



The protein balance of the honey bee worker

K Crailsheim

► To cite this version:

K Crailsheim. The protein balance of the honey bee worker. *Apidologie*, 1990, 21 (5), pp.417-429.
hal-00890873

HAL Id: hal-00890873

<https://hal.science/hal-00890873>

Submitted on 11 May 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

The protein balance of the honey bee worker

K Crailsheim

Institut für Zoologie an der Karl-Franzens-Universität, Universitätsplatz 2, A-8010 Graz, Austria

(Received 6 May 1990; accepted 15 June 1990)

Summary — For honey bee workers, protein requirements and consumption, digestion, intestinal absorption and utilisation all depend on the worker's age and functional status in the system of division of labor within the colony. Workers at the stage of nurses play a pivotal role not only in protein digestion but also in its distribution. They are well equipped for digestion of pollen, are rich in protein and produce large amounts of jelly, which they distribute to almost all members of the colony.

protein metabolism / nutrition / digestion / polyethism / *Apis mellifera*

INTRODUCTION

The honey bee is one of the few insects whose commercial importance lies in being cultivated, not exterminated. This led to intensive investigations of its nutritional needs. Many papers dealt with the types of food foraged by the colonies, the influence of artificial food or the fate of food-stuffs taken up by bees. Nevertheless, many questions concerning protein metabolism remain unanswered, due to the complexity and plasticity of the social system of the bees.

Temporal polyethism of the worker caste depends primarily on the age of the insect; the division of labor is also influenced by the availability of food inside and outside the hive, on the actual needs of the colony (Rösch, 1925, 1930; Lindauer, 1952; Ribbands, 1952; Free, 1965; Michener, 1969; Seeley, 1982; Winston and Neilson Punnet, 1982; Kolmes, 1985) and on season (Merz *et al*, 1979). Hormone levels influence the

beginning of different age-dependent tasks (Robinson, 1987; Robinson *et al*, 1989). In the last few years more and more genetically determined controlling mechanisms have been investigated (Calderone and Page, 1988; Frumhoff and Baker, 1988; Robinson and Page, 1988; Kolmes *et al*, 1989; Robinson *et al*, 1989; Rothenbuler and Page, 1989).

This review describes the protein metabolism of workers with special attention paid to their age and function.

PROTEINACEOUS NUTRIENTS

When worker larvae hatch, they are provided with jelly from the brood food glands of the nurses (Haner and Rembold, 1964; Rembold, 1974) until they reach an age of 3.5 d. Afterwards, they receive a mixture of jelly, honey and pollen; this latter food mixture has less protein but more carbohydrates (von Rhein, 1956; Kunert and Crailsheim, 1987). Approximately 1/3

of the dry weight of the jelly fed at first consists of high molecular weight proteins (Habowsky and Shuel, 1959; Patel *et al*, 1960). The percentage of protein in the food for the young larvae depends not only on the age of the larvae but is also subject to seasonal fluctuations: during 1 summer a decline from more than 50% (protein of dry weight) to less than 30% was demonstrated in 2 colonies investigated in parallel (Kunert and Crailsheim, 1987).

Soon after emergence, young bees start to consume honey and beebread from an area near their brood nest (Lotmar, 1938; Lindauer, 1952; Dietz, 1969; Haydak, 1970; Hagedorn and Moeller, 1967). Beebread is the pollen that has been collected by the foragers and prepared for storage in the combs. It is mixed with secretion from workers (Pain and Maugenet, 1966; Maurizio, 1969; Herbert and Shimanuki, 1978) and contains various microorganisms (Pain and Maugenet, 1966; Gilliam *et al*, 1989). The nutritive value of pollen stored in this manner is higher than that of laboratory-stored pollen (Beutler *et al*, 1949; Van Der Vorst and Jacobs, 1980). In addition, the newly emerged workers receive food from older bees in trophallactic interactions; Perepelova (1928) observed that bees of 1–3 d of age were fed by others whose age ranged from 7–14 d. Free (1957, 1959) found that the young bees also received food from workers of other ages; however, he also confirmed that nurses of 7–14 d were predominant in the group of food providers. From about the third day of life, the number of receiving and giving interactions are similar in number, but generally there is rather more transfer from old to young bees than in the opposite direction (Free, 1957). As the age class from 7–14 d provides the protein for the larvae, it could be assumed from the above findings that newly emerged bees receive not only hon-

ey but also protein-rich products of the nurse's brood glands.

These trophallactic transfers also occur when bees are caged (Free, 1957; Korst and Velthuis, 1982; Moritz and Hallmen, 1986). Such laboratory experiments were used to prove the transfer of proteinaceous food between nurses and 1 d-old bees (Crailsheim, 1990a). This transfer was also shown in small free-flying colonies (Crailsheim, 1990b).

Pollen is the main natural source of protein for bees; nectar collected by bees contains rather low concentrations of amino acids (Baker and Baker, 1973). The pollen consumption of the workers increases until they reach the age when they become nurses (Lotmar, 1938; Lindauer, 1952; Pain, 1961), after which pollen consumption tends to decrease.

During the winter when there is no food available outside the hive, bees consume the stored food, *ie* honey and beebread. Beebread is important for surviving the winter as well as for the start of colony repopulation in the spring (Wille *et al*, 1987).

One way of obtaining protein is cannibalism. In certain situations, such as lack of proteinaceous food, workers may eat eggs and brood (Haydak, 1935; Myser, 1952; Newton and Michl, 1974). This food can lengthen the lifespan of the cannibal bees themselves (Weiss, 1984), or it can be used for brood rearing or feeding the queen (Weiss, 1984; Webster and Peng, 1987; Webster *et al*, 1987).

QUALITY OF PROTEIN

That pollen is necessary for a colony has been known for a long time (Eckert, 1942). Two authors laid down the essential foundations for all subsequent work on protein requirements and metabolism in the early

fifties. Maurizio (1954) described the importance of pollen for the colonies and Groot (1953) described the nutritional value of various sources of protein for the colonies as well as the requirements of individual bees. He classified the following 10 amino acids as being essential for the growth of the honey bee: arginine, histidine, lysine, tryptophan, phenylalanine, methionine, threonine, leucine, isoleucine and valine; the largest requirements were for leucine, isoleucine and valine.

