

Acute Colorectal Ischemia after Aortic Surgery: Pathophysiology and Prognostic Criteria

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Acute colorectal ischemia is a rare though potentially lethal complication of aortic surgery. We reviewed our recent experience with 16 cases in order to analyze its causative and prognostic factors. The incidence was 2.8%, and the inferior mesenteric artery was occluded in all cases. All patients also had severe occlusive disease of at least two of the hypogastric or deep femoral arteries. Hypoperfusion due to arterial ligation, prosthetic occlusion or embolism was responsible in half the cases. Ischemia and perfusion due to aortic cross-clamping or perioperative hemorrhage were involved in the rest of the cases. Postoperative mortality was 31%. The mortality was lower for partial, nontransmural necrosis, and for elective operations. Recurrent intestinal ischemia, transmural necrosis, surgery for ruptured aneurysm, intestinal hemorrhage and pulmonary edema were associated with a higher mortality rate. All patients with anuria or extrarenal epuration and hepatic cytolysis died. Although reconstruction of the inferior mesenteric artery might lessen the incidence of postoperative colonic ischemia due to hypoperfusion, the role of oxygen free radicals should be investigated in humans, in order to afford colonic protection against the consequences of ischemia-reperfusion. (*Ann Vasc Surg* 1992;6:111–118).

KEY WORDS: Aortic surgery; colorectal ischemia; occlusive disease; oxygen free radicals; bacterial translocation.

Acute colorectal ischemia occurs infrequently after reconstructive aortoiliac surgery. Its prevalence is between 0.3% and 10% [1–3], whereas the

clinical incidence is 2% and coloscopic incidence is 6% [4]. Mortality, on the other hand, varies between 50% and 100% [2,3,5,6]. Recently, new concepts concerning bacterial translocation, i.e. systemic dissemination of bacteria usually found in the intestinal tract [7,8], and cellular toxicity of oxygen free radicals [9,10], have led to a better understanding of the pathophysiology of acute colorectal ischemia. The reservoir of enterobacteria found in the intestinal tract could play a major role in multiorgan failure usually observed in intensive care patients and could contribute to secondary multiorgan failure observed after acute colorectal ischemia. The role of oxygen free radicals in the genesis of ischemia reperfusion lesions of the intestinal tract is well

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established in the animal [11]. Studies have yet to be conducted in humans.

The objectives of this retrospective study were to: 1) define the trigger factors of acute colorectal ischemia, and in the light of experimental studies, delineate the pathophysiological mechanisms; 2) enumerate the clinical and biological signs of ischemia as well as of bacterial translocation; and 3) analyze the causes and prognostic criteria of mortality.

MATERIAL AND METHODS

Between May 1985 and December 1988, 566 patients underwent reconstructive aortoiliac surgery. Sixteen of these patients (2.8%) had postoperative acute colorectal ischemia. Fifteen patients were men. Mean age was 66.8 years (range 54–82 years). The diagnosis was established by colonoscopy or at laparotomy. Colonoscopy findings included mucosal paleness, purpuric spots, submucosal hematoma, mural ulcerations, or slough. Histopathological studies were performed routinely during colonoscopy to detect findings consistent with acute colorectal ischemia such as arteriolar microthrombosis and mural lesions, including peeled-off or ulcerated mucosa. At laparotomy the absence of pulsations in the mesenteric marginal artery, the presence of purpuric spotting or gangrene, and the transmural aspect of slough were noted. Indications for initial surgery included elective or emergency infrarenal abdominal aortic aneurysmectomy. The procedures performed (aorto-aortic, aortobiliac or aortobifemoral prosthetic bypass) were noted, as well as the clinical signs usually associated with acute colorectal ischemia, including hypo- or hyperpyrexia, diarrhea, bleeding per anum, pain in the left lower quadrant, intestinal obstruction, and shock. Modifications of serum phosphorus, alkaline phosphatases, transaminase enzymes—including aspartate aminotransferases (ASAT) or alanine aminotransferases (ALAT), creatine phosphokinases (CPR), lactic acid dehydrogenase (LDH)—and bacteremia of intestinal origin were noted for each patient.

