Secondary Dysfunction of the Intestinal Barrier in the Pathogenesis of Complications of Acute Poisoning

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Abstract—The last decade has been marked by an exponential increase in the number of publications on the physiological role of the normal human gut microbiota. The idea of a symbiotic relationship between the human organism and normal microbiota of its gastrointestinal tract has been firmly established as an integral part of the current biomedical paradigm. However, the type of this symbiosis varies from mutualism to parasitism and depends on the functional state of the host organism. Damage caused to the organism by external agents can lead to the emergence of conditionally pathogenic properties in the normal gut microbiota, mediated by humoral factors and affecting the outcome of exogenous exposure. Among the substances produced by symbiotic microbiota, there are an indefinite number of compounds with systemic toxicity. Some occur in the intestinal chyme in potentially lethal amounts in the case they enter the bloodstream quickly. The quick entry of potential toxicants is prevented by the intestinal barrier (IB), a set of structural elements separating the intestinal chyme from the blood. Hypothetically, severe damage to the IB caused by exogenous toxicants can trigger a leakage and subsequent systemic redistribution of toxic substances of bacterial origin. Until recently, the impact of such a redistribution on the outcome of acute exogenous poisoning remained outside the view of toxicology. The present review addresses causal relationships between the secondary dysfunction of the IB and complications of acute poisoning. We characterize acute systemic toxicity of such waste products of the normal gut microflora as ammonia and endotoxins, and demonstrate their involvement in the formation of such complications of acute poisoning as shock, sepsis, cerebral insufficiency and secondary lung injuries. The principles of assessing the functional state of the IB and the approaches to its protection in acute poisoning are briefly considered.

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Basic terms and abbreviations: Bacterial exotoxins—toxic proteins and peptides released by living bacteria into the external environment; Brain—blood barrier (BBB)—a set of structural elements
separating the central nervous system (CNS) and blood; Endogenous substance—a compound produced by the cells of the host or its symbiotic microflora; Endotoxemia—accumulation of toxic endogenous substances in blood, including toxic low molecular weight metabolites of bacterial and/or tissue origin, endotoxin, and bacterial exotoxins; Endotoxicosis—a set of clinical manifestations of endotoxaemia; Endotoxin—the sum of outer cell wall lipopolysaccharides of gram-negative bacteria; Endotoxicemia—endotoxin accumulation in blood; Intestinal barrier (IB)—a set of structural elements separating the intestinal chyme and blood; Low molecular weight bacterial metabolites—substances of bacterial origin with a molecular weight of less than 1500 Da; Primary intestinal barrier dysfunction—preformed (inborn or acquired) chronic intestinal barrier disorder; Secondary intestinal barrier dysfunction—damage to the intestinal barrier caused by external exposure; Toxicant—any substance that causes damage to the organism in a non-mechanical way when exposed at a given dose; Xenobiotic—a foreign substance

INTRODUCTION

The legal analysis of the causes of a patient’s death implies a choice out of four possibilities: untimely, incorrect, inadequate medical care, or lack of its connection with the outcome of the disease [1]. One of the sources of error in this choice is the unpredictability of complications of acute diseases, so vividly manifested during the recent COVID-19 pandemic [2]. Unpredictability is equally inherent to complications of acute poisoning: acute circulatory disorders, sepsis, secondary acute lung injuries and acute cerebral insufficiency develop not in all patients. The reasons for such an individual variability require further detailization.

As we have shown previously, the probability of fatal complications in acute poisoning depends on the functional state of the vascular endothelium [3]. Some of its damaging substances are produced by the normal gut microbiota [4]. Based on the concept that the human body is an ecosystem with the gut microbiota being its integral constituent [5, 6], these substances are considered hereinafter as endogenous. In healthy people, the ability of such substances to cross the enterohematic or intestinal barrier (IB), a system of structural elements separating the intestinal chyme from blood, is insignificant, and their latent leakage leads to a number of chronic diseases [4, 6]. In mild exogenous poisoning, the effects of endogenous substances of intestinal origin can be overshadowed by the direct toxic effect of the xenobiotic. However, their quick and massive entry into the bloodstream in severe exogenous poisoning can aggravate the state of the organism and be involved in the pathogenesis of complications.

Direct or indirect damage to the IB by a xenobiotic, hereinafter referred to as its secondary dysfunction, can provoke the entry of metabolites and cellular bacterial components into the bloodstream in doses affecting the outcome of acute exogenous poisoning. The composition of the mixture of these substances a priori depends on that of the gut microbiota, which is individually variable [5] and, according to some estimates [7], even unique. This variability may underlie the stochasticity of complications of acute poisoning.

There is a large body of data characterizing the effect of xenobiotics on the IB. There are also numerous data on the involvement of toxicants of intestinal origin in the pathogenesis of critical states of the organism. This allows tracking for the first time the relationship between secondary dysfunction of the IB and some complications of acute poisoning. Such a relationship is the subject of the present review.

The aim of the review was to reveal the ways to prevent complications of acute poisoning through the identification of the role of secondary dysfunction of the IB in their pathogenesis.

SOURCE SELECTION ALGORITHM

The search for data characterizing the relationship between complications of acute poisoning and secondary IB dysfunction revealed an obvious novelty of this subject matter. Therefore, this review only provides basic information, formulates the concept, but is in no way systematic. It mainly cites the publications of fundamental importance that came out over the last 10 years or earlier. The main source of information was the PubMed database. The following queries were
used: “acute poisoning AND (gut microbiota OR intestinal microflora OR gut microflora OR metabolome OR human gut microbiome OR gut barrier OR leaky gut syndrome OR intestinal barrier OR gut–brain axis OR gut–liver axis)”. No conference abstracts were used for analysis.