Thus, the nutritional value of the pollen for bees is primarily defined by its absolute and relative content of essential amino acids. Accordingly, pollen from different plants has a different nutritional value for bees. This could be demonstrated by measuring such diverse parameters as the growth and development of the fat body (Maurizio, 1954), the development of hypopharyngeal glands (Maurizio, 1954; Standifer, 1967), lifespan (Maurizio, 1954; Wahl, 1963; Schmidt *et al*, 1987) or brood rearing (Wahl, 1963; Herbert *et al*, 1970). Caged bees show the ability to distinguish between different kinds of pollen. They prefer protein-rich pollen (Schmidt and Johnson, 1984) and react to the presence of phagostimulants in pollen (Schmidt, 1985).

Less important for bees in nature than for artificial feeding of bees is the fact that the age of pollen (Dietz and Haydak, 1965; Hagedorn, 1968) and the way it was collected – *ie* whether the pollen was collected by humans or by bees (Maurizio, 1954) – influences its value for bees.

INTESTINAL TRANSPORT, DIGESTION AND ABSORPTION

Pollen is ingested either fresh or from bee-bread. There is no digestion of protein in the crop, as salivary and hypopharyngeal

glands do not produce proteolytic enzymes (Arnold and Delage-Darchen, 1978). Amino acids from nectar do not penetrate the crop wall (Crailsheim, 1988a). Pollen grains are quickly transported through the crop to the proventriculus which regulates the passage of all nutrients into the midgut. In agreement with their intensive pollen consumption, the worker caste has the largest proventriculus as compared to the drones and to the queen (Pabst and Crailsheim, 1990). Transport of liquids is regulated according to requirements (Crailsheim, 1988b, c); solid food, *ie* pollen, is compressed into bolusses by the proventriculus which are then transported through the stomodeal valve, a flexible pipe (Trappman, 1923) into the midgut (Whitcomb and Wilson, 1929; Bailey, 1952; Schreiner, 1952; Barker and Lehner, 1972; Klungness and Peng, 1984a; Peng and Marston, 1986). There, the pollen bolusses are surrounded by peritrophic membranes. They do not leave the tubule of peritrophic membranes (endoperitrophic space) but pass through the anterior intestine into the rectum (Schreiner, 1952). The time required for passage through the midgut depends on the amount of foodstuffs in this section, on the age of the bee and on the way the bees are kept; the length of time from ingestion until pollen, found in the rectum ranged from a few h up to more than 1 d. The long duration is necessary because pollen is difficult to digest, due to its tough wall (Barker and Lehner, 1972; Klungness and Peng, 1984b; Peng *et al*, 1986). Some types of pollen leave the midgut without being utilized completely (Peng *et al*, 1985). For a pollen mixture, offered in a formulated diet, an apparent digestibility of 77% was shown by Schmidt and Buchmann (1985).

The peritrophic membranes form compartments for digestion (Moritz and Crailsheim, 1987) as also described for other

insects (Terra *et al*, 1979; Terra and Ferreira, 1981; Terra, 1988, 1990) and contain immobilized aminopeptidases (Peters and Kalnins, 1985). The chemical composition of the membranes depends on the age of the bees (Pabst *et al*, 1988).

Upon entering the midgut, pollen is thought to be broken down by osmotic shock (Kroon *et al*, 1974) and/or by the effects of proteolytic enzymes. In another bee (*Melipona quadrifasciata*), osmotic shock is less pronounced and only causes an opening of the pores on the pollen grains (Martinho, 1975). Gut proteolytic enzymes are only present to a limited extent in the last pupal stage and in emerging bees, but they increase rapidly in the first hours of imaginal life (Moritz and Crailsheim, 1987). Pavlovsky and Zarin (1922) were the first to investigate extensively the proteolytic enzymes in the gut of the honey bee; they found several types. Giebel *et al* (1971) described 4 endopeptidases with different properties. Dahlman *et al* (1978) only found 3 endopeptidases, 1 trypsin-like and 2 chymotrypsin-like. Grogan and Hunt (1980, 1984) first proved differences in gut proteolytic activity between house bees (those who did not yet forage) and foragers; Moritz and Crailsheim (1987) showed age-dependence and a close correlation with soluble protein content in the midgut. Trypsin-like and chymotrypsin-like enzymes showed their maximum activity at pH 8.5 and around pH 8 respectively. The highest levels of proteolytic activity were found in the nurse bees; this activity declined when bees started to forage. Low levels were found in bees in winter; the same population of bees showed a slight increase when bees again began to forage and to perform brood-care tasks in spring. Proteolytic activity in the gut of workers is also influenced by the exigency of the bees' social system. Neither a queen nor open brood was necessary for normal de-

velopment of intestinal proteolytic activity when bees lived in colonies, but proteolytic levels were much lower if bees were caged, even in groups and with sufficient proteinaceous food. The composition of the diet also influenced proteolytic levels (Crailsheim and Stolberg, 1989).

Only little proteolytic activity was found in the midgut tissue, somewhat more in the ectoperitrophic space and most in the endoperitrophic space where pollen is transported (Moritz and Crailsheim, 1987; Jimenez and Gilliam, 1989). The latter authors postulate a countercurrent flow in the endo- and ectoperitrophic spaces as described for other insects (Terra and Ferreira, 1981; Terra *et al*, 1985; Terra, 1988, 1990). Such a flow enables the enzymes to be utilized optimally and explains why only small amounts of liquid pass from midgut into the rectum of bees (Crailsheim, 1988b).

