Impact of nutrition on canine behaviour: current status and possible mechanisms

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Each year, millions of dogs worldwide are abandoned by their owners, relinquished to animal shelters, and euthanised because of behaviour problems. Nutrition is rarely considered as one of the possible contributing factors of problem behaviour. This contribution presents an overview of current knowledge on the influence of nutrition on canine behaviour and explores the underlying mechanisms by which diet may affect behaviour in animals. Behaviour is regulated by neurotransmitters and hormones, and changes in the availability of their precursors may influence behaviour. Tryptophan, the precursor of serotonin, may affect the incidence of aggression, self-mutilation and stress resistance. The latter may also be influenced by dietary tyrosine, a precursor to catecholamines. As diet composition, nutrient availability and nutrient interactions affect the availability of these precursors in the brain, behaviour or stress resistance may be affected. PUFA, especially DHA, have an important role as structural constituents in brain development, and dietary supply of n-3 and n-6 PUFA could modify aspects of the dopaminergic and serotonergic system and, consequently, cognitive performance and behaviour. Finally, persistent feeding motivation between meals can increase stereotyped behaviour and aggression and decrease resting time. This feeding motivation may be altered by dietary fibre content and source. At present, few studies have been conducted to evaluate the role of nutrition in canine (problem) behaviour through the above mentioned mechanisms. Studies that explore this relationship may help to improve the welfare of dogs and their owners.

Dogs: Food: Nutrients: Behaviour

Introduction

The domestic dog (Canis familiaris) is believed to have evolved from the grey wolf (C. lupis) as a separate species at least 15 000 years ago and it is thought to be the first animal species to be domesticated by humans¹,². At the present time, as a result of selective breeding, approximately 400 distinct dog breeds are recognised worldwide, representing a large variation in body size and weight, with the latter ranging from 1 to 90 kg. Initial functions of dogs such as hunting, shepherding and guarding have diminished gradually in importance in favour of the dog’s role as a companion to humans³. Though most human–dog relationships are fulfilling, each year a large number of animals are abandoned by their owners or relinquished to animal shelters⁴. Aggression toward people and animals, running away, destructive behaviour, disobedience, house soiling and excessive barking are unwanted behaviours that make owners relinquish or abandon their dogs⁵. Although only 20 % of the dogs in the US shelters are assigned by their owners for euthanasia⁶, a further 40 % of dogs admitted are euthanised⁷. Of the sheltered dogs that are purchased by new owners, approximately 20 % are returned to shelters⁶,⁷ and a large proportion of these animals are euthanised⁴. The number of dogs and cats euthanised annually in the USA is estimated to be between 5 and 17 million⁸,⁹, with 3–6 million as a result of behaviour problems¹⁰. Strategies that combat problem behaviours in dogs will greatly benefit animal welfare. The behaviour of individual dogs is controlled by numerous factors and from studies in humans it can be derived that nutrition plays a role also. For example, diets rich in vitamins and minerals may decrease anti-social behaviour in schoolchildren¹¹ and supplementation of vitamins, minerals and essential fatty acids.

Abbreviations: DHA, docosahexaenoic acid; EPA, eicosapentaenoic acid; GLP-1, glucagon-like peptide-1; ISF, insoluble fibre; LNAA, large neutral amino acids; ME, metabolisable energy; PUFA, polyunsaturated fatty acid; PYY, peptide tyrosine tyrosine; SF, soluble fibre; VFA, volatile fatty acids.

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decreased anti-social behaviour, including violence, of young adult prisoners. Dietary effects on behaviour have been investigated for anti-social aspects, but also for behavioural changes related to ageing and, in this, dogs have been used as a model for humans. Dogs develop similar cognitive deficits and neuropathology as can be seen in ageing humans and elderly suffering from dementia. Milgram and co-workers initiated a series of experiments with young and aged beagle dogs to study dietary interventions on age-related cognitive decline. Results showed that canine food enriched with antioxidants and mitochondrial cofactors decreased the rate of cognitive decline in aged beagle dogs under laboratory conditions and improved age-related behavioural changes in older pet dogs held in home situations (for reviews, see Roudebush et al. and Zicker). These findings demonstrate clearly that canine behaviour can be influenced by dietary components.

The present review presents an overview of our current knowledge on the influence of dietary macronutrient composition on the behaviour of dogs and explores the underlying mechanisms by which diet may affect behaviour. Findings from food–behaviour studies in dogs and other mammals are integrated to assess in what way problem behaviour in dogs may be reduced through dietary means.

**Effects of dietary amino acids and protein content on behaviour**

After ingestion, proteins are enzymically degraded and absorbed in the small intestine mainly as tripeptides, dipeptides and free amino acids. After hydrolysis of the peptides in the enterocytes, the free amino acids are transported through the portal vein to the liver. Amino acids are important constituents required for the synthesis of enzymes and other proteins, and used as precursors for the synthesis of neurotransmitters and hormones. For example, serotonin, catecholamines, acetylcholine and histamine are metabolites from tryptophan, tyrosine, choline and histidine, respectively. These neurotransmitter precursors (except for choline) are amino acids and are natural dietary constituents. Behaviour results from signal detection, transmission and processing in the (central) nervous system, which is accomplished and modulated by chemical messengers such as neurotransmitters and hormones. Changes in neurotransmitter precursors such as tryptophan and tyrosine are, therefore, likely to influence behaviour. The amount and timing of food intake, diet composition and digestibility are all factors that determine the availability of different amino acids, i.e. precursors of chemical messengers. Consequently, such factors may influence behaviour. The effects of tryptophan and tyrosine on behaviour will be discussed as these could be relatively potent modulators; for similar reports on choline, histidine and threonine, we refer to Young.