STRUCTURE AND FUNCTION OF THE NORMAL INTESTINAL BARRIER

The IB comprises of mucin, the intestinal mucosal epithelium [8], symbiotic epithelium-associated microorganisms [9], and the endothelium of submucosal blood and lymphatic capillaries [10]. Substances that escaped absorption by blood and lymphatic capillary plexuses in the intestinal wall have to overcome additional structural elements of the IB on their way to blood: the layer of smooth muscle cells, the mesothelium of visceral and parietal peritoneal sheets with the fluid-filled peritoneal cavity in between, and the endothelium of blood and lymphatic capillaries of the parietal peritoneum.

The mucin layer, 150 μm thick, consists of hydrated glycoproteins and separates epithelial cells from the aggressive environment of the intestinal luminal space. Mucin production is impaired in mucosal ischemia, which is most dangerous for the colon with its high density of bacterial colonization. Normally, symbiotic aerobic bacteria oust the pathogenic microflora from the apical surface of entero- and colonocytes, while providing them with essential substances [5]. The main function of the 20-μm epithelial cell monolayer is the selective absorption of substances from the chyme.

Due to the presence of small-intestinal villi and large-intestinal crypts, the absorptive surface area of the gastrointestinal mucosa reaches 200 m² [8]. Normally, substances are mainly absorbed from the intestinal chyme transepithelially. The proportion of paracellular transport is presumably proportional to that of intercellular contacts on the luminal surface of the intestine, which is estimated at 0.1% [11]. Intercellular contacts are of two types, tight and adherens junctions, and consist of actin; they provide mechanical strength to the epithelium by connecting the plasma membranes of neighboring cells to the intracellular cytoskeleton [8]. Intercellular contacts serve as an object of damaging effect of a number of xenobiotics [12], against which the proportion of paracellular transport increases.

A part of the chyme substances that overcome the intestinal epithelium, enter the submucosal network of blood and lymphatic capillaries and then proceed further to the basins of the portal vein and thoracic ducts, which is the main pathway under normal conditions. The other part, having passed through the visceral and parietal peritoneal sheets, as well as a 0.25-mm fluid-filled peritoneal cavity between them, enters lymphatic vessels of the thoracic duct basin or the blood capillary network of the inferior vena cava basin. This process, transperitoneal diffusion, is facilitated by the absence of the large-intestinal diffusion barrier in the form of a continuous longitudinal muscular layer [13]. Transperitoneal diffusion of ammonia [14] and endotoxin has been shown experimentally [15]. The substances involved in transperitoneal diffusion avoid presystemic metabolism in the liver on their way to the general blood flow. In portal hypertension, the role of transperitoneal diffusion may increase due to the delayed delivery of substances of intestinal origin to the portal vein basin [16]. Transperitoneal diffusion serves as a target for detoxification therapy with a peritoneal dialysis [17].

Normally, the IB is impermeable to intestinal bacteria: DNA of *Escherichia coli* and *Bacteroides* was detected in blood plasma only against the IB injury [18]. Substances to be removed from the organism penetrate from the chyme into blood via passive diffusion [19]. Their absorption is approximated [20] by the equation of the Fick’s first law of diffusion for membranes: \( J = D \cdot (C_i - C_o) \), where \( J \) is a diffusion flux density (mol \( \times \) m\(^{-2} \) \( \times \) s\(^{-1} \)), \( D \)—membrane permeability coefficient (m \( \times \) s\(^{-1} \)), \( C_i \) and \( C_o \)—concentrations of a substance on the epithelial and endothelial sides of the membrane, respectively (mol \( \times \) m\(^{-3} \)). Bioavailability of toxic products of the gut microbiota increases with values of the multipliers \( (C_i - C_o) \) and \( D \). The former is limited by the content of these substances in the chyme, the latter by the state of the IB. Presumably, absorption increases with increasing hydraulic pressure of the chyme due to smooth muscle spasm and (or) the intensification of gas produc-
The routes of diffusion of substances from the intestinal chyme into lymph and blood are depicted schematically in Fig. 1.

**NORMAL GUT MICROBIOTA**

Prokaryotes of more than four hundred species inhabit the gastrointestinal tract (GIT) of a healthy human. The total number of their cells exceeds the number of cells of the host organism, while their mass is estimated at 0.3 % of the body weight [22]. The density of bacterial colonization in the lower (distal) GIT is higher than in its upper (proximal) division (Table 1).

Samples of gastric contents taken in fasting people are practically sterile. In hypochlorhydria, there are more bacteria in the gastric lumen, and they are mainly represented by obligate anaerobes, streptococci, lactobacilli, neisseria and staphylococci. At pH > 5.0, the microbial composition of gastric contents is indistinguishable from that of the small intestine. In 2/3 people aged 51–60 years, the ammonia-producing Gram-negative bacterium *Helicobacter pylori* is detected in the mucin layer of the gastric mucosa. In *H. pylori*-infected individuals, the pH of gastric contents increases, which adversely changes the composition of distal gut microbiota [23].

In the duodenum and jejunum, bacterial vegetation is counterbalanced by their rapid removal due to secretion, motility and bactericidal effect of bile, with Gram-positive cocci (*Streptococci, Peptococci*) and bacilli (*Lactobacilli, Bifidobacteria*) being predominant. The microbiota of the

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**Table 1. The content of bacteria in the chyme of the human gastrointestinal tract (GIT)**

| GIT division       | Concentration, mL⁻¹ | Volume, mL | Number of bacteria |
|--------------------|---------------------|------------|--------------------|
| Stomach            | ≤ 10³* [5]           | 250–900**  | ≤ 10⁷*             |
| Duodenum and jejunum| 10³–10⁴**           | 400**      | ≤ 10⁷              |
| Ileum              | 10⁸**               | 400**      | ≤ 10¹¹             |
| Large intestine    | 10¹¹**              | 400**      | ≤ 10¹⁴             |

*—At pH < 3.0; **—[22].
terminal ileum is similar to that of the cecum due to reflux from the latter [5]. In the colonic chyme, bacteria account for an average of 27% of its dry weight [22]. Anaerobic bacteria (predominated by *Escherichia coli, Bacteroides fragilis, Lactobacilli and Bifidobacteria*) in the colon are 1000 times as numerous as aerobic bacteria [5]. Anaerobes are represented by Gram-positive (*Bifidobacteria, Eubacteria, Propionibacterium*) and Gram-negative (*Fusobacterium, Enterobacteriaceae*) bacteria. The colonic mucosa is inhabited not only by symbiotic bacteria, but also by conditionally pathogenic urease-expressing bacteria from the family *Enterobacteriaceae* [24].

