INTRODUCTION

Acute pneumonia (AP), described by Hippocrates two and a half millennia ago, is one of the oldest nosologies in medicine. However, a long history of popularity and study of AP does not prevent it from being an important and urgent problem so far. "Pneumonia is a common illness affecting approximately 450 million people a year and occurring in all parts of the world." It is a major cause of death among all age groups resulting in 4 million deaths (7% of the world’s total death) yearly. In addition, AP is the only non-specific inflammatory process of all known nosologies, which develops in the vascular system of the small circle of blood circulation. One of the leading cases of tissue transformation in the area of inflammation is a vascular reaction, which has a stereotypical sequence, while the vessels of the small circle of blood circulation are an integral part of the regulation of systemic blood flow. The combination of these reasons reflects the importance of a comprehensive study of the vascular bed of the lungs to understand the mechanisms of development and manifestation of AP.

MATERIALS AND METHODS

Studies were conducted in the Clinic of Pediatric Surgery at The State Institute for Advanced Training of Doctors (Novokuznetsk, USSR) in 1976–1985. During this period, the department treated 994 children aged 4 months to 14 years with various forms of community-acquired pneumonia and its purulent-destructive complications. Many patients from this number were selectively hospitalized in our department in the initial period of the most aggressive forms of AP immediately at diagnosis. The reason for hospitalization of the last group of patients is that the surgical
clinic was the only place of intensive care in our region with a population of about 2 million people. This group of patients was characterized by high mortality (>10%) and rapid development of pleural complications. Unsatisfactory results of traditional treatment (massive doses of antibiotics, oxygen supply, methods of bronchial drainage, and intravenous infusion) and a large concentration of very severe patients are forced to look for effective solutions.

The work began with a review of theoretical ideas about the nature of AP, its causes and mechanisms of development. For this purpose, known scientific data from certain biomedical rules and laws were used. To clarify some of the nuances of the disease and obtain objective evidence of the measures taken, special studies and experiments were conducted. One of the fragments of this work is presented below.

Rentgenoanatomical studies of lung specimens. A total of 56 whole lung anatomical preparations were taken for this study during the autopsy of children who died of pneumonia. Separate contrast vessels of the lungs (arteries or veins) followed by X-ray examination were performed on 47 samples (in 9 cases bronchi were contrasted). This type of research was carried out as follows. The procedure was carried out immediately after the autopsy and extraction of the specimen, which was not subjected to any additional effects. The polychlorinated vinyl catheter was inserted into the initial part of the corresponding vessel and strictly fixed in this position by an external silk ligature. The fixed catheter had a cannula at the distal end, to which a syringe with a liquid contrast agent was attached. Contrast agents were used, which are usually used in the clinic for diagnostic purposes by intravenous (eg, “urographin”). The contrast was injected a slight pressure on the plunger of the syringe until the first signs of resistance. X-ray specimen was performed immediately after contrast injection.

Pathological transformation in lung tissue ranged from inflammatory infiltration to necrosis and suppuration, and the extent of these changes was necessarily assessed by standard macro- and micro-scopic studies.

Since these studies, there have been no radical changes in the solution of the AP-problem, and no such tests have been conducted. Therefore, the results of research may be of interest to specialists in this field.

RESULTS AND DISCUSSION

The inflammatory process, regardless of localization, is accompanied by five classical signs reflecting the severity of tissue damage. These signs of inflammation, which were described by Celsus and Galen centuries ago, are well known because they remain at the heart of our understanding of this pathology. AP is a classic inflammatory process and has all of the above symptoms-heat, pain, redness, swelling, and loss of function. It is easy to note that among the presented signs of inflammation the fifth sign (loss of function) is of the greatest importance, since the specificity and severity of its clinical manifestations just depend on the localization of the process.

In this regard, it should also be noted the well-known fact that AP, as a rule, occurs without the appearance of the second classical sign of inflammation (pain), due to the lack of pain receptors in the lung tissue itself. The appearance of pain syndrome in AP usually indicates the participation of pleural sheets in the inflammatory process. However, other types of receptors, in particular, baroreceptors, which are in the area of inflammation, cannot avoid the irritating effects of inflammatory tissue transformation, and this fact can significantly affect the regulation of blood circulation in the body.

Well-known scientific facts about inflammation allow us to recall that the inflammatory transformation of tissues occurs due to the vascular reaction, which is based on successive stages of changes in blood flow, blood filling, and permeability of the vascular wall. In the case of AP, especially its aggressive forms with lobar organ damage, the anatomical picture of these stages has long been described and is well known. These anatomical stages are described in detail in the lobar forms of AP without destructive changes: Congestion, red hepatization, gray hepatization, and resolution.