**Findings and mechanisms in different mammals**

**Tryptophan.** A diet high in tryptophan has been shown to reduce mouse killing by rats, reduce aggression in vervet monkeys, enhance exploratory behaviour in female silver foxes and reduce self-injurious behaviour in rhesus monkeys. In contrast to the observed reductions in aggression in some experimental conditions, dietary supplementation of tryptophan has also been shown to increase territorial aggression in male mice. Dietary tryptophan may also influence the resistance or tolerance to stress and, therefore, change the behavioural stress response. Koopmans et al. reported enhanced recovery after social stress as measured by lower plasma cortisol and noradrenaline concentrations in pigs fed a surplus of dietary tryptophan compared with pigs fed diets containing a ‘normal’ concentration of tryptophan. In addition, supplementation of dietary tryptophan reduced plasma cortisol concentrations during a stress-inducing mental arithmetic task in healthy stress-vulnerable humans. It was, therefore, suggested by Markus et al. that tryptophan supplementation above normal dietary concentrations could improve the ability of an individual to cope with stress. The effects of dietary tryptophan on stress resistance involve different pathways. In rats a variety of stressors, such as immobilisation, foot shock, and hypothermia, increase brain tryptophan and serotonin turnover. Depressed humans show decreased plasma tryptophan concentrations in comparison with normal subjects. It appears that initially stressors stimulate serotonin turnover, which over time may deplete serotonin (precursor) supplies and result in decreased serotonin (precursor) concentrations.

Quantitatively the most important pathway for tryptophan metabolism, after protein synthesis, is the kynurenine pathway which is responsible for over 90% of tryptophan catabolism. In humans, normally 1% of the available tryptophan is converted to serotonin which is mainly present in the gastrointestinal tract. The first and rate-limiting step in the synthesis of serotonin is the hydroxylation of tryptophan to 5-hydroxytryptophan by the enzyme tryptophan hydroxylase (Fig. 1). Tryptophan hydroxylase is normally about half saturated with tryptophan. Consequently, an increase in tryptophan in the brain, which increases serotonin synthesis and serotonergic neurotransmission, can maximally double serotonin synthesis. The second step in the synthesis of serotonin is the decarboxylation of 5-hydroxytryptophan to serotonin which is stored in vesicles in the nerve terminal were it is held before release. When serotonin is released into the synaptic cleft, serotonin can bind to different subtype receptors (for reviews, see Barnes & Sharp and Hoyer et al.). Via binding to these different receptors, serotonin can produce many different effects on post-synaptic cells influencing various parts of the brain involved in controlling a variety of physiological functions including hormone releases, cardiovascular functioning, pain, appetite, and in general mood and behaviour. Tryptophan transport across the blood–brain barrier and metabolism is in part affected by animal factors such as breed, sex, social status, age, activity and level of arousal. The availability of dietary tryptophan to the brain is largely dependent on the composition of the ingested diet. Tryptophan is found in nearly all protein-containing foods where it is found in a lower concentration compared with the other large neutral amino acids (LNAA) tyrosine, phenylalanine, leucine, isoleucine...
and valine\textsuperscript{45}. For access into the brain, tryptophan shares the same carrier as other LNAA for transport across the blood–brain barrier\textsuperscript{34}. Central tryptophan concentrations can either be increased by increasing plasma tryptophan or by lowering plasma concentrations of LNAA\textsuperscript{34,46}. As tryptophan is normally present in only small concentrations in dietary protein compared with other LNAA, the consumption of a meal high in protein will decrease the ratio of tryptophan to other LNAA\textsuperscript{47} and thereby potentially lower serotonin synthesis.

The fraction of unbound tryptophan as compared with that bound to albumin is another factor that may influence tryptophan availability to the brain\textsuperscript{48}. In mammals, approximately 80–90\% of all tryptophan molecules in the blood are bound to serum albumin\textsuperscript{49}. It has been suggested that the majority of the albumin-bound tryptophan is available for passage across the blood–brain barrier\textsuperscript{46,50}, but possibly the concentration of circulating free tryptophan may be especially important\textsuperscript{48}. According to Chaouloff\textsuperscript{48}, three factors affect circulating free and bound tryptophan concentrations: (i) the rate of lipolysis because blood non-esterified fatty acids displace tryptophan from its binding to albumin\textsuperscript{51}; (ii) the activity of tryptophan 2,3-dioxygenase, the rate-limiting enzyme in tryptophan detoxication through the kynurenine pathway – activation (inactivation) of this enzyme decreases (increases) circulating free tryptophan may be especially important\textsuperscript{52}; (iii) uptake into peripheral and central tissues. Carbohydrate-induced insulin rises facilitate the uptake of most LNAA into skeletal muscle, but not tryptophan bound to albumin\textsuperscript{53,54}. Consequently, the ratio of tryptophan relative to LNAA increases. This results in a competitive advantage of tryptophan over LNAA for uptake at the blood–brain barrier. However, as little as 2–4\% of the energy of a meal as protein seems to prevent this increased availability of tryptophan\textsuperscript{31,55}.

**Tyrosine.** In rats, a high-tyrosine diet prevents adverse behavioural and neurochemical effects (for example, immobility during a swim test, depletion of brain noradrenaline) of various acute stressors including hypothermia\textsuperscript{56}, restraint and tail-shock\textsuperscript{57–59}. Human studies also suggest beneficial effects of tyrosine under conditions of stress (for reviews, see Lieberman\textsuperscript{60} and Young\textsuperscript{17}).

Tyrosine, which can be synthesised from phenylalanine, is the direct precursor for the catecholamines dopamine, noradrenaline and adrenaline\textsuperscript{32}. Dopamine can be synthesised from tyrosine in neurons in two steps. The first and rate-limiting step is the conversion of tyrosine to dihydroxyphenylalanine by the enzyme tyrosine hydroxylase. In rats, central tyrosine hydroxylase is approximately 75\% saturated with tyrosine\textsuperscript{33}. In the second step, dihydroxyphenylalanine is decarboxylated to dopamine which can be used as an endproduct (neurotransmitter) in neurons or further converted to noradrenaline or adrenaline\textsuperscript{61}. Like tryptophan, tyrosine competes with other LNAA at the blood–brain barrier for entry into the brain\textsuperscript{34} and is taken up into skeletal muscle under the influence of insulin\textsuperscript{53,54}. In diets, tyrosine is typically available in much higher concentrations compared with tryptophan and high-protein meals will typically raise tyrosine concentrations in the brain, but will lower the concentration of tryptophan\textsuperscript{32}. Catecholamines play a key role in a variety of behavioural, neuroendocrine and cardiovascular responses during stress\textsuperscript{60}. Increases in

