Rodent models of pulmonary embolism and chronic thromboembolic pulmonary hypertension

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1. Introduction

Pulmonary embolism (PE) is the third most common cardiovascular disease [1]. It is associated with high in-hospital mortality and the development of acute and chronic complications [2]. Moreover, observational studies have shown a trend towards an increase in the number of PE cases per year over time [3]. According to Keller et al. [4], the incidence of PE in Germany from 2005 to 2015 increased from 85 to 109 cases per 100 000 people per year. Similar data were obtained from the Danish National Patient Register, in which there was an increase in the incidence of PE between 2004 and 2014, from 45 to 83 cases per 100 000 people per year [5]. This trend may be accounted for by an increase in life expectancy, a greater prevalence of risk factors for PE, and a significant improvement in PE detection. In addition, PE can be accompanied by a number of complications. One of the most serious complications is chronic thromboembolic pulmonary hypertension (CTEPH), which is characterized by incomplete lysis of thromboemboli and a sustained increase in pulmonary artery pressure. The five-year survival rate of patients with CTEPH receiving only oral anticoagulant therapy, with a mean pulmonary artery pressure of more than 30 mmHg, does not exceed 10% [6].

The treatment and prevention of PE, as well as its complications, require high-tech approaches aimed at improving patient prognosis. In this regard, experimental studies aimed at clarifying the pathogenesis, improving diagnostic methods, and identifying new drug targets for prevention and treatment of PE and its complications represent an unmet clinical need. Considering the current trends in preclinical research, the use of rodent models, such as mice, rats, hamsters, and rabbits is becoming more widespread. The use of these animal species is ethically acceptable and economically feasible.

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The literature describes a large number of models used to induce PE. All available approaches to modeling PE can be divided into three main groups:

- Induction of thromboembolism
  - Induction of thrombus formation in vivo
  - Injection of blood clots prepared in vitro
  - Introduction of non-thrombogenic particles
  - Surgical procedures.

A separate direction in this area is the modeling of CTEPH. Due to the complex pathogenesis of this condition, as well as the high fibrinolytic activity of blood plasma in rodents, modeling this pathology is difficult and currently does not have an optimal solution.

The choice of animal species and the approach to modeling PE are primarily determined by the objectives of the specific study. This article provides an overview of the main approaches to the modeling of PE and discusses the advantages and disadvantages of each method. This study analyzed publications from 1978 to 2021, in which PE was induced in rodents.

2. PE modeling using thrombosis induction in vivo

For PE modeling by inducing endogenous thrombus formation, animals are injected intravenously with agonists causing platelet aggregation or substances that trigger the coagulation cascade. Table 1 summarizes the studies performed using these models. This group of models is most often used to study the anticoagulant [7, 8, 9] and antiplatelet [10, 11, 12] activity of pharmacological substances, and to investigate the pathogenesis of various forms of blood coagulation disorders [13, 14, 15]. Mice are most commonly used for this purpose [16, 17, 18, 19]. This makes it possible to form large study groups and thus partially compensates for the low reproducibility of these models. Occasionally, rabbits [20] are used for this purpose.

Collagen is often used for the formation of endogenous blood clots either on its own [7, 21] or in combination with epinephrine [14, 17, 22] or norepinephrine [11]. The doses of collagen and epinephrine administered varied within significant limits: dose range for collagen varied from 12.5 μg/kg [23, 24] to 100 mg/kg [25], while for epinephrine it varied from 0.075 μg/kg [24] to 1 mg/kg [26].

Generally, the administered dose of thrombus inducers was determined by calculating the target mortality in the control group. For example, in a study by Crikis et al. [7], isolated intravenous administration of collagen at a dose of 0.075 μg/g was used to model PE in mice. The mortality rate of animals in the control group 1 h after collagen injection was 53%. Ryu et al. [22] induced CTEPH in mice by intravenous administration of collagen (500 μg/kg) in combination with epinephrine (50 μg/kg). Fifteen minutes after drug administration, the mortality was 82.9 ± 10.7%.

Other substances could also be used to induce clot formation in vivo after systemic administration. These include adenosine diphosphate (ADP) [12, 18, 27], thrombin [9, 28, 29], high molecular weight poly-phosphate [13], thromboxane A2 receptor agonist U46619 [24], neutrophil elastase [27, 30], arachidonic acid [18], tissue factor [8], or a combination of tissue factor with phospholipids and calcium chloride [13].

Katsumata et al. used ADP as an agent for modeling PE [27]. Here, mice were injected once intravenously with ADP at a dose of 280 mg/kg. The mortality rate in the early stages after introduction of the thrombogenic agent was 43%. It should be noted that isolated use of ADP is rare. The dose of ADP administered varies from 280 mg/kg to 40 mg/100 g [18, 27]. An example of such an isolated application of ADP is a study by Katsumata et al. [27], in which mice were injected once intravenously with ADP at a dose of 280 mg/kg. The mortality rate in the early stages after introduction of the thrombogenic agent was 43%.

Hsu et al. [18] used three different prothrombotic agents simultaneously to model PE in mice: ADP, collagen, and arachidonic acid were used at a dose of 40 mg/100 g, 2.5 mg/100 g, and 9 mg/100 g of animal weight, respectively. With intravenous administration of these substances, the death rate from PE was 90% during the first 3 min of observation. The authors indicated differences in the effectiveness of various standard drugs affecting thrombus formation, and the test substance to influence the survival rate of animals after PE. In this study, aspirin and indomethacin have been shown to reduce mortality from PE caused by collagen and arachidonic acid but do not affect survival outcomes using the ADP model. Heparin had no effect on any of the models used. Such differences should be considered when choosing a model to test new drugs that affect thrombus formation.

To increase the efficiency of thrombus formation, a combination of ADP with other prothrombotic agents, such as neutrophil elastase, is occasionally used [27]. Two non-standard approaches to modeling PE were described by Banno et al. [13]. In the first approach, human re-combinant tissue factor containing phospholipids and calcium was used as a prothrombotic agent; in the second approach, high molecular weight polyphosphate was used. Polyphosphate is a linear polymer of inorganic phosphates that acts as a natural negatively charged surface that activates the blood coagulation system. Doses of both substances were selected based on 20% survival 20 min after PE in the control group of mice. The authors did not provide comparisons with other available PE models. However, it is likely that the choice of non-standard prothrombotic agents was determined by the aim of the research, which was to study the effect of protein S mutation on venous thromboembolism. Thus, the researchers were primarily interested in the coagulation cascade during thrombus formation, which is facilitated by the agents used in the modeling.

In contrast to the above-described models utilizing non-localized formation of clots in the circulation followed by embolism, there were occasional attempts of PE modeling more closely mimicking the clinical scenario, implying formation of localized thrombi in the limb veins with subsequent displacement and obstruction of pulmonary artery branches. An example of such an approach is a study by Shaya et al. [31], in which the factor V Leiden paradox was investigated; the incidence of deep vein thrombosis was proportionally greater than that of PE in these individuals. To simulate this phenomenon, a piece of filter paper impregnated with 1.8% ferric chloride was applied to the femoral vein. The exposure time was 5 min. After 2 h, the size of the thrombus in situ was assessed, as was the severity of PE.

The above models are characterized by significant variability in the obtained results, which should be considered when planning the sample size. Owing to this, a vast majority of such experiments are performed on readily available mice. Among the advantages, the high pathophysiological accuracy of such models should be pointed out, as this makes it possible to study pharmacological substances with sufficient fidelity to correct the coagulation and aggregation properties of blood.

2.1. Administration of thrombi prepared in vitro

The protocol for modeling PE using thrombotic masses prepared in vitro assumes the initial sampling of animal blood and fabrication of native or modified thrombi of a given size and number, followed by their intravenous administration to the same animal. In some studies, allogeneic [32, 33] or even xenogeneic blood [34, 35, 36] has been used for ex vivo preparation of thromboemboli. These models are produced in animals of various species, including mice [37, 38, 39], rats [37, 40, 41], hamsters [35, 42, 43], and rabbits [44, 45, 46] (Table 2).

