Thermal and digestive constraints to foraging behaviour in marine mammals

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While foraging models of terrestrial mammals are concerned primarily with optimizing time/energy budgets, models of foraging behaviour in marine mammals have been primarily concerned with physiological constraints. This has historically centred on calculations of aerobic dive limits. However, other physiological limits are key to forming foraging behaviour, including digestive limitations to food intake and thermoregulation. The ability of an animal to consume sufficient prey to meet its energy requirements is partly determined by its ability to acquire prey (limited by available foraging time, diving capabilities and thermoregulatory costs) and process that prey (limited by maximum digestion capacity and the time devoted to digestion). Failure to consume sufficient prey will have feedback effects on foraging, thermoregulation and digestive capacity through several interacting avenues. Energy deficits will be met through catabolism of tissues, principally the hypodermal lipid layer. Depletion of this blubber layer can affect both buoyancy and gait, increasing the costs and decreasing the efficiency of subsequent foraging attempts. Depletion of the insulative blubber layer may also increase thermoregulatory costs, which will decrease the foraging abilities through higher metabolic overheads. Thus, an energy deficit may lead to a downward spiral of increased tissue catabolism to pay for increased energy costs. Conversely, the heat generated through digestion and foraging activity may help to offset thermoregulatory costs. Finally, the circulatory demands of diving, thermoregulation and digestion may be mutually incompatible. This may force animals to alter time budgets to balance these exclusive demands. Analysis of these interacting processes will lead to a greater understanding of the physiological constraints within which the foraging behaviour must operate.

Keywords: digestion; thermoregulation; foraging; diving; marine mammals; bioenergetics

1. INTRODUCTION

Scientists often refer to animals performing in an ‘optimal’ manner, either referring to how they allot their time to different behaviours, or the performance of their physiological systems. In truth, what an animal does and how it does it is the result of a series of compromises. Time is finite, and an animal rarely faces the luxury of having only a single objective to fulfil. Similarly, increasing the performance of a given physiological system will almost inevitably detract from other concurrent requirements. Like time, energy, blood and nutrients are also finite. With that in mind, an animal’s observed foraging behaviour may only appear logical (or more importantly, predictable) if it is viewed as a concession to a number of concurrent demands.

Marine mammals face distinctive environmental conditions that can translate into unique physiological challenges. The following review examines the potential physiological constraints (or concurrent demands) to foraging behaviour in marine mammals.

We present a novel framework that centres around three broad processes: prey acquisition; prey consumption; and thermoregulation. The framework allows us to review the effects and complex physiological interactions that shape the foraging behaviour at the individual level. We briefly review the scientific research that contributes to this paradigm, although the scope of the task precludes a complete synopsis of any single section. More importantly, we have placed past research in a novel larger context, and highlighted gaps in our current knowledge base that should be addressed in future studies. Our aim is to help direct future research with a view to improving quantitative foraging/energetic models.

There is a stark dichotomy in the general approaches to foraging models between scientists studying terrestrial mammals and those concentrating on marine mammals. Studies of terrestrial foraging have primarily approached the problem as an attempt by the animal to optimize energy and time in the acquisition of resources (for review, see Pyke 1984). This approach has resulted in a vast realm of theory that has largely been derived from initial optimal foraging models based on marginal value theorem (Charnov 1976; Stephens & Krebs 1986).

In contrast, studies of the foraging strategies of marine (diving) mammals have primarily focused on...
the physiological constraints that limit prey acquisition. In particular, these studies have been predominantly concerned with the way that oxygen reserves limit the time available below the surface or at depth to actively pursue and capture prey. The role of oxygen storage/usage in limiting foraging time is an overwhelmingly unique, almost defining, aspect of marine mammal foraging behaviour. Therefore, it should not be surprising that this constraint has been the focus of most of the research on marine mammal foraging. However, there are other physiological constraints that define foraging patterns in marine mammals which are often neglected. In this paper, we will discuss the interaction of three physiological processes that impose limitations on foraging patterns: the physiological demands of prey acquisition; prey processing; and thermoregulation.

Within these aforementioned physiological processes, we have defined three general types of potential conflict. First, there may be an absolute time restriction if the activities are mutually exclusive. Second, increased energy costs of one activity may also increase the costs (or decrease the efficiency) of another. Conversely, increased costs in one parameter may actually decrease the costs of another. Third, physiological changes required to maximize one process may be incompatible with other processes. Therefore, animals have to potentially prioritize among activities and/or manage with suboptimal performance in several aspects of their physiology.

Certainly, the physiological interactions of digestion, thermoregulation and foraging are not unique to marine mammals. Marine mammals, after all, are anatomically and physiologically similar to terrestrial mammals in most respects. For example, while Williams et al. (2001) discerned allometric differences in the size of the small intestine between marine and terrestrial carnivores, there is no indication that the digestive physiology and biochemistry of marine mammals are fundamentally different from those of their terrestrial equivalents. And while marine mammals live in an environment with greater potential for heat loss (the specific heat capacity of water is 25 times that of air), many terrestrial mammals live in equally thermally challenging environments. However, the physiological adaptations used by marine mammals to maximize underwater foraging times potentially impact on their ability to digest prey and maintain thermoneutrality, and vice versa (figure 1). While these interactions may not represent over-riding physiological constraints to foraging behaviour in the same manner as the balanced usage of oxygen stores, we will illustrate how they are significant parameters that should be addressed in models of marine mammal foraging behaviour and energy budgets.

2. LIMITS ON FOOD INTAKE
The ability of a marine mammal to meet its energy demands with sufficient prey is dependent on a number of limiting factors. In the framework of this paper, we have divided these factors into constraints on the processes of prey acquisition and processing.

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Prey acquisition is limited by the amount of time that an animal can spend foraging and the proportion of that foraging time which can be spent at depth pursuing prey (figure 1 no. 1). Foraging time is impacted by the intrusion of other incompatible behaviours. As an obvious example, the time spent ashore (to nurse, mate, or for thermoregulatory considerations) directly decreases the potential foraging time. Within that potential foraging period, the amount of time that an animal can spend actively pursuing prey is also limited by the proportion of time it can spend at depth, minus the time ‘wasted’ diving to depth and recovering at the surface (inter- and post-dive intervals). These aspects of foraging efficiency are closely tied to the animal’s aerobic dive limit (ADL), which is determined by physiological capacity and aerobic requirements (see §2a).

Food intake is further limited by aspects of prey processing. While additional prey may be available for capture, an animal can only consume a finite amount over a specific time, given the anatomical and biochemical limitations of the digestive system itself. The rate at which an animal can process prey is primarily limited by its consumption capacity (initial intake) and digestion time (passage rate through the system; figure 1 no. 2).

Finally, there are several avenues of interaction between prey acquisition and processing. There are indications that circulatory adjustments required to maximize diving and digestion may be, to a degree, incompatible (figure 1 no. 3; see §2c). Additionally, the metabolic requirements of digestion may impinge on the foraging efficiency by decreasing ADLs (figure 1 no. 4).