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**Fig. 1.** Effects of dietary characteristics on tryptophan uptake by the central nervous system and synthesis of serotonin from brain tryptophan (adapted from Grimmett & Sillence\textsuperscript{185} with modifications). \(\rightarrow\), Factors that may ultimately decrease brain tryptophan; 5-HTP, 5-hydroxytryptophan; NEFA, non-esterified fatty acids.
brain tyrosine have little or no effect on catecholamine synthesis\textsuperscript{17}, but the situation may be different during stress when brain noradrenaline turnover increases and noradrenaline concentrations decrease\textsuperscript{57,63}. An enhanced noradrenergic activity is part of a normal adaptive stress response\textsuperscript{64}. In stressed rats (tail-shock), ingestion of a high-tyrosine diet reversed the post-stress decline in brain noradrenaline and attenuated behaviour changes, i.e. decreased locomotion, standing on hind legs, hole-poking in a novel open field\textsuperscript{57}. This suggests that a high-tyrosine diet may be beneficial during severe stress, as it prevents depletion of the substrate required for catecholamine synthesis in times of high catecholaminergic activity and demand.

**Findings in dogs**

Studies on the effects of tryptophan or tyrosine on behaviour in dogs seem to be limited to one. DeNapoli et al.\textsuperscript{65} formulated diets with high or low protein content (approximately 310 or 190 g crude protein/kg, respectively) and with or without tryptophan supplementation (1-45 g/kg) in order to provide varying tryptophan contents and tryptophan:LNAA ratios (Table 1). Each of the four diets was fed in random order for 1 week to thirty-three privately owned dogs that displayed a high territorial aggression, dominance aggression or hyperactivity. There was no effect of dietary protein or tryptophan content on the behavioural scores within each group of problem behaviour. However, when the groups of dogs were analysed as one study population a lower territorial aggression score was obtained for dogs fed the high-tryptophan diet compared with dogs fed the low-tryptophan diet, but only when fed a low-protein diet. In addition, dogs fed the high-protein diet without tryptophan supplementation showed a higher dominance aggression score compared with dogs on the other dietary treatments.

Three studies in literature have reported that low-protein diets decreased aggression in dogs, though these were not performed under controlled experimental conditions. In a study with seven aggressive golden retrievers held at in-home living conditions, incidences of aggression as reported by their owners immediately decreased after the introduction of a low-protein diet (15–18% of total energy).\textsuperscript{66} Unfortunately, neither the composition of the experimental diet nor the composition(s) of the diet(s) before the dietary intervention were reported. The reduction in aggressive incidences, however, was only sustained in three dogs; two dogs deteriorated again in their behaviour and contact was lost with the remaining two clients. In another study, twelve dogs that exhibited either high territorial aggression, dominance aggression or hyperactivity and fourteen control dogs were fed each of three diets varying in protein content (180, 250 and 310 g crude protein/kg DM) for 2 weeks at in-home living situations\textsuperscript{59}. The low-protein diet and medium-protein diet decreased territorial aggression scores compared with the high-protein diet. No effects of dietary protein content in dogs with dominance aggression or hyperactivity were found. Additional behavioural analysis of the group of dogs demonstrating territorial aggression revealed that five of these dogs showed dominance-related

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**Table 1. Effect of dietary protein and tryptophan (TRP) content on canine behaviour**

<table>
<thead>
<tr>
<th>Authors</th>
<th>Dogs and design</th>
<th>Diets*</th>
<th>Results</th>
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<tbody>
<tr>
<td>Mugford\textsuperscript{66}</td>
<td>Seven aggressive golden retrievers at in-home living situations. Measurements were not reported</td>
<td>15–18% protein of total dietary energy based on approximately 20% meat and 80% boiled rice</td>
<td>(a) Territorial aggressive dogs showed lower territorial aggression scores when fed diets 1 and 2 compared with diet 3; the remaining five territorial aggressive dogs fed the low-tryptophan diet, but only when fed a low-protein diet. In addition, dogs fed the high-protein diet without tryptophan supplementation showed a higher dominance aggression score compared with dogs on the other dietary treatments.</td>
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<tr>
<td>Dodman\textsuperscript{10}</td>
<td>Twelve territorial aggressive, twelve dominance aggressive, twelve hyperactive and fourteen control dogs (age 1 year) fed each diet (Latin square) at maintenance level at in-home living situations for 7 d. Each day, owners scored their dogs for territorial aggression, dominance aggression, excitability and fearfulness</td>
<td>(a) No changes in behaviour within each behaviour group for any dietary treatment. (b) When all dogs were combined, territorial aggression scores were higher for dogs fed diet 1 compared with diets 2 and 3. (c) When all dogs were combined, territorial aggression scores were higher for dogs fed diet 1 compared with diets 2 and 3.</td>
<td></td>
</tr>
<tr>
<td>DeNapoli et al.\textsuperscript{65}</td>
<td>Eleven territorial aggressive, eleven dominance aggressive and eleven hyperactive dogs (age &gt; 1 year) fed each diet (at random) at maintenance level at in-home living situations for 14 d. Each day, owners scored their dogs for territorial aggression, dominance aggression, excitability and fearfulness</td>
<td>(a) No changes in behaviour within each behaviour group for any dietary treatment. (b) When all dogs were combined, dominance aggression scores were higher for dogs fed diet 1 compared with diets 2 and 3.</td>
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\textsuperscript{*} Values are presented on a DM basis. LNAAs, large neutral amino acids (tyrosine, phenylalanine, leucine, isoleucine, valine).
Effects of dietary lipids on behaviour

Lipids have various functions, such as constituents of cellular membranes, precursors for chemical messengers (for example, steroid hormones) and their use as an energy source or stored in the body as adipose tissue. After adipose tissue, the central nervous system has the greatest concentration of lipids. The structural constituents in the grey matter of the brain and retinal tissues in mammals are derived from dietary linoleic acid (18:2n-6) and α-linolenic acid (18:3n-3). Both are polyunsaturated fatty acids (PUFA) and can be metabolised to long-chain PUFA by sequential alternating enzymic desaturation and elongation. Linoleic acid can be metabolised to arachidonic acid (20:4n-6) which can be further metabolised to docosapentaenoic acid (22:5n-6). The enzymic desaturation and elongation of α-linolenic acid yields eicosapentaenoic acid (EPA) (20:5n-3) which can be further metabolised to docosahexaenoic acid (DHA) (22:6n-3).

