its own estimate of food profitability. The second set of factors involves various environmental features, (1) that may influence the duration of a foraging cycle such as the impact of temperature on walking/flying speed of foragers, or (2) that may alter the trail lifetime such as the absorbency of the substrate (Detrain et al., 2001; Detrain and Deneubourg, 2002; Jeanson et al., 2003). Likewise, a longer path toward a feeding site (Goss et al., 1989) or a longer food distance from the nest (Detrain et al., 1999) may reduce the rate of trail reinforcement and lead to a remote food source losing out in its competition with other resources. In these latter cases, one food source or one path can be selected without any modulation of the recruiting signal by the individuals. In the next sections, we consider only the first set of factors that act upon recruitment dynamics by reviewing how an ant or a honeybee actually fine-tunes its communication in order to select the best food sources.
4.1 FOOD SELECTION AND TUNING OF TRAIL PHEROMONE BY ANTS

In the case of trail-laying ants, if a rich source (1 M sucrose solution) is discovered at the same time as a poor one (0.1 M sucrose solution), the food source with the highest sugar content is selected and exploited by most of the foragers (Fig. 6A). There is no evidence of any qualitative change in the chemical content of recruitment trails laid by ant scouts depending on food quality. Information about food quality is “coded” only through quantitative changes in the trail intensity: more frequent trail spots will be laid...
by recruiting ants to more concentrated sugary liquid sources (see e.g. Hangartner, 1969; Hölldobler and Wilson, 1990; Detrain et al., 1999).

Experimental data show that this trail modulation, as a function of food quality, is rather weak. For instance, in Lasius niger, the average number of trail marks laid towards a 1 m versus a 0.1 m sugar solution is only of 1.5 higher (Beckers et al., 1992a, 1993). Then, as predicted by Eq. (1), at the trail bifurcation point, the likelihood of an individual choosing the more concentrated trail is only slightly higher than that of it choosing the less marked trail path (3–4% higher for the first steps of recruitment). The slight impact of differences in signal intensity on the choice by individual ant (Fig. 4) contrasts with the clear-cut collective choices that ultimately emerge at the colony level (Fig. 6A). Indeed, based on numerical simulations, there is always a higher percentage of foragers that orient themselves towards the sweetest source, this rich food source drawing more than 80% of ants in nearly all cases. Hence, due to the non-linearity of the response to chemical trails, even a slight modulation of the recruiting signal drives the whole colony to a nearly unanimous choice of the most profitable food source.

Besides food quality, foragers can also convey information about food size. The decision of each scout to lay a trail is ruled by simple criteria such as the ability to ingest a wanted volume of liquid food (Mailleux et al., 2000, 2003b). The prey resistance to traction is another criterion used by foragers to estimate prey size: for small prey, foragers lay only a weak trail whereas, for a prey item too large to be individually retrieved, ants dash back to the nest laying a more continuous chemical trail and trigger an intense food recruitment (de Biseau and Pasteels, 1994; Detrain and Deneubourg, 1997). Such decision criteria are highly functional as they incidentally take into account multiple sources of information, such as food size, food weight or current number of ants cooperating in food retrieval (Detrain et al., 1999; Detrain and Deneubourg, 2002). They also act as individual thresholds above which a scout decides to recruit nestmates so that, at the colony level, there will be a higher percentage of trail-laying foragers directed towards large food sources.

Hence, due to the autocatalytic nature of chemical recruiting signals, ants have a high potential to draw most of the foraging force towards the most profitable food source (Beckers et al., 1990, 1993; Sumpter and Beekman, 2003; Portha et al., 2004). Even slight changes in the trail-laying behavior of a few foragers can boost the reinforcement of one foraging path by the succeeding ants and can ultimately lead to the exploitation of the most profitable food by the whole colony.

