Nutrition in the perioperative period

Module 17.1

Metabolic Responses to Surgical Stress

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Learning Objectives

- Understand how the body reacts to injury and surgery;
- Have knowledge about insulin resistance in surgery;
- Have knowledge about how metabolic changes in surgery are related to surgical complications;
- Have knowledge about ways to reduce the catabolic response to elective surgery and how this affects recovery;
- Have knowledge about treatments that may impact the metabolic response to elective surgery;
- Know about patients with special metabolic problem.

Contents

1. The metabolic response to injury
   1.1. Insulin actions under normal conditions
   1.2. Insulin resistance in surgical stress
2. Complications after surgery
   2.1. Metabolism and complications
   2.2. Treating insulin resistance
3. Special metabolic risk groups
   3.1. The malnourished patient
   3.2. The patient with diabetes
   3.3. The patient with cancer
4. Measures that disturbs homeostasis
   4.1. Oral bowel preparation
   4.2. Preoperative fasting
   4.3. Pain control
   4.4. Multimodal metabolic approach
5. Summary

Key Messages

- Surgery and injury initiate a series of reactions causing catabolism;
- Insulin resistance is key to the metabolic response to surgery;
- Insulin resistance is closely related to many surgical complications and to postoperative fatigue and delayed recovery;
- Several simple treatments in the perioperative period can help reduce the catabolic reactions and this will enhance recovery and reduce complications;
- Fasting overnight is obsolete and patients should be given clear fluids to drink up until 2 hours before anaesthesia, preferably a carbohydrate rich drink;
- Avoiding unnecessary oral bowel preparation and good pain control are other important measures to support adequate metabolic and nutritional care.
1. The metabolic response to injury

Injury and surgery immediately spark a series of stress responses in the body. The most important reactions involve the release of stress hormones and cytokines. The magnitude of these reactions is related to the amount of stress inflicted. With greater stress, increasingly strong reactions cause more marked catabolic reactions. Central to all these reactions and the subsequent metabolic situation is the loss of the normal anabolic actions of insulin, i.e. the development of insulin resistance (1), Figure 1. Excessive catabolic reactions are generally not beneficial for the body and a state of catabolism with continuous breakdown of muscle tissue and loss of energy stores prolongs the time to recovery. Hence a key aspect of enhancing recovery after surgery is related to minimising the negative metabolic effects by reducing the catabolic responses and have the patient return to balanced metabolism as quickly as possible again. Key to this process is the maintenance of proper energy and protein balance. Thus, nutrition in the perioperative care is central for recovery.

Figure 1. Insulin resistance after surgery. The relative change in insulin sensitivity (postoperative/preoperative insulin sensitivity x 100, as determined using the hyperinsulinaemic normoglycaemic clamp technique) in relation to the magnitude of the surgical trauma (the size of the operation). From Thorell et al (1).

1.1. Insulin actions under normal conditions

Insulin is the most important anabolic hormone in the body. Insulin regulates glucose metabolism keeping it within very tight limits in the healthy man. Insulin ensures that glucose levels are normalised shortly after food intake by activating rapid glucose uptake and storage in muscle and fat, along with glucose loading in the liver as glycogen. This uptake is stimulated via specific glucose transporters, GLUT 4, that are activated by insulin. These transporters secure active and rapid uptake of glucose in these organs. Most other organs and cells have a transient increase of glucose uptake after carbohydrate intake. This uptake uses other transporters that act upon the prevailing glucose level. Since plasma glucose rises transiently after a meal, these organs also have a transient increase in uptake. This balance and interaction between these two different ways of regulating glucose uptake is important for daily glucose control, but is equally
important for the development of some complications in surgery, as will be outlined below.

Insulin also controls protein metabolism, primarily by reducing protein breakdown in the muscle, but also by supporting protein synthesis in the presence of amino acids. Insulin also controls fat metabolism by stimulating formation of triglycerides and blocking their breakdown.

As outlined above, in insulin sensitive cells, primarily muscle and fat, insulin acts via specific receptors on the cell surface. Inside these insulin sensitive cells specific signalling pathways are activated securing anabolic reactions such as glycogen storage and protein synthesis in muscle or the blocking of lipolysis in fat cells.