Findings and mechanisms in different mammals

There is ample scientific literature available in which the effects of both dietary deficiency and supplementation of PUFA on animal performance in cognitive or behavioural tests are evaluated (for reviews, see Wainwright and McCann & Ames). For example, the learning ability of rodents decreased when fed n-3 fatty-acid-deficient diets and increased when fed DHA-supplemented diets compared with rodents fed diets adequate in n-3 fatty acid concentrations. Other studies, however, did not find affects of dietary n-3 PUFA manipulation on learning performance as tested with a Morris water-maze in rats or mice. Dietary PUFA seem to affect animal cognition but can also cause behavioural changes. Rats fed n-3 PUFA-deficient diets showed increased aggression scores in a resident intruder test and increased expression of stress-related behaviours during several stress tests compared with male rats fed adequate amounts of n-3 PUFA. Similarly, anxiety was found to be increased in mice fed a diet deficient in n-3 PUFA, though others did not observe any effects of dietary PUFA on anxiety in mice or rats.

The dopaminergic and serotonergic systems in the brain are known to play important roles in learning, emotions, and impulse control, which makes it tempting to assume that the effects of PUFA on behaviour run through these systems. Indeed, both systems are known to be influenced by PUFA. Rats deficient in n-3 PUFA compared with rats fed diets with α-linolenic acid showed a reduction in dopamine concentration in the frontal cortex and an increase in dopamine concentration in the nucleus accumbens but no effects in the striatum. In the frontal cortex of these animals the rate of dopamine synthesis and breakdown mediated by monoamine oxidase have been linked to the reduced dopaminergic storage pools. Changes in dopamine concentrations were followed by changes in number of D2 receptors. n-3 PUFA-deficient rats had a lower number of D2 receptors in the frontal cortex but higher in the nucleus accumbens. Rats fed diets supplemented with EPA and DHA had an increased dopamine concentration and D2 binding possibly as a result of a reduction in monoamine oxidase activity in the frontal cortex compared with rats fed adequate amounts of PUFA.

As for dopamine concentrations, frontal cortex serotonin concentrations were increased in rats fed diets supplemented with n-3 PUFA. In line with this, serotonin in the frontal cortex was reduced in piglets fed n-3 and n-6 PUFA-deficient formula for 18 d from birth compared with piglets fed formula supplemented with linoleic acid and α-linolenic acid and/or arachidonic acid and DHA. The findings in the frontal cortex may not extrapolate to other brain areas. For example, in the hippocampus of 2-month-old rats fed an n-3 PUFA-deficient diet extracellular basal serotonin concentrations were increased. This was probably due to reduced storage pools not due to decreased activity of monoamine oxidase. Such effects of n-3 PUFA deficiency on serotonin concentrations are not found in all studies (for example, Delion et al.).
In addition to the observed changes in the dopaminergic and serotonergic systems in different brain regions, physical properties (for example, fluidity, permeability) of cerebral membranes may also mediate dietary effects on cognition and behaviour\textsuperscript{79}. For example, chronic dietary deficiency in n-3 PUFA resulted in low concentrations of n-3 PUFA in the rat brain\textsuperscript{26,77} whereas diets high in EPA and DHA resulted in high concentrations of EPA and DHA in the brain of rats\textsuperscript{25,85,100,101}. In addition, dietary α-linolenic acid deficiency induces a more pronounced reduction in DHA concentrations in the frontal cortex than in the striatum and cerebellum\textsuperscript{72,91}. Besides changes in brain PUFA compositions, dietary PUFA may alter properties of the neuronal membrane, such as the activity of membrane-bound enzymes, receptors and ion channels\textsuperscript{102}. These alterations may affect neurological functioning and may, therefore, also contribute to the observed changes in cognitive functioning and behaviour.

\textit{Findings in dogs}

To the authors’ knowledge, there are at this moment no scientific articles available regarding the influence of n-3 or n-6 PUFA deficiency or enrichment on canine behaviour or cognitive performance. Since DHA is essential for the development and function of the brain and retina\textsuperscript{73}, its supply may affect neurological development in puppies. For example, low dietary concentrations of DHA during the gestation or lactation of bitches and dry diets for puppies depressed their retinal sensitivity\textsuperscript{103,104}. Although the immediate connection between the cellular effects of DHA and visual sharpness and cognitive abilities in receiving dietary DHA still needs more support\textsuperscript{72}, studies seem to emphasise the importance of DHA in the diet of bitches during gestation until weaning and the diet of puppies in order to ensure optimal neurological development. At present, there is no recommended allowance for DHA for both bitches in gestation and lactation or puppies, but the recommended allowance for α-linoleic acid is 3.35 g/1000 kJ (0.8 g/1000 kcal) ME\textsuperscript{68}. A diet high in α-linolenic acid fed from breeding throughout lactation increased α-linolenic acid concentration in milk but failed to do this for DHA\textsuperscript{105,106}. In a recent study, puppies converted α-linolenic acid to DHA during the first month of weaning but little conversion of α-linolenic acid to DHA occurs after weaning\textsuperscript{106}. It seems that the capacity of puppies to synthesise DHA from dietary α-linolenic acid or other n-3 fatty acid precursors is active for only a short time during the neonatal period and is decreased thereafter. The amount of dietary α-linolenic acid for sufficient synthesis of DHA and the amount of DHA required for optimal neurological development in puppies still remain to be determined. Whether the provision of sufficient DHA for optimal neurological development in dogs also results in changes in the dopaminergic and serotonergic systems and subsequent effects in cognitive abilities or behaviour in later life remains to be confirmed.