There is less information about the absorption of substances through the midgut wall of the honey bee than for other insects (Turunen, 1985). The transport of sugars was shown to be passive by simple diffusion (Crailsheim, 1988d). Subsequent *in vivo* experiments demonstrated the concentration dependence of leucine transport as well as inhibition of this transport by the structurally similar amino acid isoleucine, but not by glycine, arginine or glutamic acid (Crailsheim, 1988a). This finding of a rather specific carrier mechanism could be confirmed by *in vitro* experiments which furthermore proved that the transport of leucine is not energy-dependent (Haszonits and Crailsheim, 1990). *In vivo* as well as *in vitro* experiments had previously shown different transport kinetics in bees in summer and in winter; transport being faster in summer. These results are in good agreement with data about protein metabolism. Morphological studies offer an explanation for the lower transport capacity

for amino acid in winter. Freeze-etch studies showed a lower number of particles in the plasma membrane of midgut epithelial cells in winter bees (Pabst and Crailsheim, 1987); such particles are described in mammals as being carriers or membrane-bound enzymes.

PROTEIN UTILISATION AND TURNOVER

Large amounts of protein are required during larval development; 1 larva needs about 50 mg jelly to reach a weight of 35 mg (von Rhein, 1956). During about 3.5 d, the high protein content of a 3 mg-larva is reduced from nearly 19% (of fresh weight), to about 9% (Kunert and Crailsheim, 1987).

If the nursing bees themselves lack proteinaceous food, they can continue to nurse larvae, but only for about 1 week. All larvae fed after this period were found to be unable to reach maturity and were removed by the bees. Protein-deprived nurse bees had lower nitrogen levels in all 3 body sections, as compared to nurses before the onset of protein deprivation. The largest reduction (~11%) was seen in the abdomen. Haydak (1935) concludes from these results that the nurses use their own tissue protein for rearing the larvae. Larvae that were nursed deficiently but reached maturity, developed into dwarfed adults (Jay, 1964).

After emergence, bees require protein for imaginal tissue differentiation. They consume a great deal of pollen and receive proteinaceous food from other workers. Upon emerging, they have a nitrogen content of 1.74 mg; this increases to 2.65 mg in 21–38 d-old bees (Haydak, 1934). A comparison of protein- or nitrogen-content-values of emerging workers taken from different authors with different methods is giv-

en by Kunert and Crailsheim (1988). The protein content at emergence depends on the availability of food outside the hive during the nursing season; the difference between bees nursed during bad pollen harvesting conditions and between those nursed during good conditions amounts to more than 13%.

At the moment of emergence, bees which were destined to survive the winter had a higher protein content than summer bees (Kunert and Crailsheim, 1988). They are reared during late summer and autumn, a period with less pollen available outside the hive but with considerable amounts of beebread inside. Furthermore, much fewer larvae are reared compared to the period from May to July.

During the hive-bound period, the most remarkable protein increase was found in the heads; nurses had about twice the amount found in newly emerged bees (Haydak, 1934). This is due to the system of work allotment in the honey bee community. Workers at the age of nurses have well-developed hypopharyngeal glands (Krathky, 1931; Moskovljevic-Filipovic, 1952; Groot, 1953; Maurizio, 1954; Beams *et al*, 1959; Fluri *et al*, 1982; Mortiz and Crailsheim, 1987; Fergusson and Winston, 1988; Suzuki, 1988; Crailsheim and Stolberg, 1989; Knecht and Kaatz, 1990) which are superbly equipped for synthesis of protein-rich secretions (Brouwers, 1982; Takenaka and Kaatz, 1987; Webster *et al*, 1987; Webster and Peng, 1987; Knecht and Kaatz, 1990; Crailsheim, 1990a). The secretion products of these glands are then distributed throughout the colony. The development of these glands is stimulated by a signal from the brood (Huang and Otis, 1989; Huang *et al*, 1989) and suppressed by juvenile hormone (Jaycox *et al*, 1974; Fluri *et al*, 1982; Liu, 1989) at least at high dosages (Sasagawa *et al*, 1989). Workers with the highest rates of protein

synthesis also exhibit the highest protein levels (4.3%) in the haemolymph; this is reduced to 1.7% in foragers (Sinizki and Lewtschenko, 1971). In winter the protein levels are again higher than in summer; this is due to a lower titer of juvenile hormone (Fluri *et al*, 1982). In summer, when workers stop producing jelly upon reaching the age of foragers, the shrunken hypopharyngeal glands produce other proteins, mainly enzymes (Simpson *et al*, 1968; Halberstadt, 1980; Sasagawa *et al*, 1989). The correct development of hypopharyngeal glands depends on proper nutrition (Soudek, 1927; Free, 1961; Maurizio, 1962; Standifer, 1967; Herbert *et al*, 1977) and on the existence of a social system (Crailsheim and Stolberg, 1989). When Schmidt and Buchmann (1985) offered such a proper pollen-containing diet to a small colony in a flight cage, they determined an average need of 3.07 mg N for 1 bee of the colony per 28 d. This corresponds to 19.2 mg protein.

The fat body is the general storage organ in the bee larva and contains large amounts of protein. Its composition changes with age (Krajewska and Hryniewiecka-Szyfter, 1988). Subsequently, it provides material for the newly developing imaginal tissues in pupae (Schnelle, 1923). After emergence, the protein content of the fat body increases until the worker reaches the nurse stage in summer. Protein content is then lower in foragers and highest in bees that survive the winter (Shehata *et al*, 1981; Fluri and Bogdanov, 1987; Southwick, 1989). Winter bees have higher protein levels in the haemolymph and more protein in their hypopharyngeal glands than foragers in summer (Fluri *et al*, 1982). Thus, both compartments function, together with the fat body, as protein storing organs. Many proteins circulating in the haemolymph are produced by the fat body (Lensky and Rakover, 1983).

Injections of radioactively labelled amino acids or feeding of labelled protein enable us to determine the site and rate of protein synthesis as well as catabolism. The high activity of hypopharyngeal glands was demonstrated *in vivo* with tracer methods (Webster and Peng, 1987; Webster *et al*, 1987; Knecht and Kaatz, 1990; Crailsheim, 1990a). Ninety min after an injection of ^{14}C -phenylalanine, nurses had more than 25% of the radioactivity in the protein fraction of their caput, less than 10% in the thorax and more than 25% in the abdomen. Foragers showed similar results in the thorax and abdomen, but contained somewhat less than 10% in the caput. More than half of the tracer found in the protein fraction of the head of nurses was in the hypopharyngeal glands after 6 h (Crailsheim, 1990a).