1.2. Insulin resistance in surgical stress

With any major injury, such as a major operation, the actions of insulin are overrun as a result of the release of the stress hormones (glucagon, catecholamines, cortisol and growth hormone) and the inflammatory reactions mediated by cytokines. In response to stress, amino acids, free fatty acids and glucose are released to the bloodstream from various tissues. Substrate metabolism is also changed and the body starts to consume fat over glucose. Following medium to large size operations such as colorectal surgery these reactions are reversible with the treatment of exogenous insulin. This was shown nicely in studies by Brandi et al who illustrated that if insulin is infused in sufficient amounts to normalize glucose to normal levels, the rest of metabolism is also normalized (2). In these patients nutrition was provided as total parenteral nutrition during the course of the study. Thus, when nutrition is provided and insulin action is reinstated, protein breakdown is normalised, and free fatty acid levels and substrate oxidation return to normal once the effects of insulin on metabolism are reinstituted. From a clinical point of view it seems that infusion of sufficient insulin to normalise glucose levels can be used as an end point target to achieve these reactions.

2. Complications after surgery

2.1. Metabolism and complications

In recent years it has become evident that the changes in metabolism and the excessive catabolism are involved in the development of many of the common complications occurring after surgery. Hyperglycaemia is one such cause of complications. The development of surgical hyperglycaemia has many similarities to that described for hyperglycaemia in diabetic patients (3). Table 1. However, while insulin resistance is progressive and slow in the development and progression of diabetes, in surgery insulin resistance is rapid in its onset but transient and has passed within weeks in most cases.

Table 1. Insulin resistance and glucose metabolism in diabetes type 2 and in the postoperative patient

<table>
<thead>
<tr>
<th>Factor</th>
<th>Surgery</th>
<th>Diabetes type 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose level</td>
<td>Elevated</td>
<td>elevated</td>
</tr>
<tr>
<td>Glucose production</td>
<td>Elevated</td>
<td>Elevated</td>
</tr>
<tr>
<td>Peripheral glucose uptake</td>
<td>Reduced</td>
<td>Reduced</td>
</tr>
<tr>
<td>GLUT 4 activation in muscle</td>
<td>Reduced</td>
<td>Reduced</td>
</tr>
<tr>
<td>Glycogen formation</td>
<td>Reduced</td>
<td>Reduced</td>
</tr>
</tbody>
</table>

Hence some of key characteristics of hyperglycaemia are an increase in glucose production, a relative reduction in glucose uptake in the periphery and the loss of activation of glucose transporters and glycogen storage in response to insulin stimulation in muscle and fat. The same changes also occur in type 2 diabetes. With insulin
resistance the main mechanism for glucose uptake into the large depots in muscle and fat is blocked. This, along with the increase in glucose production contributes to the sustained elevation of glucose. Instead of glucose uptake in the normal depots, glucose uptake is markedly increased in the organs and cells that take up glucose in relation to the prevailing glucose level, including blood cells, renal cells, endothelial cells and neural cells. With glucose levels increased, these cells have no immediate mechanism of blocking glucose uptake in response to this rapidly developing stress. In addition, these cells have no storage capacity for glucose. This leaves glycolysis as the only metabolic pathway remaining for the elevated glucose inflow. When glycolysis is overloaded, this eventually causes problems for these cells. With massive glucose inflow to the mitochondria the oxidative capacity is eventually overrun and oxygen free radicals are produced. This may cause changes in the cell metabolism that ultimately results in changes in gene expression and signalling. These reactions occur in many cells such as endothelial tissue, kidney, nerve cells and blood cells. These are also the key cells involved in many of the most common complications such as cardiovascular complications, renal failure, neuropathy and infections. While the details of these mechanisms are explored more in depth in diabetic complications, there are several similarities between those in diabetes (3) and in surgical stress.