Concerning commercial dog food, it seems likely that in dogs deficiencies of PUFA are rare as long as fat oxidation during process and storage of the food is limited\textsuperscript{107}. Levels of PUFA, particularly the n-3 family, are nowadays higher in commercial dog food compared to foods of several years ago\textsuperscript{108} (Delton-Vandenbroucke \textit{et al.}, 1998). However, the amount and ratio between n-6 and n-3 fatty acids may differ considerably between commercially available diets. The n-6:n-3 fatty acid ratio of twelve commercial dry dog foods was found to differ considerably, ranging from 17:1 to 5.1\textsuperscript{109}.

\textbf{Effects of dietary carbohydrates on behaviour}

Feeding of mammals is a discontinuous process in which periods of food consumption are interspersed with periods of non-eating\textsuperscript{110}. Food intake behaviours are controlled by feelings of hunger\textsuperscript{111} and satiety\textsuperscript{110}, but may be modulated by psychological and social factors\textsuperscript{112}. Numerous central and peripheral signal molecules are involved in the regulation of eating (for reviews, see Bray\textsuperscript{113}, de Graaf \textit{et al.}\textsuperscript{114} and Strader & Woods\textsuperscript{115}). The rate and site of degradation of nutrients largely determines the postprandial physiological state of an animal and in this way the extent and duration of satiety and, therefore, behaviour. There is a wide variety of carbohydrates with different physical and chemical properties. These properties can affect the rate and site of degradation of these carbohydrates\textsuperscript{116}. In single-stomached animals, degradable carbohydrates may be digested with endogenous enzymes in the first part of the gastrointestinal tract, or fermented by micro-organisms that colonise predominantly the last part of the gastrointestinal tract. Products derived from digestible carbohydrates are mainly monosaccharides. The digestion of starch and absorption of monosaccharides are primarily responsible for the fluctuations in the postprandial blood glucose concentrations that subsequently may modify tryptophan availability in the brain when protein intake is low (see section on Findings and mechanisms in different mammals: Tryptophan, and influence mood in at least humans (for a review, see Benton\textsuperscript{117}). The indigestible carbohydrates are often referred to as dietary fibre, which contains non-starch polysaccharides, resistant starch and non-digestible oligosaccharides. The fermentation endproducts of dietary fibre are volatile fatty acids (VFA; acetic, propionic and butyric acid), lactate, alcohol and the gases methane, hydrogen and carbon dioxide\textsuperscript{118}. Apart from the fermentability, other physical and chemical properties of dietary fibre include solubility, ability to bind water and affect viscosity, and possible interactions with the digestion and absorption of starch, protein and fat. In addition, the duration of satiety experienced by animals between meals may be affected by carbohydrates, which in turn may reduce the behavioural side effects of a high feed motivation.

\textit{Findings and mechanisms in different mammals}

The effects of dietary carbohydrate sources (i.e. fibrous ingredients) on animal behaviour have been relatively well studied especially in pigs, where non-lactating sows were fed energy-restricted diets in order to prevent excessive lipid deposition and reduced reproduction performance. Commonly diets for sows are formulated to meet the daily nutrient requirements for maintenance and reproduction. However, the latter may not result in a sufficient level of...
amounts of feed it has been suggested that hunger is most shown to be more active compared with sows fed large diets high in fibrous ingredients (sugarbeet pulp, oat hulls, soybean hulls, wheat bran) can be fed resulting in an increased time of sows laying down\(^2\), increased resting time, less time spent on foraging and aggression\(^3\) and reduced posture changes 8 and 10 h after feeding\(^4\). The latter authors compared sows fed a high- and a low-fermentable carbohydrate diet (for further examples, see Meunier-Salaün et al.\(^5\)). The relationship between dietary fibre content and stereotyped behaviour has also been documented in horses. A large survey among trainers of race horses in Sweden revealed a negative correlation between the amount of roughage provided and the incidence of stereotyped behaviour (cribbing or wind-sucking, weaving, box-walking) or wood-chewing in horses\(^6\). Wood-chewing may be related to a ‘fibre deficiency’ in the diet and represent attempts to increase dietary fibre intake\(^7\)–\(^9\). The effect of fibrous ingredients on behaviour is not generic for all fibre sources; for example, solvent-extracted coconut meal and soyabean hulls as a dietary fibre source do not appear to affect physical activity in pigs\(^10\), whereas sugarbeet pulp silage does\(^1\). Since sows which are fed low amounts of feed were shown to be more active compared with sows fed large amounts of feed\(^3\) it has been suggested that hunger is most likely the cause of the increased physical activity\(^1\).

The variety in physical and chemical properties of different fibrous ingredients results in differences between these fibres in creating and maintaining satiety and preventing feelings of hunger. The biological mechanisms behind the satiating properties of dietary fibre are still not fully understood, but several dietary fibre characteristics seem to be important. First, fibres with a high water-binding capacity may increase the volume and weight of the gastric contents when liquids are available. The weight or volume may stimulate stretch receptors that can induce gastric signals of satiation\(^1\)–\(^3\). Second, gastric emptying can be affected either directly by dietary fibres high in intragastric viscosity\(^4\) or indirectly through the stimulation of the release of glucagon-like peptide-1 (GLP-1) (a potent inhibitor of gastric emptying\(^5\)). Stimulation of GLP-1 production can be mediated through carbohydrate fermentation in the distal part of the gastrointestinal track\(^6\), or through the production of VFA (mainly acetate) which stimulates the release of peptide tyrosine tyrosine (PYY)\(^7\)–\(^9\). The effects of GLP-1 and PYY in delaying gastric emptying are referred to as the ‘ileal brake’ mechanism which results in a moderate and stable flow of nutrients from the stomach into the small intestine\(^1\). A decrease in postprandial gastric-emptying rate will, consequently, prolong gastric distension and gastric signals of satiation\(^1\)–\(^3\). This mechanism was studied by Moran et al.\(^1\) in rhesus monkeys where intramuscular injections of PYY reduced gastric emptying and resulted in a decrease in food intake. In addition, there are indications that PYY in the brain reduces appetite in humans\(^1\)–\(^3\), although this is still a subject for debate\(^4\). Third, fibrous dietary ingredients may increase small-intestinal transit time\(^5\), possibly also by stimulation of PYY which is found to suppress intestinal motility\(^6\). An increase in small-intestinal transit time: (i) prolongs contact between nutrients and intestinal receptors involved in maintaining satiety\(^7\) and postpones feelings of hunger\(^8\); (ii) results in the slowing down of starch digestion and subsequent absorption of glucose, thereby maintaining more stable postprandial glucose and insulin concentrations in the blood\(^9\). A transient decline in blood glucose level preceded meal initiation in rats\(^10\) and humans\(^1\)–\(^3\), and caused a delay in the decrease in blood glucose concentrations. This may prolong satiety and postpone hunger and meal initiation (for a review, see Campfield & Smith\(^1\)). Finally, fermentation of carbohydrates may yield VFA which leads to a higher level of satiety by (i) PYY-mediated reduction of gastric emptying rate\(^1\) and (ii) becoming a source of energy (mainly acetate) at times when glucose supply from the small intestine is decreasing, which stimulates longer-term satiety\(^1\)–\(^3\),\(^4\),\(^5\).