(a) Prey acquisition

The amount of time a marine mammal can spend at depth actively pursuing prey during a single dive, the frequency of those dives and the inter- and post-dive recovery periods are all related to its ADL. The ADL is defined as the amount of time that an animal can remain submerged without relying on anaerobic metabolism (i.e. without accumulating lactate; Kooyman et al. 1980, 1983). It is not that an animal is unable to dive beyond this physiological point, but there are important repercussions. Primarily, increased surface intervals are required to clear the circulating lactic acid levels from the blood. Aerobic diving limits are often viewed as a major determinant of diving ability (Costa et al. 2001); therefore, they represent a critical constraint to foraging behaviour (Castellini 1991). There is a growing body of literature that has examined the aspects of physiology which define an animal’s ADL. Despite the obvious logistical difficulties in monitoring blood lactate levels in a diving marine mammal, such biochemical changes have been examined in both the laboratory and the field in several species (see review by Costa et al. 2001). As a generalization, it appears that marine mammals only rarely approach (let alone exceed) their ADLs when foraging in the wild.

In lieu of actual measures of ADL, several studies have relied upon the concept of calculated aerobic dive limits (cADL). This value is derived by estimating the animal’s total internal oxygen stores (including haemoglobin stores in the blood, myoglobin stores in the muscle tissues and, in some cases, pulmonary stores in the lungs) and metabolism during the dive (the rate at which oxygen stores are consumed).

There are several problems with these estimates. First, they assume that all of the oxygen stores in the body are used completely (and evenly) before anaerobic metabolism begins. In truth, selective vasoconstriction will ensure that certain tissues will become hypoxic before all oxygen in the body is depleted. Indeed, low partial pressures are required before the substantial amounts of oxygen stored in myoglobin can be released from muscle stores (Davis et al. 2004). Second, there are very few direct measures of diving metabolic rate in marine mammals. These are usually restricted to captive animals and may not be comparable to animals foraging in the wild. As a result, there appears to be a discrepancy between cADLs and the behaviour of marine mammals in the wild, with some studies reporting a substantial proportion of observed diving bouts exceeding cADLs (Costa et al. 2001) and others reporting most foraging dives far below the cADL (Croll et al. 2001). Obviously, more reliable data would come from studies that have directly measured lactate levels in diving mammals, but the challenges in such research are considerable.

Physiologically based modelling of diving behaviour must not only take into account the behaviour observed during single foraging dives, but also explain the larger patterns of repeated dives with short surface intervals that are characteristic of most marine mammal foraging. This has led to a body of work that has produced models of optimal dive times (Houston & Carbone 1992) and (inversely) optimal breathing patterns (Kramer 1988). As succinctly stated by Green et al. (2005), foraging models of diving homeotherms, in a sense, have been primarily concerned with viewing oxygen rather than prey as the limited or patchy resource that defines behavioural patterns.

Our ability to test models of foraging behaviour and prey acquisition in marine mammals is further hampered by our general inability to differentiate between non-foraging, and successful and unsuccessful foraging dives. Although emerging technologies hold promise of actual measures of foraging success (e.g. Andrews 1998; Hooker et al. 2002; Austin et al. 2006), the current link between diving behaviour and actual foraging usually has to be inferred.

(b) Prey processing

Foraging models usually equate rates of prey acquisition with rates of energy and nutrient intake, i.e. prey processing. Rarely is there any consideration of the ability of the animal to process its food within a given time frame. Digestive constraints represent a finite limitation in the rate of energy intake. Digestive capacity (sometimes referred to as ‘consumption capacity’) is the amount of food that an animal can process over a sustained period. It is affected by the maturity of the animal, the size of its digestive system, the type of digestive system, the digestibility of prey, the flexibility of the digestive system to different prey types and availability, and levels and type of concurrent activity.

The importance of digestive capacity as a physiological bottleneck in the chain of resource acquisition and assimilation is not a novel concept.
The frequency of feeding episodes is another way that digestive capacity can impact foraging behaviour. Most animals do not forage continuously; there is evidence that fragmented foraging periods may increase maximum intake levels (Zynel & Wunder 2002). However, this also means that animals must fulfil their daily energy requirements within a restricted period each day. Similarly, if animals are unable to forage every day (e.g. due to other required behaviours or lack of prey availability), then the amount of prey they have to consume to maintain overall energetic homeostasis must be proportionally increased. Restricted maximum foraging times will contribute to increased required food intake rates during the periods of foraging that approach the digestive capacity of an individual animal (see study by Rosen & Trites 2004 detailed below).

It is recognized that the digestive capacity has important ecological implications (Weiner 1992; Kersten & Visser 1996), including potentially explaining the proximate physiological cause of a metabolic ceiling (Peterson et al. 1990). However, most measures of digestive capacity are restricted to avian species (e.g. Zwarts et al. 1996; Kvist & Lindstrom 2003; Zharikov & Skilleter 2003; Van Gils et al. 2005), with only a few for carnivorous mammals (Kirkwood 1983; Zynel & Wunder 2002). This narrow focus on avian digestive capacity may be due to the fact that, in general, birds consume food which is abundant but high in indigestible material and thus requires long processing times (Bednekoff & Houston 1994), and that they have limited capacity for energy storage due to flight considerations. Therefore, many birds may live near the edge of their digestive capabilities.

Whatever the reason, there is a paucity of information on maximum digestive capacity among mammalian carnivores, and there exist only a few estimates for marine mammals. Kastelein published several reports of digestive capacity (measured as maximum sustained food intake) of captive marine mammals (Kastelein et al. 1990b, 1994, 1995). One paper (Kastelein et al. 1990a) reports the food intake of seven Steller sea lions (Eumetopias jubatus) (ranging in age from 1 to 16 years) as between 4 kg (the youngest) and 26 kg (the oldest male) after a day of fasting. However, this consumption was not sustained and the data are not available to determine the relationship between animal mass and digestive capacity.

In a study of young captive Steller sea lions, the digestive capacity was estimated by measuring the maximum food intake rates with varying levels of energy density of prey and foraging opportunities (Rosen & Trites 2004). The animals were normally maintained on herring at a ration equivalent to 5–7% of their own body mass. In the experiment, they were offered either high-energy herring or lower-energy capelin, free of human interference or performance demands.

The primary aim of the experiment was to determine the capacity (and inclination) of these animals to increase food intake to compensate for differences in prey quality over short (5–10 days) periods of time. Based on differences in gross energy content, the sea lions would have had to consume 83% more capelin than herring. In fact, the sea lions consumed on average 89% more capelin during the trials when food was offered ad libitum for 7 h each day (figure 2).

In order to ensure that the animal’s digestive capacity was being measured, it was necessary to increase the rate of food intake required to fulfil energy requirements. Rather than increase the individual animal’s daily energy requirements, the amount of time available to consume the required food was decreased. In one of the treatment combinations, the sea lions were offered food every other day (for 7 h), rather than daily. Theoretically, the sea lions should have consumed twice the amount of herring to maintain the same average intake levels over 2 days, and 236% more capelin, compared to trials when herring was offered every day. Although the sea lions increased their intake rates under these conditions, they appeared unable to consume sufficient prey to maintain their usual energy intake levels (figure 2). More interestingly, there appeared to be a maximum level of food intake equivalent to 16–20% of the animal’s own body mass (figure 3). Behavioural observations appeared to confirm this physical limitation.

