Tissue fuel and weight loss after injury

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In physiological and metabolic studies it is common to refer to a mythical 'average' or '70 kg' man. It is useful to consider how this average adult individual would respond to injury as judged by the literature on injury of the past decade, with particular attention to the ideas which pertain to the supposed energy and fuel demands of this individual. The most common feature of the injured person is a period of prompt weight loss, the rate of which is roughly proportional to the severity of the injury. Extra demands are placed on vital organs. Nitrogen excretion is increased, reflecting protein breakdown. These postinjury changes are consistent with the concept that injury produces large increases in the level of resting metabolism. Such increases in the demand for tissue fuel occur at a time of reduced calorie intake, hence a large weight loss is to be expected as the body consumes its own tissue for fuel. If body protein is considered to be a primary energy reserve of the body, then extra nitrogen excretion would naturally follow the breakdown of protein to meet these extra fuel needs. Several authors have suggested calorie expenditures of 5,000 to 6,000 calories per day (approximately three times normal) following a major injury such as an extensive burn (Artz and Reiss, 1957; Moore, 1959). Extreme weight loss after injury presumably reflects the loss of both protein and adipose tissue in order to meet such large energy demands. In addition to the increased oxidation of protein and fat, there is some decrease in tolerance to exogenous glucose and perhaps a tendency to hyperglycaemia. This has been termed the 'diabetes of injury' and has been interpreted as a reduction in the capacity to oxidize glucose.

Cuthbertson divided the response to injury into an 'ebb and a flow' phase (Cuthbertson, 1942). The 'ebb' period was considered the period of diminished vitality and metabolism with circulatory deficiency as its central feature and occurring immediately after the injury. The 'flow' period was considered the period of increased metabolism or 'traumatic inflammation' which was considered to begin in 24 to 48 hours after the injury and was assumed to correspond to an inflammatory defence and removal of necrotic tissue. This flow phase of convalescence was observed to be accompanied by an increase in blood sugar and body temperature, as well as increases in pulse and respiration and probably increased activity of the hypothalamic-pituitary-adrenocortical axis with depletion of body protein.

Moore (1953) subdivided the 'flow' phase of convalescence after injury into an early or catabolic phase lasting three to seven days after elective operation and a longer period after major injury and sepsis, and then a 'turning point' of a few days with a reduction in nitrogen excretion and weight loss, followed by weeks of anabolism with restoration of protein stores, adipose tissue, and general vitality. The metabolic response in the posttraumatic period is based on the type and extent of the injury and the previous health of the patient. As an approach to grading the severity of trauma, a 'scale of 10' has been proposed (Moore and Ball, 1952). The repair of a hernia would be regarded as 1 or 2 on this scale, whereas the extreme injuries such as major burns would qualify as 9 or 10. In the middle ranges would be ordinarily elective abdominal surgery, such as subtotal gastrectomy or colectomy.

Weight Loss after Injury

Mild to moderate degrees of weight loss are so common that careful efforts to monitor weight loss are often lacking in hospital care. It is
assumed that tissue function is not restricted by mild weight loss and that the loss can be restored without difficulty as convalescence progresses. At the opposite extreme is the prompt but non-specific worry of the clinician when faced with the patient having undergone extreme weight loss. Studley was among the first to call attention to some of the implications of weight loss in preoperative patients (Studley, 1936). In a study of 46 patients operated upon for chronic ulcer disease he found that the percentage of preoperative weight loss was the only factor that correlated with operative mortality rates. The mortality rate was 33% in those patients who had lost more than 20% of body weight before operation. In those who lost less than 20% of their normal weight the mortality was 3.5%. Between the lack of concern with lesser degrees of weight loss and the worry over extreme loss is a large and ill-defined range of weight loss whose development is only slowly noticed, where the composition of tissue loss is not clear, and the metabolic significance remains uncertain.