A classic example of PE modeling using in vitro-prepared thromboemboli is the modeling protocol used by Tang et al. [47]. In this study, changes in gene expression in the pulmonary artery in the subacute period (7 days) after PE were studied. Rabbits were used, and 0.5 ml of blood was preliminarily collected from the marginal ear vein.
Table 1. PE modelling by blood clots induction in vivo.

| № | Animal species | Blood clots agents | Substance/action | Aim of study | Administration vessel | Study duration | CTEPH pathway possibility | Authors |
|---|----------------|--------------------|------------------|-------------|-----------------------|--------------|--------------------------|---------|
| 1 | Mice           | Collagen + epinephrine | Collagen = 0,8 μg/g; Epinephrine = 60 μg/kg | Studying of cathesatin role in blood clots | Inferior vena cava | 30 min | – | [100] |
| 2 | Mice           | Thrombin           | 20 U             | Studying of gelsolin role in PE prevention | Tail vein | 15 min | – | [101] |
| 3 | Mice           | Collagen + epinephrine | Collagen = 400 μg/kg; Epinephrine = 60 μg/kg | Studying of functional role of platelet isoform CD45 | retro-orbital sinus | Acute study (After respiratory distress < 300 s) | – | [102] |
| 4 | Mice           | Collagen + epinephrine | Collagen = 400 μg/kg; Epinephrine = 60 μg/kg | Definition of functional role of ELMO1 peptide in platelets | Intravenous (n/d) | Acute study (After respiratory distress) | – | [103] |
| 5 | Mice           | Application of ferric chloride (III) (soaked filter paper) on femoral vein Times of experiment: 0.5 μL; Collagen = 64 μg/kg; Epinephrine = 5.4 μg/kg | Size of filter paper - 1 x 2 mm; FeCl3 concentration-1.8%; Exposure time - 5 min | Studying of generation rate, stability and embolization risk in case of factor V Leiden (FVL) variant | Femoral vein | 2 h | – | [31] |
| 6 | Mice           | Collagen + epinephrine | Collagen = 280 μg/kg; Epinephrine = 29 μg/kg | Studying of inhibitor tyrosineproteinokinase MER effectiveness in platelets inactivation and blood clots prevention | Intravenous (n/d) | 15 min | – | [104] |
| 7 | Mice           | 1) Thromboplastin 2) Collagen + epinephrine | Thromboplastin = 5 μL; Collagen = 64 μg/kg; Epinephrine = 5.4 μg/kg | Studying of platelets TL-1 receptor role of immune origin hemorrhage | Intravenous (n/d) | 30 min | – | [105] |
| 8 | Mice           | Collagen/2MeSADP + epinephrine | Collagen = 400 μg/kg; 2MeSADP = 20 μg/kg; Epinephrine = 30 μg/kg | Studying of platelets activation mechanisms in case of hemostasis | retro-orbital sinus; Inferior vena cava | Acute study (After respiratory distress) | – | [106] |
| 9 | Mice           | Collagen + epinephrine | Collagen = 430 μg/kg; Epinephrine = 20 μg/kg | Studying of glucose metabolism effect on platelets function regulation | Intravenous (n/d) | 25 min | – | [14] |
| 10 | Mice          | Collagen + epinephrine | Collagen = 500 μg/kg; Epinephrine = 50 μg/kg | Studying of thrombin inhibitor anticoagulant potency in PE prevention | Tail vein | 15 min | – | [107] |
| 11 | Mice          | Thromboplastin    | 7.5 μL/g | Studying of the vaccination anticoagulant properties to factor XI in thrombosis prevention | Intravenous (n/d) | 3 min | – | [8] |
| 12 | Mice          | Collagen + epinephrine | Collagen = 0,6 μg/g; Epinephrine = 0.2 μg/g | Study of the thromboxane receptor antagonist antithrombotic properties in PE prevention | Tail vein | Acute study (After respiratory and heartbeat distress) | – | [17] |
| 13 | Mice          | Collagen + epinephrine | Collagen = 500 μg/kg; Epinephrine = 50 μg/kg | Studying of the antithrombotic and anticoagulant activity of alkaloids Scolopendra subspinipes mutilans | Tail vein | 15 min | – | [108] |
| 14 | Mice          | 1) Tissue factor (human recombinant) with phospholipids and calcium 2) High molecular polyphosphate | Till 80% mortality after 20 min after administration | Assessment of protein S mutation role in the development of venous blood clots in PRGS1 knockout mice | Inferior vena cava | 20 min | – | [13] |
| 15 | Mice          | Collagen + epinephrine | Collagen = 0,5 μg/kg; Epinephrine = 0,06 μg/kg | Studying of the limiglydol antithrombotic properties in PE prevention | Tail vein | 1 day | – | [109] |

(continued on next page)
Table 1 (continued)

| №  | Animal species | Blood clots agents | Aim of study | Administration vessel | Study duration | CTEPH pathway possibility | Authors |
|----|----------------|--------------------|--------------|-----------------------|---------------|-------------------------|---------|
| 16 | Mice           | Splenectomy followed after 1 month narrowing of the inferior vena cava lumen, by ligation | Studying of splenectomy role in CTEPH formation | – | 1–28 days | CTEPH +/- (after 28 days pressure in PA wasn’t measured) | [92]    |
| 17 | Mice           | Thrombin           | 1500 IU/kg body weight | Tail vein | 15 min | – | [28]    |
| 18 | Mice           | Collagen + epinephrine/ collagen + epinephrine + amyloid β | Collagen – 1000 μg/kg; Epinephrine – 10 μg/kg; Amyloid β – 2,5 mg/kg | Tail vein | 15 min | – | [15]    |
| 19 | Mice           | Thrombin           | 1250 IU/kg body weight | Tail vein | 15 min | – | [9]     |
| 20 | Mice           | 1. Collagen + noepinephrine 2. Thrombin | 1. Collagen – 500 μg/kg; noepinephrine – 80 μg/kg; 2. Thrombin - 2000 IU | Intravenous (n/d) | 15 min | – | [11]    |
| 21 | Mice           | Thromboplastin     | 1–1,33 mg/mice | Tail vein | 30 min | – | [110]   |
| 22 | Mice           | Collagen           | Collagen – 0,4 mg/kg | Tail vein | 1 h | – | [21]    |
| 23 | Mice           | Collagen + epinephrine | Collagen – 80 mg/kg; Epinephrine – 1 mg/kg | Tail vein | 15 min | – | [26]    |
| 24 | Mice           | Collagen           | 0,075 μg/g body weight | external jugular vein | 1 h | – | [7]     |
| 25 | Mice           | Collagen + epinephrine | Collagen – 500 μg/kg; Epinephrine – 50 μg/kg | Tail vein | 15 min | – | [22]    |
| 26 | Mice           | Neutrophil elastase + collagen-induced arthritis | Neutrophil elastase – 5 U/kg, 2 times per day administration during 3 days | Tail vein | 4 days after last injection | – | [30]    |
| 27 | Mice           | Collagen + epinephrine/U46619 | Collagen – 12,5 μg/kg; Epinephrine – 0,075 μg/kg; U46619 – 0,2 mg/kg | Intravenous (n/d) | 15 min | – | [23]    |
| 28 | Mice           | Thromboplastin     | Thromboplastin – 65 μL/kg or 140 μL/kg | Jugular vein | 30 min | – | [111]   |
| 29 | Mice           | Collagen III type + epinephrine | Collagen – 100 mg/kg; Epinephrine – 90 μg/kg | Jugular vein | 8 min | – | [25]    |
| 30 | Mice           | Collagen + epinephrine | Collagen – 800 μg/kg; Epinephrine – 60 μg/kg | Jugular vein | 5 min | – | [112]   |
| 31 | Mice           | ADP/NE + ADP       | NE – 10 μg/mouse; A₁D₄ – 28 mg/100 g | Intravenous (n/d) | Acute experiment (n/d) | – | [27]    |

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Subsequently, thrombotic masses were formed under sterile conditions at room temperature for 45 min, from which embolizing particles 5-mm long were prepared by cutting. Blood clots were suspended in 10 ml of normal saline and injected into the femoral vein. An additional 5 ml of saline was used to flush the catheter and advance the thromboemboli.

In the majority of studies, tubes of corresponding diameter were used for the formation of thromboemboli of given sizes. A thrombotic mass formed inside each tube, and the resulting thrombotic mass was cut to the desired length, giving it the characteristics of the desired naturally occurring thromboembolus [48, 49].

To accelerate the formation of thrombotic masses in vitro in a number of studies, thrombin [33, 50, 51], tissue factor [49], and calcium chloride [50, 51, 52] were used. As mentioned above, rodent blood has a high fibrinolytic activity, which leads to rapid lysis of exogenously administered blood clots. In order to overcome this problem, fibrinolysis was prevented by adding a 5 ml of saline containing 10 mg/kg of U46619, a thromboxane A2 receptor antagonist.

In order to observe embolus formation, the PE pathway was generated using a combination of thrombin, epinephrine, and collagen. These agents were used in various doses, as shown in Table 1.