As suggested previously, hunger is most likely the cause for the observed behavioural effects seen in sows\(^3\). Hunger or appetite is correlated with the peripheral concentration of ghrelin\(^1\), a twenty-eight amino acid peptide synthesised predominantly in the stomach\(^1\)–\(^3\). For example, a rise in blood ghrelin concentration is associated with meal initiation in humans\(^4\). Supplementation of short-chain oligofructose (average degree of polymerization of 4.5) in a diet for 3 weeks decreased energy intake and lowered ghrelin concentrations in rats compared with rats fed the control diet without fructan supplementation. However, rats fed a diet supplemented with long-chain oligofructose (average degree of polymerization of 25.0) showed a decrease in energy intake but not in ghrelin concentrations compared to rats fed the control diet\(^5\). It is suggested that the lower blood ghrelin concentrations may contribute to a decrease in appetite during fasting\(^6\). Whether these results were accompanied with changes in behaviour (for example, food-seeking behaviour) requires further investigation. Fig. 2 shows the effects of dietary fibre on satiety.

**Findings in dogs**

‘When we are considering how a dog is behaving, we really should be considering what is inside the stomach’ (Mugford\(^1\), p. 1046). Despite this statement, little additional research has been conducted on the association between canine behaviours and satiety or feeding motivation between meals. To the authors’ knowledge, three studies have investigated the effects of dietary fibre on satiety and feeding motivation in dogs of which only one also studied canine behaviour and another measured ad libitum food intake of dogs fed diets varying in fibre source and content (Table 2). Butterwick & Markwell\(^1\) fed overweight dogs (> 115 % ideal body weight) six different moist diets varying in type and amount of fibre on an energy-restricted basis (45 % restriction of calculated maintenance energy requirements; ME (kJ) = 461 × body weight (kg)\(^1\)–\(^3\)). The four experimental high-fibre diets formulated to vary in soluble fibre (SF) and insoluble fibre (ISF), i.e. (g/kg DM) 40–8 SF and 13–6 ISF; 112–5 SF and
37.5 ISF; 35.7 SF and 202.4 ISF; 24.8 SF and 310.6 ISF, were compared with two dry control diets (36.5 SF and 14.6 ISF; 45.5 SF and 15.2 ISF). The authors found no differences in time spent at behaviours related to feeding motivation (i.e. cumulative time spent at feeding bowl and number of visits to bowl 30 min after feeding, intake of a meal (canned diet) provided 3 h after introduction of the test diets) between dogs fed the different diets. In contrast, Jewell & Toll\textsuperscript{162} did find effects of fibre content on the satiety of dogs. Dogs with \textit{ad libitum} access to dry diets with a medium or high crude fibre content (135.5 and 223.4 g/kg DM) decreased total ME intake compared with dogs that had \textit{ad libitum} access to low-crude fibre diets (16.3 and 16.4 g/kg DM). When dogs were offered a subsequent meal, 30 min after the end of the last meal, energy and DM intake were lower in dogs fed the high-fibre diet compared with dogs consuming the low-fibre diet\textsuperscript{162}. Similarly, Jackson \textit{et al.}\textsuperscript{163} observed that a high-fibre content in dry diets reduced energy intakes in dogs. These authors fed dogs in the morning either a diet high in total dietary fibre (26.7 SF; 263.7 ISF g/kg as fed) or low in total dietary fibre (18.1 SF; 123.2 ISF g/kg as fed) followed 6 h later by \textit{ad libitum} access to a diet containing 23.2 SF; 123.5 ISF g/kg as fed. Average energy intake over the day was lower (kJ/kg body weight) in the dogs fed the high-fibre diet compared with dogs consuming the low-fibre diet\textsuperscript{162}. Similarly, Jackson \textit{et al.}\textsuperscript{163} observed that a high-fibre content in dry diets reduced energy intakes in dogs. These authors fed dogs in the morning either a diet high in total dietary fibre (26.7 SF; 263.7 ISF g/kg as fed) or low in total dietary fibre (18.1 SF; 123.2 ISF g/kg as fed) followed 6 h later by \textit{ad libitum} access to a diet containing 23.2 SF; 123.5 ISF g/kg as fed. Average energy intake over the day was lower (kJ/kg body weight) in the dogs fed the high-fibre diet compared with the energy intake of the dogs fed the low-fibre diet in the morning (273 ± 332 kJ (65.3 ± 79.4 kcal)/kg body weight). The difference in average daily energy intake was the result of the energy intake in the morning since there were no significant differences observed in intake of the diet provided in the afternoon between the high-fibre (181 kJ (43.2 kcal)/kg body weight) and low-fibre (197 kJ (47.2 kcal)/kg body weight) groups. These latter two studies showed that high levels of fibrous dietary ingredients in dogs can increase satiety and reduce energy intake. This, however, was not confirmed in a study by Butterwick & Markwell\textsuperscript{161}. The latter may be due to the energy restriction and the large differences in DM content of diets between studies. Energy restriction will result in an increased feeding motivation in dogs to a level that nullifies the possible effects of fibre on satiety\textsuperscript{163}. DM content of the moist diets fed to dogs in the study of Butterwick & Markwell\textsuperscript{161} ranged between 132 and 168 g/kg whereas Jewell & Toll\textsuperscript{160} and Jackson \textit{et al.}\textsuperscript{162} fed dry diets with a DM content between 908 and 923 g/kg. On an energy basis, intake of a diet with a high DM content or high energy density will result in lower weight of the digesta in the stomach compared with a diet with similar nutrient composition but lower DM content. A low dietary DM content will therefore have a higher weight of digesta in the stomach and will stimulate stretch receptors which affect satiety in dogs\textsuperscript{164}. Finally, food intake in g DM/kg body weight was found to be lower in dogs with \textit{ad libitum} access to a diet with 15 g short chain fructo-oligosaccharides/kg DM compared with dogs with \textit{ad libitum} access to a diet with 60 g cellulose/kg DM\textsuperscript{165}. The authors suggested that satiety between diets was altered because of the differences in fermentability of the fibre sources included in the diets. Unfortunately, no measurements were made in this study to elucidate possible mechanisms underlying their observed difference in food intake.