The extent of weight loss which has been reported as a result of partial starvation provides some background for considering weight loss after injury. The nutritional literature contains many reports of the weight loss of civilians, which reached 10-20% of body weight when subjected to the food restrictions of wartime. Famine conditions and wartime prison life have been reported to cause 15-30% weight loss without being lethal. The cachexia of late starvation, such as in certain concentration camp victims, was associated with the loss of 30 to 45% of body weight. Krieger (1920) attempted to show that a lethal degree of weight loss existed from starvation alone and suggested that 40% loss was lethal in acute starvation and 50% in semi-starvation. Keys and others (Keys, Brožek, Henschel, Mickelsen, and Taylor, 1950) reviewed this concept and emphasized that such figures could only represent rough approximations at best. It appears that underlying organic disease usually decreases the extent of weight loss from starvation which can be survived, and there is some reason to believe that acute surgical conditions further decrease the tolerance to prolonged weight loss.

The weight loss after injury is usually due to a combination of decreased dietary intake and a variable increase in basal energy expenditure. The extent of weight loss has been shown by many workers to be influenced by the patient's sex, body build, the preoperative nutritional status, the degree of injury, and the presence of complicating factors such as infection. Our experience with adult surgical patients suggests that loss of weight between 5 and 10% occurs after mild, uncomplicated forms of injury such as elective operation. Intermediate levels of weight loss, from 10 to 20%, occur within four to six weeks after more severe injury, particularly if involving multiple fractures or major sepsis. If there is continuing inflam-

mation, such as active peritonitis, draining fistulae, or invasive infection from an infected burn surface, the loss of body weight will reach 20 to 30%. Severe tissue wasting will be evident unless obscured by water retention. Weight loss beyond 30% represents precachexia with increasing trouble from weakness, decubitus ulcers, and joint contractures. Death is often associated with bronchopneumonia.

Composition of the Adult Body

An approximate idea of the normal composition of the adult body is necessary to understand the weight loss which occurs after injury. Classic information about the chemical composition of the human body has depended upon the analysis of postmortem tissues. In the last few decades methods have been developed which permit estimates of the body composition of the living subject. These techniques have largely been directed toward the calculation of the body content of fat or water. Fat is an anhydrous tissue with a lower density than lean tissue. Therefore, if the density of the whole body can be measured by immersion (Buskirk, 1961) or by pneumatic means in a chamber (Siri, 1961) an estimation of body fat can be made. Another method which is more feasible for the study of the ill or injured patient involves an injection of a tracer material which is distributed throughout the body water and allows the calculation of body content of the given material from dilution of the tracer (Moore, 1946).

In order to convert the value for total body water to the value of fat-free tissue in the body, the percentage of water in lean tissue must be known. The studies of Pace and Rathburn (1945) have shown that fat-free tissue normally contains close to 73% water by weight. Thus the fat-free body weight obtained from isotope dilution can be subtracted from total body weight to arrive at fat content. The factor of 73% water in fat-free tissue probably has some variation in disease and injury. No methods are currently available for the direct determination of the amount of protein or skeleton in the body weight (Baker, 1961). But evidence from many lines of study suggest that the body composition presented in Fig. 1 is a reasonable approximation of that which is present in an average, healthy adult male weighing 70 kg. The total mineral content of the body, largely found in the skeleton, represents 3 to 5% of the total body weight. This amount is not only small but also relatively static when compared with the changes occurring elsewhere in the body. The remaining 95% of the body can be conveniently regarded in two phases: the aqueous phase, amounting to approximately 55%, and the organic phase, amounting to approximately 40% of total body weight. The organic and the aqueous phases of body composition will be discussed separately, but it is important to remember that the metabolic
Fig. 1  The approximate proportions of aqueous and organic phases are shown for the body composition of an average 70-kg adult male. The small fraction of body weight (3 to 5%) shown for minerals is largely that of the skeleton.

machinery or 'body cell mass' is a combination of organic materials together with intracellular water and electrolytes. In terms of body composition the body cell mass (Moore, Olesen, McMurray, Parker, Ball, and Boyden, 1963) appears to include tissue glycogen and a major portion, but not all, of the total body protein. Therefore it is the body cell mass which is constantly converting oxygen and foodstuffs to carbon dioxide, water, work, and heat. It seems probable that the amount of average turnover of this body cell mass bears a fundamental relationship to alveolar ventilation, cardiac output, and resting metabolic expenditure.