### Table 1 (continued)

| No. | Animal species | Blood clots agents | Aim of study | Administration vessel | Study duration | CTEPH pathway possibility | Authors |
|-----|----------------|---------------------|--------------|----------------------|----------------|--------------------------|---------|
| 32  | Mice           | Collagen + epinephrine | Collagen – 12.5 μg/mouse, Epinephrine – 0.075 μg/mouse | Investigation of the combination of nitroaspirin and clopidogrel antithrombotic properties in the PE prevention | Intravenous (n/d) | 2 min | – | [113] |
| 33  | Mice           | Collagen + epinephrine/U46619 agonist to α2 thromboxane receptors | Collagen - 12.5 μg/kg; Epinephrine – 0.075 μg/kg; U46619 – 0.2 μg/kg | Studying of the NO-releasing statin antithrombotic properties | Intravenous (n/d) | 15 min | – | [24] |
| 34  | Mice           | Collagen + epinephrine/collagen | Collagen – 25/12/6/3 μg; Epinephrine – 1 μg | Studying of the α2/1 integrin role in the platelet aggregation and thrombus formation | Right Jugular vein | 3 min | – | [19] |
| 35  | Mice           | Thrombin           | It was counted as a 90% mortality dose | Studying of low molecular weight heparin anticoagulant action mechanisms | Tail vein | 15 min | – | [114] |
| 36  | Mice           | Thrombin           | 1250 IU/kg body weight | Investigation of the protein C anticoagulant properties in blood clots prevention | Tail vein | 15 min | – | [29] |
| 37  | Mice           | Collagen + epinephrine | collagen – 400 μg/kg; epinephrine – 50 μg/kg | Study of the glycoprotein IIb/IIIa antagonist antithrombotic properties in the PE prevention | Tail vein | 15 min | – | [16] |
| 38  | Mice           | ADP                | 300 μg/g body weight | Assessment of the drug antithrombotic properties | Tail vein | 10 min | – | [12] |
| 39  | Mice           | Collagen + epinephrine | collagen – 25 μg; epinephrine – 1,5 μg | Evaluation of the antithrombotic drugs effectiveness in the PE prevention | Tail vein | 3 min | – | [10] |
| 40  | Mice and rabbits | Mice: human thrombin/collagen + epinephrine; Rabbit: thrombin/ADP/platelets activating factor | Mice: human thrombin 1250 IU/kg, collagen 1,25 mg/kg, epinephrine 75 μg/kg
Rabbits: thrombin 10/20 IU/kg or ADP 20 μg/kg or platelets activating factor – 50 ng/kg. | Studying of the anticoagulant and antithrombotic properties of defibrotide | Mice – tail vein; Rabbits – marginal auricular vein
Mice – 15 min; Rabbits – 5 min | – | [20] |
| 41  | Mice           | Thrombin           | Thrombin (0, 25, 50, 75 IU/mouse) | Studying of the thrombomodulin anticoagulant properties in the PE prevention | Tail vein | 15 min | – | [115] |
| 42  | Mice           | Thrombin           | 3,8–15 IU/mouse | Studying of the thrombomodulin role in the PE pathogenesis | Tail vein | 1 h | – | [116] |
| 43  | Mice           | ADP/collagen/ arachidonate | ADP – 400 mg/kg; Collagen – 25 mg/kg; Arachidonate – 90 mg/kg | Studying of the Agrimonial pilosa extract antithrombotic properties in the PE prevention | Intravenous (n/d) | 3 min | – | [18] |

**Note:** ADP — adenosinediphosphate, i/v — intravenous administration, IU — international unit, PE — pulmonary embolism, CTEPH — chronic thromboembolic pulmonary hypertension, 2MeSADP — 2-methylthio-adenosine-5'-diphosphate, n/d — no data, NO — nitrogen oxide (II), NE — neutrophil elastase, U46619 — 9,11-dideoxy-11a,9a-epoxymethanoprostaglandin F2α.
## Table 2. Administration of thrombi prepared in vitro.

| No | Animal species | Embolizing particles | Material | Label | Size | Dose | Preparation conditions | Aim of work | Vessel of sampling/ administration | Study duration | Possibility of CTEPH formation | Authors |
|----|----------------|----------------------|----------|-------|------|------|------------------------|-------------|-----------------------------------|---------------|----------------------------------|---------|
| 1  | Mice           | Autologous thrombi  | –        | 1 × 1 mm | 30 pcs | –    | –                      | Studying of miR-106b-5p role in PE | Blood sampling – tail vein | 7 days | – [117]                          |         |
| 2  | Mice           | Xenogenic fibrinous thrombi (human plasma) | $^{125}$I-labeled fibrin | 12.5 μL | 20 pcs | in vitro addition of CaCl$_2$ and thrombin | Study of α2-antiplasmin contribution to fibrinolytic insufficiency | TE administration – jugular vein | 4 h | – [39]                          |         |
| 3  | Mice           | Autologous fibrinous thrombi | – | 1 × 1.2 mm | 30 pcs | in vitro addition of CaCl$_2$ and thrombin | Study of mesenchymal stem cells use effectiveness in acute PE | Blood sampling – tail vein; TE administration – jugular vein | 1 day | – [118]                          |         |
| 4  | Mice           | Xenogenic fibrinous thrombi (human plasma) | $^{125}$I-labeled fibrin | 1.5–5 μm | – | – | – | Study of t-PA, tenecteplase and reteplase fibrinolytic capacity in PE | TE administration – jugular vein | 1 h | – [119]                          |         |
| 5  | Mice           | Fibrinous thrombi    | $^{125}$I-labeled fibrin | – | – | – | – | Study of pulmonary vascular activity regulation using tissue plasminogen activator via NMDA receptors | TE administration – jugular vein | 1 h | – [120]                          |         |
| 6  | Mice           | Autologous fibrinous thrombi | – | 1.2 × 1 mm | 30 TE | in vitro addition of CaCl$_2$ and thrombin | Study of the number and function of endothelial precursors from the bone marrow in acute PE in mice. | Blood sampling – tail vein. TE administration – jugular vein | 1 h, 1 day, 2 days | – [51]                          |         |
| 7  | Mice           | Autologous fibrinous thrombi | $^{125}$I-labeled fibrin | – | 25 μL | – | – | Study of fibrinolytic properties during adenoviral transfection of the plasminogen activator gene in mice with varying degrees of tPA deficiency | TE administration – jugular vein | 16 h | – [38]                          |         |
| 8  | Mice           | Xenogenic thrombi from modified RBC (rats RBC) | – | – | – | Chlorpromazine and glutaraldehyde were added | New model development | Blood sampling – blood rats; TE administration – tail vein | 15 min | – [36]                          |         |
| 9  | Mice & rats    | Fibrinous thrombi    | $^{125}$I-labeled fibrin | 3–5 μm | – | in vitro addition of CaCl$_2$ and thrombin | Study of the tPA fibrinolytic activity associated with RBC in the thrombosis prevention | – | 1 h | – [37]                          |         |
| 10 | Rats           | Autologous thrombi + tranexamic acid + carrageenan | – | – | 32 ± 5 Administration was repeated after 4 and 11 days from the 1st administration | – | Study of the aseptic inflammation role in the CTEPH formation | Blood sampling – external jugular vein. TE administration – jugular vein | 32 days | CTEPH + [90]                      |         |

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Table 2 (continued)