The mechanisms behind the observed effects of dietary fibre on inducing and maintaining satiety in humans and pigs (see previous section) have in part been also observed in dogs. Stimulation of stretch receptors through infusion of liquids or filling a balloon with water placed in the stomach reduced sham feeding in dogs, indicating that stimulation of stretch receptors induces satiety in dogs\textsuperscript{164}. Gastric emptying was reduced in dogs as fibre (for example, psyllium, guar gum) content and viscosity of the meal increased\textsuperscript{166}, which will prolong gastric distension.

\textbf{Fig. 2.} Effects of dietary fibre (DF) on satiety. \textit{(- - -)}, Factors that may ultimately increase the residence time of digesta in the designated segments of the gastrointestinal tract; WBC, water-binding capacity; VFA, volatile fatty acids; GLP-1, glucagon-like peptide-1; PYY, peptide tyrosine tyrosine.
and gastric signals of satiation. In addition, a study of Bueno et al.\textsuperscript{123}, in which dogs were fed different fibre sources (wheat bran, cellulose, guar gum), both gastric emptying and intestinal transit time were affected with the effect depending on the fibre source included.

A delay in gastric emptying and thus an increase in intestinal transit time by dietary fibre (alginate) results in more stable blood glucose concentrations as observed by Jewell \& Butterwick \& Murray et al.\textsuperscript{66} On day 7, one of two diets was provided 75 min after first meal. On day 14, the other diet was offered 75 min after first meal. After 14 d, the experimental design was repeated but each group of dogs received the other of the two diets Study 2, identical as study 1 but with different diets

<table>
<thead>
<tr>
<th>Authors</th>
<th>Dogs and design</th>
<th>Dietary fibre content or source*</th>
<th>Results</th>
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<tr>
<td>Jewel &amp; Toll\textsuperscript{66}</td>
<td>Study 1, two groups of fifteen beagle dogs were assigned to one of two dry diets fed once per d at maintenance level for 14 d</td>
<td>(1) 16 g CF/kg (study 1)</td>
<td>(a) Energy intake of all dogs was lower than energy on offer</td>
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<td>On day 7, one of two diets was provided 75 min after first meal. On day 14, the other diet was offered 75 min after first meal. After 14 d, the experimental design was repeated but each group of dogs received the other of the two diets Study 2, identical as study 1 but with different diets</td>
<td>(2) 136 g CF/kg (study 1)</td>
<td>(b) Dogs fed diets 2 and 4 had lower daily energy intake than dogs fed diets 1 and 3, respectively</td>
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<td>(3) 16 CF/kg (study 2)</td>
<td>(c) Energy intake of the second meal 75 min after first meal was lower when dogs were fed diets 2 and 4 compared with dogs fed diets 1 and 3, respectively</td>
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<td>(4) 223 g CF/kg (study 2)</td>
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<td>Butterwick &amp; Markwell\textsuperscript{159}</td>
<td>Six obese terrier dogs (&gt; 115% of ideal BW) were fed each of the six wet diets (6 x 6 Latin square) at 45% of maintenance level for 12 d. Number of visits to the bowl and cumulative time spent at the bowl were observed for 30 min from the start of the meal. On day 7 and 10, 8 and 11, or 9 and 12, dogs had ad libitum access to a wet diet that was provided 180 min after the first meal and food intake was measured</td>
<td>(1) 7 g CF/kg; 41 g SF/kg; 14 g ISF/kg</td>
<td>(a) No differences between diets in daily energy intake</td>
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<td>(2) 13 g CF/kg; 113 g SF/kg; 38 g ISF/kg</td>
<td>(b) No differences between diets in observed behaviours</td>
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<td>(3) 143 g CF/kg; 36 g SF/kg; 202 g ISF/kg</td>
<td>(c) No differences between diets in food intake of the second meal 180 min after first meal</td>
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<td>(4) 149 g CF/kg; 25 g SF/kg; 311 g ISF/kg</td>
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<td>(5) 15 g CF/kg; 37 g SF/kg; 15 g ISF/kg</td>
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<td>(6) 8 g CF/kg; 46 g SF/kg; 15 g ISF/kg</td>
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<td>Jackson et al.\textsuperscript{161}</td>
<td>Two groups of fifteen miniature schnauzers and toy poodles were assigned to one of two dry diets fed in the morning at 50% of daily intake and had ad libitum access to a control diet in the afternoon (approximately 6 h later) for 8 d</td>
<td>(1) 95 g CF/kg; 27 g SF/kg; 264 g ISF/kg</td>
<td>(a) Dogs fed diet 1 had lower morning and daily energy intake/kg BW than dogs fed diet 2</td>
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<td>(2) 20 g CF/kg; 18 g SF/kg; 123 g ISF/kg</td>
<td>(b) There was no difference in food intake of diet 3 in the afternoon between dietary treatments</td>
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<td>Howard et al.\textsuperscript{163}</td>
<td>Twenty-eight adult beagle dogs were stratified by BW and assigned at random to one of four dry diets with ad libitum access for 35 d</td>
<td>(1) 60 g cellulose/kg</td>
<td>(a) No differences between diets in DM intake per d</td>
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<td>(2) 15 g FOS/kg</td>
<td>(b) Dogs fed diet 2 showed lower DM intake/d per kg body weight compared with dogs fed diet 1</td>
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<td>(3) 60 g beet pulp/kg</td>
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<td>(4) 60 g beet pulp/kg; 20 g gum talha/kg; 15 g FOS/kg</td>
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CF, crude fibre; BW, body weight; SF, soluble fibre; ISF, insoluble fibre; FOS, fructo-oligosaccharides.