Recent studies have shown that the main pathways for insulin signalling are disturbed after surgical stress and this blocks the normal anabolic actions of insulin in muscle cells (4, 5). This is also true for fat cells where pathways of insulin are disturbed while pathways enhancing inflammation are enhanced (6). Muscle is also affected by surgical stress, and fatigue is a very common postoperative problem. This fatigue can be explained by a combination of disturbed intracellular glucose metabolism and protein catabolic reaction causing muscle protein breakdown and at the same time limitations of energy in the form of glucose and glycogen. The heart is another muscle that is vulnerable to stress and metabolic disturbances and insulin resistant states. Insulin has also been shown to be a key hormone for tissue healing and hence a state of insulin resistance is accompanied with lower healing capacity (7).

Many of the complications in surgical stress are similar to those occurring in patients with diabetes, Table 2.

<table>
<thead>
<tr>
<th>Cells involved</th>
<th>Surgery occurring within days</th>
<th>Diabetes type 2 developing over years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood / immune cells</td>
<td>Infections</td>
<td>infections</td>
</tr>
<tr>
<td>Renal</td>
<td>Renal failure</td>
<td>Renal failure</td>
</tr>
<tr>
<td>Endothelial</td>
<td>Cardiovascular</td>
<td>Cardiovascular</td>
</tr>
<tr>
<td>Nerve</td>
<td>Polyneuropathy</td>
<td>Polyneuropathy</td>
</tr>
<tr>
<td>Muscle</td>
<td>Fatigue</td>
<td>Fatigue</td>
</tr>
</tbody>
</table>

Thus, it is not only the changes occurring in postoperative glucose metabolism that are similar to those found in diabetes. The pattern of complications and the affected cells are also the same in many instances. While in diabetes the disease progresses slowly and complications usually develop over years, the change in surgery induced glucose metabolism is established within minutes, and the complications usually occur within the first week of surgery.

2.2. Treating insulin resistance

Studies of postoperative patients with moderate stress (APACHE II around 10-15) have shown that controlling glucose levels with insulin impacts outcome by reducing the development of some of more common complications in the surgical ICU (8, 9). In addition, observational studies in patients undergoing colorectal surgery and treated in
surgical wards where a lower glucose level was maintained had fewer complications than those with only slightly higher levels (approximately 1 mmol/l) (10).

Recently, the focus in postoperative metabolism has been on glucose. However, there is an abundant literature from earlier years showing that negative protein balance is also detrimental for recovery after surgery. Protein balance is also under strong influence of insulin. Hence, in insulin resistant states, protein balance becomes negative, mainly because of increased protein breakdown occurs in muscle, but also lack of protein supply. The main effect of insulin is by reducing protein breakdown in muscle, while protein synthesis is mainly stimulated by insulin in the presence of amino acids (11). In stress induced insulin resistance, treatment with insulin can counteract protein losses (2, 12) and support tissue healing (13). Smaller experimental studies in man clearly suggest that retaining insulin action is a key to anabolism and is likely to play a role in the avoiding complications after surgery. In larger clinical studies this notion has been supported in a large single centre randomized trial of patients after mainly thoracic surgery (8). These patients were given a combination of enteral and parenteral nutrition, and when given insulin to normalize glucose to 4.5 – 6.0 mmol/l, the authors reported a marked reduction in complications that affected cells sensitive to hyperglycaemia and a marked reduction in mortality.

In a follow up large multicentre trial of patients under greater stress a similar treatment resulted in the opposite effect with a slightly but significantly higher mortality (14). These seemingly conflicting findings are likely to be explained by some main differences between the trials. The first investigation studied patients under less stress than the second study. This is obvious from the roughly 3 times higher mortality rate in the second trial. In situations of increasing stress, the effect of insulin eventually vanishes and may even be counter productive (9). Secondly, in the second trial, the patients received fewer calories than in the first trial while given large doses of insulin. This may also have contributed to the differences, since insulin is normally only active at times of substrate availability. The effect of providing the calories along with insulin is still not investigated on any large scale, but this may be an important piece of the puzzle in understanding the pathophysiology of surgical stress and how to best deal with it. Lastly, the extent to which the protocol was followed differed between the two trials. In the first one performed in a single unit, the protocol compliance was very good and the variation in glucose levels was substantially less than in the second trial. This may also help explain differences.