Thus rough estimates of maximum food intake can be integrated with data on available foraging times and ranges of prey energy densities to estimate a variety of...
parameters, including the minimum frequency and energy density of prey needed to sustain an animal. It can also be used to estimate minimum energy deficits that arise from scenarios where an animal is unable to ingest sufficient prey to meet its energy requirements. Unfortunately, estimates of maximum food intake are rarely integrated into models of individual foraging behaviour to set realistic endpoints to food intake variables.

Digestive capacity may be more problematic in younger animals, given their naturally higher metabolic rates due to growth. Larger body size also allows animals to fast longer, given that the metabolic requirements scale intraspecifically to body mass $^{<1.0}$. A similar discrepancy of scaling exists between metabolic requirements and gut capacity, the latter of which scales isometrically with mass (Peters 1983; Schmidt-Nielsen 1984). This suggests that smaller animals have a smaller digestive capacity relative to energy requirements when compared with larger animals. It is unclear whether this scaling discrepancy exists on a developmental basis—although work has been done on the ontogeny of specific aspects of digestive physiology (e.g. Buddington & Diamond 1989; Debray et al. 2003), almost no work has been published on changes in maximum food intake with age. However, their relatively higher energy requirements imply that young animals are generally living on the edge, energetically speaking.

But what are the implications of these relationships, particularly in regard to foraging behaviour? First, younger animals may be under pressure for greater selectivity in prey types. Although many marine mammals are often categorized as 'generalist' feeders (often based solely on the observation of a varied diet), they may be more selective than is typically appreciated. The innate physiological limitation in food intake may require that animals bypass lower-quality prey resources and continue to 'search' for prey which can more readily fulfill their energetic requirements. Like many animals, this suggests that resource quality (gross nutritional value) may be as important as resource availability in driving prey selection. Whelan & Brown (2005) provide a review of the relationship between prey quality, prey selection and digestive constraints, but clearly more work needs to be done on the interactions between prey selection and digestive capacity in marine mammals.

(c) Trade-offs between prey acquisition and processing

There are at least three potential conflicts between prey acquisition (diving) and processing (digestion): time-budget restrictions; interactive energetic costs; and incompatible circulatory changes. In practice, it is very difficult to clearly differentiate these potential conflicts from observations of foraging behaviour alone, and the interactions between these demands are probably more complicated than we fully appreciate to date.

Time-budget restrictions may often be the result of physiological incompatibilities between diving and digestion, such that the time devoted to processing prey impinges upon time available for acquiring more prey. This disjunction may result from incompatible circulatory adjustments required to maximize the efficiency of diving and digestion (figure 1 no. 3). Given the scientific focus in marine mammal physiology on the adaptations they posses to maximize their diving ability, it is somewhat surprising that the results of these circulatory changes—specifically the potential for conflicting demands in circulatory patterns—have been largely overlooked.

Figure 2. Food intake of captive Steller sea lions with different combinations of prey type and feeding opportunities. Average daily food intake ($\pm$ s.d.) relative to the 'baseline' amount of fish consumed during the daily herring phase of the 'satiation' study. Dotted lines represent 'expected' values required to maintain an equivalent gross energy intake based on relative energy content of herring and capelin and daily or alternate day prey availability. Asterisk indicates significant differences between observed and expected consumption levels. Figure modified from Rosen & Trites (2004).

Figure 3. Ad libitum food intake of captive Steller sea lions. The frequency of relative daily food intake (expressed as percentage of body mass) for five captive Steller sea lions in all the treatment combinations during 'satiation' trials (Rosen & Trites 2004).
Many early studies of marine mammal physiology specifically studied the ways that circulatory changes were employed to limit metabolism and, therefore, increase aerobic dive times. Irving et al. (1935) and Scholander (1940) helped define the classical ‘diving response’ which included the three pillars: apnea; bradycardia; and peripheral vasconstriction. Scholander (1964) contended that vasconstriction turned marine mammals into ‘heart, lung and brain machines’. Initial studies with restrained animals suggested an ‘absolute’ response of peripheral vasostriction and bradycardia (Scholander’s ‘dive reflex’). Subsequent studies suggested that some of the previously observed adaptations were by-products of the forced nature of the submergence. Later studies on free-diving animals suggested that the dive response was, in fact, graded to the length of the dive (Castellini 1991) and may not be apparent in shorter shallow dives (Kooyman 2002). Still, it would appear that the demand for peripheral vasconstriction during longer foraging dives may be incompatible with the high rates of blood flow required for digestion and assimilation processes.

One of the most comprehensive studies on circulatory changes during marine mammal diving was conducted by Zapol et al. (1979). They studied changes in blood flow in a variety of systems in a Weddell seal (Leptonychotes weddelli) during forced submergence. Although changes in blood flow in the digestive system were not specifically measured, the overriding vasconstriction observed throughout the seal—with the notable exception of the brain—adds weight to the theory that diving and digestion demand incompatible circulatory adjustments (figure 1 no. 3). This potential conflict is discussed further in §4c.

However, even if foraging dives are restricted to depths where there is no circulatory conflict between diving and digestion, there will be an inherent metabolic conflict (figure 1 no. 4). The various processes of digestion (mechanical and chemical breakdown, assimilation) are known to increase the rates of oxygen consumption (known as heat increment of feeding (HIF) or specific dynamic action; Blaxter 1989). Increased metabolism due to digestion will impact foraging ability by increasing metabolic overhead, thereby decreasing available aerobic dive time. At some point, the cost of digestion could possibly decrease the energetic efficiency of foraging to the point where it was no longer profitable.

The potential problems of incompatible circulatory demands and increased foraging costs have led to the (often assumed) disjunction between foraging and digestive activities. In the most extreme expression of this scenario, animals would concentrate solely on prey capture until their immediate gut capacity had been reached. At this point, the animal is forced to rest rather than continue foraging (Crocker et al. 1997). Drift dives, characterized by episodes of languid, non-powered ascents or descents (Crocker et al. 1997; Biew et al. 2003) have been hypothesized to function as a period of inter-foraging digestion (as well as an opportunity to purge metabolic waste; Crocker et al. 1997).

Of course, this division between foraging and digestion does not necessitate that an animal should fill its stomach and then completely digest the contents before resuming foraging. A host of other factors (including prey availability, indirect costs of foraging, other required behaviours, flexibility in digestive capacity and processing times) help determine when an animal should switch between digestion and foraging. Some studies have clearly demonstrated that periodic foraging patterns increase maximum consumption capacity (Zynel & Wunder 2002).

The relationship between digestion and foraging is probably best characterized as a physiological ‘compromise’ in the sense that neither process is functioning at maximum efficiency. However, another rarely considered factor—the anatomical ability of marine mammals to consume their prey at depth—may necessitate a complete separation of digestion and foraging. Animals are ultimately limited in the size of the prey they can consume at depth by their gape size. Returning to the surface to consume prey alters foraging costs through increased energy and time costs, and potential increases in predation risk (e.g. killer whales, Orcinus orca). Nonetheless, it is common to observe seals coming to the surface to tear apart large fish such as salmon. The decision to forage on prey that has to be brought to the surface to be consumed represents another clear incompatibility between foraging and diving behaviour.