4.2 FOOD SELECTION AND TUNING OF RECRUITMENT DANCES BY HONEYBEES

As for trail-laying ants, the recruitment activity within the hive is adjusted to both increased and decreased food profitability: honeybees will allocate
more recruits to resources that have a higher pay-off for the colony. If food profitability is gauged as low by one forager, it will abandon the resource, whereas if it estimates it as high, the forager may decide to recruit nestmates and to tune the intensity of recruiting signals according to the food profitability (see e.g. von Frisch, 1967; Seeley and Towne, 1992; Seeley, 1994, 1995; Seeley et al., 2000; De Marco and Farina, 2001; Afik et al., 2008). Mainly quantitative changes in the recruitment signals occur. Indeed, the strength of dances inside the hive intensifies with the profitability of the flower patch. The number of waggle dances ($D_j$) as well as their relative duration ($d_j$) can then be viewed as indicators of the overall value of the nectar source, such as the distance to the flower patch or the sweetness of the nectar. A higher occurrence of dances coding for a rich resource ($D_j d_j$) slightly increases the likelihood of a bee being recruited towards this source (Fig. 4). This tuning of dances results in a cascade of positive feedback at the colony level so that, at the stationary state, more than 60% of the flow of foragers will be focused on the sweetest flower patch in around 80% of cases (Fig. 6B). However, in the remaining cases, the collective choice of the most profitable food source is less clear-cut than for ants, since 20–40% of honeybees go on exploiting the poorer source. These predictions have been supported by experiments showing that the hive is actually able to focus its foraging effort on the food source with the highest sugar concentration, i.e. with the highest energy reward (von Frisch, 1967; Seeley, 1995; Dyer, 2002).

### 4.3 Redirecting Collective Choice in a Changing Environment

Ants and honeybees share the same potential to select the best food source when several resources are discovered simultaneously (Fig. 6). However, hives and ant nests differ greatly in their ability to redirect their choice depending on the temporal sequence of food discoveries.

In the case of ants, if a rich source (1 M sugar solution) is discovered after recruitment is well underway to a first discovered but poorer source (0.1 M), then the poor source will remain the most exploited while the rich one will be neglected (Figs. 2B and 7). Theoretical simulations predict that the poor source will keep attracting the majority of foragers in all cases (Fig. 8A). The difficulty ant colonies have in redirecting their foraging activity towards more profitable sources has been experimentally confirmed in several mass-recruiting species (Beckers et al., 1990; Traniello and Robson, 1995). Similar to ants, scent-marking stingless bees may be “trapped” in the exploitation of the first discovered food source (Schmidt et al., 2006a). When stingless bees experience an increase of sugar concentration over time, the newcomers at the richer source never outnumber the bees that have already exploited the poor source and reinforced the scent trail (Biesmeijer and Ermers, 1999). The difficulty of both ants and stingless bees to redirect foraging towards more profitable food sources could be
due to the chemical nature of their communication. Indeed, chemical recruiting signals elicit responses, which are strongly non-linear and highly sensitive to slight differences in signal intensities. The new trail to the richer source will never succeed in overcoming the ongoing recruitment to the first discovered food source despite its lower profitability. Likewise, the selection rate of a food source is correlated to the number of workers

FIG. 7 Flexibility of foraging choices in ants and honeybees. In ants, the first exploited food (source 1 at $t_1$) will remain preferentially foraged at $t_2$ even a richer food (source 2) has been introduced. Conversely, honeybees that exploit the first discovered food (source 1 at $t_1$) are able to shift their activity at $t_2$ towards a more profitable resource (source 2) that has become available.
FIG. 8 Collective response to the delayed introduction of a new richer food source in ants and honeybees. (A) Frequency distribution of simulations as a function of the fraction of ants exploiting the rich source (after 120 min; 1000 simulations). When the recruitment is well-established to the first poor source, a richer source is introduced at $t = 30$ min (arrow). Even at 120 min, the poor source remains selected in all cases. (A, inside box) A simulation showing the dynamics of the fraction of foragers at the poor (dotted line) and at the rich (solid line) food source. Only a few individuals were occasionally observed at the rich source. (B) Frequency distribution of simulations as a function of the fraction of bee foragers exploiting the rich source (after 360 min, 1000 simulations). In honeybees, due to the slow growth of recruitment, foragers have access to the first poor source for a longer time before a new richer source is introduced at $t = 60$ min (arrow). At 360 min, the rich source is exploited by more than 60% of the honeybee foragers in nearly all cases. (B, inside box) A simulation showing the progressive shift of bees from the first flower patch (dotted line) towards the more profitable patch (solid line).
already exploiting it and this is likely to increasingly reinforce the scent trail (Pasteels et al., 1987; Goss et al., 1989; Sumpter and Beekman, 2003; Sumpter and Pratt, 2003).

Conversely, in the case of honeybees, the hive is expected to be able to select the most profitable nectar source, even if poorer food sources have been previously discovered. If between two nectar sources, the sugar concentration of one suddenly increases, the number of foragers exploiting the other source decreases, as experimentally shown by Seeley et al. (1991). As predicted by theoretical simulations, the delayed introduction of a sweeter source induces a reorientation of the honeybee foraging at the expense of the first discovered source (Figs. 7 and 8B). Over time, the sweetest food source will attract more foragers, will reach the same level of exploitation as the first food source and ultimately will become preferentially foraged by honeybees. In the majority of cases, 60–80% of bee foragers will end up redirecting and focusing their activity towards the rich food source (Fig. 8B). The high flexibility of honeybee foraging patterns is due to the proportional relation between occurrence/duration of dances and the number of recruited foragers. Provided that inactive workers are still available within the hive and ready to engage themselves in foraging, they can be recruited towards a richer flower patch by dancers coding for this new target. The reorientation towards more profitable resources may be facilitated by the phenomenon of cross-inhibition between forager groups (described by Seeley, 1995). Indeed, upon entering the hive, bees loaded with nectar of lesser value experience longer search times for nectar receivers, which decrease the rate of dances coding for this poor source.