Somewhat over half of the organic material in the body is shown as fat. A body fat content of 17 kg represents 24% of the total body weight. It is recognized that fat can vary from 10 to over 40% of total body weight. The fat content of healthy adults normally increases with age from approximately 10% at the age of 20 years to 25 or 30% at the age of 55 years. Women are usually somewhat fatter than men. Carbohydrate is present in the human body in small amounts only, approximately 300 g of glycogen being present in liver and muscle and in addition 10 to 20 g of glucose in the extracellular water. Therefore the total carbohydrate provides less than 1,500 calories and amounts to less than 1% of total body weight. A protein content of 11 kg in the representative adult male is an estimate based on analyses of individual tissue and calculations by difference, utilizing many human studies, of the content of body fat. The amount of skeletal material and lean tissue and perhaps a small amount of structurally important fat appear to be genetically controlled for the individual. But the total amount of fat is a variable segment of body composition, since it reflects the voluntary alterations of energy balance.

Loss of body weight following operative or physical injury is so common that the metabolic dynamics involved and the potential consequences that may ensue have failed to receive serious consideration. It was not until Cuthbertson's pioneering observation (1932) of the excessive excretion of urinary nitrogen, sulphur, and phosphorus by patients who had sustained fractures of long bones that any significant analysis of posttraumatic weight loss was attempted. When an average 70-kg man loses 15 kg, the various segments of his composition do not share equally in this weight loss (Moore, McMurray, Parker, and Magnus, 1956). The total body water shows small decreases in absolute amount, while the relative amount of water may increase. The extracellular space shrinks with acute weight loss. By contrast, chronic weight loss is associated with intracellular depletion. Acute illness and trauma are accompanied by the accumulation of water and salt. Water accumulates in excess of sodium, resulting in hypotonicity. This response is of particular importance in understanding the body composition of the non-obese patient who has lost more than 10% of his body weight from a major injury with the associated partial starvation. The principal decrease occurs in the intracellular portion of the body, mainly represented by protein and cell water, together with a significant loss of fat. The extracellular space shows a relative, or even an absolute, increase in volume. The body behaves as though protection from chronic weight loss can be provided by enlarging the extracellular space as the body cell mass shrinks. The composition of the blood volume is an excellent indicator of the change in total body composition, since the decrease of the body cell mass is reflected by a decrease in red cell volume, and the abnormal enlargement of the extracellular water is evidenced by an abnormally large plasma volume. Because these changes are in opposite directions, the total blood volume is often equal to the normal amount for this person in health.

Any metabolic behaviour which is thought to occur after injury must account for the extent of the weight loss which is commonly observed. The ideas from the literature indicate that the weight loss is explicable on the basis of injury causing a large increase in energy expenditure, and therefore body substance, particularly protein, is torn down to provide the extra fuel which is required. This formulation is based on a large volume of data about weight loss and nitrogen loss after injury which in general shows a correlation between the rates of nitrogen loss and weight loss. Maximum rates of loss of both nitrogen and total weight occur together (excluding transient diuresis), and it is rare for sustained weight gain to begin before a positive nitrogen balance has been achieved. Such observations are entirely consistent with the idea that protein serves as a major energy reserve during injury and sepsis. Thus measurements are needed to determine what are the increased calorie needs after injury or sepsis, and what calorie contribution is made by the protein breakdown which can be shown to occur.
Energy Requirements after Injury

Because of the central role of energy requirements after injury in determining the amount of body tissue required for fuel and also the caloric contribution of tissue protein to these energy demands, our studies began with efforts to measure the resting metabolic expenditure after various forms of injury and infection. No chamber calorimeters have been designed for the direct measurement of body heat loss which would allow appropriate professional care of the critically ill patient. Therefore we have turned to indirect calorimetry as a way of measuring the amount and composition of each type of foodstuff being metabolized. This involves measuring the intake of oxygen, the output of carbon dioxide, and the excretion of nitrogen. The clinical importance of measuring metabolic expenditure under nonbasal conditions had led to efforts at simplification, but the assumptions and restrictions of the modified methods have made them undesirable for the accurate study of acutely injured patients. Certain techniques require a degree of isolation of the patient which prevents close professional care while the patient is being studied. Therefore a system of continuous measurement of gas exchange in acutely ill patients has been designed for use without a tight-fitting mask or mouthpiece or other attachment to the upper airway. It involves a closed system using a rigid, transparent head canopy (Fig. 2) which is practical and comfortable for long study periods (Kinney, Morgan, Domingues, and Gildner, 1964). A lightweight plastic neck seal provides a leak-proof connection to the canopy, which is ventilated with a continuous stream of conditioned air. The air leaving the canopy is passed through pipes in the wall to a gas analyzer, located in another room, where flow and gas concentrations are continuously monitored. A unique calibration system allows routine measurements of gas exchange to be reliable within \( \pm 5\% \), and often within \( \pm 3\% \).