Sea otters (Enhydra lutris) process much of their prey at the surface. However, sea otters must also contend with a rather unique temporal conflict between foraging and thermoregulation. Rather than relying on a blubber layer, the sea otter’s incredibly thick pelage serves as the primary thermal buffer (Morrison et al. 1974), aided at lower temperatures by vasconstriction (Costa & Kooyman 1982). This results in two conflicting behavioural requirements. First, sea otters must consume tremendous amounts of food (equivalent to 20–25% of its own body mass daily), partly to maintain thermoneutrality, and therefore spend a large portion of their day foraging. However, to maintain the low thermal conductance of their pelage, they must also spend a large portion of their day grooming at the surface (Kenyon 1981). This grooming behaviour is also, in itself, a surprisingly energetically expensive behaviour (Yeates et al. 2005), adding to the animal’s required daily food intake.

Clearly, the behavioural and physiological demands of digestion and foraging interact in a number of ways, not all of which are clearly understood. However, this complex relationship may prove central to constructing accurate bioenergetic and behavioural models of marine mammal foraging.

3. CONSEQUENCES OF INSUFFICIENT FOOD INTAKE

Important to the discussion of physiological constraints to foraging are the subsequent effects of changes in body mass and composition which may result from an animal’s inability to obtain energetically adequate levels of prey. Tissue catabolism is the direct result of an inability to obtain or process sufficient food to fulfil energy requirements. During certain periods, the usage of internal energy reserves is a predetermined strategy.
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(i.e. part of their natural life history), but at other times, it is the result of unexpected energy deficits. To understand the consequences and limitations of this strategy, it is important to consider the subsequent effects on physiology, foraging costs and thermoregulatory capabilities.

(a) Changes in body composition
When an animal is unable to satisfy its energetic requirements from external sources (figure 1 no. 5), it must catabolize internal tissues. The anatomical source of mass loss can be almost as important as the degree of tissue loss. The preferential usage of either lipid or protein sources under certain conditions not only results in changes in relative body condition (figure 1 no. 6), but also reflects physiological ‘decisions’ resulting from numerous conflicting requirements and constraints.

Lipid is generally regarded as the preferred tissue for catabolism as it provides greater energy per mass (approx. 39 kJ g⁻¹) than protein (approx. 18 kJ g⁻¹; Schmidt-Nielsen 1997). Additionally, lipids provide greater amounts of metabolic water than protein (107 and 41 g per 100 g tissue, respectively) that may also be lacking from external sources. According to general theories of fasting physiology, an animal should primarily use the lipid stores to fulfill additional energy requirements, except for a small amount of protein catabolism required for gluconeogenesis for the central nervous system (Øritsland 1990). Only when lipid stores are depleted should substantial protein usage occur, since protein degradation is the proximate mechanism for death via organ failure (Cherel et al. 1992).

It is assumed that most mammals with predictable seasonal periods of food restriction (owing to either food scarcity or behavioural demands that preclude foraging) will display protein-sparing physiological adaptations. Certainly, there is a trend for mammals that experience such seasonal changes in energy balance (i.e. predicted periods when they must catabolize tissues) to have a substantial discrete hypodermal blubber layer in addition to lipid reserves situated around certain internal organs (e.g. Scholander et al. 1950; Pond 1978; Leader-Williams & Ricketts 1982; Adamczewski et al. 1995). The blubber layer of marine mammals fulfills the (sometimes contrary) dual roles of insulation and energy reserve (except animals such as sea otters, polar bears (Ursus arctos) and, to a lesser extent, fur seals where pelage is the primary insulative organ). In field studies, the extent of subcutaneous lipid stores has been taken as an indication of a pinniped’s energy state, health status or overall ‘condition’ (Renouf et al. 1993; Naess 1998; Pitcher et al. 2000; Tierney et al. 2001).

The animal’s lipid reserves are often expressed as a proportion of its total body mass, yielding a simple (or perhaps simplistic) body condition index.

However, the relative extent of a diving mammal’s blubber reserves also affects the animal’s thermoregulatory capabilities. Decreases in insulation due to tissue catabolism will potentially increase thermal costs (figure 1 no. 7), leading to further imbalances in the animal’s energy budget (figure 1 no. 8), and eventually to further depletion of the lipid reserve. Depletion of the primary thermoregulatory organ can potentially lead to an escalating discrepancy between increasing energy requirements and decreasing energy reserves.

Energy requirements may also increase due to changes in buoyancy resulting from a reduction in lipid reserves. Changes in buoyancy directly affect the cost of diving (and even the animal’s gait; figure 1 no. 9) which not only decreases foraging efficiency by decreasing ADLs but also potentially affects its energy balance (figure 1 no. 12) and, in turn, further depleting its lipid reserves.

Therefore, while primarily using lipids for fulfilling energy requirements (versus non-lipid tissues, primarily ‘core’ protein structures) is preferential from the perspective of maximizing energy yield per gram of tissue and sparing vital protein structures, there are additional considerations that may mitigate this strategy. Using greater core tissues will help maintain the thermal layer and preserve buoyancy, thereby limiting the cost of foraging (and limiting further energy deficits). However, while this strategy may save immediate locomotor and thermal costs, it may also have significant short- and long-term health costs.

There is a great deal of data on the patterns of tissue usage in pinnipeds. In general, they follow general mammalian fasting theory and primarily use lipids. Although there are a substantial number of studies, almost all are on phocid seals (see review by Castellini & Rea 1992). Few studies have been undertaken on otariid seals which generally have smaller relative lipid reserves than phocid seals. Studies of other marine mammals have concentrated on describing observed seasonal and developmental changes in lipid stores (Lockyer et al. 1985; Lockyer 1993; Naess 1998; Dunkin et al. 2005).

In general, phocid seals follow the mammalian strategy of employing protein-sparing metabolic strategies during natural fasts. Otariids appear to follow a similar strategy, although their relatively smaller lipid layers may preclude such extreme metabolic adjustments. Beauplet et al. (2003) found that 56% of mass loss was derived from lipids and only 10% from proteins in naturally fasting sub-Antarctic fur seal (Arctocephalus tropicalis) pups (the remainder being metabolic water). Similar results were reported for naturally fasting Antarctic fur seal pups (Arnould et al. 2001b) and adult males holding territories (Boyd & Duck 1991).

Further investigations have suggested that the pattern of tissue catabolism in marine mammals is more complicated, reflecting shifts in energetic priorities. In experimentally fasted/food-restricted captive Stellar sea lions, approximately 60% of mass loss could be attributed to lipid sources (Rosen & Trites 2005; Kumagai et al. 2006). However, the relative rates of lipid versus protein tissue usage were dependent on both season and food quality (lipid/protein content). When on short-term restricted feedings, the proportion of total mass loss comprising lipid varied significantly between seasons, ranging from 24 to 107% (figure 4). In addition, a greater proportion of the total mass loss was derived from lipid stores while consuming a low-lipid/high-protein diet than while consuming a high-lipid/low-protein diet, except during the summer (a natural period of high growth) when the trend was substantially
Figure 4. Changes in Steller sea lion body mass and composition during food restriction. Mean ± s.d. of rate of total body mass loss (black bars, averaged across diets) during 9 days of restricted feeding of herring or pollock in seven captive Steller sea lions (n=7 for each season). Mass loss was significantly greater in the winter trials. The contribution of lipid loss to total mass loss is also shown separately for herring (grey bars) and pollock (white bars) diets. There was a significant season × diet interaction—more of the mass loss was derived from lipid stores while consuming pollock than while consuming herring, except in the summer when the pattern was reversed. More than 100% lipid loss in the herring group in the summer suggests that they were gaining lean body mass while losing overall body mass. Figure from data in Kumagai et al. (2006).