The composition of the normal gut microbiota depends on diet [25], age, antibacterial drug intake, and a number of uncontrollable conditions [5]. The production of toxic substances in the intestine is promoted by the predominance of *Proteobacteria* and *Fusobacteria* over *Bacteroidetes* while suppressing *Lactobacilli* and *Bifidobacteria* [26]. The number of microorganisms in the GIT depends on its motility. The normal gastrointestinal transit time is 10–48 min for the stomach, 2.5–4.0 h for the small intestine [27], and 25–40 h for the large intestine [22]. The duration of chyme passage through the large intestine determines the greatest contribution of its microbiota to the production of toxic substances [5].

**TOXIC PRODUCTS OF NORMAL GUT MICROBIOTA**

The normal intestinal microbiota produces substances both essential and toxic for the host [28]. The validity of the hypothesis formulated by I.I. Mechnikov more than a century ago on the ability of substances produced by the gut microbiota from nutrients to cause systemic pathological processes under certain conditions [29] is now fully proven. From 2011 to 2021, the annual number of publications available under the keywords “gut barrier”, “gut–brain axis”, “gut–liver axis”, “gut microbiota”, “intestinal microflora”, “gut microbiota AND metabolome”, “intestinal barrier”, “leaky gut syndrome” and “human gut microbiome” has grown exponentially. On the PubMed.gov Web site, it increased by 7, 16, 17, 18, 23, 23, 30, 34, 37, 40, and 44 times, respectively, during this time, whereas in previous years, the growth was linear.

Toxic substances of bacterial origin with systemic toxicity include bacterial exotoxins, endotoxin, and products of bacterial transformation of food proteins, amino acids, amino alcohols and phospholipids (ammonia, amines, phenols, heterocycles). Some amines, heterocycles and phenols are converted in the liver into secondary toxicants (trimethylamine-N-oxide, indoxyl sulfate, p-cresyl sulfate) involved in the pathogenesis of cardiovascular and other chronic diseases. Ammonia and endotoxin, asymptptomatically circulating in the blood of a healthy individual in trace amounts, can exhibit acute systemic toxicity at higher concentrations, as discussed below.

**Ammonia.** No less than 2/3 of the ammonia produced in the organism is of intestinal origin. In enterocytes, the main mechanism of its formation is a glutaminase reaction, while in the colon, it is derived due to the metabolic activity of bacteria: deamination of amino acids and nitrogenous bases, as well as hydrolysis of the urea diffusing from blood to the luminal surface of the mucosa [30]. The ureolytic activity of microorganisms associated with the colonic mucosa accounts for the formation of half of the ammonia of intestinal origin [24].

From the intestinal chyme, ammonia enters the general blood flow through both the portal vein and the liver, as well as via transperitoneal diffusion [14]. In a healthy individual, about 4 g of ammonia is delivered into blood (mainly portal) from the intestine daily [31]. Its level in the blood of hepatic veins depends positively on that in the portal blood, while the latter on the ammonia content in the chyme [32]. Due to presystemic metabolism of ammonia, its content in hepatic venous blood is 2–3 times lower than in portal blood [33] and an order of magnitude lower than in mesenteric venous blood of the large intestine [5].

In arterial blood plasma, the normal concentration of ammonia ([NH₃] + [NH₄⁺]) is 30 μM [34]. Its elevation is associated with neurotoxicity, first described in the laboratory of I.P. Pavlov [35]. At ([NH₃] + [NH₄⁺]) values in blood plasma up to 100 μM, hyperammonemia is asymptomatic; at 100–200 μM, vomiting, ataxia,
Irritability and hyperactivity are noted, while at values more than 200 μM, there ensue convulsions and coma [36]. At an average colonic chyme volume of 0.4 L [22] and an ammonia content of 5.7–39.0 mmol/L [37], the colonic ammonia pool is 2.3–15.6 mmol/L. If this amount were evenly distributed in 5 L of blood, 2 L of lymph, and 0.4 L of chyme at the same time instant, the plasma ammonia concentration would reach 311–2108 μM, i.e., the values far above the threshold of coma.

Ammonia enters cells in an unionized form, as NH₃. Considering the ammonia basicity constant (pKa = 9.15 at t = 37°C), the proportion of ammonia present as NH₃ in blood plasma with a pH of 7.36 is 1.6%. The pH of the cytoplasm is lower, making NH₃ to diffuse into cells even when the total blood level of both forms of ammonia is normal [30]. In metabolic acidosis or gaseous alkalosis, an increased difference between pH values in blood plasma and the cytoplasm intensifies ammonia influx into cells along the NH₃ concentration gradient [38]. This explains the absorption of ammonia from blood by the brain, the accumulation of osmotically active glutamine therein, and brain edema with increasing ammonia levels in brain tissue [39]. In this case, neurotoxic effects of ammonia are possible even in the absence of hyperammonemia.

In addition to neurotoxicity, ammonia has endothelial toxicity, as demonstrated both in vitro and in vivo. In cultured brain capillary endothelial cells, it induces oxidative stress, the accumulation of nitrogen oxide (NO) and arachidonic acid peroxidation products [40, 41], destroys extracellular matrix, increases plasma membrane permeability to a fluorescein-isothiocyanate dextran derivative, causes the expression of the transmembrane carrier of arginine, the NO synthase substrate [40]. Endothelial cell growth medium treated with ammonium salt causes swelling of cultured astrocytes, which shows the involvement of the vascular endothelium in the development of brain edema during hyperammonemia [42]. In phagocytes, ammonia activates Toll-like type receptors 4, which enhances the inflammatory response to endotoxin and formation of free oxygen and nitrogen radicals, causes swelling and increased permeability of the cerebrovascular endothelium [42].