| No | Animal species | Embolizing particles | Material | Label | Size | Dose | Preparation conditions | Aim of work | Vessel of sampling/ administration | Study duration | Possibility of CTEPH formation | Authors |
|----|----------------|----------------------|----------|-------|------|------|------------------------|-------------|----------------------------------|---------------|----------------------------------|---------|
| 11 | Rats           | Autologous thrombi   | –        | 1 x 2 mm | 30 TE | –    | –                      | Urokinase effect on PE formation | Blood sampling – tail vein; TE administration – jugular vein | 6 h         | –                               | [121]   |
| 12 | Rats           | Autologous thrombi + tranexamic acid | – | 1 x 3 mm | Administration was repeated after 4 and 7 days from the 1st administration | – | Study of transcription factor O-1 and apoptosis role in remodeling of pulmonary artery branches in CTEPH | Blood sampling – orbital vein; TE administration – jugular vein | 1, 2 and 4 weeks | CTEPH + | [86] |
| 13 | Rats           | Autologous thrombi   | –        | 1.1 x 2 mm | 25 TE | Incubation on water bath 70°C, 10 min | Evaluation of low molecular weight heparin effect on intimal hyperplasia in acute PE | Blood sampling – tail vein; TE administration – jugular vein | 1, 4, 7, 14 and 28 days | CTEPH + | [85] |
| 14 | Rats           | Autologous thrombi + tranexamic acid | – | 1 x 3 mm | 40 TE; Administration was made 2 times with 4 days interval | – | Study of the tissue factor and transcription factor O-1 expression in CTEPH modelling | Blood sampling – orbital vein; TE administration – jugular vein | 1, 2 and 4 weeks | CTEPH + | [53] |
| 15 | Rats           | Autologous thrombi + tranexamic acid | – | 1 x 3 mm | Administration was made 2 times with 4 days interval | – | Study of the tissue factor and autophagy role in the remodeling of pulmonary artery branches in CTEPH | Blood sampling – orbital vein; TE administration – jugular vein | 1, 2 and 4 weeks | CTEPH + | [87] |
| 16 | Rats           | Autologous thrombi + tranexamic acid | – | 3 mm x 1 mm | 15 emboli; Administration was made 3 times with 2 weeks interval | 0.2 ml of blood, 37°C till night | Study of the brinolysis substance Rg1 effect on fibrous myocardial remodeling in CTEPH | Blood sampling – tail vein; TE administration – jugular vein | 4 weeks – preparation time; + 4 weeks of observing | CTEPH + | [88] |
| 17 | Rats           | Autologous thrombi   | –        | ~2 mm³ | 15-20 pcs | – | Study of the aspirin effects on ERK and PI3K/Akt signaling pathways in PE | Blood sampling – orbital vein; TE administration – jugular vein | 3 days | – | [41] |
| 18 | Rats           | Autologous thrombi   | –        | 1.1 x 3 mm | 0.5 ml (18 mg/100 g) | in vitro addition of CaCl₂ and thrombin | Chimeric antibodies to MCP-1 suppress the development of acute pulmonary hypertension after PE | Blood sampling – carotid; TE administration – jugular vein | 1, 4 and 8 h | – | [52] |
| 19 | Rats           | Allogenic fibrinous thrombi | 125I labeled fibrin | 10–100 μm | 75 μL/kg | in vitro addition of CaCl₂ and thrombin | Study of the SMTP substance fibrinolytic properties after PE | TE administration – tail vein | 20 min | – | [40] |
| 20 | Two rats lines: Sprague-Dawley & Copenhagen | Autologous thrombi + tranexamic acid | – | SD – 1.4 mm diameter; Cop. – 1.14 mm diameter | SD – 18 mg/100 g; Cop. – 5 mg/100 g | in vitro addition of CaCl₂ and thrombin | PE modelling with fibrinolysis suppression and stimulation of coagulation on different rats’ species | Blood sampling – jugular vein. TE administration – jugular vein | 1 and 5 days | – | [50] |

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| №  | Animal species | Embolizing particles | Material | Label | Size | Dose | Preparation conditions | Aim of work | Vessel of sampling/administration | Study duration | Possibility of CTEPH formation | Authors |
|----|----------------|----------------------|----------|-------|------|------|-------------------------|-------------|--------------------------------|----------------|---------------------------------|---------|
| 21 | Rats           | Allogenic thrombi    | –        | –     | 10 mm × 1 mm | 3 TE | in vitro addition of thrombin | Study of plasma proteome changes in acute PE | TE administration – jugular vein | 1, 8, 24, 48 h | –                               | [48]    |
| 22 | Rats           | Allogenic thrombi    | –        | –     | 10 × 1 mm | 3 TE | in vitro addition of thrombin | Study of paracrine factors in acute PE | Blood sampling – from another rats (cardiac puncture), TE administration – jugular vein | 1, 8, 24 and 48 h | –                               | [33]    |
| 23 | Rats           | Xenogenic fibrous thrombi (rabbit plasma) | –        | –     | –    | –    | –                       | Study of targeted thrombolysis of monoclonal antibodies chimeric molecules to the pulmonary artery endothelium and urokinase. | TE administration – jugular vein | 2 h                               | [34]    |
| 24 | Rats           | Allogenic fibrous thrombi | 129I-labeled fibrin | 10–100 μm | –    | –    | in vitro addition of CaCl₂ and thrombin, Thrombi were frozen by liquid nitrogen before blending | Study of the surfactin C effect on the fibrinolytic capacity of plasma caused by the plasminogen activation in PE | TE administration – tail vein | 80 min                           | –       | [122]                           |
| 25 | Rats           | Allogenic thrombi    | –        | –     | –    | –    | –                       | Study of RV and LV dysfunction in massive acute PE | Blood sampling – donor femoral artery, TE administration – jugular vein | 30 min                           | –                                 | [32]    |
| 26 | Rats           | Allogenic thrombi    | 129I-labeled fibrin | 1,47 × 12 mm | 1 TE | –    | in vitro addition of CaCl₂ and thromboplastin | Study of thrombolytic properties of Desmodus Rotundus salivary plasminogen activator in PE | Blood sampling – carotid (rat donor); TE administration – jugular vein | 1 h                               | –       | [49]    |
| 27 | Rats and hamsters | Rat – allogenic fibrous thrombi; hamsters – xenogenic fibrous thrombi (human plasma) | –        | –     | –    | –    | 2 models comparison, Evaluation of t-PA effect | TE administration – inferior vena cava | 2,5 h                           | –                                 | [123]   |
| 28 | Hamsters       | Xenogenic fibrous thrombi (human plasma) | 129I-labeled fibrin | – | 50 μL | – | – | Study of thrombolytic and antiplatelet properties of a recombinant chimeric molecule containing an activated platelet agonist and single-chain urokinase. | TE administration – jugular vein | 1,5 h                           | –                                 | [35]    |

(continued on next page)
| No | Animal species | Embolizing particles | Material | Label | Size | Dose | Preparation conditions | Aim of work | Vessel of sampling/administration | Study duration | Possibility of CTEPH formation | Authors |
|----|----------------|----------------------|----------|-------|------|------|------------------------|-------------|--------------------------------|----------------|---------------------------------|---------|
| 29 | Hamsters       | Xenogenic fibrinous thrombi (human plasma) | ¹²⁵I-labeled fibrin | – | 25 μL | – | Study of the recombinant t-PA anticoagulant activity with a Lys296-Gly302 deletion | TE administration – left jugular vein | 1.5 h | – | [54] |
| 30 | Hamsters       | Xenogenic fibrinous thrombi (human plasma) | ¹²⁵I-labeled fibrin | – | 25 μL | – | Study of thrombolytic and antiplatelet properties of a conjugated molecule consisting of u-PA and monoclonal antibodies to platelets | TE administration – jugular vein | 1.5 h | – | [43] |
| 31 | Hamsters       | Autologous fibrinous thrombi | ¹²⁵I-labeled fibrin | – | 25 μL | – | Comparative study of bolus thrombolytic properties and continuous administration of a chimeric plasminogen activator molecule (t-PA/u-PA) | TE administration – jugular vein | 1.5 h | – | [42] |
| 32 | Rabbits        | Autologous thrombi | – | 3 × 10 mm | Mean number of thrombi 4.9 ± 1.2 (before shocked) | – | Study of sodium nitroprusside and diltiazem effect on the sympathetic nervous system activation. | Blood sampling – femoral vein. TE administration – pulmonary artery (catheterized through the femoral vein) | 2 h | – | [96] |
| 33 | Rabbits        | Autologous thrombi | – | 3 × 10 mm | Mean number of thrombi 4.9 ± 1.2 (before shocked) | – | Study of the sodium nitroprusside effects on hemodynamics in massive PE with cardiogenic shock | Blood sampling – femoral vein. TE administration – pulmonary artery (a catheter passed through the femoral vein) | 2 h | – | [57] |
| 34 | Rabbits        | Autologous thrombi | – | 1 × 3–4 mm | Thrombi from 1 ml of blood | Incubation on water bath 70 °C, 10 min | Study of the nitric oxide inhalation effect on troponin I and blood clotting factors during massive PE | Blood sampling – auricular vein. TE administration – jugular vein | 120 min | – | [46] |
| 35 | Rabbits        | Autologous thrombi | – | 1–2 mm | Thrombi from 1 ml of blood | – | Assessment of hemodynamic changes during acute PE | Blood sampling – jugular vein. TE administration – jugular vein | <1 h | – | [56] |
| 36 | Rabbits        | Autologous thrombi | – | 0.8–1 mm | Thrombi from 1 ml of blood | – | Assessment of hemodynamic changes during acute PE | Blood sampling – jugular vein. TE administration – jugular vein | 5 min | – | [55] |
Table 2 (continued)