Another consideration with respect to patterns of tissue catabolism is that the depletion of lipid stores may be further constrained by the morphological function they serve in swimming. In cetaceans, blubber functions not only in buoyancy control, but also in streamlining and as structural components (Pabst et al. 1999). This can lead to preferential lipid usage at different sites of the body. For example, in harbour porpoises (Phocoena phocoena), the thorax appears to be the primary site for lipid deposition and depletion (Koopman et al. 2002). Although substantial lipid masses are located at the tailstock, these do not appear to be depleted during periods of undernutrition. This suggests that the blubber at the tailstock is required for maintaining locomotory capabilities and streamlining.

(b) Changes in swimming costs

As previously described, most studies of marine mammals indicate that they primarily lose lipid mass rather than protein mass during periods of negative energy balance. Two potential results of such a strategy would be a decrease in thermoregulatory capacity (figure 1 no. 7; see §3c) and an alteration in the animal’s buoyancy (figure 1 no. 9). While there has been substantial research on the energetic consequences of the former, only recently has consideration been given to the latter.

It has been well documented that buoyancy affects diving performance (Lovvorn & Jones 1991). Therefore, it is not surprising that there is also evidence that buoyancy influences behavioural decisions about swimming in marine mammals. Efficient locomotion is particularly important for breath-holding divers, as increased locomotor costs will not only incur direct energetic costs (figure 1 no. 12) but also limit dive time (and decrease foraging efficiency) through increased oxygen depletion rates (figure 1 no. 11; Skrovan et al. 1999; Sato et al. 2003).

Change in gait (active strokes or gliding) is one strategy that animals use to optimize locomotion. Selection of a specific gait depends partly on the relative buoyancy of the animal in the medium (Williams 2001). An animal’s buoyancy naturally decreases with water depth, largely through compression of pulmonary air stores (Williams et al. 2000), a physical characteristic that many divers appear to take advantage of, as evident through changes in gait with depth and individual density. However, the energetic consequences of changes in buoyancy induced by changes in lipid stores are not clear. Most work to date has concentrated upon changes in diving behaviour with changes in buoyancy.

A common characteristic in marine mammal swimming biomechanics is alternating between active stroking and gliding, during either ascent or descent portions of their dives. The use of gliding behaviour during diving has become loosely categorized as ‘drift diving’. As an example, right whales (Eubalaena spp.) must use strong strokes at the surface to counteract their positive buoyancy, but they can conversely take advantage of this buoyancy to power glides on their ascent (Nowacek et al. 2001). Sperm whales (Physeter macrocephalus) and bottlenose dolphins (Tursiops truncatus) also glide more during portions of dives when buoyancy aids their movement (due to natural changes in buoyancy with depth), and whales that glide more during ascent glide less during descent (and vice versa; Skrovan et al. 1999; Miller et al. 2004).

The plasticity of this change in diving behaviour is evident by comparing swimming strategies of individuals with different body densities. Sato et al. (2003) observed that fatter Weddell seals descended with a stroke-and-gliding method, while thinner individuals prolonged their gliding phase. Gliding appeared to be a more efficient locomotor technique, as surface intervals between dives were less for gliding than for stroking animals.

In grey seals (Halichoerus grypus), body fat (hence density) was demonstrated to be related to dive characteristics—although all animals were negatively buoyant, those with higher fat mass (relatively more buoyant) had slower descent rates (Beck et al. 2000). These findings would suggest a cost to greater lipid stores, except that the more buoyant seals also had slower ascent rates, contrary to predictions. Similarly, in a study of northern elephant seals (Mirounga angustirostris) which had their buoyancies experimentally altered, less buoyant individuals had faster descent rates, but there was no relationship between buoyancy...
and rate of ascent (suggesting that ascents were all powered; Webb et al. 1998). In contrast, Biuw et al. (2003) found that fatter southern elephant seal (Mirounga leonina) pups had higher ascent and slower descent rates during the drift portion of dives. They even constructed a model that reasonably predicted measured lipid content by dive profiles alone. Some, but not all, male New Zealand fur seals (Arctocephalus forsteri) have been observed to use drift dives—the ones that did were mostly larger males (Page et al. 2005).

Changes in buoyancy are not just a by-product of changes in body composition. A number of diving vertebrates use lung capacity to alter buoyancy to maximize dive performance (e.g. Lovvorn & Jones 1991; Sato et al. 2002; Hays et al. 2004; Miller et al. 2004; Wilson & Zimmer 2004). However, other physiological considerations (e.g. nitrogen narcosis, decompression sickness, oxygen toxicity) may limit the amount of air or time during which this strategy may be used (Moore & Early 2004; Hooker et al. 2005). Again, it should be noted that actual changes in the cost of diving with changes in body composition (through either changes in buoyancy or gait) has rarely been measured. In theory, increases in the cost of diving and swimming through direct energetic cost (figure 1. no. 12) and increased metabolic overhead for aerobic dives (figure 1 no. 11) can have substantial impacts on observed short- and long-term foraging behaviour.

(c) Changes in thermoregulatory capacity

The previously discussed tendency for marine mammals to preferentially metabolize stored fat to offset energy deficits can potentially reduce valuable insulation to the point of increasing the energetic cost of thermoregulation (figure 1, no. 7). This can contribute to the onset of a downward spiral of reduced body condition and increased energy deficit, until thermal balance can no longer be maintained (figure 1, no. 8).

The challenge of balancing the usage of the blubber layer for thermoregulation and for an energy source is especially relevant to young animals which, given their higher surface area to volume ratio compared with larger adult animals, would be expected to have higher rates of heat loss in water. Additionally, maintaining thermal neutrality may be a relatively greater energetic burden to juveniles which are also faced with the energetic costs of anabolism required to continue growing and complete ontogeny. As juveniles mature, their ability to thermoregulate also influences the energetic costs associated with the development of swimming and diving skills necessary for their primarily aquatic existence (Donohue et al. 2000).

The extent to which changes in body condition affect thermoregulatory abilities and costs has been directly measured primarily in pinnipeds, and mostly in young animals. This focus is partly due to logistical concerns, and also due to the fact that this age class potentially faces the greatest thermal challenge owing to high surface-to-volume ratios and high energetic requirements for growth.