Hyperammonemia causes endothelium-dependent NO-mediated dilation of cerebral arterioles, which is involved in the development of brain edema and increased intracranial pressure in acute hepatic failure [38]. Intravenous administration of ammonium acetate to rabbits increased the permeability of their blood–brain barrier (BBB) to polyethylene glycol-400 [43]. An increased BBB permeability was also noted in patients with chronic hepatic encephalopathy [44]. Thus, the intensification of the ammonia entry into the brain induces oxidative stress, inflammatory injury, increased BBB permeability, brain edema, cerebral blood flow dysregulation, and impaired neurological functions.

Endotoxin. The content of lipopolysaccharides of Gram-negative bacteria, cumulatively known as “endotoxin”, in the colonic chyme of an adult individual is close to 2.5 g/L [45]. In blood plasma it is nine orders of magnitude lower, 9 ng/L, which is the threshold of inflammatory activation of macrophages and endotheliocytes. Moderate endotoxemia is observed in periodontitis, whereas in diabetes mellitus, liver cirrhosis, Alzheimer’s disease, and sepsis, plasma endotoxin levels can reach 500 ng/L [46]. In blood, endotoxin is present in the active free form or as complexes with proteins, usually undetectable in laboratory tests [17].

Endotoxin has a pronounced endotoxicity, which is manifested by oxidative stress, glycocalyx destruction, leukocyte adhesion, vasospasm, thrombosis [47] and increased BBB permeability [48]. Endotoxin content in blood, typical for sepsis, entails a “cytokine storm” and septic shock [49]. Lipopolysaccharides of Escherichia coli have a more pronounced proinflammatory effect compared to those of Bacteroides dorei and Bacteroides vulgatus [50]. Endotoxemia leads to thromboxane-dependent pulmonary vasoconstriction and pulmonary hypertension [51] combined with systemic vasodilation [52, 53]. The effect of endotoxin on endothelial permeability manifests itself in an increased permeability of the BBB both for endotoxin itself and for other substances [4, 54], noncardiogenic pulmonary edema [55], liver [56, 57] and renal [58] dysfunction. In endotoxemia, the ability of the liver to incorporate ammonia into urea is impaired [59]. The systemic toxicity
profiles of ammonia and endotoxin are presented in Table 2.

Table 2. Publications documenting the systemic toxicity of ammonia and endotoxin at concentrations exceeding the physiological norm

| Systemic toxicity manifestations       | Ammonia          | Endotoxin          |
|--------------------------------------|------------------|--------------------|
| Systemic inflammation                | [42]             | [47, 49, 52]       |
| Circulatory disorders                | [38]             | [47, 52]           |
| Coagulopathy                         | –                | [47]               |
| Pulmonary gas exchange disorders     | –                | [47, 51, 55]       |
| Renal failure                        | –                | [58]               |
| Hepatic failure                      | –                | [56, 57]           |
| Acute cerebral insufficiency         | [38, 42–44, 60]  | [4, 54]            |

Table 3. Some markers of intestinal barrier permeability

| Substance      | Molecular weight, Da | Molecule, diameter, nm | log P**          | Number of hydrogen bonds | References*** |
|----------------|----------------------|------------------------|------------------|--------------------------|---------------|
|                |                      |                        |                  | donor | acceptor |                   |
| Paracellular type of permeability prevails |                      |                        |                  |                  |               |
| PEG 400*       | 380–420              | 0.53                   | −1.43 (0.61)     | 2   | 8        | [61]           |
| PEG 4000*      | 3800–4200            | 1.00                   | −0.93 (0.98)     | 2   | 80       | [61]           |
| Erythritol     | 122.1                | 0.86                   | −3.16 (0.35)     | 4   | 0        | [62]           |
| Lactulose      | 322.3                | 1.33                   | −3.92 (0.50)     | 8   | 3        | [62–64]        |
| Sucrose        | 342.3                | 1.31                   | −3.65 (0.43)     | 8   | 3        | [62]           |
| EDTA (pH 7)    | 292.2                | 1.45                   | −4.43 (1.30)     | 4   | 6        | [62]           |
| Dextran        | 3000–20 000          | 5.5–10.0               | −                  | 3n+2| 2n+1     | [64]           |
| Transcellular type of permeability prevails |                      |                        |                  |                  |               |
| Mannitol       | 397.6                | 0.89                   | −4.96 (0.38)     | 6   | 0        | [61, 63, 65]   |
| Sucralose      | 397.6                | 1.26                   | 0.88 (0.65)      | 5   | 5        | [64]           |

*—At the molecular weight specified in column 2; **—log P_{octanol/water} (determination error); ***—literature source documenting the use of the substance for IB permeability evaluation.

EVALUATION OF INTESTINAL BARRIER PERMEABILITY

For an integral evaluation of the IB permeability, the levels of substances injected into the stomach are determined in blood or urine. Ideal permeability markers are biochemically inert substances that overcome the IB via passive diffusion. Of the substances determined in urine, preference is given to those that are not reabsorbed in the renal tubules. In the case of increased excretion of these substances with urine, increased IP permeability is postulated for other, similar in size, molecules [11]. Molecular weight, hydrophilic-lipophilic (log P_{octanol/water}), and the number of hydrogen bonds formed by the molecule are a priori indicators of the predominance of paracellular or transcellular transport of the test substance. The latter route is promoted by moderate hydrophilicity (log P_{octanol/water} is within the range from 0 to 5) and the ability of the molecule to form no more than five donor or acceptor hydrogen bonds. The molecular weight and effective
molecular diameter of the substance are used to evaluate the pore size in the IB (Table 3).