The type of surgical treatment performed in acute colorectal ischemia (intestinal resection, cholecystectomy, prosthetic thrombectomy) and the different surgical complications observed (recurrent ischemia of the small intestine or colon, colonic stricture, pancreatitis, acute cholecystitis) were listed. The medical complications usually seen during severe infective states included the adult respiratory distress syndrome (ARDS), defined as severe hypoxemia associated with bilateral pulmonary edema and low wedge pulmonary artery pressure; moderate acute tubular disease defined as clearance of creatinine between 5 and 30 ml/min; severe acute tubular disease, defined as clearance of creatinine less than 5 ml/min; moderate hepatic cytolysis defined as high transaminase titers, which were less

than three times the normal value; severe hepatic cytolysis, defined as transaminases more than three times normal values; upper digestive tract hemorrhage caused by acute gastroduodenal ulceration requiring blood transfusion. Prognostic criteria for each of these elements were established.

The delay between the vascular procedure and the onset of colonic ischemia was calculated. The cause of death was noted. The simplified index of severity (SIS) as established by Legall and associates [12], which has been widely used and validated in several pathologies in intensive care, was calculated to classify patients according to their degree of severity. This score was based on 15 variables including age, necessity for artificial ventilation, cardiac rate, respiratory rate, systemic arterial pressure, temperature, diuresis, Glasgow neurological score, natremia, kalemia, base deficit, glycemia, blood urea, white blood cell count, and hematocrit. Statistical analysis of comparisons of percentages was made with the Chi-square test, using Yates' correction for small samples. A *p* value less than 0.05 was considered to be significant.

RESULTS

Clinical findings for all patients are shown in Tables I and II. Ischemic colitis was observed in 12 of 358 cases (3.4%) after cure of infrarenal aorta aneurysm and in four of 208 cases (1.9/100) after surgery for aortoiliac atherosclerotic occlusive disease. Isolated aortic clamping was found as the only promoting factor in eight cases. Cholesterol embolism (*n* = 2), perioperative hemorrhagic shock (*n* = 3), ligation of a patent inferior mesenteric artery (*n* = 1) represent the other causative factors in this series. Two rare etiologic factors were found in our patients: Patient 2 presented an extensive thrombosis of an aortobiliac graft secondary to severe thrombocytopenia due to heparin; Patient 6 had undergone ligation of a previously patent inferior mesenteric artery which led to colonic vascular compromise after transverse colectomy performed for lymphoma. In two cases, the inferior mesenteric artery had been revascularized with satisfactory follow-up arteriographic findings: outcome was favorable without further surgery.

The diagnosis of colonic ischemia was established during surgery for ruptured aneurysm of the infrarenal aorta in three cases, by colonoscopy in eight cases, and by exploratory laparotomy in five cases. Ischemia was located in the left colon in all cases. Necrosis was transmural in six of 16 cases (37%). Clinical signs are listed in Table III. Of 16 patients two had systemic bacterial translocations, one and four days after aortic surgery, and the other, seven days after aortic surgery. In both cases, inoculation was multiple: organisms included *Bacteroides fra-*

TABLE I.—Clinical Findings

Case no.	Sex, age, indications for vascular surgery	Arteriographic findings	Operation	Mechanism of acute ischemia	Localization of ischemia and diagnostic delay	Medical complications	Surgical treatment	Outcome
1	Male, 76 years elective infrarenal aorta aneurysmectomy		Aortoiliac prosthetic graft	Clamping	Right and left colon D12	ARDS hemodialysis digestive hemorrhage	+ D15 total colectomy cholecystectomy	Death + D70 multiple organ failure persistent peritonitis
2	Male, 58 years elective infrarenal aorta aneurysmectomy	Celiac axis SMA, IMA normal	Aortobiiliac prosthetic reimplant IMA	Thrombocytopenia due to heparin	Small intestine Right and left colon D2	Moderate tubular disease	+ D6 thrombectomy graft and SMA	Favorable
3	Male, 64 years elective infrarenal aorta aneurysmectomy	Stenosis celiac axis SMA normal IMA occluded	Thrombo- endarterectomy aortic tube	Cholesterol embolism	Small intestine right and left colon liver, gallbladder D2	ARDS moderate tubular disease severe cytolysis	+ D2 exploratory laparotomy + D3 second look	Death + D3 multiple organ failure
4	Female, 74 years ruptured infrarenal aorta aneurysm	Stenosis celiac axis SMA normal IMA occluded	Aortic tube	Hemorrhagic shock	Left colon gallbladder D5	ARDS hemodialysis severe cytolysis	+ D5 Hartmann + D13 cholecystectomy peritonitis	Death + D62 infective pulmonary pathology
5	Male, 65 years ruptured, infected aortic		Aortobiiliac prosthetic graft	Hemorrhagic shock	Left colon D0		+ D1 Hartmann cholecystectomy	Favorable
6	Male, 60 years aortoiliac atherosclerosis	Celiac axis normal stenosis SMA IMA normal	Aortobiiliac prosthetic graft	Ligation IMA	Left colon D3		+ D28 left colectomy for cicatrical colonic stricture	Favorable
7	Male, 64 years aortoiliac atherosclerosis	Iliac axis normal SMA normal IMA occluded	Aortobifemoral prosthetic graft	Clamping	Right and left colon rectum D6	moderate cytolysis	+ D7 total coloprotectomy cholecystectomy + D9 closure + D20 fistule rectal stump	Favorable
8	Male, 79 years fissurated infrarenal aorta aneurysm		Aortic tube	Clamping	Left colon D8		Coloscopic surveillance	Favorable