Being flexible and able to shift easily to a more rewarding food patch provides obvious benefits to the insect society in terms of optimal exploitation of environmental resources. One may question why ants differ from bees by displaying such poor flexibility in their foraging patterns. The answer depends in part upon whether there is a selective advantage in being aggregated to concentrate the colony’s efforts on a single site, or whether it is better to distribute colony workforce more widely. While the amount of flower nectar collected by the hive results from the simple summation of foraging journeys by crop-loaded bees, the food amount retrieved to the ant nest is more variable. Indeed, it closely depends on the ability of workers to cooperate in the killing of prey or to collectively retrieve large food items. Besides, while cooperative defense in bees is restrained to the hive surroundings (Guzmannovoa and Page, 1994; Millor et al., 1999; Hunt et al., 2003), ants benefit from cooperative defense of food sites and their aggressiveness toward aliens increases with the number of surrounding nestmates (see e.g. Pontin, 1961; Levings and Traniello, 1981; Lumsden and Hölldobler, 1983; Czechowski, 1984; Detrain and Pasteels, 1992; Sakata and Katayama, 2001). Concentrating the foraging effort on a few
patches gives the ant colony an edge when faced with competitors, and limits the investment in defending food sources with the highest rates of return. Moreover, within a large group of cooperating ant workers, specialized castes and task-partitioning become possible, which may contribute to an increase in global foraging efficiency (Detrain and Pasteels, 1991; Ratnieks and Anderson, 1999; Anderson et al., 2002).

As regards honeybees, due to dance recruitment, scouts having discovered more profitable flower patches can easily redirect the hive activity towards these new resources. The flexibility of the dance recruitment, coupled to an intense exploration, allows the hive to maintain a long-term foraging success even when flowering patches are ephemeral and variable in richness. Dance-language recruitment is presumably well-adapted to the resource variability experienced by the hive (Seeley and Visscher, 1985; Sherman and Visscher, 2002).

All these examples have questioned whether and how insect societies make collective decisions in order to exploit the most profitable food sites. There are striking similarities in the mechanisms allowing ants and honeybees to reach such a collective decision: no individual visits all food sources and none compares them nor decides which one is the best. Instead, accurate collective decisions arise from the competition between positive feedback loops and from the tuning — even if slight — of recruiting signals towards the various food alternatives. The different pools of foragers “compete” to recruit more of their comrades towards each food location. However, the amplifying potential of recruitment strongly differs between ants and honeybees with a higher level of non-linearity associated with trail pheromone signals than with dances. Such a difference has a deep impact on the properties of patterns emerging at collective level. The high non-linearity of chemical trails will make the occurrence of bifurcation more likely, i.e. collective choice of one food source, when identical food sources are offered simultaneously. On the other hand, the non-linearity of the response to chemical recruitment in ants impedes the reallocation of foragers when food profitability changes.

5 Randomness and foraging: adaptive tuning of error in ants and honeybees

In the previous sections, the recruitment process was described in an idealized way so that every recruited worker succeeds in reaching the food target. However, for a long time, biologists have been puzzled by the following paradox: impressive structures and consensus emerge at the society level, while individuals show random, highly variable and, in some cases, not very efficient behavior [see e.g. Grasse stimergy theory (1959) about the building behavior of termites]. The duality between the
stochasticity of individual behavior and the determinism of the colony response is not peculiar to insect societies and occurs in other biological and non-biological systems (Nicolis and Prigogine, 1977). To solve this paradox, many researchers have argued that social insect behavior is less erratic than it appears, or have considered randomness as an unavoidable side-effect of animal behavior. Insect societies are expected to be able to afford more easily such a variance because of the large number of interacting workers (Oster and Wilson, 1978). If one ant fails, another will succeed: some inefficiency due to randomness in individual behavior is compensated by the reliability of the whole system, which will ultimately reach the goal. However, stochastic events should not be reduced to side-effects since they may have more creative and adaptive consequences (Deneubourg et al., 1983). At first sight, the most accurate system of communication may seem the best one, and this is certainly the case when only one food source is present and located at the same place for a while. However, natural environments are more complex, with several sources that may fluctuate or be simultaneously available to the ants. The “lost” recruits that wander around the foraged sites may then discover new food sources of higher profitability. Hence, when multiple food sites are available, there is an optimal error level which minimizes the time needed for discovering food sources and which maximizes foraging efficiency.