Much work on the thermal effects of changes in blubber reserves on energy requirements has been completed on phocid seal pups (e.g. Irving & Hart 1957; Worthy 1991; Muelbert & Bowen 1993). Phocid pups typically undergo a long post-weaning fast on land. Besides facing the immediate paradox of having to use blubber as an energy source yet conserve it as an insulative layer, ‘weaners’ must also consider the future impact that blubber depletions would have on survivorship during subsequent foraging trips. For example, northern elephant seal pups exhibit differing patterns of energy usage depending on body mass and body composition at the end of weaning. Fatter pups use more fat and spare proportionally more protein than leaner pups. This individual variation in energy usage in elephant seal pups may ensure that leaner pups become fatter (as a proportion of total mass) by maintaining a higher level of protein catabolism, thus enabling all pups to have sufficient lipid stores for thermoregulation during their first foraging trip (Noren et al. 2003; Noren & Mangel 2004). Northern elephant seal pups that have sufficient lipid stores at the end of the post-weaning fast were shown to be able to remain thermally neutral in 4°C water (Noren 2002).

Similar results have been shown for harbour seal pups through energetic modelling exercises (Harding et al. 2005). Model results suggest that the cost of thermoregulation varies dramatically with body size. Smaller pups (17–32 kg) were estimated to require more than three times the extra energy to compensate for heat loss over the winter months as compared with larger (heavier) seals. The model predicts that the first year of winter pup survival is strongly related to pup masses measured in autumn, and that pups which do not reach at least 26 kg are likely to suffer the effects of cold stress at the onset of winter water temperatures (Harding et al. 2005).

Fewer studies have measured how body condition affects the thermal capabilities of otariids. A study of Antarctic fur seals (Arctocephalus gazella) found that both pups and yearlings displayed elevated resting metabolic rates in cold water (0.6°C); in fact, the lower critical temperature for both age groups was estimated to be surprisingly high, 14.4°C. Both total body mass and percentage of lipid mass predicted thermoregulatory abilities in both age groups (Rutishauser et al. 2004).

While examining the ontogeny of thermoregulatory capabilities in northern fur seal (Callorhinus ursinus) pups, Donohue et al. (2000) noted that whole-body metabolic rates of post-moult pups were equivalent to those of smaller pre-moult pups measured in water. In other words, the mass-specific metabolic rates of pre-moult pups were higher than post-moult pups. This observation, coupled with stable body temperatures in post-moult pups, was thought to reflect increased thermoregulatory capabilities of post-moult pups compared with the greater thermal costs to pre-moult pups across a wide range of water temperatures (5–20°C). Decreasing mass-specific metabolic rates as the pups matured suggested energetic savings due to larger body size, increased lipid stores (from 14–18 to 34–39% as pups mature) and insulation provided by the post-moult pelage. Similarly, the dependence of sub-Antarctic fur seals on their pelage for reducing thermoregulatory costs is evident in the decrease in field metabolic rate from pre- to post-moult pups, along with the corresponding increases in lipid reserves (from 10.4 to 37.1%, respectively; Beaufort et al. 2003).
There are fewer direct measures of changes in thermoregulatory capabilities with changes in lipid mass in cetaceans. On an interspecific basis, small cetaceans from temperate climes have greater lipid reserves than more tropical species (Worthy & Edwards 1990). Loss of lipid in the former may have greater energetic consequences, given their colder environment (and hence greater potential for heat loss), although tropical species may actually live closer to their lower critical temperatures (Hampton & Whittow 1976). Emaciated bottlenose dolphins display a significant rise in the conductivity of their blubber layer, suggesting that the lipid content (quality) of the blubber layer changed rather than the actual depth (quantity; Dunkin et al. 2005).

Bioenergetic modelling has provided a vehicle to examine some of the constraints on thermoregulation that are otherwise difficult to collect on free-ranging animals (Hokkanen 1990; Boily 1995; Kvadsheim et al. 1997). For example, Roscow (2001) modelled the degree of heat loss in Steller sea lions under a variety of aquatic and terrestrial environmental conditions, from which costs of thermoregulation were estimated for animals over a range of body sizes (50–1000 kg) and body conditions (gauged by blubber depth, 1–5 cm). In general, under all environmental conditions, as the thickness of the blubber layer decreased the energy required for thermoregulation increased, so that sea lions with poor body condition (1 cm in blubber depth) experienced the highest thermoregulatory costs. For juvenile and adult sea lions (100 and 500 kg, respectively) at rest, critical blubber depths for three water temperatures (−2, 4 and 8°C) were modelled. Critical blubber depth (below which a thermoregulatory cost is incurred) was 3 cm for adults, when modelled with average winter water temperatures (4°C), and 2 cm with summer water temperatures (8°C). Juvenile sea lions were predicted to need slightly thicker blubber (3.5 cm) to avoid thermal costs at these water temperatures. In cold water (−2°C), critical blubber depth was never attained for either juveniles or adults with up to 5 cm of blubber depth. In air (0°C), critical blubber depth was approximately the same (between 1.25 and 1.5 cm) for juveniles and adults. Like many models, the results are limited by quality of physiological data currently available for this species, and further studies are required to refine the sensitivity of the model (e.g. Hoopes et al. 2004).

Despite the uncertainties involved in modelling thermoregulation, the depletion of a marine mammal’s lipid reserves can have serious thermoregulatory consequences. To survive in the face of limited food intake, the animal must adopt a series of physiological and behavioural responses that limit or reverse the continuing cycle of depletion of insulation and resulting increasing energy deficit.

4. ENERGY AND CIRCULATORY TRADE-OFFS BETWEEN FORAGING AND THERMOREGULATION

The effect of depletion of the blubber layer on thermoregulatory costs only becomes important when it impinges on the minimum insulative layer required for thermal neutrality. The thickness of this minimum layer is difficult to ascertain, as it relies on many physiological and environmental factors, many of which have not been adequately explored. In fact, few measures of thermoneutral zones (or even lower critical temperatures) exist for marine mammals (Liao 1990; Williams et al. 1993; Hansen et al. 1995). Comparisons are complicated by the observation that pinnipeds have two effective lower critical temperatures—one on land and another in the water—that can differ substantially (Irving & Hart 1957), and the impact of activity and digestion on ‘effective’ thermoneutral zones (see below).

Although some species have relatively ‘skimpy’ lipid reserves, others have significantly more than required for thermoregulation alone, what Watts et al. (1993) coined ‘thermoregulation below the zone of irrelevance’. As an extreme example, Hokkanen (1990) observed that the blubber layer of the bowhead whale (Balaena mysticetus) was sufficient to maintain thermoneutrality in liquid nitrogen. However, whether a marine mammal has ‘overabundant’ insulation or not, there are significant interactions between thermoregulation and foraging that must be considered.

For those marine mammals that also live in the terrestrial environment, individuals must often balance the costs and benefits of thermoregulation and foraging behaviour. Pinnipeds must find a compromise between remaining on land where thermal costs may be lower than in the water (but at the expense of receiving no energy input), and spending time in the water foraging where they have greater opportunities to overcome energy debts through food intake (but may expend more energy in thermoregulation). An extreme example of the spatial discord between thermoregulatory and foraging considerations may be demonstrated by seasonal baleen whale migration from productive high-latitude feeding grounds to low-latitude breeding grounds. This movement has traditionally been explained as a required disjunction between areas optimal for foraging and those for thermoregulation in young calves (Brodie 1975), although the hypothesis is not universally accepted (Corkeron & Connor 1999).