The evaluation of IB permeability using these markers proceeds from the assumption of their preferential absorption in a certain segment of the GIT. Sucrose is used to assess the permeability of the gastric wall, and sucralose is used to assess that of the colonic wall. However, it is possible that the site of absorption changes under pathological conditions or the influence of therapy. For example, lactulose and mannitol are absorbed mainly in the small intestine, because in the colon they are quickly broken down by microorganisms [65]. However, against the suppression of the latter by antibiotics, these markers can accumulate in the chyme of the large intestine and be absorbed therein.

For the selective evaluation of mucosal epithelial permeability, isolated loops of the intestinal segment are consecutively filled with a marker solution, ligated and incubated in a buffer solution. The marker accumulation rate is considered to be an indicator of epithelial permeability, since circulation and hence the vascular endothelium, do not function under these conditions [61]. The barrier function of GIT vascular endothelium can be selectively evaluated using dyes (Evans blue, fluorescein) or compounds labeled with radioactive isotopes [66]. Intravital microscopy [67] and ex vivo microscopy of GIT tissues provide a valuable supplement to a direct investigation of the transport of substances across the IB [68].

### INTESTINAL BARRIER DAMAGE IN ACUTE POISONING

**Direct damage to the IB by xenobiotics.** Many xenobiotics and their metabolites damage the IB directly. Local action of irritants and cauterizing substances triggers a cascade of pathological processes, including protein denaturation, tissue necrosis, release of inflammatory mediators, and blood plasma transudation [69]. Ethanol [70], arsenic, salts of heavy metals [69, 71], opioids [72], some mycotoxins [73] exhibit direct entero-toxicity. Its intensity in non-steroidal anti-inflammatory drugs made them a means for experimental modeling of acute gastroenteritis [74]. Dextran sulfate and 2,4,6-trinitrobenzene sulfonate are used to experimentally simulate acute colitis [75]. Some mycotoxins cause inflammatory damage to the IB by enhancing the effect of endotoxin thereupon [76]. T-2 mycotoxins and deoxynivalenol increase IB permeability to polyethylene glycol-4000 [77].

Direct enterotoxicity is possible not only by ingestion, but also through other routes of entry of toxicants into the organism. The small intestinal and, to a lesser extent, gastric and colonic epithelium is a typical cell renewal system with a high proliferative activity. This makes it sensitive to substances that cause reproductive or interphase death of enterocytes. Differences in enterotoxic effects of such substances are determined only by the level of target cell differentiation. Adriamycin induces apoptosis of enterocyte at position 4–5 from the stem cell located at the base of the inter villar crypt, isopropyl methanesulfonate, nitrogen mustard and actinomycin D at position 6–7; fluorouracil, milener, cyclophosphamide and cycloheximide at position 7–9, vincristine and hydroxyurea at position 10–11 [68]. At a sufficient dose, all these toxicants cause denudation of the small intestinal epithelium. In rats with acute cyclophosphamide intoxication, IB permeability to methylene blue, mannitol and lactulose increases [63]. Enterotoxicity of cytostatic drugs in hematopoietic stem cell transplantation is one of the factors that limit patient survival [78].

An increased IB permeability against the background of its chemical damage has been demonstrated experimentally. In rats administered ammonium acetate through the gastric probe, intraperitoneal injection of cyclophosphamide accelerated ammonia and glutamine accumulation in blood and the brain [79] and provoked acute neurological disorders, such as opisthotonus and apnea, characteristic of the effect of ammonium salts at much higher doses [80, 81]. At the same time, isolated administration of the ammonium salt did not affect animal behavior significantly (Fig. 2).

In the above examples, direct damage to the IB by cyclophosphamide metabolites (aldocyclophosphamide, acrolein, phosphoramid mustard, etc.) promoted the entry of ammonia from the chyme into blood at a rate at which hyperammonemia exceeded the seizure threshold.
IB damage in acute systemic hypoxia. In critical states of the organism, acute intestinal hypoxia can be caused by a disruption of external respiration and circulation. In acute severe poisoning, such states include exotoxic shock, while in the absence of respiratory support—respiratory depression, neuromuscular blockade, and bronchoobstructive syndrome. Due to “centralization” of blood circulation in such patients, the GIT experiences deeper hypoxia than the “vital organs”.

Aerobic type of energy metabolism of enterocytes predisposes them to the breakdown of ATP resynthesis in acute hypoxia [82]. This is evidenced by intestinal damage during high-altitude hypoxia, as manifested in inflammation, ulceration and bleeding that exacerbated altitude sickness [83]. Normobaric hypoxia provoked an increase in IB permeability in rats during treadmill running [84]. IB permeability increases in hemolytic anemia [85] and acute blood loss [86]. Uncoupling of oxidative phosphorylation in coloocytes impairs their resistance to *Escherichia coli* [87]. Mice with more active oxidative phosphorylation in the colonic mucosa are more resistant to the local toxic effects of dextran sulfate or trinitrobenzene sulfonate [88]. Hypoxia reduces mucin production in the intestine, which entails inflammatory alterations in the mucosa. The submucosal capillary plexus in the small intestine is more developed than in the large intestine, which accounts for a higher sensitivity of the latter to ischemia [89].

IB damage due to smooth muscle spasms. Gastrointestinal smooth muscle spasm is a characteristic symptom of poisoning with cholinesterase inhibi-
tors and serotonergic drugs, but can also be a reaction to circulatory “centralization”, or a result of the effect of serotonin produced by Enterobacteria on smooth muscles [90]. Serotonergic stimulation of the colon is accompanied, in addition to spasms of its own smooth muscles, by arterial spasm and local circulatory hypoxia [91]. Microcirculatory disorders in the intestine can damage it in the same way as it occurs under systemic hypoxia. The hypothesis of the “muscle spasm–ischemia–pain” triad as a factor of increased IB permeability in irritable bowel syndrome has been put forward [90]. In acute poisoning, the IB reaction to such changes may be identical.