| № | Animal species | Embolizing particles | Material | Label | Size | Dose | Preparation conditions | Aim of work | Vessel of sampling/ administration | Study duration | Possibility of CTEPH formation | Authors |
|---|----------------|----------------------|----------|-------|------|------|------------------------|-------------|----------------------------------|--------------|-------------------------------|---------|
| 37 | Rabbits        | Autologous thrombi   | –        |   | 5 mm | Thrombi from 0.5 ml of blood | – | Study of gene expression changes in the pulmonary artery during the subacute period (7 days) after PE | Blood sampling – marginal auricular vein. TE administration – jugular vein | 7 days | – | [47] |
| 38 | Rabbits        | Autologous thrombi   | –        |   | 2–4 mm diameter, 7–10 mm length | 2 pcs in vitro addition of thrombin | Changes in tissue factor expression in rabbits during acute PE | Blood sampling – marginal auricular vein. TE administration – jugular vein | 3, 8 and 24 h | – | [44] |
| 39 | Rabbits        | Autologous thrombi   | –        |   | 2 × 20 mm | – | Study of FIIa enzyme fibrinolytic activity from the Agkistrodon acutus venom in acute PE | TE administration through a catheter in the RV | 2 h | – | [45] |
| 40 | Rabbits        | Autologous thrombi   | –        |   | – | Thrombi from 10 ml or 1 ml of blood | – | Comparison of the traditional iodine-containing contrast effectiveness and contrast agent based on the liposomal pool of blood in PE | Blood sampling – auricular vein. TE administration – enteral jugular vein | 270 min | – | [58] |

Note: RV — right ventricular, TE — thromboemboli, PE — pulmonary embolism, CTEPH — chronic thromboembolic pulmonary hypertension, CaCl₂ — calcium chloride, ¹²⁵I — iodine-125 labelled, MCP-1 — monocyte attractant protein-1, SD — Sprague-Dawley, Cop. — Copenhagen, t-PA — tissue plasminogen activator, u-PA — urokinase plasminogen activator.

Inhibitors such as tranexamic acid have been used to suppress the fibrinolytic activity [50, 53].

The activity of the fibrinolytic system may vary depending on the genetic background of animals of the same species. For instance, Copenhagen rats were found to have lower fibrinolytic activity than Sprague-Dawley (SD) rats [50]. An intravenous administration of ex vivo formed thromboemboli with tranexamic acid caused a concomitant inhibition of fibrinolysis, resulting in a significant increase in systolic pressure in the right ventricle (RV) (45–55 mmHg) immediately after the intravenous administration of the thromboemboli, followed by normalization of the systolic pressure in the RV within 24 h in SD rats and within 5 days in Copenhagen rats. Lysis of thromboemboli after 24 h and 5 days was 95% and 97% in SD rats, and 70% and 87% in Copenhagen rats, respectively. Thus, even in Copenhagen rats, the degree of thromboemboli lysis was high enough to sustain elevated pressure in the RV for more than 5 days, which leads to the conclusion that such an approach is inappropriate for modeling persistently increased pressure in the RV and in the development of CTEPH.

Although whole blood is most commonly used to form thromboemboli ex vivo, in some cases, only certain components are used, for example, blood plasma from which fibrin clots are formed. Fibrin clots are more suitable for preclinical studies on the effectiveness of new fibrinolytic modifications [38, 42]. Moreover, such thromboemboli can be better standardized in size and provided with isotope labeling, as shown by Murciano et al. [37].

In addition to the introduction of the label, other modifications of thromboemboli can be used to achieve the required properties. For example, to study new approaches to the treatment of PE, which are not associated with an effect on platelets, a model was proposed to modify erythrocytes with chlorpromazine, which has antihemolytic properties, and glutaraldehyde, which causes fixation of erythrocytes [36]. This modification made red blood cells more resistant to lysis.

In mice, rats, and hamsters, this type of model is most often used to test new thrombolytic [35, 37, 38] and anticoagulant [54] substances. A number of studies on changes in the level of paracrine factors in PE have been carried out to reveal the genetic basis of pathological processes that occur in the acute and subsequent periods of PE [33, 48, 53].

In rabbit models, changes in gene expression in the pulmonary artery and lung tissue were studied at different periods after PE [44, 47], changes in hemodynamics during acute PE were assessed [55, 56], drugs were tested for the correction of hemodynamic disorders [46, 57] and new contrast agents were tested for computed tomography [58].

This type of model is characterized by a significantly greater standardization than PE models reproduced by the formation of endogenous thrombi. The use of radioactive labels such as ¹²⁵I-labelled fibrin allows a detailed assessment of the distribution of blood clots in the bloodstream and their fate [35, 37, 38, 42]. Among the disadvantages of these models, it is worth noting the technical complexity and need for repetitive surgical interventions while using autologous blood clots.

2.2. Administration of non-thrombotic particles

Modeling of PE with non-thrombotic particles, consisting of natural or synthetic polymers, is most often performed in rats and rabbits, while the approaches to modeling and goals differ greatly between these species (Table 3).
Table 3. Administration of particles non-thrombotic nature.

| №  | Animal species | Embolizing particles | Material | Size | Dose | Aim of work | Administration vessel | Study duration | CTEPH pathway possibility | Authors |
|----|----------------|----------------------|----------|------|------|-------------|----------------------|---------------|--------------------------|---------|
| 1  | Rats           | Sodium alginate microspheres | 180 ± 28 µm | 9367 ± 551 per animal | CTEPH modelling | Tail vein | 8 times at 4-day intervals | CTEPH + | [93] |
| 2  | Rats           | Polystyrene microspheres | 26 µm | 0,75 ml/kg | Studying of the RAS role in PE severity | Jugular vein | 5 h | – | [124] |
| 3  | Rats           | Microspheres Sephadex G50 | 300 µm | 12 mg/kg | Role of cyclophilin A-CD147 in RV injury and dysfunction after PE | Right femoral vein | 6, 12, 24, 48, and 72 h | – | [125] |
| 4  | Rats           | Polystyrene microspheres | 25 ± 1 µm | 1,3 × 10⁶/100 g of body weight | Studying of HIF-1α expression and its correlation with pulmonary artery and RV remodeling in PE | Right jugular vein | 12 weeks | CTEPH + | [60] |
| 5  | Rats           | Polystyrene microspheres + inhibitor of tyrosine kinase receptor VEGF (SU5416) | 85 µm | 97 × 10³/100 g of body weight | CTEPH modelling | Tail vein | 3 weeks/6 weeks | CTEPH + | [91] |
| 6  | Rats           | Polystyrene microspheres collagen-fibrin coated, suspended in thrombin solution | 45 µm | 3 administrations were done: 1 × 10⁵/1 g of body weight (1st, 2nd administrations), 0,75 × 10³/1 g of body weight (3rd administration) | CTEPH modelling | Tail vein | 9 days after last administration | CTEPH +/-(short observation time) | [61] |
| 7  | Rats           | Polystyrene microspheres | 25 ± 1 µm | 1,95 × 10⁶/100 g of body weight; Administration was done during 10 min | Studying of the stimulant soluble guanylate cyclase effect on RV after pulmonary embolism | Right jugular vein | 5 hours/18 h | – | [63] |
| 8  | Rats           | Polystyrene microspheres | 25 µm | 1,8 × 10⁶/100 g of body weight | Studying of PE pathogenesis | Right jugular vein | 18 h | – | [126] |
| 9  | Rats           | Polystyrene microspheres | 25 µm | 1,8 × 10⁶/100 g of body weight | Studying of the increased expression of type 2 arginase role on vasoconstriction in pulmonary embolism | Right jugular vein | 18 h | – | [65] |
| 10 | Rats           | Polystyrene microspheres | 25 µm | 1,3 or 1,6 × 10⁶/100 g of body weight | Investigation of the soluble guanylate cyclase stimulator effect in pulmonary embolism | Right jugular vein | 5 h | – | [127] |
| 11 | Rats           | Polystyrene microspheres | 26 µm | 1,5 mL/kg | Studying of the Rho-kinase role in PE pathogenesis | RV | 6 h | – | [128] |
| 12 | Rats           | Polystyrene microspheres | 25 ± 1 µm | 1,3 or 1,65 or 2,0 × 10⁶/100 g of body weight | Studying of the plasma proteins relation with the development of pulmonary hypertension after pulmonary embolism | Jugular vein | 18 h | – | [66] |