Experience with this equipment in the study of over 200 patients with various surgical conditions has allowed the establishment of ranges of increase in resting metabolic expenditure relative to the predicted basal metabolic expenditure of an individual of a given size, sex, and age. The normal range for average adult males is considered to be \( \pm 10\% \) of the predicted normal value from BMR tables, as shown in Figure 3. Uncomplicated elective operations are followed by no significant
alteration outside of the ± 10% of the preoperative values. Major skeletal injury commonly results in increases of 10 to 20% above normal for one to three weeks, followed by a reduction to values of 10 to 20% below normal as a reflection of the tissue depletion which has occurred by that time. Infection in a major serous cavity usually produces increases of 15 to 50% above normal, and some elevation above normal will remain as long as inflammation is present, despite tremendous tissue depletion and weight loss. The only form of injury that has sustained levels of hypermetabolism above these ranges is the extensive burn, where the metabolism may remain from 40 to 100% above normal for weeks at a time.

These levels of increased metabolism following injury are doubtless of great significance to the body, but do not reach the levels of hypermetabolism (5,000 to 6,000 Cal/day) which have been suggested in the surgical literature. The measured increase in energy expenditure is not large enough alone to account for the serious, prolonged weight loss in the range of 400 to 900 g per day which is known to occur after major injury. Therefore it becomes increasingly important to gain an understanding of the daily balance between protein and non-protein sources of body fuel, and to examine the associated water loss contributing to this rate of weight loss. This can be done with four or five measurements of gas exchange each waking day when calculated in terms of a resting gas exchange for each 24 hours. This is combined with the nitrogen excretion value to calculate the daily resting non-protein calorie expenditure using the table of Zuntz and Schumberg as corrected by Lusk (1928) as in Figure 4. In order to measure the caloric expenditure of patients out of bed, a portable head canopy was designed which allowed for continuous gas exchange while sitting, standing, or walking under ward conditions. Such measurements in normal volunteers revealed remarkable similarities in the percentage increase of caloric expenditure above resting values. These increases were in the range of 7% for sitting, 17% for standing, and 145% for walking in the hospital corridor at a rate of 1.2 to 1.5 miles per hour. The caloric expenditure during walking was 0.039 Cal/min/kg (Long, Kopp, and Kinney, 1969). Detailed activity sheets were recorded by the nurses for periods up to 23 days on six patients. It was found that during a 24-hour period the patient was supine 50 to 80% of the time and spent less than 10% of the day walking. When these activity figures were converted to calories it was found that the pre-operative activity was approximately 20% greater than calories. The corresponding value dropped to only 5% for the first few days after operation, and then gradually returned to 20% by 12 to 14 days.

**Tissue Composition of Weight Loss**

A study was undertaken to examine the contribution of tissue components of the weight loss that occurred in 10 patients as a result of uncomplicated operation (Kinney, Long, Gump, and Duke, 1968). Changes in daily weight were measured on a hydraulic bed scale, protein content by nitrogen balance, and fat content by indirect calorimetry. The total daily caloric balance was determined by subtracting the total metabolic expenditure from the caloric intake. The non-protein caloric balance was the difference between the total caloric balance and the calories associated with protein loss. The non-protein caloric balance was then divided by a factor of 9.3 to determine the daily loss or gain of body fat. The nitrogen balance for each day was converted to dry protein using a factor of 6.25. In order to calculate the hydrated lean tissue equivalent, a moisture content of 75% was used.