**IB damage in gastrointestinal stasis.** Gastrointestinal stasis is a potentially fatal complication in intensive care unit (ICU) patients [92]. It is one of the specific manifestations of opioid (cholinergic antagonist and serotonin agonist) toxicity [11]. It has been observed in a rat barbiturate coma model [93], as well as in acute poisoning with alkylating toxicants. Figure 3 shows the position of radiological barium sulfate shadows in rats that received barium sulfate suspension through the gastric probe immediately after intraperitoneal (i.p.), subcutaneous (s.c.) or intragastric (gavage) administration of cyclophosphamide at a dose of 1000 mg/kg [94]. A sharp suppression of propulsive activity is quite evident in all GIT divisions, being most pronounced after intragastric infusion of a toxicant.

In gastrointestinal stasis, bacterial vegetation is not compensated by their removal, due to which the type of host—normal gut microbiota relationships becomes no longer mutualistic and approaches parasitism. Not only the total luminal concentration of ammonia produced by bacteria ([NH$_3$] + [NH$_4^+$]) but also the pH of the chyme, and hence the proportion of ammonia repre-
presented in a highly penetrant NH₃ form, increase [37]. The cytotoxic effect of NH₃ reduces the life cycle of colonocytes [95] and mucin production [96], which can damage the IB and increase the bioavailability of substances produced by colonic microflora.

Considerable gas production, characteristic of gastrointestinal stasis, can intensify the diffusion of substances through the IB. It increases the barometric pressure in the intestinal lumen and, according to Dalton’s first law, the partial vapor pressure of volatile substances. According to Dalton’s second law, it increases the tension of the corresponding gases dissolved in the chyme. Damage to the IB, as well as an increased tension of volatile substances in the chyme, are potential factors that increase their bioavailability in gastrointestinal stasis.

**Inflammatory damage to the IB.** Damage to the EGB by an exogenous toxicant promotes endotoxin entry into blood. It stimulates the secretion of proinflammatory cytokines (TNF-α, IFN-γ and IL-6) by phagocytes, leading to both systemic and local inflammation [97]. Cyclophosphamide administration to pygmy pigs at a dose of 50 mg/kg increased TNF-α and IL-6 serum levels and IB permeability [98]. Intestinal mucositis leads to bacteremia in hematopoietic stem cell recipients during cytostatic therapy [99]. Submucosal inflammation and leukocytoclastic vasculitis accompanied by mucosal ulceration have been reported in rats administered with mercuric chloride [100]. Acute inflammation of the colon provokes inflammation in other organs as well [101], which may promote the entry of toxic substances of intestinal origin from blood.

Thus, in acute severe poisonings, there are prerequisites for an increase in the IB permeability for metabolites and cellular components of the intestinal microflora. The mechanisms of secondary IB dysfunction in acute poisoning include direct damaging effects of xenobiotics or their metabolites, gastrointestinal stasis, smooth muscle spasm, gastrointestinal inflammation and hypoxia. Along with secondary IB dysfunction, the delivery of toxic substances from the intestinal chyme into the general blood flow in acute poisoning can be promoted by the excessive growth of the intestinal microbiota and increased gas production. All these synergistic factors lead to endotoxemia.

**ENDOTOXEMIA IN ACUTE POISONING**

Endotoxemia is an elevated blood level of biologically active substances produced in the organism. Proceeding from the ideas of the organism as an ecosystem with the intestinal microbiota as its constituent [5, 6], the compounds produced by the latter are classified as endogenous substances.

The blood level of medium molecular weight (MMW) substances is used as an integral index of endotoxemia [102]. In patients with uncomplicated poisoned with psychopharmacological drugs, blood levels of such substances were by 32% higher than in healthy donors, and those of endotoxin—10 times higher [103]. After intestinal lavage, the blood level of MMW substances decreased by 18 %, endotoxin—by half, while without lavage, these indices of endotoxemia continued to grow. Thus, in poisoning with psychopharmacological drugs, the colonic chyme is a source of endotoxemia.

The issue of which specific substances are involved in endotoxemia formation under acute poisoning is of interest. One of them is endotoxin, whose increased blood level has been shown in severe acute poisoning with colchicine [97], verapamil [104], and ethanol [105]. Manifest endotoxemia accompanies patients’ preparation for hematopoietic stem cell transplantation, during which cytostatic drugs are administered at potentially lethal doses. Endotoxin accumulation in blood in such a treatment leads to a systemic inflammatory response [106, 107].

In acute exogenous poisonings, endotoxemia is paralleled with hyperammonemia [108, 109]. Listed below are xenobiotics, in acute poisonings with which it has been documented (to exclude liver injuries from the causes of hyperammonemia, specific hepatotoxic agents are not omitted):

- acetazolamide [110];
- calcium hopantenate [111];
- clozapine [112];
- cocaine [113];
- cyclophospham [79];
- cyclopropane [114];
– cyanides [115];
– ethanol [116];
– diethyl ether [117];
– fluoroacetate [118];
– 5-fluorouracil [119];
– homocysteine thiolactone [120];
– lead salts [121];
– methanol [122];
– oxygen [123];
– sodium thiopental [93];
– trimethoprim [124];
– valproates [125].

Thus, endotoxemia in acute poisoning involves endotoxin and ammonia, while changes in blood levels of other components of the intestinal metabolome require further investigation. The role of endotoxin and ammonia in the pathogenesis of acute complications is characterized by the information provided below on endotoxemia in the corresponding syndromes of various etiologies.

ENDOTOXEMIA IN CRITICAL STATES OF AN ORGANISM

Critical states of an organism are endotoxemia patterns, a constellation of clinical manifestations of endotoxemia. The role of endotoxin and ammonia in the pathogenesis of various critical states is discussed below, depending on whether the symptoms of these states are reproduced by endotoxin or ammonia administration.