One may question how an insect society finds the optimal strategy between a high accuracy of communication allowing immediate food exploitation and a low accuracy allowing new food discoveries. In ants, randomness is intimately related to the ability of foragers to orient and to follow a chemical trail. Recruits that leave the nest may lose the trail and never reach the food source. Hangartner (1967, 1969) showed that orientation along a trail is mainly due to osmotropotaxis, meaning that the ant perceives different concentrations in trail pheromone between its left and right antennae. For a given concentration in trail pheromone, the number of ants that are still present on the trail decreases exponentially as a function of the length of trail already followed (Pasteels et al., 1986; Detrain et al., 1988; Calenbuhr and Deneubourg, 1992; Calenbuhr et al., 1992). The mean length of trail followed by ants increases with trail concentration, and the ants’ orientation may become impaired above a certain concentration of trail. Since the amount of trail pheromone determines the width of its active space, it determines the probability of foragers staying within the pheromonal tunnel during their sinuous walk to reach the food source. Hence, an ant colony can easily modulate the accuracy of recruitment, by varying the quantity of trail pheromone deposited. If the food source is of high quality, a strong trail and its large active space will make its loss less likely by foragers and will guide nearly
all recruits to the feeding site. If the food source is of poor quality, a weak trail may result into more frequent exits from the active space, may lead to a more diffuse spatial distribution of workers and may promote an extensive search for new food sources over the foraging area (Deneubourg et al., 1983; Edelstein-Keshet et al., 1995). Similarly, at the beginning of recruitment, when the trail amount is still weak, the fraction of ants losing the trail is high and new food sources are likely to be discovered. The error level decreases as recruitment proceeds due to trail reinforcement by new recruits. The coupling of randomness in trail following behavior and competition between recruitment trails appears as a powerful strategy for optimizing foraging success.

An adaptive tuning of error has also been suggested as an interpretation of variations in the accuracy of honeybee dances (Towne and Gould, 1988). Information about food location differs in its level of accuracy as a function of the distance from the nest. At short distances (a few decameters), the dance is circular and contains no directional information. When food sources are remote, i.e. hundreds of meters from the hive, a bee recruiter is potentially able to code spatial information with a high level of accuracy through the direction of waggle runs. However, one curious feature of bee dancers is that their consecutive waggle runs may be produced with some directional scatter. A bee dancing for a less remote food source will perform relatively less precise waggle runs so that recruits will be distributed across several flower patches instead of being focused on where the communicating bee has been foraging (Weidenmuller and Seeley, 1999). The high-energy costs associated with flight may have favored a higher accuracy of communication about more remote food location. It is, however, suggested that imprecision in dance communication is caused primarily by physical constraints such as the ability of dancers to turn around quickly enough when the advertised site is nearby (Beekman et al., 2005; see also Tanner and Visscher, 2006). The accuracy of directional information of food sources also changes according to the size of the resource. When the target of recruitment is a large flower patch, the dances show a lower directional accuracy and a higher angular divergence of consecutive waggle runs than in dances coding for a more definite spot in space such as a nestbox. This tuning of the dances’ information content, which spreads recruits over a certain area, can be viewed as an adaptation to the typical spatial configuration of the recruitment targets (Weidenmuller and Seeley, 1999).

The modulation of the amount of trail pheromone laid by ants, as well as the fine-tuning of the angular divergences of bee dances, can both be viewed as striking adaptations for balancing the trade-off between an efficient exploitation of food and new food discoveries around the nest.
6 Spatial patterns

6.1 Bee lines and trail networks

In both ants and honeybees, the spatial distribution of foraging routes follows that of food resources: routes leading to highly rewarding food patches will be preferentially foraged, whereas routes leading to poor or exhausted resources will be neglected. There are, however, several major differences in the spatial foraging patterns displayed by ant colonies or hives. As a whole, the pattern of foraging routes displayed by the hive appears as a juxtaposition of straight lines (beelines) between the nest and the food sources (Fig. 9B). These beelines effectively translate information encoded in the waggle dance. Even though some local deviations may occur due to the imprecision of the communication or to physical obstacles, newly recruited bees fly directly from the hive to the vicinity of a food source. Then, in the final stages of their flight, recruits proceed to search for the exact location of the flowers, their flight may become sinuous and their spatial information may be completed by odor and visual cues that help in the localization of exploited flowers (Tautz and Sandeman, 2003; Riley et al., 2005; Arenas et al., 2007). This searching behavior close to the flower patch accounts for the fact that the arrival of recruits at the source is often later than would be expected for a direct flight between the hive and the food source.