ARDS = adult respiratory syndrome; SMA = superior mesenteric artery; IMA = inferior mesenteric artery

TABLE II.—Clinical Findings

Case no.	Sex, age, indications for vascular surgery	Arteriographic findings	Operation	Mechanism of acute ischemia	Localization of ischemia and diagnostic delay	Medical Complications	Surgical treatment	Outcome
9	Male, 67 years elective infrarenal aortic aneurysmectomy	Celiac axis normal SMA normal IMA occluded	Aortoiliac prosthetic graft	Clamping	Right and left colon D2		+ D2 Total colectomy + D4 recto operation for ischemia of ileostomy	Favorable
10	Male, 75 years elective infrarenal aortic aneurysmectomy	Celiac axis normal SMA normal IMA occluded	Thrombo- endarterectomy of interrenal aorta, aortobiliac prosthetic graft implantation IMA	Clamping	Left colon D4		Coloscopic surveillance	Favorable
11	Male, 63 years elective infrarenal aortic aneurysmectomy	Stenosis celiac axis SMA normal IMA occluded	Aortic tube aorto left renal, PTFE graft	Cholesterol embolism	Left colon D2	Hemodialysis severe cytotoxicity intestinal hemorrhage	+ D2 Hartmann cholecystectomy	Death + D2 Necrotizing pancreatitis
12	Male, 60 years aortoiliac atherosclerosis	Celiac axis normal SMA normal IMA occluded	Aortobifemoral prosthetic graft	Clamping	Left colon D7		+ D8 Hartmann	Favorable
13	Male, 71 years elective infrarenal aortic aneurysmectomy	Celiac axis normal SMA normal stenosis IMA	Aortobiliac prosthetic graft	Ligation IMA	Left colon D0	Moderate cytotoxicity	Coloscopic surveillance	Favorable
14	Male, 82 years ruptured infrarenal aortic aneurysmectomy		Aortobifemoral prosthetic graft	Hemorrhagic shock	Left colon D0	Hemodialysis moderate cytotoxicity	+ D6 Hartmann	Death + D14 left and right colonic necrosis
15	Male, 82 years ruptured infrarenal aortic aneurysm	Celiac axis normal SMA normal IMA normal	Aortobifemoral prosthetic graft	Clamping	Left colon and rectum D3	ARDS moderate tubular disease moderate cytotoxicity digestive tract hemorrhage	+ D3 Hartmann + D4 Abdomino- perineal resection cholecystectomy	Favorable
16	Male, 66 years elective infrarenal aortic aneurysmectomy	IMA occluded	Aortic tube	Clamping	Left colon D6	Moderate tubular disease moderate cytotoxicity	+ D6 Hartmann + cholecystectomy + D10 small intestinal fistula	Favorable

ARDS = adult respiratory syndrome; SMA = superior mesenteric artery; IMA = inferior mesenteric artery

TABLE III.—Clinical signs of acute colorectal ischemia

	Patients	Percent
Temperature > 38°5 < 36°5	11	69%
Diarrhea	10	62%
Shock	9	56%
Left iliac fossa	7	44%
Intestinal obstruction	6	38%
Rectal bleeding	5	31%
Bacteremia	2	12%

gilis associated with *Clostridium perfringens* in one, and *Escherichia coli* and *Clostridium perfringens* in the other. **Coloscopy** was performed in eight patients (50%) and in all cases led to the suggestion of the correct diagnosis. **Biopsies** were obtained in six of these patients and **always confirmed** the diagnosis. Blood was drawn for analysis between the evening before and the third day after diagnosis. Results of biologic abnormalities are reported in Table IV.