* Values are presented on a DM basis except for the data of Jackson et al.\textsuperscript{161}, which are as-fed.

Although dogs have a relatively small and simple large intestine, dogs are capable of fermenting a significant quantity of dietary non-digestible carbohydrates\textsuperscript{172}. Moreover, the faecal microflora of dogs were found to give similar in vitro organic matter disappearance results compared with the microflora from humans, pigs and horses\textsuperscript{173}. The latter indicates that differences between these species in carbohydrate fermentation capacity are probably dependent on factors other than the microbial population. The extent of fermentation in the gastrointestinal tract in an animal largely depends on the time available for microbial fermentation\textsuperscript{173–175}. In dogs, a transit time through the total gastrointestinal tract between 20 and 35 h is considered normal\textsuperscript{176}. The large-intestinal transit of digesta can take up to 90% of the total gastrointestinal transit time\textsuperscript{175,178}, presenting a considerable time for large-intestinal microflora to ferment undigested substrates entering from the ileum. The VFA produced can be used by the hindgut bacteria for protein synthesis, resulting in an increase in microbial mass, or absorbed in the large intestine. The contribution of large-intestinal VFA absorption towards the total energy maintenance requirements of dogs has been reported to be approximately 2–7%\textsuperscript{179,180}. However, the latter authors did...
not to provide information on the way these values were derived. In addition, the effect of production and absorption of acetate as an energy source for body tissues on postprandial satiety remains to be investigated. The work of Poutau et al. on a method to evaluate acetate production and metabolism using stable isotopes may be the starting point for further exploration of the importance of carbohydrate fermentation in the gastrointestinal tract and satiety in dogs.

To our knowledge, there is no information available in the scientific literature regarding possible influences of dietary fibre on ghrelin concentrations and behaviour in dogs. However, when dogs are fed one scheduled meal per day, ghrelin concentrations increase before and decrease rapidly after the meal to remain relatively constant throughout the rest of the day, which may indicate little potency of ghrelin concentrations to affect canine behaviour throughout the day.

Nowadays, dry extruded diets for dogs may contain 30% or more carbohydrates of which starch is the major component. Moreover, the non-digestible carbohydrate fraction in diets can also make up a considerable proportion. As mentioned previously, fibres differing in physical and chemical properties have diverse physiological responses in animals. Nutrient digestion as well as transit time through the gastrointestinal tract may be influenced by the amount and source of fibre included in canine diets. In the case of a reduction in nutrient digestibility when fibres are included, it is necessary to increase the concentration of some nutrients in order to ensure that the nutrient requirements of the animals are met. Future canine research on the behavioural effects of dietary fibre should account for the fact that different breeds may respond differently (in terms of satiety). Gastric emptying rate is inversely related to body weight in dogs of different sizes. Moreover, large-breed dogs have a longer large-intestinal transit time and increased apparent total dietary fibre digestibility, which may increase the production and the use of VFA but may increase gastrointestinal discomfort as a result of enhanced fermentation activity.

The degree of satiety in animals such as pigs has been shown to affect behaviour, including aggressive and stereotyped behaviour. Although likely, it is up till now unknown whether canine behaviour can be affected by degree of satiety and further research is required. Assuming that behaviours in dogs are more favourable during times of satiety than during times of hunger as observed in pigs (for example, aggression), specific dietary fibres through their potential to prolong satiety may assist in preventing unwanted canine behaviours.

Conclusions

The present contribution provides an overview of current knowledge on the influence of dietary macronutrient composition on canine behaviour. It can be concluded that little research has been conducted in this field although research in other species indicates that there is potential to modify behaviour in dogs through nutrition. There is evidence that dietary composition can modulate animal and human behaviour through different mechanisms. Dietary protein may contain the precursors tryptophan and tyrosine for the respective neurotransmitters serotonin and catecholamines. Since bioavailability of both tryptophan and tyrosine in the brain are dependent on the dietary protein content and amino acid composition, dietary composition may have an impact on the behaviour and wellbeing of dogs under specific circumstances (for example, stress). However, before application and extrapolation of the evidence found in mostly rodent laboratory studies into commercial canine diets is undertaken, research is required to identify the optimal and safe dietary inclusion level in combination with behavioural tests to study the magnitude of effects on (problem) canine behaviour. The n-3 PUFAs have an important role in the development of the brain, and the supply of essential fatty acids such as DHA could affect aspects of the dopaminergic and serotonergic system and, consequently, cognitive performance and behaviour as observed in rodents. Most canine studies and dietary n-3 PUFAs have been mainly focused on the effect of maternal intake of different dietary n-3 PUFAs during gestation and lactation on n-3 PUFAs in the milk and/or n-3 PUFAs intake on retinal function of puppies. It would be of interest to examine the DHA required for optimal neurological development and whether this leads to alterations in cognitive abilities or behaviour later in life of dogs. In the literature, studies have been reported which show that, depending on the physical and chemical properties, certain dietary fibres induce satiation or prolongation of satiety after a meal. However, there have been no studies conducted in which the effect of dietary fibre on physiological satiety parameters, behaviour (for example, activity) and/or feeding motivation were studied in dogs. If dietary fibre has short-term effects that result in prolongation of satiety and a reduction of hunger between meals, it may help to prevent unwanted canine behaviours and also promote long-term weight control.

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