Thermal considerations may modify short-term foraging behaviour. For example, potential heat loss increases with swimming speed due to increased costs of convective heat loss (Hoopes et al. 2004), and changes in water temperature with depth will increase the thermal gradient that the animal must combat.

It is important to note that thermoregulation and foraging are not always conflicting energetic demands. For pinnipeds, decreasing air temperatures and increasing precipitation and wind will result in environmental conditions under which thermal costs are lower in the water than hauled out on land. Additionally, the hypometabolism exhibited during diving may help offset the costs of thermoregulation (Costello & Whittow 1975; Castellini et al. 1992; Hurley & Costa 2001; Hastie et al. 2007).

A great deal of heat is also generated through muscular activity during behaviours such as diving and swimming. Similarly, heat is produced during the mechanical breakdown and biochemical absorption of food. Thus, swimming and digestion create heat ‘by-products’ that can potentially be used to offset costs of
thermoregulation rather than simply expended as metabolic ‘waste’ (figure 1 no. 10 and no. 13).

In addition to energy trade-offs, physiological adjustments in circulation may be inconsistent between thermoregulation, foraging and digestion. For example, vasoconstriction that promotes diving capabilities may limit heat dissipation to the skin surface, thereby impinging upon thermoregulatory capabilities (figure 1 no. 10). Similarly, vasoconstriction during diving that limits blood flow to the digestive system will decrease the animal’s ability to simultaneously process and acquire prey (figure 1 no. 3).

(a) Substitution of heat from activity

The idea that animals should minimize their energetic waste partly led to the suggestion that animals should divert ‘excess’ heat produced from activity to substitute for the heat required for thermoregulation (figure 1 no. 10). The importance of heat generated by muscular activity for offsetting thermoregulatory costs has been demonstrated in a range of homeotherms (Webster & Weathers 1990; Bevan & Butler 1992; Zerba & Walsberg 1992; Girardier et al. 1995; de Leeuw et al. 1998; McNamara et al. 2004; Kaseloo & Lovvorn 2006), including semi-aquatic mammals (Kruuk et al. 1994; MacArthur & Campbell 1994; Campbell & MacArthur 1998; Campbell et al. 2000). Studies have suggested that the level of thermal substitution varies with the levels of activity (heat production) and the rates of potential heat loss (Williams 1986; Hind & Gurney 1997; Kaseloo & Lovvorn 2005).

The benefits of using heat from muscular activity to offset thermoregulatory costs are only beneficial if the animal is not thermal neutral. Within the thermoneutral zone, the energy produced for basic metabolic processes is (by definition) sufficient to maintain core body temperature. The thermal balance (net heat flow) is a product of the external environment, the insulative properties of the animal and the level of internal heat generation. While the high thermal conductivity and heat capacity of water present unique challenges to marine mammals, their hypodermal blubber layer (or in the case of sea otters and some fur seals, their exceedingly dense pelage) usually provides sufficient insulation to limit heat loss during ‘normal’ physiological and environmental conditions (§3c).

Thermal substitution of heat from activity may not be possible in many cases. There is a tendency for some diving marine mammals to actually decrease body temperature during diving (Scholander et al. 1942; Kooymans et al. 1971). Whittow (1987) suggested that the observed decreases in core temperature indicated that metabolic production during dives was actually lower than at rest (although see Kooymans et al. 1980; Hill et al. 1987). This hypothesis is supported by the studies that have observed decreases in metabolic rate during diving (Costello & Whittow 1975; Castellini et al. 1992; Hurley & Costa 2001; Sparling & Fedak 2004; Hastie et al. 2007). A lack of excess heat production would obviously make any type of thermal substitution less probable.

However, hypometabolism during diving is not universal among marine mammals (and often only exhibited in deep long dives), and core temperatures have not always been observed to decline with submergence time. Increases in body temperatures during diving and swimming have been observed in a number of pinnipeds (McGinnis & Southworth 1967; Ohata et al. 1972) and cetaceans (Hampton et al. 1971; Whittow et al. 1974; Hampton & Whittow 1976). It has even been suggested that small cetaceans can only maintain internal body temperatures through the substitution of heat generated through activity (Parry 1949; Hampton & Whittow 1976).

Unfortunately, changes in core temperatures cannot definitively indicate whether thermal substitution is occurring, as it only indicates the end product of the combined demands of environmental load, internal heat production and dissipation ability. Studies that document changes in total metabolic costs across different (internal and external) thermal conditions would provide more convincing evidence.

(b) Substitution of heat from digestion

HIF is the increase in the rate of oxygen consumption (and resulting metabolic heat production) that follows the ingestion of a meal. This phenomenon is thought to result from both the mechanical and biochemical processes of digestion (Blaxter 1989). HIF has been quantified in numerous species consuming a variety of meal types, including several species of marine mammals (for review, see Rosen & Trites 1997). HIF can comprise a significant proportion (more than 20%) of gross energy intake.

It has long been suggested that homeotherms might be able to use the increase in heat production from digestion to offset concurrent thermoregulatory costs (figure 1. no. 13; Rubner 1902; Kleiber 1975; Lavigne et al. 1982). Unfortunately, empirical evidence to support this hypothesis is equivocal (Robbins 1993; Rosen & Trites 2003). While some studies of terrestrial mammals have provided evidence that HIF can at least partially substitute for thermoregulatory costs (Simek 1975; Masman et al. 1988; Chappell et al. 1997; Jensen et al. 1999), others have indicated little or no interaction between these two bioenergetic parameters (figure 5; Klaassen et al. 1989; Campbell et al. 2000).

Several variables may affect the likelihood of thermal substitution occurring. First, more heat produced through digestion will mean a greater source and greater chance of detecting thermal substitution. In general, high protein diets will induce greater HIF (Jobling 1983), which has also been demonstrated among marine mammals (Rosen & Trites 1997). Second, a higher potential for heat loss (such as through increased thermal gradient, conductivity of the medium) should result in greater thermal substitution. Therefore, one might expect that the tendency for substitution would be greatest for aquatic mammals, given their potential for losing heat to their environment. However, while there is evidence to support thermal substitution among some aquatic mammals (Costa & Kooymans 1984), other experiments have failed to support this hypothesis (MacArthur & Campbell 1994; Campbell et al. 2000; Rosen & Trites 2003; Williams et al. 2004b). The apparent inconsistency in results between studies that have and have not reported thermal substitution raises
Figure 5. Theoretical interactions between HIF and thermoregulatory costs. (a) In one scenario (thermal substitution), the total cost of metabolism due to digestion and thermoregulation remains constant across environmental temperatures, even below the animal’s lower critical temperature. (b) In an alternate scenario (thermal independence) total metabolism increases as temperature decreases due to increased thermoregulatory costs regardless of digestive heat production.

interesting questions for comparative physiologists. Uncertainty whether thermoregulation and HIF are cumulative or offsetting costs also presents a potentially large source of error when calculating energy budgets or constructing bioenergetic models.