The half-life of total body protein was shortest in foragers (11–13 d), was about 50% longer in nurses and was about 4 times longer in bees in winter that were taken from the outside of a wintercluster (Crailsheim, 1986). The reduction of metabolism in winter bees (Omholt and Lonvik, 1986) is a useful adaptation to the lessening of the duties of these bees. Many of them remain almost motionless during long periods on the outside of the cluster and therefore should not need much protein turnover. Nurses in summer that have to do the work of digestion for a large part of the colony need much protein and have a faster turnover. Their enzymatic equipment is adapted to this level of protein metabolism.

Foragers are highly active flyers and cover on average 21.5 km per d (Neukrich, 1982) which requires the metabolic equivalent of more than 6 mg sugar solely for flight (Heran and Crailsheim, 1988); such an amount of metabolic work is certainly bound to increase protein turnover. Accordingly, they are able to absorb more

leucine and probably more other amino acids than winter bees (see previous section).

Honey bees are known to use carbohydrates for their energy requirements during flight (Nachtigall *et al.*, 1989). Nevertheless, a considerable fraction of ^{14}C from ^{14}C -amino acids was shown to be exhaled as $^{14}\text{CO}_2$ (Webster *et al.*, 1987; Crailsheim, 1988a, 1990a) in bees that were resting or walking, showing that amino acids are ultimately broken down and utilized for energy metabolism. The latter function in other situations than flight requires further investigation.

ACKNOWLEDGMENTS

I acknowledge the critical reading of the manuscript by A. Buccics and the help with the literature by B. Schöffmann. Experiments carried out at the Department for Zoology in Graz were supported by the Fonds zur Förderung der Wissenschaftlichen Forschung, project nos 5674 and 7069.

Résumé — Le métabolisme des protéines chez l'ouvrière d'abeille. Il existe, au sein de la caste de ouvrières, des différences dans le métabolisme des protéines dues à des facteurs tels que l'alimentation, l'âge des abeilles, leur fonction au sein de la colonie.

Les abeilles récemment écloses consomment du pollen et du miel en quantités croissantes et reçoivent également de la nourriture larvaire de la part des congénères de la ruche plus âgées. Leur teneur en protéines s'élève, pendant cette période, puis reste à un niveau élevé jusqu'à la fin de leur vie. La teneur en protéases intestinales et la taille des glandes hypopharyngiennes augmentent continuellement.

A l'âge de 8 j, dans une colonie moyenne et dans des conditions normales de miellée, elles occupent la fonction de nourrices, pour laquelle elles consomment d'énormes quantités de pollen. Elles sont alors abondamment pourvues de protéases intestinales et le transport des acides aminés de l'intestin dans l'hémolymphe est réalisé par des mécanismes spécifiques. Ce transport – qui a été étudié pour la leucine – se passe de la même façon chez les nourrices et les butineuses, mais il est plus lent chez les abeilles de la grappe hivernale. Les glandes nourricières des nourrices sont bien développées et produisent une nourriture larvaire riche en protéines, distribuée au couvain et à la reine, mais aussi aux ouvrières de tous âges. Quand les abeilles passent au stade de butineuses, leurs glandes hypopharyngiennes diminuent et changent de fonction.

La circulation des protéines corporelles des nourrices est semblable à celle des butineuses pendant la période de miellée, mais elle est beaucoup plus rapide que chez les abeilles d'hiver.

métabolisme des protéines / nutrition / digestion / polyéthisme / *Apis mellifera*

Zusammenfassung — Der Proteinstoffwechsel von Arbeiterinnen der Honigbiene. Innerhalb der Kaste der Arbeiterinnen bestehen Unterschiede im Eiweißstoffwechsel, die von Faktoren wie Ernährung, individuelles Alter und Funktion innerhalb der Stockgemeinschaft beeinflusst werden. Frisch geschlüpfte Bienen fressen in zunehmendem Maße Pollen und Honig und werden auch von älteren Stockgenossinnen mit Futtersaft gefüttert. Sie erhöhen ihren Eiweißgehalt während dieser Zeit, der dann bis zu ihrem Lebensende auf hohem Niveau bleibt. Der Gehalt an intestinalen Proteasen und die Größe der Hypopharynxdrüsen nimmt kontinuierlich zu.

Im Alter von etwa 8 Tagen haben sie in einem durchschnittlichen Volk bei normalen Trachtverhältnissen die Funktion von Ammen, in der sie sehr große Mengen an Pollen fressen. Sie sind hierzu reichlich mit intestinalen Proteasen ausgestattet, und der Transport von Aminosäuren aus dem Darm in die Hämolymphe wird durch spezielle Trägermechanismen bewerkstelligt. Dieser Transport – untersucht für die Aminosäure Leucin – geschieht bei Ammenbienen und bei Sammlerinnen in ähnlicher Weise, bei Bienen die in der Wintertraube leben jedoch langsamer. Die Futtersaftdrüsen der Ammen sind groß und produzieren eiweißreichen Futtersaft, der an die Brut, die Königin, aber auch an Arbeiterinnen aller Altersstufen verfüttert wird. Beim Übergang zur Ausflugsfähigkeit werden die Hypopharynxdrüsen kleiner und ändern ihre Funktion.

Der Umsatz des Körperproteins von Ammenbienen ist während der Trachtzeit dem von Flugbienen ähnlich, er geschieht aber wesentlich rascher als bei Arbeiterinnen, die im Winter leben.