(continued on next page)
| №  | Animal species | Embolizing particles | Aim of work | Administration vessel | Study duration | CTEPH pathway possibility | Authors |
|----|----------------|----------------------|-------------|----------------------|---------------|--------------------------|---------|
| 13 | Rats           | Polystyrene microspheres | 24 ± 1 μm, 2.0 × 10³/100 g of body weight | Investigation of inflammation in case of RV injury in PE | Right jugular vein | 1 day–6 weeks | – [62] |
| 14 | Rats           | Microparticles Sephadex G50 | 300 μm, – | Investigation of atorvastatin effects in PE | Tail vein | 24 h | – [72] |
| 15 | Rats           | Polystyrene microspheres | 25 ± 1 μm, 2.08 × 10⁶/100 g of body weight | Investigation of inflammation in case of RV injury in PE | Right jugular vein | 20 h | – [64] |
| 16 | Rats           | Microparticles Sephadex G50 | 300 μm, 9 mg/kg | Investigation of doxycycline hemodynamic effects in PE | Left femoral vein | 1 h | – [71] |
| 17 | Rats           | Polystyrene microspheres | 24 ± 1 μm, 0.05 ml/100 g of body weight | Studying of the thromboxane synthase and cyclooxygenase inhibition effects in pulmonary embolism | Left jugular vein | 17 h | – [129] |
| 18 | Rats           | Polystyrene microspheres | 24 ± 1 μm, 1,3/1,95 (× 10⁶/kg) | Studying of pulmonary embolism and following pulmonary hypertension binding, associated with the development of asptic inflammation | Jugular vein | 18 h | – [59] |
| 19 | Rats           | Latex microspheres | 24 ± 1 μm, 0.125 ml per 100 g of body weight | Repeated administrations were done with 1 min interval till aimed pressure in pulmonary artery | Left jugular vein | 40–45 min | – [68] |
| 20 | Rats           | Latex microspheres | 25.7 ± 5.8 μm, – | Development of PE model with a controlled increase in pulmonary artery pressure | Pulmonary artery | 24 h | – [67] |
| 21 | Rats           | Carbon fiber composite microspheres | 15 ± 5 μm, 0.013/0.01 g per 100 g of body weight | Studying of myocardium ultrastructural changes in the early stages after PE | Subcutaneous vein | 1 h/24 h for doses 0.013/0.01 g per 100 g of body weight, respectively | – [69] |
| 22 | Rabbits        | Graphite microspheres | 5 μm, 1 cubic centimeter (cc.) per pound of body weight | Studying of the sympathetic nervous system role in vasospasm in the acute period of pulmonary embolism | – | 15 min | – [70] |
| 23 | Rabbits        | Gelatine sponge | 4 × 4 × 10 mm or 2 × 4 × 10 mm, 4 pcs | Dual-energy computed tomography testing for pulmonary embolism | Femoral vein | 7 days | – [76] |

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In rats, polystyrene microspheres are most often used as embolizing particles [59, 60, 61, 62]. The standard diameter of microparticles is 23–26 μm [59, 63, 64]. For example, Watts et al. [65] used polystyrene microspheres with a diameter of 25 μm in the form of a 10% suspension (1.3 × 10⁶ microspheres/ml). Microspheres were injected into the right jugular vein at a dose of 1.8 × 10⁶ per 100 g in anesthetized male SD rats weighing 350–400 g. According to the data of wire myography, a significant decrease in relaxation of the vascular rings of the pulmonary artery in response to acetylcholine exposure was observed in experimental animals compared to controls.

A similar protocol was used by Zagorski et al. [66], where male SD rats weighing 375–450 g were injected with a suspension of polystyrene microspheres 25 ± 1 μm in 0.01% Tween-20 to a final concentration of 13 × 10⁶ microspheres per ml. Three different doses of microspheres (1.3 × 10⁶/100 g, 1.65 × 10⁶/100 g, and 2.0 × 10⁶/100 g) were injected into the jugular vein of an anesthetized animal. The study showed a dose-dependent increase in systolic pressure in the RV at 2 h and 18 h post-embolization.

In addition to polystyrene microspheres, latex [67, 68], carbon fiber [69] graphite [70] and Sephadex [71, 72] microparticles have also been used in rats.

In a study by Rieger et al. [67], latex microparticles were used to model PE in rats. Anesthetized male Wistar rats weighing 300–350 g were injected with latex microspheres of 25.7 ± 5.8 μm in diameter through a catheter placed in the pulmonary artery until a mean pulmonary artery pressure of 35 mmHg was reached.

In a number of studies, Sephadex microspheres (Pharmacia Biotech, Germany) were used. They are gel filtration resins obtained by cross-linking dextran with epichlorohydrin [71, 72]. Palei et al. [71] performed experiments on male Wistar rats (240–280 g) by injecting them with Sephadex G50 microspheres at a dose of 9 mg/kg and an embolizing particle size of 300 μm, to simulate PE. The authors indicated in pilot studies that various doses of microspheres were tested in the range of 5–15 mg/kg. The 9 mg/kg dose was chosen because it induced a significant systemic arterial hypotension but did not lead to death. Sixty minutes after embolization, the mean arterial pressure decreased by 25.5 ± 4.4 mmHg.
Models based on the use of non-thrombotic emboli are also used in rabbits. However, the materials used to fabricate emboli in rabbits differ significantly from those used in rats. This difference is in part due to the larger size of the pulmonary artery in rabbits, thus allowing the intravascular delivery and positioning of emboli. For example, intravascular delivery of a silicone balloon resulting in obstruction of a branch of the pulmonary artery is well monitored by X-ray or magnetic resonance imaging. It is often used to investigate new protocols for these diagnostic methods and to test contrast agents [73, 74, 75].

In addition, gelatin sponge has been used as an embolizing particle in rabbits. The size of these embolizing fragments is standardized at 4 × 4 × 10 mm or 2 × 4 × 10 mm [76, 77, 78, 79, 80].

In a study by Zhang et al. [80] while testing the method of dual-energy lung computed tomography for the diagnosis of PE, a model of embolization of the pulmonary artery branches in rabbits using a gelatin sponge was used. New Zealand white rabbits weighing 2–3 kg were used in this study. Four emboli from a gelatin sponge measuring 4 × 4 × 10 mm or 2 × 4 × 10 mm were injected into the right femoral vein of each rabbit. Two hours after embolism, emboli in the lungs were detected in 18 out of 20 rabbits in the experimental group. In the remaining two rabbits, embolizing particles were found in the inferior vena cava.

Thus, rats and rabbits are the main species in which models with non-thrombotic particles are used. Most of the studies simulating PE in rats were devoted to the study of cardiac and vascular remodeling after PE, as well as drugs for the treatment of PE, which did not target blood coagulation or fibrinolytic systems. A specific niche for the use of this type of model in rabbits is the study of X-ray and magnetic resonance images and testing of new contrast agents.

The main advantages of models using non-thrombotic particles are high standardization and repeatability of the results obtained, and simplicity of modeling. However, artificial particles cannot fully simulate important processes occurring during thromboembolism, such as partial or complete biodegradation and release of biologically active substances: endothelin, catecholamines, platelet-derived growth factor, fibrin degradation products.

### 2.3. Surgical techniques for modeling of PE

Pulmonary artery occlusion can be achieved not only by embolization with thrombotic or non-thrombotic particles, but also by using surgical techniques. A number of studies have described the ligation of one of the pulmonary arteries as a model of PE [81, 82, 83] (Table 4). Since the left lung in the rat consists of only one lobe (in contrast to the four lobes in the right lung) and is accompanied by a significantly lower perfusion volume, the left pulmonary artery is preferred for modeling. This is also facilitated by the more convenient surgical access to the left pulmonary artery.

All studies with this type of model were conducted in rats. This model has a number of significant drawbacks: it poorly reflects the pathophysiology of the thromboembolic process and is irreversible. The experiment with the pulmonary artery ligation also requires specific surgical experience. In this regard, the main purpose of using this type of model is to study the remodeling of the RV after PE [81].

### 3. Modeling of CTEPH

CTEPH is one of the most severe complications of PE and is characterized by a persistent increase in pulmonary artery pressure caused by partial or complete lack of thromboemboli lysis. The high fibrinolytic ability of rodent blood plasma significantly complicates the process of CTEPH modeling. Therefore, in addition to PE, factors leading to CTEPH should be modeled using repeated injection of embolizing particles, stabilization of natural thromboemboli to prevent their lysis, suppression of neoangiogenesis and stimulation of aseptic inflammation.

Most researchers agree that the isolated use of native thrombi prepared in vitro in healthy small laboratory rodents cannot cause the formation of CTEPH [50, 84] because the blood plasma of rodents has great fibrinolytic activity.