Shock is an acute state of inadequate tissue blood supply, with hyperammonemia being one of its biochemical markers. Plasma and brain tissue ammonia levels are manifold elevated in rabbits and dogs when modeling heat or insulin shock [126]. In patients with septic shock, venous blood ammonia levels were 80% higher, while in those who died eventually of septic shock, 2.2 times higher compared to patients with sepsis but without shock [127]. In patients with hemorrhagic shock, ammonia levels in arterial blood plasma were twice as high as in patients with hemorrhage but without shock [128]. In severe trauma, venous blood ammonia levels were twice as high as in controls, while in deceased patients, it was by 70% higher than in survivors [129]. Plasma ammonia levels in patients admitted to the ICU because of cardiac arrest was 4.8 times higher than that of patients with spontaneous circulation [130]. Hyperammonemia not only accompanies shock but is also involved in its pathogenesis. The administration of ammonium salts to rabbits reproduced key manifestations of heat and insulin shock [126].

In experiments, shock is also reproduced by endotoxin administration [131]. Endotoxin accumulation in blood is characteristic of septic [132], traumatic and hemorrhagic [133] shock.

IB damage is one of the mechanisms of endotoxemia in shock. When modeling hemorrhagic shock in rats, the animals were losing claudin-3, a protein of intestinal epithelium tight junctions, with urine [134]. Using MMW substances as endotoxemia markers, it has been shown that, in hemorrhagic shock, they accumulate first in intestinal tissues, then in portal blood, and finally in carotid artery blood [135].

Thus, hyperammonemia and endotoxinemia are characteristic of endotoxemia that accompanies shock and is involved in the development of its different types.

Sepsis is a systemic inflammatory response to endotoxemia and bacteremia, which most frequently complicate acute poisoning in alcohol addicts [105]. Sepsis developed in 6.4% of the methanol poisoning victims [136]. Endotoxemia constantly accompanies sepsis [137], while bacteremia has only been observed in 67% of such patients [138]. Endotoxin accumulation in blood occurs in sepsis caused by Gram-negative, as well as Gram-positive bacteria and fungi, due to IB damage [139].

Clinical manifestations of sepsis are reproduced by endotoxin administration to animals [140]. In endotoxemia characteristic of sepsis, the blood coagulation cascade is activated, resulting in disseminated microvascular thrombosis [141], which leads to kidney [10], heart and liver [142] damages, multiple organ failure [132], and pulmonary edema [143].

Ammonia is also involved in the pathogenesis of multiple organ failure in sepsis. Its plasma level in patients with this diagnosis, measured on admission to the emergency department, correlated positively with the likelihood of developing multiple organ failure within the following
Secondary acute lung injuries in acute poisoning include non-cardiogenic pulmonary edema, shock and wet lung syndromes. The involvement of endotoxin in the pathogenesis of these conditions follows from endotoxemia that accompanies all of them [144] and endotoxin-induced damage to alveolar–capillary (air–blood) barrier cells both in vitro [145] and in vivo [146]. Endotoxin administration to rats caused acute lung injury resembling such in COVID-19 and manifesting itself as an alveolar cavity hemorrhage, cytokine storm, coagulopathy, and pulmonary hypertension [147]. *Escherichia coli* endotoxin caused hemodynamic disorders in the pig lungs: its administration into the right atrial cavity increased blood pressure in the pulmonary artery [53].

In portocaval anastomosis, hyperammonemia is the predictor of pulmonary hypertension [148]. In patients with severe acute pancreatitis, the development of a lethal complication, pulmonary edema, is caused by excessive bacterial colonization of the small intestine [149].

Thus, endotoxemia accompanies secondary acute lung injuries under acute poisoning and is involved in the development of pulmonary hypertension. Ammonia is another marker of endotoxemia, promising for the prediction of secondary acute lung injuries in acute poisoning.

Acute cerebral insufficiency is a syndrome resulting from diffuse brain damage. It is observed in severe poisoning with substances having different mechanisms of action, and manifests itself in impaired consciousness and mental confusion, motor disorders, accelerated catabolism, acute respiratory and circulatory disorders [150].

Some manifestations of acute cerebral insufficiency may be associated with increased ammonia and/or endotoxin delivery to the brain. Hyperammonemia is characteristic of acute liver failure complicated by brain edema and increased intracranial pressure [151]. Intracranial pressure, neurological disorders, and mortality are positively related to serum ammonia levels in patients with extrahepatic hyperammonemia [45]. Plasma and brain tissue ammonia levels were significantly elevated in dog and rat models of diabetic coma [126]. Endotoxin increased the sensitivity of piglets to acute cerebral hypoxia. *Escherichia coli* endotoxin administration to piglets increased neuronal loss and metabolic disturbances, as well as increased the probability of brain death followed by carotid artery occlusion [152]. Endotoxin administration to ferrets exacerbated brain damage due to carotid artery occlusion [153]. Endotoxin potentiated brain edema during altitude hypoxia [154].

Thus, the available data indicate that endotoxin and ammonia produced by the normal gut microbiota are involved in the pathogenesis of abnormal conditions that can complicate acute poisoning, such as shock, sepsis, secondary acute lung injuries, and acute cerebral failure.

SECONDARY DYSFUNCTION OF THE INTESTINAL BARRIER AND STOCHASTICITY OF COMPLICATIONS OF ACUTE POISONINGS

The phenomenon of an altered tolerance of acute damaging exposures against the background of preformed chronic pathology has long been used in functional diagnostics of the latter. This phenomenon also occurs in acute poisoning. Xenobiotic-induced damage to the IB may be superimposed on chronic pathological conditions, in which it is damaged, such as intestinal dysbiosis, irritable bowel syndrome, Crohn’s disease, nonspecific ulcerative colitis, etc. This can
exacerbate endotoxemia in acute exogenous poisoning. Such an overlap, shown schematically in Fig. 4, may contribute to the individual variability in the severity of their complications.