This group lost approximately 6% of their starting weight within 10 days after operation. Fat provided 75 to 90% of the calories while protein accounted for the remainder, as shown in Figure 5. The contribution of water to this weight loss was found to be in excess of the predicted quantity for both men and women. The magnitude of the water loss did not become evident until the second four-day period after operation which suggested a definite antidiuretic influence over the first three days. Continued water loss was observed in many patients during the second postoperative week while protein restoration was under way. The calories derived from this weight loss were unexpectedly low in the men (1,584 Cal/kg) as a result of excessive water loss and a relatively high protein to fat utilization. From such observations it is evident that weight loss cannot be used to predict caloric expenditure and hence is not an appropriate guide to postoperative nutrition.

Another group of surgical patients was studied to determine the proportion of calories supplied by protein after varying degrees of injury (Duke,
Jørgenson, Broell, Long, and Kinney, 1970). Patients undergoing elective operation were similar to the normal subjects with protein supplying an average of 15% of the caloric expenditure. Protein supplied approximately 20% of calories in the tissue fuel of six patients with major trauma and sepsis, while a similar increase in nitrogen excretion of four patients with major burns averaged only 14% of the resting caloric expenditure, since the latter was elevated to an average of nearly 60%. Therefore, protein (or amino acids) provides only 12 to 22% of the caloric expenditure, even in the forms of injury

where the nitrogen excretion per day is significantly increased. Thus, it seems that the nitrogen loss after injury must be occurring for reasons other than merely to provide extra calories.

Our findings are in agreement with those of many other investigators who have noted that the rate of weight loss tends to parallel the extent of negative nitrogen balance. The apparent correlation between weight loss and nitrogen loss can be explained partly on the basis of both the amount of tissue used for fuel and the proportion of fat to lean tissue in the mixture. Fat is an anhydrous form of fuel with approximately 9 Cal/g, while lean tissue has only approximately 1 Cal/g (Kinney, 1959). This is because body protein exists in the body with approximately 3 parts of water, hence the 4 Cal/g of dry protein becomes 1 Cal/g of hydrated lean tissue when body protein is degraded following injury. Therefore, the weight loss associated with major injury or sepsis will have a relatively high proportion of hydrated lean tissue in the fuel mixture when considered by weight, whereas, when considered by calorie contribution, the hydrated lean tissue contribution is only 12 to 22% of the entire resting caloric expenditure. Thus, the extreme weight loss of injury and sepsis is not due to the huge needs for fuel in the metabolic furnace, but rather the body is meeting modest increases in energy demands with a low energy fuel that requires more fuel (and hence has a faster weight loss) to meet whatever calorie deficit exists.

The idea that body protein is the major, or even a good, source of calories whenever the body is faced with large increases in resting metabolism following injury is not correct. Hence, explanations for the nitrogen loss after injury must be sought in alterations of intermediary metabolic pathways other than the final fuel mixture for oxidation.

Intermediary Metabolism after Injury

The pathways of intermediary metabolism between the content of the three major foodstuffs in the adult male body are shown in abbreviated form in Figure 6. From the previous discussion it appears that the tricarboxylic acid cycle, shown in the lower right corner of the diagram, has only modest increases in its need for two-carbon fuel after injury and sepsis, and that protein or amino acid breakdown is a poor source of such fuel. Fat from adipose tissue stores appears to be mobilized and oxidized without difficulty to meet whatever demands occur for extra calories as a result of injury. Therefore, one is left with the question as to the meaning of the extra nitrogen excretion seen after injury and sepsis.

The pathways of intermediary metabolism in Fig. 6 are deliberately oversimplified to emphasize that most amino acids yield carbohydrate

![Diagram of metabolic pathways](http://jcp.bmj.com/)

**Fig. 5** The components of postoperative weight loss over 10 days are expressed as percentages of fat, protein, and water. The percentage of total calories derived from fat approximates 80 to 90% of the total despite contributing only 10 to 30% to the weight loss. Reproduced with kind permission of the publisher from Lusk (1928).

![Diagram of metabolic pathways](http://jcp.bmj.com/)

**Fig. 6** An abbreviated diagram of the major pathways of intermediary metabolism which interconnect the three foodstuffs. See text for discussion of nitrogen loss (largely urea) after injury which is probably more related to mechanisms of gluconeogenesis than to the total demands for two-carbon fuel.
intermediates or are ‘glucogenic’ on deamination. It appears that while fatty acids can provide two-carbon fragments readily for general tissue fuel, they cannot be used to provide a net gain of carbohydrate intermediates, glycogen, or circulating glucose (Coleman, 1969). Therefore, in an actual or threatened situation of energy deficit, fat stores can be drawn upon for calories, but protein represents the only sizeable reserve for carbohydrate intermediates—something which is also critical for survival.