Nevertheless, in some studies, the isolated use of thrombi or in combination with tranexamic acid provided the necessary effect in the form of a prolonged increase in pulmonary artery pressure. For example, in a study by Zhou et al. [85] a single in vitro intravenous injection of prepared autologous thrombi into SD rats led to an increase in the mean pulmonary artery pressure on the 28th day after modeling. Before administration, thrombotic masses were treated for 10 min over a water bath at 70°C; the size of the embolizing particles was 1.1 mm × 2.0 mm. The effectiveness of tranexamic acid as a fibrinolysis inhibitor for stabilizing exogenously introduced thrombi and the formation of CTEPH is controversial, with some studies confirming its value. For example, a series of studies demonstrated the possibility of using this approach for CTEPH modeling [53, 86, 87]. Experiments were performed on SD rats. Blood was collected from the orbital vein using a tube with an inner diameter of 1 mm. Thrombotic masses were formed at room temperature for 24 h. Cylindrical autologous blood clots were cut to achieve 3-mm length thrombi and suspended in saline containing tranexamic acid at a dose of 200 mg/kg of animal body weight. The suspension was injected into the jugular vein of the anesthetized animal at the rate of 0.2 ml/min. The second injection was performed 4 days after the first injection. Hemodynamic analysis, molecular genetics, and histological studies were carried out at 1, 2, and 4 weeks after modeling. There was a significant increase in the level of mean pulmonary artery pressure and pulmonary vascular resistance in comparison with healthy animals, and a significant increase in these parameters over time.

Li et al. [88] also used a combination of exogenously prepared autologous blood clots with tranexamic acid. During the simulation, rats received three intravenous injections of autologous thromboemboli at 2-week intervals. Each injection included the administration of fifteen 3 × 1 mm embolizing particles suspended in 2 ml of saline at an administration rate of 0.5 ml/min. During the entire simulation period, tranexamic acid was injected intraperitoneally at a dose of 12.5 mg/kg per day. Four weeks after modeling according to the authors’ data, there was a significant increase in systolic pressure in the RV according to cardiac catheterization data, as well as an increase in the ratio of the RV area to the area of the left ventricle according to histological examination.

A similar protocol was used in the previous study by Runyon et al. [50], in which thromboemboli were prepared in vitro in combination

### Table 4. Surgery methods of PE modeling.

| No. | Animal species | Surgery procedure | Aim of work | Study duration | CTEPH pathway possibility | Authors |
|-----|----------------|-------------------|-------------|---------------|--------------------------|---------|
| 1   | Rats           | Permanent ligation of left pulmonary artery | Studying of pulmonary artery occlusion effect on right ventricular afterload | 10 min | – | [81] |
| 2   | Rats           | Permanent ligation of left pulmonary artery | Studying of CTEPH development mechanisms | 2 weeks/5 weeks | CTEPH + | [83] |
| 3   | Rats           | Permanent ligation of left pulmonary artery | Modeling post-obstructive pulmonary hypertension | 2 weeks | CTEPH +/– (short observing time) | [82] |

Note: CTEPH — chronic thromboembolic pulmonary hypertension.
with tranexamic acid and did not result in a persistent increase in systolic pressure in the RV in two rat strains. This was associated with rapid lysis of the thromboemboli. Thus, CTEPH models based on single or multiple administrations of prepared ex vivo thromboemboli in combination with tranexamic acid in some cases can give the desired result, although this has not been confirmed by all authors who have used this technique.

Recurrent mechanical obstruction of pulmonary artery branches with thromboemboli is typically considered to be a major mechanism of CTEPH development. However, other mechanisms are also discussed. In particular, chronic vascular inflammation in genetically predisposed individuals might contribute to extensive remodeling of pulmonary vascular bed and increased pulmonary vascular resistance [89]. In one of the studies, an attempt was made to reproduce CTEPH by repeated administration of autologous thrombi in combination with tranexamic acid against the background of a systemic inflammatory response induced by the administration of carrageenan at a dose of 20 mg/kg [90]. In support of the inflammatory theory of CTEPH pathogenesis, the authors observed a more significant increase in pulmonary artery pressure and a more intense remodeling of the pulmonary arteries in the group of animals treated with carrageenan.

As discussed above, blood plasma in rodents is characterized by an outstandingly high fibrinolytic activity, which, despite several previously described successful studies, decreases the utility of thromboembolic models of CTEPH. Therefore, non-thrombotic emboli are much more commonly used for CTEPH modeling. For example, Liu et al. [60] intravenously administered polystyrene microspheres of 25 ± 1 μm in size at a dose of 1.3 × 10^7/100 g body weight in rats. At subsequent observation points (3 days and 1, 2, 4, 8, and 12 weeks), there was a consistent increase in the mean pressure in the pulmonary artery, the relative thickness of the media in the distal branches of the pulmonary artery, and the severity of RV hypertrophy.

Arias-Loza et al. [61] used embolizing microspheres with a complex structure; a fibrin-collagen coating was applied to polystyrene microspheres with a diameter of 45 μm to ensure better adhesion between the particle surface and vascular endothelium. The resulting microspheres at a dose of 1 × 10^8 microspheres/g of animal weight and thrombin at a dose of 0.0027 U/μl were injected into the tail vein of anesthetized male Wistar rats. With a one-week interval, the injection of the embolizing mixture was repeated twice, and during the last injection, the dose of microspheres was reduced to 750 microspheres/g of animal weight. Transthoracic echocardiography (TTE) was performed 7 days after the last injection of microspheres, and the hemodynamics of the right heart chambers were studied and histological examination was performed after 9 days. The authors noted a significant increase in systolic pressure in the RV compared with the control, an increase in the level of brain natriuretic peptide, and a decrease in RV systolic function according to TTE data. It should be noted that the presented model reflects pulmonary hypertension against the background of the subacute stage of PE. A longer observation period is required for final CTEPH formation.

Another hypothesis of CTEPH pathogenesis suggests that a decrease in the intensity of angiogenesis after embolization is of great importance in its formation. Normally, the activation of angiogenesis after repeated episodes of embolism can play an important compensatory role since it leads to a decrease in pulmonary vascular resistance. In the study, it was difficult to ensure decreased angiogenesis; however, a solution was found in a study by Neto-Neves et al. [91], in which CTEPH was modeled against the background of administration of the tyrosine kinase blocker SU5416 to rats, which is capable of suppressing the postreceptor signaling of vascular endothelial growth factor. In this study, modeling was carried out on male SD rats weighing 420 g. It was observed that the combined single intravenous administration of 85 μm polystyrene microspheres at a dose of 97 000/100 g of body weight and SU5416 lead to a stable increase in systolic pressure in the RV within 6 weeks after embolization. Attention was drawn to the fact that the isolated use of microspheres at the indicated dose did not lead to a significant increase in systolic pressure in the RV.

Frey et al. [92] indicated a possible approach to modeling PE with the further development of CTEPH. This study examined the role of splenectomy in the development of CTEPH. In the course of the study, splenectomy was performed in anesthetized mice, and 1 month after that, a ligature was applied to the inferior vena cava below the left renal vein, causing narrowing of the vessel (residual blood flow corresponded to the 5-0 prolene thread diameter). On the 28th day, the thrombi that formed below the ligature in the group without splenectomy had almost completely resolved, while thrombus lysis in splenectomized animals was significantly delayed. The authors pointed out that the increase in thrombus volume after splenectomy was associated with platelet activation, and the subsequent delay in thrombus resolution was due to inhibition of thrombus recanalization. Thus, although this study did not directly simulate PE, it disclosed a possible approach for the development of new models, reflecting one of the particular clinical variants for CTEPH formation.

Considering the clinical scenario of CTEPH, one may conclude that it is critical to ensure a delayed but controlled pattern of degradation of emboli after repetitive embolic occlusion of segmental/subsegmental branches of the pulmonary artery to develop a successful model of CTEPH. This goal has not been completed by previous studies [93] because the majority of thrombotic emboli are dissolved soon after their administration, while synthetic emboli, such as polystyrene, are not degradable. To circumvent this problem, we suggest the use of partially biodegradable sodium alginate microspheres for modeling CTEPH. Sodium alginate is a natural biopolymer derived from brown algae with high biocompatibility. In previous studies, this polymer was actively used to encapsulate cells of various origins, including beta cells of the Langerhans islets [94] and stem cells [95] with their subsequent implantation into the body. In our study, acellular sodium alginate microspheres have been repeatedly administered in the tail vein in rats for induction of CTEPH. It has been shown that 6 weeks after the last injection of microspheres, there was a decrease in exercise tolerance according to the treadmill test, a significant increase in systolic pressure in the RV and endothelin-1 levels, enlargement of the pulmonary trunk and outflow tract of the RV according to TTE, and remodeling of the pulmonary artery branches according to the histological examination data (Figure 1).