A major feature of honeybee lines is that the trajectory of each forager is individual. Before leaving the hive, each recruit has received, through the recruitment dance, the whole information needed to locate a flower patch. Each recruit exiting the hive then takes a flying route of its own. Ideally, this route should connect the hive entrance to the food patch in the straightest way possible in order to minimize the energy expenditure associated with flight. Hence, the spatial foraging pattern of the hive can be seen as a bundle of hundreds of bee lines that radiate out of the hive entrance and that are more concentrated in the angular zones leading to exploited flower patches. The spatial distribution of beelines is thus expected to be highly flexible: it should reflect changes in the direction coded in waggle runs and should closely follow changes in the spatial distribution of flower patches (Fig. 9B).

In ants, the spatial configuration of the foraging trails is quite different from that of beelines. Instead of being the mere juxtaposition of individual routes, trails often organize themselves into a network. Depending on the species, ants minimize the traveling time (Dussutour et al., 2006) and/or the energy costs (Denny et al., 2001) so that the final pattern of the trail network is deeply influenced by a set of spatial factors as well as by the history of the foraging activity (e.g. food discoveries, level of area marking, etc.) (Deneubourg et al., 1990; Traniello et al., 1991; Fourcassie and
Deneubourg, 1994; Fourcassié et al., 2003). For instance, if a new food source becomes available close to a source that is already foraged by ants, scouts will be more likely to discover it, they will lay a trail and, through path integration, they will follow a rather straight trajectory towards the nest. On their homeward journey, scouts may cross a well-established trail leading to another exploited food patch and may then decide to join the main stream of workers going back to the nest. Over time, successive visits of the foragers to the new food source will lead to the emergence of a new lateral trail branched onto the main foraging trail (Hölldobler and Wilson, 1970, 1990; Hölldobler and Möglich, 1980; Detrain et al., 1991; Quinet and

FIG. 9  Schematic representation of ant trails and beelines. Larger circles stand for larger food sources or larger flower patches. (A) Ant trails: Several resources may be concurrently exploited but some food sources remain neglected. Most of the foraging activity is focused on one food source. If depleted, this source becomes less foraged while another one starts attracting the foragers. A lateral trail that initially connects a food source to the nest entrance may, after a while, partially fuse with the main stream of foragers. A new trail can also originate from a previously exploited food source. (B) Beelines: The spatial distribution of beelines fits that of flower patches and radiate around the nest entrance. Flights become less frequent towards locations in which food patches become depleted. The direction of beelines is flexible and may markedly change when flowers blossom at quite different locations.
For the simple case of two close food sources, two straight independent trails leading to each food source scarcely coexist and most often merge into a common trail. As a result, the foraging activity becomes more concentrated on fewer paths as one travels from the food sites towards the nest entrance. The geometry of trail bifurcations within the network provides polarity information to foragers joining a trail, and these foragers can re-orientate themselves if they initially walk in the wrong direction (Jackson et al., 2004). At the individual level, a tree-shaped trail network may lengthen the distance traveled by one forager, since its trajectory somewhat deviates from a straight line directly connecting the food source to the nest. However, at the society level, the building of a tree-shaped trail network greatly decreases the total trail length in comparison to a “star graphs” network with one direct trail going to each food source. Several benefits are associated with tree topologies: by reducing total network length, they facilitate the maintenance of foraging pathways (i.e. by actively cleaning or building construction around them; Howard, 2001), as well as their defense against competitors or predators.

The diversity of trail networks reported for ant species is a by-product of the interactions between trail-laying ants and any environmental factors that impede access to food sources or that may influence the probability of connecting two trails, such as the spatio-temporal availability of food resources (Deneubourg et al., 1989; Franks et al., 1991; Lopez et al., 1994; Detrain et al., 2000). For instance, two trails are more likely to merge into a common trunk trail when the two sources are persistent, close to each other, and/or remote from the nest. The higher sensitivity of the trail network to group-size effects is another major difference between ants and honeybees. In honeybees, an increase in the foraging effort will simply lead to an increase in the number of concurrent beelines. Conversely, in trail-laying ants, an increase in the number of workers may lead to qualitative changes in the spatial patterns of foraging routes. For instance, there is a shift from a diffuse foraging pattern to a tree-like trail network as the number of interacting foragers increases (Detrain et al., 1991; Nicolis and Deneubourg, 1999). Such dependence on colony size is the typical signature of a self-organized process of pattern formation.