3. Special metabolic risk groups

3.1. The malnourished patient

The malnourished patient is at particular risk of complications and slower recovery (15). From a metabolic point of view these patients have substantially smaller reserves of energy and protein and they are also likely to have reduced lower immune function. It is therefore important to identify patients who are malnourished or at risk of becoming malnourished. It is advisable to inform the patient the importance of eating normal food up until the night before surgery and also to be liberal with providing nutritional supplements during the period before the operation (16). In the more extreme and complicated cases, enteral and/or parenteral nutrition is often indicated as outlined in chapter 4.

3.2. The patient with diabetes

The patient with diabetes is another patient with higher risks of complications. These patients are at risk of being catabolic from the very start if their diabetes is not under control. In addition, diabetic patients become even more insulin resistant after surgery. Some reports indicate that it may be the peak glucose value that is related to major
outcomes after surgery, and diabetic patients more often reach higher glucose levels after surgery compared with non-diabetics (17). Studies have shown that patients with diabetes under good control empty a preoperative carbohydrate drink at the same rate from the stomach as healthy volunteers. This was true for patients both on oral medication and those on insulin. The glucose level came up higher and stayed elevated longer than in the healthy controls, but from a safety point of view also these patients may be able to prepare for surgery this way if also given their ordinary morning medication (18).

3.3. The patient with cancer

Patients with cancer coming for surgery often have disturbed glucose metabolism, even if they have not been diagnosed with diabetes. This is illustrated in a recent study in colorectal cancer patients, showing that every fourth patient coming for colorectal surgery without known diabetes had an elevated HbA1c as an indicator of glucose intolerance (10). This is not all that surprising since cancer is known to cause insulin resistance. What was not known was that the patients with an elevated preoperative HbA1c also had a higher glucose level after surgery. In addition, they displayed higher CRP levels and developed more complications, in particular infectious complications.

The surgeon usually meets the patient a few weeks before the patient is to receive the operation. Many of the patients will have cancer surgery, and some will have radiation or chemotherapy, but most of them will have surgery planned for as quickly as possible. In most units the operation can be done within a few weeks. This still allows the patient to prepare metabolically and for the surgeon to institute appropriate treatments to secure that the patient is in the best metabolic and nutritional status by the time of the operation.

4. Measures that disturb homeostasis

4.1. Oral bowel preparation

Oral bowel preparation was used based on the belief that this would reduce complications by minimising the risk of faecal contamination. However, large randomized trials have shown that the use of mechanical bowel preparation before colonic surgery has no such protective effect (19, 20). Oral bowel preparation does however have major effects on fluid balance (20) and it stops the patient from having a meal in the afternoon and evening the day before surgery. This results in a prolonged period of fasting and this disturbs homeostasis. For colonic surgery, mechanical bowel preparation should not be used routinely. For rectal surgery, the information available is less clear (20).

4.2. Preoperative fasting

Preoperative fasting was first proposed in 1848 after the first anaesthetist death reported (21), and became one of the best known medical rules during the last century (22). There is now overwhelming evidence for more liberal fasting guidelines with intake of clear fluids up until 2 hour before the onset of anaesthesia and surgery (23, 24). Nevertheless, the old overnight fasting routine is still practiced in many countries. In addition to causing unnecessary discomfort for the patients, the fasted state of metabolism coming in to surgical stress has been shown not to be optimal (25). Instead of fasting, setting off daytime metabolism with a carbohydrate load has been shown to have several positive effects on outcomes after elective surgery. Many of these effects can be associated with the effect on insulin action and insulin sensitivity that a carbohydrate load can have.
The normal diurnal rhythm of metabolism can be separated into two major entities, daytime metabolism that starts with breakfast and night-time metabolism that prevails during the later phase of the night. Insulin is a key regulator of both. When we eat breakfast insulin is released and activates several mechanisms to ensure that the body stores the nutrients just consumed. Digestion is slow and takes a few hours. Therefore, the effects of insulin remain for 4-5 hours, and are usually still active by the time the next meal is consumed. During the day metabolism is dominated by storage and anabolism, all under the influence of insulin. It is only during the late night when the interval between meals is prolonged that the effects of insulin wanes. At this point other hormones are activated, mainly glucagon and cortisol. They are both anti-insulinergic and catabolic and sets metabolism in breakdown mode. This is the situation that the body comes into surgery if in an overnight fasted state.