Due to the partial and controlled biodegradation of microspheres and high biocompatibility, sodium alginate is a promising material for the manufacture of embolizing particles in CTEPH simulation. An additional advantage of using sodium alginate microspheres is the possibility of encapsulating a thrombogenic mass with the effect of a delayed release of biologically active substances secreted by platelets (serotonin, beta-thromboglobulin, thrombospordin, calcium ions, etc.) and fibrin degradation products, which makes it possible to further improve the compliance of pathogenesis with simulated pathology.

In a study, ligation of the left pulmonary artery was used as an approach to CTEPH modeling [83]. The authors noted a stable increase in systolic pressure in the RV at 2 and 5 weeks after ligation. However, it should be pointed out that this modeling approach seems to be overly mechanistic and reflects only the reduction of the vascular bed as one of the links in CTEPH pathogenesis, without taking into account the molecular mechanisms of vasoconstriction mediated through endothelial dysfunction and aseptic inflammation.

Significant obstruction of the branches of the pulmonary artery in PE is accompanied by high mortality due to acute afterload on the RV. In many patients, CTEPH is formed with recurrent PE, which leads to a significant reduction in the vascular bed of the pulmonary circulation; however, it allows the RV to adapt to a gradually increasing afterload.

In experimental CTEPH modeling, the repeated injection of embolizing particles was also often used, which increased the total volume of embolized vessels and made it possible to reduce mortality due to a gradual increase in right ventricular afterload [53, 61, 88, 93]. The frequency of administration in this approach varied from 2 [53,87] to 8 [93]. In addition, to decrease the acute afterload on the RV in PE and
prevent the development of acute heart failure, a low injection rate of embolizing particles was used [88].

Thus, at the moment there is no universally accepted CTEPH model. However, despite the complexity of modeling this pathology, over the past ten years, several successful models have been developed, which can be used for preclinical testing of new drugs and for studying the pathogenesis of this pathology. The most common features of CTEPH models are a long observation period (usually more than 4 weeks), repeated administration of embolizing particles, the use of artificial particles resistant to fibrinolysis, or active suppression of the animal’s natural fibrinolytic system.

3.1. Species-dependent differences in autonomic innervation of pulmonary vessels and their impact on PE/CTEPH modeling

When choosing the optimal PE model for a number of tasks, it is necessary to consider the specific features of the structure and regulation of the vascular bed of the pulmonary circulation. The most significant interspecies differences are related to innervation.

The pulmonary vascular bed of all mammals is innervated by three types of nerve fibers: sympathetic, parasympathetic, and sensitive. Compared to other organs, the density of innervation and reactivity to vasoactive substances of neuronal origin is highest in large vessels and decreases towards the periphery. In rats and mice, sympathetic and parasympathetic efferent perivascular axons barely reach the hilum. However, in guinea pigs and rabbits, as in humans, this innervation extends to small intrapulmonary vessels [96, 97].

Thus, rabbits and guinea pigs are the preferred species from those considered in this review for studying the autonomic nervous system. However, one should also consider the complexity of modeling certain variants of PE and especially CTEPH in these species. This requires an individual approach when choosing the type of animal for a particular study.

3.2. Evaluation methods used to describe the PE and CTEPH models

Today, taking into account the rapid development of diagnostic methods, there are many methods of verification and assessment of hemodynamics and molecular genetic abnormalities in the experimental modeling of PE and CTEPH. The entire set of assessment methods can be roughly divided into functional, morphological, molecular biology, and biochemistry. In addition, time-based survival estimates have been used in acute models [9, 17, 28] (see Table 5).

The choice of a set of diagnostic methods for a particular study depends on a wide range of parameters: the purpose of the study, the type of animals, the selected model of PE and CTEPH, and the objective capabilities of the laboratory.

The gold standard for determining hemodynamic changes after PE is catheterization of cardiac cavities with manometry; it is more often used
in rats [83, 93] and rabbits [55]. A possible expanding option for cardiac catheterization is the use of catheters with combined pressure-volume transducers, allowing the construction of a pressure-volume-loop [98]. Telemetry technologies are used to continuously record hemodynamic parameters [91].

The use of isolated heart methods [62, 91] and wire myography of the branches of the pulmonary artery [65] allows a detailed study of the functional activity of the structure under study, excluding the influence of external factors.

An important study that allows the assessment of exercise tolerance, thereby determining the functional class of heart failure, is the treadmill test [93]. Most research using this method has been conducted in rats.

Histological examination of the lungs and heart is an integral part of the overwhelming majority of studies. This allows for the most accurate description of morphological changes that occur after PE. In addition to the standard staining with hematoxylin and eosin, dyes specific to connective tissue are also often used in chronic studies: Masson’s trichrome [91], Picrosirius red [61, 62], and Azan according to Heidenhain [93].

Immunohistochemistry makes it possible to clarify the localization of PE models might be recommended for the studies aiming preclinical testing of anticoagulants, antiplatelet agents, and fibrinolytics: i) administration of in vitro prepared blood clots in rats; ii) pharmacological induction of thrombi formation in mice. The use of artificial particles in this case is impossible, and the use of rabbits in most of these studies does not meet the considerations of humanity and economic feasibility.

However, in the study of hemodynamic effects, without considering the reasons for the formation of thromboembolism, it is advisable to use artificial particles. Models of this type are easy to perform and provide a more stable hemodynamic result.

Owing to larger size and better spatial resolution, rabbit models of PE seem to be more suitable for testing new diagnostic X-ray- and magnetic resonance-based approaches. In these models, a prominent branch of pulmonary artery is either embolized by in vitro prepared blood clots in rats; ii) pharmacological induction of thrombus formation in mice. The use of artificial particles in this case is impossible, and the use of rabbits in most of these studies does not meet the considerations of humanity and economic feasibility.

CTEPH modeling is unique and complex because of insufficient understanding of the pathogenesis of the disease and the significant fibrinolytic activity of rodent plasma. Models used to reproduce CTEPH should be characterized by a persistent increase in pulmonary artery pressure and a stable reduction of the vascular bed due to emboli. The observation period in most of the analyzed publications was more than 4 weeks, which confirmed the stability of the reproduced changes in the models used. An important criterion confirming the success of modeling this pathology is a persistent increase in pulmonary vascular resistance, as well as hypertrophic and fibrotic changes in the pulmonary vessels and right ventricle according to histological examination data.

### Summary of methods used to describe the PE and CTEPH models.

| Functional measurements | Morphological measurements | Molecular biology and biochemistry |
|-------------------------|---------------------------|----------------------------------|
| Exercise tolerance testing (treadmill-test) | Histopathological/immunohistochemical examination: | Serum NT-proBNP |
| Heart catheterization and pressure measurement: | - Lung | Transcriptomic analysis of the heart and lung |
| - RV | - heart | - Serum levels of cytokines and growth factors with ELISA |
| - LV | - Computed tomography with contrast enhancement | - Analysis of cytokine and growth factor expression in the heart and lung: |
| - PA | - Perfusion scintigraphy | - Western blotting |
| - Left atrium | - Pressure-volume (PV) loop | - RT-PCR |
| - Wire myography | - Magnetic resonance imaging | - ELISA (from tissue extracts) |

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### 4. Conclusion

Based on analyses of the publications describing PE modeling presented in this article, some key conclusions can be drawn. First, it is noteworthy that there is no single optimal method for modeling PE and CTEPH. The choice of a specific model and animal species is determined based on the objectives of the proposed study. For example, two kinds of PE models might be recommended for the studies aiming preclinical testing of anticoagulants, antiplatelet agents, and fibrinolytics: i) administration of in vitro prepared blood clots in rats; ii) pharmacological induction of thrombi formation in mice. The use of artificial particles in this case is impossible, and the use of rabbits in most of these studies does not meet the considerations of humanity and economic feasibility.

However, in the study of hemodynamic effects, without considering the reasons for the formation of thromboembolism, it is advisable to use artificial particles. Models of this type are easy to perform and provide a more stable hemodynamic result.

Owing to larger size and better spatial resolution, rabbit models of PE seem to be more suitable for testing new diagnostic X-ray- and magnetic resonance-based approaches. In these models, a prominent branch of pulmonary artery is either embolized by artificial particle(s) or occluded by the inflation of intraluminal balloon.
Further improvement of experimental models of PE and CTEPH is critical for a deeper understanding of the pathogenesis of these disorders and for the preclinical development of new drugs to prevent complications and reduce morbidity and mortality.

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