The aforementioned studies all measured the interaction between thermoregulation and digestion through changes in total energy production (figure 5). An alternative avenue for an energetic interaction between digestion and thermoregulation is when HIF decreases thermoregulatory costs by helping to maintain core temperatures (irrespective of changes in total metabolism). Several studies have demonstrated the combined effects of digestion and environmental temperature on both maintaining body temperature and reducing shivering thermogenesis (Costa & Kooyman 1984; Maloney et al. 1999; Rashotte et al. 1999). This includes a strategy of delayed digestion to maximize the substitution effects of HIF during the most thermally challenging periods (Rashotte et al. 1999). However, other studies of the effect of HIF on maintaining body temperature have yielded mixed results (MacArthur & Campbell 1994; Koh & MacLeod 1999). This includes hypometabolism in certain tissues and decreased core temperatures across a variety of environmental temperatures. In addition, studies may be more fruitful by investigating the conditions that define the extent of substitution which occurs in an individual animal rather than classifying species into those that do or do not exhibit thermal substitution.

(c) Circulatory conflicts
Vasoconstriction is observed during long dives in marine mammals and serves primarily to maintain blood pressure in light of bradycardia. Vasoconstriction also results in certain circulatory changes that may impact thermoregulation while foraging. This includes hypometabolism in certain tissues and decreased core temperatures in very long dives. This section speculates on the potential circulatory conflicts between thermoregulation and diving, and the potential limitation placed on foraging behaviour by over-riding thermoregulatory concerns.

In small cetaceans (e.g. dolphins), the primary thermoregulatory concern during activity appears to be promoting heat dissipation. Surface vessel systems at peripheral sites (flippers and flukes) assist in heat dissipation during exercise and cooled returning blood is used to cool temperature-sensitive internal organs. However, vasoconstriction associated with diving physiology may limit heat dissipation abilities (Whittow 1987). This may partly explain the increases in body temperature observed during swimming in some species (Hampton et al. 1971; Whittow et al. 1974; Hampton & Whittow 1976).

Several studies have examined how bottlenose dolphins manage the seemingly incongruent demands...
of increased circulation for heat dissipation from exercise and vasoconstriction to maximize dive times (figure 1 no. 10). These animals decrease heat flow to thermal windows during submergence and increase blood flow to peripheral sites after exercise, presumably in an effort to maximize heat dissipation rates (Noren et al. 1999). These changes in heat flow were mirrored by changes in heart rate (Williams et al. 1999). These studies suggest that the majority of heat dissipation is deferred until animals surface, thereby maximizing the oxygen-sparing mechanisms associated with diving. However, this deferment of heat dissipation is limited by the animal’s ability to withstand the resulting increase in core temperatures during dives, as demonstrated in studies with (restrained) harbour seals (Phoca vitulina) (Hammel et al. 1977) and ducks (Johansen 1964).

In large cetaceans, the potential build-up of metabolic heat from muscular exercise would be particularly critical (Hokkanen 1990). The sheer allometry of exceedingly low surface area to volume ratios makes internal build-up of critical internal temperatures much more probable. Passive heat loss would be negligible in large cetaceans (Innes 1986); therefore, a greater emphasis would be placed on circulatory transport of excess heat to the surface (Hokkanen 1990). Large cetaceans have demonstrated the capacity to minimize heat build-up even during intense surface travel (Brodie & Paasche 1985), but it is unclear how circulatory changes during diving might interfere with this ability. It is also unclear whether all large cetaceans actually generate the anticipated amounts of heat in the manner of smaller cetaceans. Anatomical streamlining, adjustments in buoyancy and decreased relative speeds may prevent higher metabolic rates and the resulting thermal loads. In some cases, whales even possess particular circulatory adaptations designed to minimize heat loss during foraging (Heyning & Mead 1997).

In contrast to cetaceans, most studies of the thermoregulatory mechanisms of pinnipeds have investigated their methods for minimizing heat loss. This focus derives from their generally smaller body size (therefore, higher surface area to volume ratio) and tendency for polar distributions. Additionally, although the heat capacity of water is greater than air, the potential heat loss while hauled out in the winter is often greater than in the water (due to lower temperatures and higher convective heat loss).

Decreases in skin temperatures observed during periods of swimming and diving imply a change in circulation in agreement with vasoconstriction required to maintain blood pressure during bradycardia in many diving vertebrates (Boyd 2000). Vasoconstriction away from flippers—a major avenue of heat dissipation—serves both to maintain body temperature and to restrict aerobic metabolism, both of which extend foraging times (Gallivan & Ronald 1979; Willis et al. 2005). Although overheating can be a serious problem while on land for pinnipeds, the need for dumping heat during foraging has rarely been addressed. This is partly due to the smaller size of most pinnipeds and possibly due to their generally lower levels of heat generation during swimming and diving compared with cetaceans.

While it appears that the circulatory adjustments required to maximize dive times and minimize thermoregulatory costs in pinnipeds are complementary, this relationship is finite. The same level of bradycardia that minimizes heat loss from the flippers and skin also prevents transfer of muscle-generated heat to the core of the animal, potentially leading to hypothermia. Some pinnipeds may reduce the energy required to maintain body temperatures by reducing their defined core region (another by-product of vasoconstriction) or by lowering their defended body temperature set-point. However, such adjustments will only delay hypothermia. Therefore, longer dives may be restricted not by the ADL but by the animal’s ability to maintain viable core body temperatures.

5. SUMMARY
Mathematical optimal foraging models have been used to explore time-budgeting and diving behaviour of marine mammals in consideration of energy requirements and prey availability, distribution and quality (Boyd 1999; Arnould et al. 2001a; Thompson & Fedak 2001; Houston et al. 2003; Mori et al. 2005). Recently, models have begun to consider the full dynamic process of foraging (from behaviour through energy allocation), by individual marine mammals in variable environments (e.g. Dall & Boyd 2002; Matthiopoulos et al. 2005; Frid et al. 2006). This paper delineates and describes some of the physiological interactions that partly formulate foraging behaviour. Inclusion of these concepts may aid in the refinement of predictive foraging models. This paper also highlights areas where research is lacking, as an aid to defining and stimulating future research in these areas.

This paper specifically examined how the foraging behaviour of marine mammals may be physiologically limited by demands of prey acquisition (diving), prey processing (digestion) and thermoregulation. The ability of an animal to consume sufficient prey to meet its energy requirements is partly determined by its foraging abilities (e.g. available foraging time and diving capabilities) and digestive capacity (e.g. time devoted to digestion, maximum consumption capacity). Failure to consume sufficient prey will further impact foraging, thermoregulatory and digestive capacity through several interacting avenues. Catabolism of tissues to maintain energy balance will cause changes in the cost of thermoregulation and diving/swimming. A downward spiral of increased tissue catabolism to pay for increased energy costs may result. Conversely, the heat generated through digestion and foraging activity may help offset thermoregulatory costs. Finally, there may be trade-offs between the circulatory demands of diving, thermoregulation and digestion.

Observed foraging behaviour is an integration of a multitude of competing demands on an animal. While it is difficult to model or conduct research on all of these parameters, we can concentrate upon those that we deem most important. Physiological processes alone cannot explain diving behaviour, but they represent finite limits to an animal’s abilities. For example, as noted by Castellini (1991), “while metabolic limits define the

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outer boundaries of diving, behavioural patterns place the diving animal at certain points inside that window’.

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