6.2 SELECTION OF THE SHORTEST PATH

For walking ants that encounter large obstacles (stones, crevices, etc.) compared to their tiny size, recruitment trails not only prevent them from losing their way but also allow them to select the shortest route towards the feeding site. While the choice of one path is random over a bridge with two equal branches, most of the foragers follow the shortest path of a bridge with branches of unequal length (Goss et al., 1989; Vittori et al., 2006).
This adaptive selection of the shortest path occurs even though individual ants have no knowledge of relative path lengths, as well as no memory of the routes they have previously taken. For ants that lay a trail in both directions (from the nest to the food source and vice versa), this selection is simply due to the shorter trip duration and hence to the increased rate of trail reinforcement over the short path. In some ant species that lay a trail only during their homeward trip (Beckers et al., 1992b), the selection of the shortest path is also due to the U-turns made by workers that double back on a trail, return to the common branch point and then follow the second branch. Because when an ant is backtracking after its U-turn it does not lay down trail pheromone, the higher frequency of U-turns along the long path ultimately leads to a lower amount of pheromone over the longest branch. The adaptive selection of the shortest path thus arises “automatically” at the colony level from differences in the recruitment dynamics depending on branch length. This results from the synergistic effect of a shorter duration of foraging trips and of a lower frequency of U-turns on the rate of trail reinforcement.

It is difficult to find the counterpart for the honeybees. Firstly, the spatial information provided by the orientation of waggle dances is, by essence, the shortest flying route connecting the hive to the flower patch, since it results from a quite efficient path integration of the foraging journey by the recruiter (Collett and Collett, 2004, 2006; Collett et al., 2006). Nevertheless, in the presence of obstacles, short cuts and shorter-lasting foraging journeys may be achieved on an individual basis by honeybee foragers that are more experienced about how to reach the flower patch.

7 Conclusions

7.1 What about other group-living arthropods?

Collective decision-making can be as accurate and effective as decisions made by solitary animals, even by those famous for their well-developed capacities for information processing, such as vertebrates (Krebs and Davies, 1984). The strength of collective decision-making is that adaptive choices may arise automatically without the need for any leaders, which would have a synoptic view of all environmental opportunities. Consensus decisions can be reached even when the individuals do not know how the information they possess compares with that of conspecifics. Many examples of collective decision-making have been reported in the literature not only for ants or honeybees (Hölldobler and Wilson, 1990; Seeley, 1995; Theraulaz and Spitz, 1997; Bonabeau et al., 1997; Camazine et al., 2001), but also for other group-living organisms including bacteria, which may achieve quorum sensing and behave collectively (Miller and Bassler, 2001;
Ben Jacob et al., 2004). Likewise, complex patterns have been reported in fungal networks (Bebber et al., 2007) and slime-moulds (Shirakawa and Gunji, 2007). In this respect, group-living arthropods display amazing collective patterns and coordinate their efforts efficiently even though their social organization is less complex than that of social insects (Fitzgerald, 1995; Fitzgerald and Costa, 1999; Deneubourg et al., 2002; Costa, 2006). The choice of a common shelter or the cooperative exploitation of one resource by several individuals is then guided not only by cues related to the quality of these resources, but also by recruiting and/or arresting signals emitted by conspecifics. Hundred of examples of aggregative behavior, which rely on pheromones, have been reported for species living in very different ecological situations [e.g. Triatoma kissing bugs (Lorenzo and Lazzari, 1996); cockroaches (Miller and Koehler, 2000; Said et al., 2005); Drosophila fruit flies (Takahashi, 2006); lobsters (for a review see Wertheim et al., 2005; Horner et al., 2006)]. The active recruitment of conspecifics toward a common place can also be mechanically mediated, as has been shown in gregarious larvae of herbivorous insects, such as treehoppers, which use vibrations to orient themselves and eventually to move from low- to high-quality sites (Cocroft, 2005).

Although collective patterns of aggregation are widely reported in arthropods, there are surprisingly few experiments investigating how collective choices emerge among group members and how they are influenced by the ways conspecifics interact with each others. The use of silk impregnated with pheromone is one of the most studied long-range interactions in gregarious arthropods. Social caterpillars (Fitzgerald, 1995) and mites (Yano, 2008) may lay silk threads that attract conspecifics in a manner similar to ant trails. The tuning of the amount of pheromone released over the silk enables caterpillars to collectively select richer leaf patches as well as to reach a consensus about a common resting site at the end of a day’s activity (Fitzgerald and Peterson, 1983; Fitzgerald, 1995; Ruf et al., 2001). Moreover, like ants or stingless bees, forest tent caterpillars, Malacosoma disstria, show a lack of flexibility in their collective decision-making. Indeed, when faced with a nutritionally balanced and a less profitable, unbalanced source placed at opposite ends of a bridge, the colony of caterpillars as a whole, randomly discovers one of the two resources, goes on following the recruitment silk threads and becomes trapped for at least one day in the exploitation of the resource discovered first independently of its quality (Dussutour et al., 2007a,b).