Proteinstoffwechsel / Ernährung / Verdauung / Polyethismus / *Apis mellifera*

REFERENCES

- Arnold G, Delage-Darchen B (1978) Nouvelles données sur l'équipement enzymatique des glandes salivaires de l'ouvrière d'*Apis mellifica* (Hymenoptère, Apidae). *Ann Sci Nat Zool* 20, 401-422
- Bailey L (1952) The action of the proventriculus of the worker honeybee. *Apis mellifera* L. *J Exp Biol* 29, 310-326
- Baker HG, Baker I (1973) Amino-acids in nectar and their evolutionary significance. *Nature* 241, 543-545
- Barker RJ, Lehner Y (1972) A look at honeybee gut functions. *Am Bee J* 112, 336-338
- Beams HW, Tahmisiyan TN, Anderson E, Devine RL (1959) An electron microscope study on the pharyngeal glands of the honeybee. *J Ultrastruct Res* 3, 155-170
- Beutler R, Opfinger E, Wahl O (1949) Pollen-ernährung und Nosemabefall der Honigbiene. *Z Vergl Physiol* 32, 383-421
- Brouwers EVM (1982) Measurement of hypopharyngeal gland activity in the honeybee. *J Apic Res* 21, 193-198
- Calderone NW, Page RE Jr (1988) Genotypic variability in age polyethism and task specialization in the honey bee, *Apis mellifera* (Hymenoptera: Apidae). *Behav Ecol Sociobiol* 22, 17-25
- Crailsheim K (1986) Dependence of protein metabolism on age and season in the honeybee (*Apis mellifica carnica* Pollm.). *J Insect Physiol* 32, 629-634
- Crailsheim K (1988a) Transport of leucine in the alimentary canal of the honeybee (*Apis mellifera* L.) and its dependence on season. *J Insect Physiol* 34, 1093-1100
- Crailsheim K (1988b) Regulation of food passage in the intestine of the honeybee (*Apis mellifera* L.). *J Insect Physiol* 34, 85-90
- Crailsheim K (1988c) Intestinal transport of glucose solution during honeybee flight. In: *Biona report 6* (Nachtigall W, ed) Gustav Fischer, New York, 119-128
- Crailsheim K (1988d) Intestinal transport of sugars in the honeybee (*Apis mellifera* L.). *J Insect Physiol* 34, 839-845
- Crailsheim K, Stolberg E (1989) Influence of diet, age and colony condition upon intestinal proteolytic activity and size of the hypopharyngeal glands in the honeybee (*Apis mellifera* L.). *J Insect Physiol* 35, 595-602
- Crailsheim K (1990a) Protein synthesis in the honeybee (*Apis mellifera* L.) and trophallactic distribution of jelly among imago in laboratory experiments. *Zool Jahrb* (in press)
- Crailsheim K (1990b) Interadult feeding of jelly in honeybee colonies (*Apis mellifera* L.). *J Comp Physiol* (in press)
- Dahlmann B, Jany KD, Pfeleiderer G (1978) The midgut endopeptidases of the honey bee (*Apis mellifica*): comparison of the enzymes in different ontogenetic stages. *Insect Biochem* 8, 203-211

- Dietz A, Haydak MH (1965) Causes of nutrient deficiency in stored pollen for development of newly emerged honey bees. *Proc Int Jubilee Beekeeping 20th Congr Bucharest*, Apimondia Publ House, Bucharest, 238-241
- Dietz A (1969) Initiation of pollen consumption and pollen movement through the alimentary canal of newly emerged honey bees. *Ann Entomol Soc Am* 62, 43-46
- Eckert JE (1942) The pollen required by a colony of honeybees. *J Econ Entomol* 35, 309-311
- Fergusson LA, Winston ML (1988) The influence of wax deprivation on temporal polyethism in honey bee (*Apis mellifera* L.) colonies. *Can J Zool* 66, 1997-2001
- Fluri P, Bogdanov S (1987) Age dependence of fat body protein in summer and winter bees (*Apis mellifera*) In: *Chemistry and Biology of Social Insects* (Eder J, Rembold H, eds) J Peperny, München, 170-171
- Fluri P, Lüscher M, Wille H, Gerig L (1982) Changes in weight of the pharyngeal gland and haemolymph titres of juvenile hormone, protein and vitellogenin in worker honey bees. *J Insect Physiol* 28, 61-68
- Free JB (1957) The transmission of food between worker honeybees. *Br J Anim Behav* 5, 41-47
- Free JB (1959) The transfer of food between the adult members of a honeybee community. *Bee World* 40, 193-201
- Free JB (1961) Hypopharyngeal gland development and division of labour in honey-bee (*Apis mellifera* L.) colonies. *Proc R Entomol Soc Lond* 36A, 5-8
- Free JB (1965) The allocation of duties among worker honeybees. *Symp Zool Soc Lond* 14, 39-59
- Frumhoff PC, Baker J (1988) A genetic component to division of labour within honey bee colonies. *Nature* 333, 358-361
- Giebel W, Zwilling R, Pfeleiderer G (1971) The evolution of endopeptidases - XII. The proteolytic enzymes of the honeybee (*Apis mellifica* L.). *Comp Biochem Physiol* 38B, 197-210
- Gilliam M, Prest DB, Lorenz BJ (1989) Microbiology of pollen and bee bread: taxonomy and enzymology of molds. *Apidologie* 20, 53-68
- Grogan DE, Haunt JH (1980) Age correlated changes in midgut protease activity of the honeybee, *Apis mellifera* (Hymenoptera: Apidae). *Experientia* 36, 1347-1348
- Grogan DE, Hunt JH (1984) Chymotrypsin-like activity in the honeybee midgut: patterns in a three-year study. *J Apic Res* 23, 61-63
- Groot de AP (1953) Protein and amino acid requirements of the honeybee (*Apis mellifica* L.). *Physiol Comp Ecol* 3, 197-285
- Habowsky J, Shuel RW (1959) Separation of the protein constituents of the larval diets of the honey bee by continuous flow electrophoresis. *Can J Zool* 37, 957-964
- Hagedorn HH, Moeller FE (1967) The rate of pollen consumption by newly emerged honeybees. *J Apic Res* 6, 159-162
- Hagedorn HH (1968) Effect of the age of pollen used in pollen supplements on their nutritive value for the honeybee I. Effect on thoracic weight, development of hypopharyngeal glands, and brood rearing. *J Apic Res* 7, 89-95
- Halberstadt K (1980) Elektrophoretische Untersuchungen zur Sekretionstätigkeit der Hypopharynxdrüse der Honigbiene (*Apis mellifera* L.). *Insectes Soc* 27, 61-77
- Hanser G, Rembold H (1964) Analytische und histologische Untersuchungen der Kopf- und Thoraxdrüsen bei der Honigbiene *Apis mellifera*. *Z Naturforsch* 19B, 938-943
- Haszonits O, Crailsheim K (1990) Uptake of L-leucine into isolated enterocytes of the honeybee (*Apis mellifera* L.) depending on season. *J Insect Physiol* (in press)
- Haydak MH (1934) Changes in total nitrogen content during the life of the imago of the worker honeybee. *J Agric Res* 49, 21-28
- Haydak MH (1935) Brood rearing by honeybees confined to a pure carbohydrate diet. *J Econ Entomol* 28, 657-660
- Haydak MH (1970) Honey bee nutrition. *Annu Rev Entomol* 15, 143-156
- Heran H, Crailsheim K (1988) Energy requirements in bees (*Apis mellifera carnica* Pollm) in free flight, with and without additional load. *Veröffentlichungen der Universität Innsbruck*, 167, p 77
- Herbert EW, Bickley WE, Shimanuki H (1970) The brood-rearing capability of caged honey bees fed dandelion and mixed pollen diets. *J Econ Entomol* 63, 215-218