These stages of inflammatory changes in lobar pneumonia have long been included in the category of classical descriptions of the disease, but this description, unfortunately, does not give a sufficient idea of a very significant detail of the restructuring of the vascular bed. According to the well-known biological postulate, the anatomical shape and structure of the organ correspond to its function (and vice versa). Edema and inflammatory infiltration of tissues arising from AP eliminate the airiness of the alveoli in the affected area and disconnect it from gas exchange. The fact of such anatomical transformation also refers to the classical characteristics of the AP. However, the transformation of the architectonics of blood vessels in the area of inflammation, which play an important role in the regulation of blood flow, requires clarification.

Our rentgenoanatomical studies have yielded the following results. Contrast studies of the vessels of the small circle of blood circulation showed that the arterial sector in the early stages (infiltration) of the inflammatory process retains its shape and the corresponding diameter of the vessels. A similar pattern is shown in Figure 1.

Pathological results on arteriograms were found only at the stage of alteration. In such situations, there was a sharp depletion of vascular pattern or a complete absence of
contrast vessels, depending on the spread and localization of necrosis. An example of such changes is shown in Figure 2.

In contrast to the arteries of the small circle, the architectonics of the venous vessels of this system is exposed to visible changes already at the stage of inflammatory infiltration of pulmonary tissue. Already in this period of development of the inflammatory process in the affected area, there is a significant decrease in the intensity of vascular pattern. It should be noted that this character of changes in vascular architectonics, which was also accompanied by a smaller penetration of contrast in the veins of large caliber, little depended on the severity of tissue damage in the AP zone. At the same time, in 2 cases with massive infiltration of the lobe of the lung, a contrast study revealed large vascular formations in non-inflammatory parts at the level of small venous branches. These formations in both cases were localized in the basal part and, from our point of view, were the evidence of excessive blood filling of healthy parts of lungs and opening of shunts [Figure 3].

Unfortunately, technical difficulties did not allow to conduct an identical study of bronchial vessels, the results of which could give a very important and interesting material for understanding the mechanisms of AP development. However, the results of studies that have been conducted, allow more detailed representation of the nature of vascular transformation in the focus of inflammation and in a small circle in the development of AP. The arterial sector of the small circle has a higher tone and blood pressure compared to the venous system. These qualities of arteries allow them to maintain their architectonics and provide the possibility of blood flow almost throughout the formation of the focus of AP. Only a complete blockade of blood flow in this area ends with the loss of a certain segment of blood vessels and irreversible consequences for the vassal tissue.

The preserved delivery of blood to the zone of inflammation at the initial stages of the process occurs in the conditions of early appearance of difficulties with its outflow. The more malleable venous area is subjected to external compression as a result of progressive processes of edema and infiltration. If in the described situation to increase blood flow to the area of inflammation, in conditions of the high permeability of the vascular wall and difficulties for blood outflow, such efforts will only stimulate tissue infiltration. The emerging restructuring of the angioarchitectonics creates a kind of

Figure 1: Arteriogram of the lung with extensive inflammation in the lower lobe. You can see the uniform distribution of contrast in the vessels of the entire lung. The diameter of the arteries is the same in all parts of the lung.

Figure 2: Arteriogram of the lung with destruction lesions in the lower lobe. The contrast did not fill the peripheral vascular network in some areas of the lower lobe, which coincide with areas of necrosis.

Figure 3: Venogram of the lung with full inflammation of the upper lobe. (Subsequent anatomical and histological studies revealed no necrotic changes). Sharp depletion of venous pattern in the upper affected part of the lung (large arrow). In the lower part of the architectonics of the vessels is preserved, but against this background, relatively large vascular formations of semicircular shape are visible.
“vascular trap.” This phenomenon is one of the explanations of those clinical observations when on the background of intravenous infusions in patients with initial stages of AP occurs catastrophically rapid deterioration of the X-ray picture.\[7\]

The described mechanism of stimulation of the inflammatory changes in the lung tissue belongs to the affected area and has a local character. However, we should not forget that we are talking about changes in the blood flow in the pulmonary vascular system. It is known that this part of the circulatory system is responsible for maintaining the necessary proportions between the two circles of blood circulation. Such proportions are based on the equality of blood volumes that pass through each half of the heart, and the inverse ratio of blood pressure. Under optimal conditions, the blood pressure of a large circle of blood circulation is about 5–6 times higher than the same indicator of pulmonary blood flow, and autonomous regulation is carried out through baroreceptors and is known to physiologists as the Schwiegk’s reflex.\[8,9\]

The study of angioarchitectonics in AP indicates anatomical vascular changes in pulmonary blood flow. According to one of the postulates of biology and physiology, anatomical changes inevitably entail functional shifts, which in AP begin primarily with pathological reflexes. Elimination of reflex influence of AP-focus on systemic circulation is an important element of first aid to such patients.\[10-15\] A detailed description of all studies and clinical testing of new approaches in the treatment of AP can be found in the published monograph.\[16\]

**CONCLUSION**

Thus, the results of the study of angioarchitectonics in AP allow to understand and present in more detail the features of the mechanisms of development of the inflammatory process, as well as facilitate the preparation of a new doctrine of the disease.

**REFERENCES**


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