Group-living spiders provide another striking example of collective decision-making. The strong correlation between the amount of exploratory webs and the localization of the spiders’ population suggests that conspecifics’ draglines are used as a recruitment means for colony formation. Actually, groups of Anelosimus studiosus spiders can only follow individuals sequentially released at a common spot when draglines
are left in place on the substrate (Furey, 1998). The silk network orients the movement of individual spiders: it acts as a recruiting structure and, due to the pattern of dragline attachments, it may lead in a binary choice setup to the collective selection of only one aggregation site (Saffre et al., 1999; Jeanson et al., 2004b).

The emergence of collective patterns does not necessarily imply the emission of recruiting signals and may also be triggered by short-range physical contact. For instance, the aggregation of cockroaches under a common shelter is due to their perception of contacts with conspecifics, in which the individual probability of moving decreases with the number of neighbors resting under a shelter (Ame et al., 2004, 2006; Jeanson et al., 2005; Halloy et al., 2007). Other spatial patterns (other than binary choices) may be seen as by-products of collective choices in which the group members choose between two or more mutually exclusive actions. For instance, in groups of locusts of increasing density, a transition occurs from a disordered movement of individuals within the group to a highly aligned collective motion. This transition may be explained by mechanisms and decision rules similar to those described previously (Buhl et al., 2006).

7.2 NON-LINEARITY OF INFORMATION FLOW: THE KEY-STONE OF COLLECTIVE DECISION-MAKING

The ways individuals interact with each other and give rise to collective responses vary, depending on the species concerned and may involve chemical signals, physical contacts or both types of stimuli (Camazine and Sneyd, 1991; Seeley et al., 1991; Visscher and Camazine, 1999a,b; Visscher, 2007). Insect societies such as the hive or the ant nest are real life models that offer a unique opportunity to investigate (1) how collective structures emerge in complex biological systems made up of numerous individuals, and (2) how collective behavior may be shaped by the properties of one species’ form of communication. Due to the specificities of chemical trails versus dances, ant nests and hives differ in the amplifying potential of their recruiting process. Likewise, in other group-living arthropods, a difference in the level of non-linearity of one individual’s response to recruiting stimuli may have major consequences on the collective patterns displayed by the whole group. Usually, chemical signals are characterized by a pheromone decay rate and a receiver response curve that are both highly non-linear. Such a non-linearity of information flow – as found in ants’ chemical trails – favors the focusing of the group at one or a few sites, but restrains the flexibility of the group to shift to more profitable resources. Hence, a non-linearity of receivers’ response is expected to be found in species that benefit from cooperation in the exploitation or in the defense of resources, as well as in species exploiting resources that are rather stable in time and space. Conversely, tactile signals such as the honeybee dances
are expected to show less non-linear properties than chemical ones since the rates of contacts should be proportional to the number of interacting individuals. The individual response to contact rates may nevertheless be non-linear as found in the case of nest-moving recruitment (Pratt et al., 2002; Seeley and Visscher, 2003; Jeanson et al., 2004a; Pratt and Sumpter, 2006) or in the case of aggregation patterns in ants (Gordon et al., 1993; Depickere et al., 2004; Nicolis et al., 2005) and cockroaches (Ame et al., 2004, 2006; Jeanson et al., 2005). Whatever the type of between-nestmate interactions, a low level of non-linearity favors an even distribution of individuals among identical resources. It also allows a closer fitting between the number of animals allocated to one resource and its relative profitability (as predicted by an ideal free distribution). Finally, it facilitates higher flexibility in-group decisions when environmental resources are changing. This should benefit species that (1) exploit resources, the availability of which is highly variable in time and space, (2) that do not need to monopolize resources against competitors, and (3) that do not need to cooperatively defend food sites because individuals are well protected against predators (e.g. by defensive secretions or effective weapons, such as in honeybees).