- Herbert EW Jr, Shimanuki H (1978) Chemical composition and nutritive value of the bee-collected and bee-stored pollen. *Apidologie* 9, 33-40
- Herbert EW Jr, Shimanuki H, Caron D (1977) Optimum protein levels required by honey bees (Hymenoptera, Apidae) to initiate and maintain brood rearing. *Apidologie* 8, 141-146
- Huang ZY, Otis GW (1989) Factors determining hypopharyngeal gland activity of worker honey bees (*Apis mellifera* L.). *Insectes Soc* 36, 264-276
- Huang ZY, Otis GW, Teal PEA (1989) Nature of brood signal activating the protein synthesis of hypopharyngeal gland in honey bees, *Apis mellifera* (Apidae: Hymenoptera). *Apidologie* 20, 455-464
- Jay SC (1964) Starvation studies of larval honey bees. *Can J Zool* 42, 455-462
- Jaycox ER, Skowronek W, Guynn G (1974) Behavioral changes in worker honey bees (*Apis mellifera*) induced by injection of a juvenile hormone mimic. *Ann Entomol Soc Am* 67, 529-534
- Jimenez DR, Gilliam M (1989) Age-related changes in midgut ultrastructure and trypsin activity in the honey bee, *Apis mellifera*. *Apidologie* 20, 287-303
- Klungness LM, Peng YS (1984a) Scanning electron microscope observation of pollen food bolus in the alimentary canal of honeybees (*Apis mellifera* L.). *Can J Zool* 62, 1316-1319
- Klungness LM, Peng YS (1984b) A histochemical study of pollen digestion in the alimentary canal of honeybees (*Apis mellifera* L.). *J Insect Physiol* 30, 511-521
- Knecht D, Kaatz HH (1990) Patterns of larval food production by hypopharyngeal glands in adult worker honey bees. *Apidologie* 21 (5), 457-468
- Kolmes SA (1985) A quantitative study of the division of labour among worker honey bees. *Z Tierpsychol* 68, 287-302
- Kolmes SA, Winston ML, Fergusson LA (1989) The division of labor among worker honey bees (Hymenoptera: Apidae): The effects of multiple patriline. *J Kansas Entomol Soc* 62, 80-95
- Kost PJAM, Velthuis HHW (1982) The nature of trophallaxis in honeybees. *Insectes Soc* 29, 209-221
- Krajewska K, Hryniewiecka-Szyfter Z (1988) Histological changes in the fat body of *Apis mellifera* L. during larval and pupal development. *Bull Soc Amis Sci Lett Poznan Ser D Sci Biol* 27, 25-36
- Kratky E (1931) Morphologie und Physiologie der Drüsen in Kopf und Thorax der Honigbiene (*Apis mellifica* L.). *Z Wiss Zool* 139, 120-200
- Kroon GH, Praagh van JP, Velthuis HHW (1974) Osmotic shock as a prerequisite to pollen digestion in the alimentary tract of the worker honeybee. *J Apic Res* 13, 177-181
- Kunert K, Crailsheim K (1987) Sugar and protein in the food for honeybee worker larvae. In: *Chemistry and Biology of Social Insects* (Eder J, Rembold H, eds) J Peperny, München, 164-165
- Kunert K, Crailsheim K (1988) Seasonal changes in carbohydrate, lipid and protein content in emerging worker honeybee and their mortality (*Apis mellifera* L.). *Comp Biochem Physiol* 75B, 607-615
- Lensky Y, Rakover Y (1983) Separate protein body compartments of the worker honeybee (*Apis mellifera* L.). *Comp Biochem Physiol* 75B, 607-615
- Lindauer M (1952) Ein Beitrag zur Frage der Arbeitsteilung im Bienenstaat. *Z Vergl Physiol* 34, 299-345
- Liu TP (1989) Juvenile hormone III induced ultrastructural changes in the hypopharyngeal glands of honeybee *Apis mellifera* L. (Hymenoptera: Apidae) without and with infection by *Nosema apis* Zander (Microsporidiae: Nosematidae). *Int J Insect Morphol Embryol* 18, 73-83
- Lotmar R (1938) Untersuchungen über den Eisenstoffwechsel der Insekten, besonders der Honigbienen. *Rev Suisse Zool* 45, 237-271
- Martinho MR (1975) Beiträge zur Untersuchung der Verdaulichkeit der Pollenkörner für *Melipona quadrifasciata*, *Anthidioides lepeletier* (Hymenoptera, Apidae, Meliponinae). 25th Int Apic Congr Grenoble France, Apimondia Publ House, Bucharest, 326