A 20% glucose infusion intravenously overnight at a rate of 5 mg/kg/min or intake of 200-400 ml of a carbohydrate rich drink at a concentration of around 12% has been used to break the overnight fasted state. At these concentrations, daytime metabolism is set. This treatment initiates the activation of glucose uptake in insulin sensitive organs (mainly muscle and fat) and breaks the overnight fasted and catabolic state before the onset of the operation (26). Intake of a carbohydrate rich drink also enhances insulin sensitivity. This is probably one of the main reasons for the postoperative effect of substantially lower insulin resistance with the use of the preoperative carbohydrate loading. The carbohydrate load has effects mainly on the peripheral uptake of glucose in the first day or two (27, 28), while later the effects of a carbohydrate load is mainly to reduce endogenous glucose production (29). Both these effects will lower glucose levels. Interestingly, some of the effects remain for a very long time after surgery. This was shown in a study in which glycogen storage capacity was reduced in fasted patients up to a month after elective colorectal surgery while this was much improved with only a preoperative carbohydrate load (30). The mechanisms behind these effects on glucose and protein has recently become more clear with studies showing that the insulin signalling pathways for the major anabolic effects in muscle cells are better preserved with carbohydrate loading compared to placebo (5). The effect is likely to be due to the stimulation of these pathways before the onset of stress by the carbohydrate load.

Setting metabolism in an anabolic state before surgery using preoperative carbohydrates has several clinically relevant effects. With this preparation, protein metabolism is better maintained (31, 32), lean body mass retained (33) and muscle function in the postoperative phase better maintained (30, 32). It is not just the skeletal muscle that is affected by metabolism and carbohydrate loading, since so also is the cardiac muscle. Hence, several reports have shown that the heart functions better when carbohydrate loading is used as opposed to after surgery in the fasted state (34-36).

### 4.3. Pain control

An important part of the catabolic response is mediated by the release of the adrenal hormones cortisol and catecholamines. The release of these hormones can effectively be blocked by the use of epidural anaesthesia (37). The placement of the epidural should be such that it covers the dermatomes around T10 to achieve this effect, preferably at the level of T8-9. Importantly, it is crucial to activate the epidural before the onset of surgery to avoid the release of these potent catabolic hormones.

The epidural has several effects and one that is related to the metabolic aspects of enhanced recovery is pain relief. Pain itself causes insulin resistance (38). Avoiding pain is a key feature during the postoperative phase, and the epidural plays a key role.
4.4. Multimodal metabolic approach

Combining epidural anaesthesia and analgesia with carbohydrate loading reduce insulin resistance in two different ways (as outlined above), and a combined effect can be achieved. This combination allowed complete enteral feeding immediately after major colorectal surgery and continued for several days without any need of insulin, and still glucose levels were kept within the normal range (at approximately 6 mmol/l) Figure 2. This was possible because the two treatments almost completely blocked the development of insulin resistance. The patient in a balanced metabolic state is able to take care of glucose control with endogenous insulin release (39). This is clinically important since rigorous glucose control often requires intravenous insulin with the entailing need for continual adjustments and this is difficult to manage on regular surgical wards.

![Figure 2. Glucose after major colorectal surgery using multimodal approach to metabolic control](image)

Combining mid thoracic epidural analgesia with preoperative carbohydrates while providing complete enteral nutrition (blue line) or hypocaloric glucose achieves the same level of glucose control without any need for insulin. From Soop et al (39).

5. Summary

This module describes the metabolic responses to surgical stress and how this impacts on outcomes. It also covers proposals for treatments that support anabolism and faster recovery after surgery by providing the optimal metabolic setting for postoperative nutrition.

References