Thirteen patients (80%) underwent **reoperations**. The **ischemic colon** was **removed** in 11 (70%). In one patient, extended intraabdominal necrosis (liver, spleen, **pancreas** gallbladder, small and large intestines) without therapeutic potential was found. A second look was performed in six patients, and in four a complementary surgical ablative procedure was necessary. Of seven cholecystectomies, three were performed for **acute cholecystitis**. In Patient 2, complete restoration of the colonic vascularity was obtained after thrombectomy.

In cases of adult respiratory distress syndrome, the mean duration of mechanical ventilation was 43 days, compared to 12 days without. Overall mortality was 31% (5 of 16) with an average follow-up of 11 months. Two patients who died were in the group operated on for infrarenal aortic aneurysm. In five cases death was directly related to acute colorectal ischemia, persisting ischemia of the small or large intestine in two cases, and multiple organ failure related to sepsis other than acute colorectal ischemia in three cases (pneumonia, persistent peritonitis in spite of several iterative laparotomies, and pancreatitis with necrosis).

The mean age of patients who died was 71.8 years compared with 64.5 years for survivors (NS). The best predictive factors for mortality (Table V) were kidney tubular disease and severe hepatic cytolysis. The SSI was calculated on the day the diagnosis of acute colorectal ischemia was made, usually the day of admission to intensive care. The SSI score was 18 on an average in the group of patients who died (range 16 to 22) and 11 in survivors (range 6–15) ($p < 0.01$).

DISCUSSION

The **incidence** of acute **colorectal ischemia** in our series of aortic surgery was **2.8%**. This can be broken down to **3.4%** for **surgery of infrarenal aortic aneurysms** and **1.9%** for **aortoiliac surgery**, which is comparable to the literature. Ernst and colleagues [4], employing **routine colonoscopic** examination, found an incidence of **7.4%** and **4.3%**, respectively. Even though these two studies are not comparable, the true frequency of **acute colorectal ischemia** could be **underestimated** by **50 to 60%** in the **absence of routine colonoscopy**. Vascular visualization permits delineation of the type of colonic vascularity most threatened by aortic clamping, as is essentially the case **when the inferior mesenteric artery is occluded**. According to our study, **two trigger mechanisms** of acute colorectal ischemia can be identified: When it is necessary to **clamp the infrarenal aorta**, certain patients experience **transient ischemia** of the **colon**. In this case, colonic vascularization is restored quickly and remains identical to preoperative status.

In other patients, **ischemia** is **prolonged**, either because of perioperative **hemorrhagic shock** or **ligation** of a patent **inferior mesenteric artery** with **previous transverse colectomy** (a mechanism already described [13]) or prosthetic thrombosis due to severe thrombocytopenia secondary to heparin, or cholesterol embolism. No one mechanism can be incriminated in the onset of acute colorectal ischemia. Experimental studies have shown that **ischemia of short duration can lead to irreversible necrosis**. Conversely, in the case of initial ischemia, cellular anoxia is not the only cause of tissue lesions.

The **role of oxygen-derived free radicals** has recently been demonstrated [14,15]. These molecules are highly reactive and are generally produced in small quantities by enzymes such as xanthine oxidase in the small intestine or aldehyde oxidase in the colon [16]. The intestinal cells possess several enzymatic systems capable of eliminating these free radicals, including superoxide dismutase [10,17,20]. During intestinal ischemia, enzymes capable of synthesizing free radicals derived from oxygen and their substrata are produced in large quantities. Reperfusion, i.e. oxygen delivery, is responsible for massive production of free radicals. These mole-

TABLE IV.—Biological changes

	Less than normal	Normal	More than normal
Serum phosphorus	43%		57%
Alkaline phosphatases	0	66%	34%
Aspartate aminotransferase	0	30%	70%
Alamine aminotransferase	0	67%	33%
Lactic dehydrogenase	0	15%	85%
Creatine phosphokinase	0	38%	62%