- Maurizio A (1954) Pollenernährung und Lebensvorgänge bei der Honigbiene (*Apis mellifica* L). *Landwirtsch Jahrb Schweiz* 62, 115-182
- Maurizio A (1959) Factors influencing the life-span of bees. *Ciba Foundation Symp* 231-243
- Maurizio A (1962) Zuckerabbau unter der Einwirkung der invertierenden Fermente in Pharynxdrüsen und Mitteldarm der Honigbiene (*Apis mellifica* L) 5. Einfluß von Alter und Ernährung der Biene auf die Fermentaktivität der Pharynxdrüse. *Ann Abeille* 5, 215-232
- Merz R, Gerig L, Wille H, Leuthold R (1979) Das Problem der Kurz- und Langlebigkeit bei der Ein- und Auswinterung im Bienenvolk (*Apis mellifica* L): Eine Verhaltensstudie. *Rev Suisse Zool* 86, 663-671
- Michener CD (1969) Comparative social behavior of bees. *Annu Rev Entomol* 14, 299-342
- Moritz B, Crailsheim K (1987) Physiology of protein digestion in the midgut of the honeybee (*Apis mellifera* L). *Insect Physiol* 33, 923-931
- Moritz RFA, Hallmen M (1986) Trophallaxis of worker honeybees (*Apis mellifera* L) of different ages. *Insectes Soc* 33, 26-31
- Moskovljevic-Filipovic V (1952) The development of the pharyngeal glands of the honeybee in the normal bee colony. *Bull Acad Serbe Sci* 4, 257-262
- Myser WC (1952) Ingestion of eggs by honey bee workers. *Am Bee J* 92, 67
- Nachtigall W, Rothe U, Feller P, Jungmann R (1989) Flight of the honey bee. III. Flight metabolic power calculated from gas analysis, thermoregulation and fuel consumption. *J Comp Physiol* 158B, 729-737
- Neukirch A (1982) Dependence of the life span of the honeybee (*Apis mellifica*) upon flight performance and energy consumption. *J Comp Physiol* 146B, 35-40
- Newton DC, Michl DJ (1974) Cannibalism as an indication of pollen insufficiency in honeybees: ingestion or recapping of manually exposed pupae. *J Apic Res* 13, 235-241
- Omholt SW, Lonvik K (1986) Heat production in the winter cluster of the honeybee, *Apis mellifera*. A theoretical study. *J Theor Biol* 120, 447-456
- Pabst MA, Crailsheim K (1987) Freeze-etch study of age-dependent changes in midgut epithelial cell surfaces of the honeybee. *Eur J Cell Biol* 43, 25
- Pabst MA, Crailsheim K (1990) The proventriculus of worker honeybee pupae, adult workers, drones and queens (*Apis mellifera* L). *Zool Jahrb* 94, 271-289
- Pabst MA, Crailsheim K, Moritz B (1988) Age-dependent histochemical changes in the peritrophic membranes of the honeybee *Apis mellifera* (Hymenoptera: Apidae). *Entomol Gen* 14, 1-10
- Pain J, Maugenet J (1966) Recherches biochimiques et physiologiques sur le pollen emmagasiné par les abeilles. *Ann Abeille* 9, 209-236
- Patel NG, Haydak M, Gochnauer TA (1960) Electrophoretic components of the proteins in honeybee larval food. *Nature* 186, 633-634
- Pavlovsky EN, Zarin EJ (1922) On the structure of the alimentary canal and its ferments in the bee (*Apis mellifera* L). *Q J Microsc Sci* 56, 509-556
- Peng YS, Marston JM (1986) Filtering mechanism of the honey bee proventriculus. *Physiol Entomol* 11, 433-439
- Peng YS, Nasr ME, Marston JM (1985) The digestion of dandelion pollen by adult worker honeybees. *Physiol Entomol* 10, 75-82
- Peng YS, Nasr ME, Marston JM (1986) Release of alfalfa, *Medicago sativa*, pollen cytoplasm in the gut of the honey bee, *Apis mellifera* (Hymenoptera: Apidae). *Ann Entomol Soc Am* 79, 804-807
- Perepelova LI (1928) The nurse bees. As cited in: Free JB (1965) *Opuit Pas* 12, 551-557
- Peters W, Kalnins M (1985) Aminopeptidases as immobilized enzymes on the peritrophic membranes of insects. *Entomol Gen* 11, 25-32
- Rembold H (1974) Die Kastenbildung bei der Honigbiene, *Apis mellifica* L aus biochemischer Sicht. In: *Sozialpolymorphismus bei Insekten* (Schmidt GH, ed) Wissenschaftliche Verlagsgesellschaft, Stuttgart
- von Rhein W (1956) Über die Ernährung der Arbeitermade von *Apis mellifica* L insbesondere in der Altersperiode. *Insectes Soc* 3, 203-212
- Ribbands CR (1952) Division of labour in the honeybee community. *Proc R Soc B*, 140, 32-43