The evolution of communication in gregarious insects has been presumably steered by whether, for a particular colony sizes and habitat, a recruitment increases food collection, favors the monopolization of resources or prevents attacks by predators. Then, the payoff of one type of recruitment should vary depending on the foraging ecology of the species. In particular, recruitment mechanisms with a higher level of non-linearity seem well-adapted to species that benefit from cooperation and collective selection of a subset of resources at the expense of a lower flexibility to changing environmental conditions. In this respect, being able to shift between signals that differ in their level of non-linearity may be advantageous for one species. For example, within the same recruitment process, some ant species may change the way they communicate with each other: scouts that have discovered a new source first perform group recruitment to activate nestmates and thereafter shift towards mass recruitment by chemical trails. Group recruitment (such as tandem recruitment or honeybee dances) is characterized by a low level of non-linearity: the number of recruited foragers is largely proportional to the number of successful scouts, as the group of recruits has to be physically guided by a recruiting leader until they reach the food source (e.g. Tetramorium ant species, Verhaeghe, 1982; see also Hölldobler and Wilson, 1990; de Biseau et al., 1994). Over the course of food exploitation, mass recruitment and non-linear properties of chemical trail recruitment become prevalent, with most of the foragers being concentrated on a few food sources (Pasteels et al., 1987). This double recruitment process confers a higher flexibility to the ant nest. Any patrolling ant which has missed the
first food source and that discovers a more profitable one, can behave as a leader: it guides groups of recruits towards this new resource, counteracts the attractiveness of the chemical trail leading to the first exploited target and, as observed in honeybees, facilitates the shift of the foraging activity towards richer food sources (Beckers et al., 1990).

A higher complexity of communication is an additional means for a group-living species to improve flexibility, as well as diversity of collective patterns. In the case of foraging, recruitment signals are often a blend of molecules with different life-times, operating at different stages of food exploitation (see e.g. de Biseau et al., 1991; Detrain and Cammaerts, 1991). Another example is the synergistic effect of recruitment trail and home-range marking, the latter being ant density-dependent (Devigne and Detrain, 2002; Devigne et al., 2004; Devigne and Detrain, 2006): this means that workers are recruited to areas as a function of their frequenceation level and hence their potential interest as foraging sites. Foraging efficiency can also be improved if repellent pheromones are used to mark unrewarding areas (Giurfa, 1993; Stickland et al., 1999; Robinson et al., 2005). A challenge for future research will be to investigate to what extent the level of non-linearity of one species’ interactions matches its need for cooperative behavior or for flexibility to adapt to environmental changes.

7.3 WHAT ABOUT INDIVIDUALITIES AND THEIR IMPACT ON COLLECTIVE DECISION-MAKING?

Whatever their levels of sociality, group-living species share a common property: the potential for amplifying, at the group level, individual traits or individual preferences. For instance, when ants are given the choice on a diamond-shaped bridge between a branch with or without a wall, about 65% of the foragers will prefer to follow the walled branch in the absence of a recruitment trail. But, as recruitment proceeds, this propensity of ants will increase and will lead up to 85% of the foragers to choose the walled path (Dussutour et al., 2005a). This demonstrates that non-linearity in between-nestmate interactions may enhance even slight preferences of the individuals and lead to much more clear-cut choices by the group as a whole. Likewise, a synergy may exist between individual memory or fidelity to an area or to a given resource and the amplifying potential of recruitment; so that the specialization of a small number of recruiters will focus most of the colony in this particular sector or onto this food source. Amplifying phenomena that underlie most collective decisions are thus powerful means by which slight changes in physiological preferences (e.g. thermopreferendum values), changes in behavioral traits (e.g. thigmotaxis) or changes in individual knowledge of the environment can have deep consequences on the way a species will live as a group, shelter exclusively in
a peculiar resting site, travel over a determined foraging path, and focus its foraging effort on a given resource.

One may erroneously think about collective decision-making and idiosyncrasy as being opposite concepts (Deneubourg et al., 1999). In fact, individual memory and individual experience do not necessarily prevent the reaching of a consensus, but instead make the patterns that will emerge at the colony level more complex and diverse (Couzin et al., 2005). When “conflicts” exist between the preferences of different individuals, there are multiple possible outcomes. One is the splitting of the group with individuals possibly losing the benefits associated with being part of a large group (Krause and Ruxton, 2002) but with the group enlarging its occupation range of the environment. Second is the reaching of a consensus with a small proportion of informed individuals acting as leaders and guiding a group primarily composed of naïve individuals towards a target location. Research should now focus on how individualities may be enhanced at the group level by positive feedbacks and give rise to a multiplicity of adaptive collective patterns. As a corollary, one should question how species that benefit from maintaining group cohesion have evolved mechanisms resolving potential conflicts between different individual preferences. Part of the answer could be found in the ratio of recruiters (or informed individuals) versus naive ones, as well as in the level of non-linearity of their communication. Answers to these questions will ultimately provide new insights into the evolution of communication and cooperation in group-living animals.

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