TABLE V.—Prognostic criteria of mortality

	Deaths	Survival	CHT
Iterative intestinal ischemia	40	27.3	NS
Transmural necrosis	80	63.7	NS
Emergency vascular surgery	40	18.2	NS
Visceral failures	—	—	—
Digestive tract hemorrhage	40	0	<0.05
ARDS	60	0	<0.05
Tubular disease clearance < 5 ml/minute	60	0	<0.01
Cytolysis > 3 × N	60	9.1	NS
Shock	100	27.3	<0.05
More than 2 visceral failures	100	9.1	<0.01
Simplified severity index	18	11	<0.01
Age	71.8	64.8	NS

ARDS = adult respiratory distress syndrome

cules possess major cellular toxicity and are responsible for tissular lesions. It has been shown experimentally that dismutase superoxide and inhibitors of xanthine oxidase reduce the permeability of the intestinal wall and attenuate the intensity of histological lesions, the risk of necrosis, of perforation, and mortality [11,14,15,21–23].

Aminophylline, a substrate of xanthine oxidase, increases histological lesions [14]. In experimental models of severe and prolonged ischemia, the introduction of superoxide dismutase did not modify the permeability of the intestinal wall or the intensity of histological lesions [11]. Under these circumstances, free radicals are not a determinant in the onset of intestinal necrosis: tissular lesions are induced by arterial ligation, and in the absence of sufficient flow, necrosis ensues. This model corresponds best to the acute colorectal ischemia seen in humans after alteration of colonic vascularity, prolonged low-flow states, or cholesterol embolism, and is best named “devascularization acute colorectal ischemia”. Conversely, during isolated clamping of pathological arteries inducing less severe ischemia, the action of free radicals could be responsible for the observed lesions. This model could explain why acute colorectal ischemia can occur in the absence of altered colonic vascularity and is named “ischemia reperfusion acute colorectal ischemia.” Clear demonstration of the role of free radicals in humans is a prerequisite for evaluation of the therapeutic options for colonic protection which have been experimented in the animal.

Arteriographic evaluation of patients at risk could help decrease the incidence of acute colorectal ischemia. When performing aortic reconstruction, it seems preferable to perform inferior mesenteric artery revascularization irrespective of the results of residual pressure measured during operation. In

the absence of revascularization, postoperative hypotension can be responsible for colonic ischemia. This is the only well-established preventive surgical procedure for acute colorectal ischemia [24].

Colonoscopy is the primary investigational tool [25], as it is diagnostic and permits the procurement of specimens for histopathological confirmation. Routine use of colonoscopy could lead to earlier diagnosis. As colonoscopy is simple and not dangerous, this investigative method should be entertained routinely whenever aortic surgery is followed by diarrhea, cerebral disorders, or cardiac, respiratory, or renal failure, even when these entities occur in an isolated fashion. The reported severity of acute colorectal ischemia is most likely due to the delay or absence of proper diagnosis.

Several studies, in animals as in humans, have attempted to find sensitive and specific biological markers for colonic ischemia: In animals these are serum titers of phosphorus [26–28], urea, uric acid [27], LDH [27], CPR [27,29–32], ASAT and ALAT, alkaline phosphatases [27], hexosaminidase [33], vasoactive intestinal peptide (VIP) [34–36]. In humans, CPK [37–39], LDH [39], hexosaminidase [40] and phosphorus [28,41] have been found to be abnormally elevated. It is not known, however, whether or not these markers are specific. Results were difficult to interpret in our 16 patients. Variation of serum phosphorus is not unusual after major surgery. Heavy glucose infusion and alkalosis due to mechanical ventilation can be responsible for the hypophosphoremia observed after aortic surgery. On the other hand, hyperphosphoremia is usually seen only when kidney failure occurs. In two of our patients, however, hyperphosphoremia was seen with normal kidney function, and no cause could be found.

As often occurs after surgery, CPR and LDH were often found to be elevated. High titers of alkaline phosphatases, ASAT and ALAT, when present, are most likely indicative of hepatic failure associated with acute colorectal ischemia [42]. While these abnormalities may often constitute alarm signals, they do not contribute to the topographical diagnosis of postoperative colonic complications. Because of their pharmacological properties, elevated serum levels of hormones usually found in the colon wall could be a partial explanation for the clinical signs observed [43,44]. A prospective study of the dosage of colonic enzymes and hormones is currently underway.