The changes in carbohydrate metabolism in the later phases after injury are not as well delineated as in the shock phase before resuscitation. Several investigators have shown that there is a tendency toward a diabetic-like glucose tolerance curve following elective operation (Hayes and Brandt, 1952) and that this is more marked after severe injury (Howard, 1955). This tendency toward hyperglycaemia and glucose intolerance during convalescence following injury has resulted in such terms as ‘traumatic diabetes’ (Thomsen, 1938) and ‘diabetes of injury’ (Drucker, Miller, Craig, Jeffries, Levy, and Abbott, 1953). This term suggests a relative or absolute lack of insulin activity and presumably a decrease in the levels of blood pyruvate and lactate after injury (Drucker, Craig, Hubay, Davis, and Woodward, 1961). The work of Drucker and his associates (Drucker, Craig, Kingsbury, Hofmann, and Woodward, 1962) has suggested a partial block in the pathways leading from pyruvate to two carbon fragments during the later phases of convalescence as well as the evidence for such changes during the shock phase.

The role of glucose metabolism in injury is further confused by the growing recognition of a syndrome first reported by Evans and coworkers (Evans and Butterfield, 1951) in occasional burn patients without any diabetic history, who developed unexpected lethargy and coma and were found to have marked hyperglycaemia without acidosis. During the past decade there has been growing awareness that lethargy, coma, and even death may occur occasionally in diabetic patients without explanation and has been given the name ‘non-ketotic, hyperosmolar coma’ (Ashworth, Sacks, Williams, and Byrne, 1968). One explanation has been that a sudden increase in circulating corticosteroids could inhibit the action of insulin, but most of the recorded cases have not been on steroid therapy. There appears to be an increasing incidence of this syndrome in acute surgical patients, when receiving large amounts of intravenous glucose.

The evidence for the idea that injury inhibits the oxidation of glucose has been indirect in man or limited to special circumstances in animal or tissue studies. Tracer studies with 14C-glucose have been employed with a mathematical model developed by Dr Jordan Spencer to relate blood and breath specific activities and the rate of CO2 production to the rate of glucose turnover and oxidation in surgical patients (Spencer, Long and Kinney, 1970). To date, 35 glucose runs have been made, including 14 volunteers (Long, Spencer, Kinney, and Geiger, 1970a) or patients undergoing elective surgery. The studies have demonstrated that glucose oxidation is essentially unchanged in minor degrees of trauma which may be expected to reduce glucose tolerance and perhaps increase the resting level of blood sugar. However, in the critically ill patient with a stable circulation and significant hyperglycaemia, our measurements indicate that such a patient has some increase in glucose oxidation as well as an increased glucose turnover in the bloodstream (Long, Spencer, Kinney, and Geiger, 1970b). We have also demonstrated that the hepatic manufacture and release into the bloodstream of new glucose, which is normally inhibited by the administration of approximately 6 g of glucose per hour intravenously, is no longer inhibited by the administration of this amount of glucose to the patient after major injury. The alteration in the control system by which the liver ceases to turn off this process for glucose manufacture is of both theoretical and practical interest, and is a subject of continuing study.

At the present time, neither the utilization of fat nor the breakdown of hydrated lean tissue with excretion of the associated water and nitrogen is adequate to explain the weight loss seen after injury. The additional reason for the rapid weight loss is that water in excess of tissue fuel is being excreted. This may be associated with the normal unloading of previously retained water, or to abnormal renal function. Another factor is the tendency seen in studies of starvation (Kekwick and Pawan, 1956) and paediatric nutritional studies (Holt, 1957) where the body excretes extra water for several days after shifting from a high carbohydrate to a high fat fuel mixture regardless of whether the fat is from the diet or from tissue stores as in starvation. The reverse phenomenon appears to occur when the body returns to a high carbohydrate diet. Analysis of the tissue composition of weight loss in patients following elective operation revealed that 35 to 45% of the weight loss was water in excess of that expected from the hydration of the measured protein breakdown (Kinney et al, 1968).

References


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