- Robinson GE (1987) Regulation of honey bee age polyethism by juvenile hormone. *Behav Ecol Sociobiol* 20, 329-338
- Robinson GE, Page RE Jr, Strambi C, Strambi A (1989) Hormonal and genetic control of behavioral integration in honey bee colonies. *Science* 246, 109-112
- Rösch GA (1925) Untersuchungen über die Arbeitsteilung im Bienenstaat, I. Teil: Die Tätigkeiten im normalen Bienenstaate und ihre Beziehungen zum Alter der Arbeitsbienen. *Z Vergl Physiol* 2, 571-631
- Rösch GA (1930) Untersuchungen über die Arbeitsteilung im Bienenstaat. II. Teil: Die Tätigkeiten der Arbeitsbienen unter experimentell veränderten Bedingungen. *Z Vergl Physiol* 12, 1-71
- Rothenbuhler WC, Page RE Jr (1989) Genetic variability for temporal polyethism in colonies consisting of similarly-aged worker honey bees. *Apidologie* 29, 433-437
- Sasagawa H, Sasaki M, Okada I (1989) Hormonal control of the division of labor in adult honeybees (*Apis mellifera* L.). I. Effect of methoprene on corpora allata and hypopharyngeal gland, and its α -glucosidase activity. *Appl Entomol Zool* 24, 66-74
- Schmidt JO (1985) Phagostimulants in pollen. *J Apic Res* 24, 107-114
- Schmidt JO, Johnson BE (1984) Pollen feeding preference of *Apis mellifera* (Hymenoptera: Apidae), a polylectic bee. *The Southwest Entomol* 9, 41-47
- Schmidt JO, Buchmann SL (1985) Pollen digestion and nitrogen utilization by *Apis mellifera* L. (Hymenoptera: Apidae). *Comp Biochem Physiol* 82A, 499-503
- Schmidt JO, Thoenes SC, Levin MD (1987) Survival of honey bees, *Apis mellifera* (Hymenoptera: Apidae), fed various pollen sources. *Ann Entomol Soc Am* 80, 176-183
- Schnelle H (1923) Über den feineren Bau des Fettkörpers der Honigbiene. *Zool Anz* 57, 172-179
- Schreiner T (1952) Über den Nahrungstransport in Darm der Honigbiene. *Z Vergl Physiol* 34, 278-298
- Seeley TD (1982) Adaptive significance of the age polyethism schedule in honeybee colonies. *Behav Ecol Sociobiol* 11, 287-293
- Shehata SM, Townsend GF, Shuel RW (1981) Seasonal physiological changes in queen and worker honeybees. *J Apic Res* 20, 69-78
- Simpson J, Riedel IBM, Wilding N (1968) Invertase in the hypopharyngeal glands of the honeybee. *J Apic Res* 7, 29-36
- Sinizki NN, Lewtschenko IW (1971) Der Gehalt an Eiweiß und freien Aminosäuren in der Hämolymphe der Arbeitsindividuen der Honigbiene. *23rd Int Beekeeping Congr Moscow*, Apimondia Publ House, Bucharest, 361-364
- Soudek S (1927) The pharyngeal glands of the honeybee (*Apis mellifica* L.). As cited in: Groot de AP (1953) Czech with English summary. *Bull Éc Sup Agron* 10, 1-63
- Southwick EE (1989) Seasonal differences in honey bees. *Am Bee J* 129, 451-452
- Standifer LN (1967) A comparison of the protein quality of pollens for growth-stimulation of the hypopharyngeal glands and longevity of honey bees, *Apis mellifera* L. (Hymenoptera: Apidae). *Insectes Soc* 14, 415-426
- Suzuki K (1988) The development of hypopharyngeal glands in honey bee workers. *Bull Fac Educ Shiba Univ* 36, 96-101
- Takenaka T, Kaatz HH (1987) Protein synthesis by hypopharyngeal glands of worker honey bees. In: *Chemistry and Biology of Social Insects* (Eder J, Rembold H, eds), J Peperny, München, 166-167
- Terra WR, Ferreira C (1981) The physiological role of the peritrophic membrane and trehalase: digestive enzymes in the midgut and excreta of starved larvae of *Rhynchosciara*. *J Insect Physiol* 27, 325-331
- Terra WR (1988) Physiology and biochemistry of insect digestion: an evolutionary perspective. *Braz J Med Biol Res* 21, 675-734
- Terra WR (1990) Evolution of digestive systems of insects. *Annu Rev Entomol* 35, 181-200
- Terra WR, Ferreira C, Bastos F (1985) Phylogenetic considerations of insect digestion. Disaccharidases and the spatial organization of digestion in the *Tenebrio molitor* larvae. *Insect Biochem* 15, 443-449
- Terra WR, Ferreira C, de Bianchi AG (1979) Distribution of digestive enzymes among the endo- and ectoperitrophic spaces and midgut

- cells of *Rhychosciara* and its physiological significance. *J Insect Physiol* 25, 487-494
- Trappman W (1923) Anatomie und Physiologie des Zwischendarmes von *Apis mellifica*. *Arch Bienenkd* 5, 190-203
- Turunen S (1985) Absorption. In: *Comprehensive Insect Physiology Biochemistry and Pharmacology* (Kerkut GA, Gilbert LI, eds) Pergamon Press, Oxford, New York, Toronto, 241-277
- van der Vorst E, Jacobs FJ (1980) Comparison of colony- and laboratory-stored pollen for maintaining the life of caged honeybees. *J Apic Res* 19, 119-121
- Wahl O (1963) Vergleichende Untersuchungen über den Nährwert von Pollen, Hefe, Sojamehl und Trockenmilch für die Honigbiene (*Apis mellifica*). *Z Bienenforsch* 6, 209-279
- Webster TC, Peng YS (1987) Passage of cannibalized tissue among honey bee (Hymenoptera: Apidae) colony members. *Ann Entomol Soc Am* 80, 814-819
- Webster TC, Peng YS, Duffey SS (1987) Conservation of nutrients in larval tissue by cannibalizing honey bees. *Physiol Entomol* 12, 225-231
- Weiss K (1984) Regulierung des Proteinhaltess im Bienenvolk (*Apis mellifica* L.) durch Brutkannibalismus. *Apidologie* 15, 339-354
- Whitcomb W Jr, Wilson HF (1929) Mechanics of digestion of pollen by the adult honey bee and the relation of undigested parts to dysentery of bees. *Wis Res Bull* 92, 1-27
- Wille H, Wille M, Kilchenmann V, Imdorf A (1987) Die Pollenernährung des überwinternden und auswinternden Bienenvolkes. *Forschungsanstalt für Milchwirtschaft, Mitteilungen der Sektion Bienen* 2, 1-11
- Winston ML, Neilson Punnett E (1982) Factors determining temporal division of labor in honeybees. *Can J Zool* 60, 2947-2952
- Winston ML (1987) *The biology of the honey bee*. Harvard Univ Press, Cambridge MS