Surgery is indicated according to the aspect of the colonic mucosa and associated signs of severity. When superficial ischemia is found on colonoscopy, operation is mandatory even when only one sign generally associated with infection, such as cerebral disorders, septic shock, pulmonary edema, acute tubular disease, hepatic cytolysis, or acute gastroduodenal ulceration, is present. The surgical procedure performed depends on the condition of the colon. Ideally all necrotic colon must be resected. In our opinion, the frequency with

which acute alithiasic cholecystitis occurs mandates a cholecystectomy whenever laparotomy is indicated.

Two patients (Nos. 7 and 16) had bacterial translocation four and seven days after operation due to transmural colonic necrosis. Such bacterial translocation can occur after clamping of the supraceliac aorta, even in the absence of acute colorectal ischemia [45]. Other authors have reported that, while endotoxemia may occur [46], bacteremia is rare and occurs late.

After traversing the intestinal wall, endotoxin is transported by the mesenteric lymphatic vessels [47] and then enters the systemic circulation without being detoxified by the liver. This might explain why the manifestations of acute colorectal ischemia occur early in comparison to bacteremia. The presence of endotoxin in the peritoneum increases the permeability of the intestinal tract to bacteria [48]. Multiple organ failure is the consequence when endotoxin and then bacteria pass through the injured colonic wall [8,44,48-50]. Portal bacteremia occurs as early as eight hours after insult in animals with severe ischemia (this concerns anaerobic germs only). Systemic bacteremia subsequent to bacterial translocation is found 48 hours later and involves anaerobic germs and particularly *Escherichia coli* [42]. As hepatic cultures have been found to be positive early in the course of infection [42], the liver probably plays the role of a filter, and this might explain the delay between portal and systemic bacteremia.

The results of these studies pose the problem of antibiotic treatment once acute colorectal ischemia has been declared. Because of the delay to systemic bacteremia, appropriate early antibiotic coverage may be difficult. The usefulness of antibiotic therapy in the absence of severe infection or peritonitis has not been shown, and empirical use of antibiotics should be discouraged because of the possible risk of emergence of resistant organisms. However, since there are organisms in the portal system, some form of treatment appears to be logical.

Preoperative intestinal decontamination might be an alternative. In cats, during experimental colonic ischemia, systemic endotoxemia was decreased and death retarded when gentamicin was instilled in the intestinal tract [51]. Associated with mechanical preparation of the colon, intestinal decontamination might reduce the frequency of bacterial translocation. Bactericidal and synergistic intravenous antibiotic therapy is indicated when systemic bacteremia occurs or when surgery is performed.

As shown by the study of predictive factors for mortality, the severity of this disease is well reflected by its consequent medical complications. Acute respiratory distress, anemia, or hepatic cytolysis are negative signs for outcome. The simplified severity index seems to be adapted to the evaluation of acute colorectal ischemia, as the difference between those patients who survive and

those who die was statistically highly significant. Mortality due to acute colorectal ischemia in our institution was 80% in 10 patients between 1968 and 1976, 46% in 13 patients between 1976 and 1977 [6,52], and 31% in the present series. This improvement is related to progress in the treatment of multiple organ failure in intensive care units and aggressive surgery as soon as multiple organ failure occurs. Indeed, only removal of all diseased colon is capable of eliminating the responsible foci.

CONCLUSION

Prevention of colonic ischemia is dependent on preoperative arteriographic findings which permit selection of patients whose colonic vascularity seems to be at risk. The inferior mesenteric artery should be revascularized whenever possible. Early diagnosis of colonic ischemia is primary. Although there are no specific clinical or biological signs associated with colonic ischemia, the indication for colonoscopy must be entertained in the presence of postoperative complications of aortic surgery. When the mucosa only is involved, surgery is indicated if general signs of sepsis are present. In our opinion, operation is mandated even when only one sign of sepsis is present, or when more than the mucosal layer is involved on colonoscopy, even when this finding is isolated. Bacterial translocation is rare and occurs late. When necrosis is not transmural, elective surgery is associated with less mortality than on the average. Repeated intestinal ischemia, transmural involvement, emergency surgery for ruptured aneurysm, intestinal hemorrhage, and pulmonary edema are associated with higher mortality. Severe kidney tubular disease or hepatic cytolysis always lead to death. Evidence that free radicals play a role in this pathology may be a step towards preventive perioperative tissular protection of the colon. The search for specific biological markers might help decrease the incidence and improve diagnostic possibilities as well as the prognosis of acute colorectal ischemia.

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