**INTESTINAL BARRIER PROTECTION IN ACUTE EXOGENOUS POISONINGS**

The intestinal chyme is an aggressive environment for the damaged intestinal mucosa, and its local damaging effect increases with intestinal dysbiosis [95]. Therefore, the planned measures on IB protection should be aimed at optimizing intestinal microbiota composition and increasing the local resistance of the mucous membrane to toxicants of bacterial origin. The main approaches to optimizing intestinal microbiota composition are prebiotic, metabiotic, probiotic and autoprobiotic therapies.

Prebiotics are nutrients that selectively promote the vegetation of host-beneficial members of the gut microbiota whose therapeutic effects can be exemplified by gut microbiota recovery in mice and guinea pigs with antibiotic-associated dysbiosis [155]. Metabiotic therapy, a variant of prebiotic therapy, implies the use of metabolites of functional microorganisms. Metabolites of *Lactobacillus plantarum* accelerated gut microbiota recovery in mice with antibiotic-associated dysbiosis by 5965 times and in guinea pigs by 308 times. The prebiotic drug Stim bifid (a complex of fructooligo- and fructopolysaccharides), when applied on the same experimental models, accelerated gut microbiota recovery by 7118 and 1047 times, respectively [155].

Probiotics are drugs derived from functional microorganisms referring to collection strains. A classic example of probiotic therapy is the use of *Lactobacillus bulgaricus* proposed by I.I. Mechnikov [29]. Preparations of live *Lactobacilli*, *Enterococci*, *Bifidobacteria*, and *E. coli* are widely used for this purpose [156]. Anaerobic Gram-positive bacteria—*Lactobacilli*, *Bifidobacteria*, *Propionobacteria*—suppress conditionally pathogenic intestinal microorganisms, as well as the production of ammonia, amines, substrates for the synthesis of trimethylamine-N-oxide, indoxyl sulfate, p-cresyl sulfate, which have both systemic and local toxicity [45]. Official microbiological preparations containing probiotic microorganisms are applicable as a source thereof. At the same time, such microorganisms are unable to be involved in the formation of a “microbial–tissue complex”, as well as in long-term colonization of the human intestine [155]. In this regard, autoprobiotic therapy, based on the application of certain indigenous bacteria isolated from the organism and returned to it after cultivation and accumulation in vitro, seems to be promising. The therapeutic efficacy of *Enterococci* [157] and *Bifidobacteria* [158], used in this way, has been shown in experimental dysbiosis.

Low-toxicity drugs, suitable for long-term application, are also promising for boosting the local intestinal resistance to toxic substances of bacterial origin. It is reasonable to search for them among the drugs having adaptogenic properties. The beneficial properties of plantain juice as a remedy to correct acute hyperammonemic and neurotoxic effects of cyclophosphamide have been experimentally proven. Against the background of just a single intragastric administration of an official preparation of plantain juice to rats at a dose equivalent to 47 mL of that for humans, the hyperammonemic effect of cyclophosphamide administered at a supralethal dose (2.1 LD50) was 1.5 times lower, the average hang time during which the rats could hold out on a wire grid in the four limb hanging test increased by 20%, while the average life expectancy of these animals increased by 30% [159].

Promising means of emergency prevention of IB lesions are the preparations able to form a protective film on the surface of the mucous membrane. Such substances can be pectins whose adhesive properties find use in hydrocolloid wound plasters [160]. When pectin is administered by ingestion, a protective film of calcium pectates forms on the mucous membranes [75], preventing substances produced by gut microflora from getting into blood. Gelatin tannate has a similar ability [161]. The protective effect of these substances is mainly expected during the drug transit in the small intestine: from 10 min to 5 h post ingestion.

The condition necessary for maintaining the intestinal barrier function in severe acute poisoning is the elimination of intestinal hypoxia and
inflammation, as confirmed by the anti-inflammatory effect of hyperbaric oxygenation in sepsis [162]. Clinical trials of oxygen-enriched gas mixtures containing inert gases, nitrogen oxide or hydrogen at low concentrations also show a lot of promise; the properties of these additives can modulate the effects of oxygen therapy. Xenon, nitric oxide and hydrogen have neuroprotective and cardiotonic properties, while argon and hydrogen sulfide have neuroprotective properties only [163]. Anti-inflammatory drugs can be supplementary to oxygen therapy: in experimental sepsis modeling, both IB dysfunction and endotoxemia were prevented by hydrocortisone administration [164].

CONCLUSION

1. In a healthy person, the intestinal chyme contains products of bacterial origin, whose amount and systemic toxicity profile, when losing the gradients of their concentration between the chyme and blood, ensure the formation of critical states of the organism, such as shock, sepsis, secondary acute lung injuries, and acute cerebral insufficiency.

2. Severe acute exogenous poisoning creates prerequisites for secondary intestinal barrier dysfunction and endotoxemia of intestinal origin.

3. Endotoxemia, including endotoxinemia and hyperammonemia, is involved in the formation of complications of acute poisoning, as manifested in critical states of an organism listed in paragraph 1.

4. Planned measures to protect the intestinal barrier from damage in the following acute poisoning should be aimed at optimizing the composition of gut microbiota and suppressing its production of cytotoxic substances, as well as boosting local intestinal resistance, by prescribing low-toxicity drugs intended for a long-term administration.

5. The main approaches to optimizing gut microbiota composition are prebiotic, metabiotic, probiotic and autoprobiotic therapies. Low-toxicity preparations having adaptogenic properties and suitable for long-term application are promising for increasing local intestinal resistance to toxic substances of bacterial origin.

6. Emergency prevention of secondary intestinal barrier dysfunction involves the use of drugs shielding the mucous membrane from the chyme and eliminating intestinal hypoxia and inflammation.

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AUTHORS’ CONTRIBUTION

Ju.Ju.I.—conceptualization, structuring the article, formulating its generalizations and conclusions; T.V.S.—participation in writing all sections of the article, searching and analyzing the scientific literature; V.L.R.—participation in writing the section “Intestinal barrier damage in acute poisoning”, searching and analyzing the scientific literature; O.A.V.—participation in searching and analyzing the scientific literature, preparing the text and list of references.

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