Basic models and approximation for the engineering description of the kinetics of the oxide layer of steel in a flow of heavy liquid metal coolant under various oxygen conditions

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Abstract

The article presents the results of corrosion processes, kinetics and changes in the oxide layer modeling using MASKA-LM software complex. The complex is intended for a numerical simulation of three-dimensional non-stationary processes of mass transfer and interaction of impurity components in a heavy liquid metal coolant (HLMC: lead, lead-bismuth). The software complex is based on the numerical solution of coupled three-dimensional equations of hydrodynamics, heat transfer, formation and convective-diffusive transport of chemically interacting components of impurities.

Examples of calculations of mass transfer processes and interaction of impurity components in HLMC, formation of protective oxide films on the surfaces of steels are given to justify the coolant technology.

Keywords
coolant, corrosion, films, HLMC, heavy liquid metal coolant, impurities, lead, lead-bismuth, mass transfer

Introduction

With the advent of liquid metal coolants necessary for nuclear power plants with fast neutron reactors, the key elements of the safety justification were attributed to the problems associated with maintaining the quality of the coolant. Maintaining the necessary physicochemical regime of the coolant (or coolant technology) has turned from an important but additional operation into one of the main components in substantiating the design and operating parameters of the reactor. There are two main tasks of the heavy liquid metal coolant (lead or lead-bismuth) technology:

• ensuring the cleanliness of the coolant and the surfaces of the circulation circuit to maintain the design thermohydraulic characteristics with long service life (several decades when the reactor plant operates at a power of up to 100%);
• prevention of corrosion and erosion of structural materials with long service life (several decades, when the reactor plant is operating at a power of up to 100%).

It should be noted that the main processes of the lead-bismuth and lead coolant technology have a common physicochemical nature. In the lead-bismuth eutectic, lead exhibits the dominant chemical activity. Considering also that the construction materials for both coolants are identical, the methods and means of the coolant technology are also similar in the principle of operation.

Therefore, understanding the process of generation of the oxide layer of steel in a flow of heavy liquid metal coolant is highly important for justification of circuit integrity and reactor safety.

Kinetics of corrosion processes of steel components in HLMC is determined by the associated processes of hydrodynamics and the interaction of dissolved components, primarily such as oxygen and iron. The justification and appropriate modeling of these processes is an important component for substantiating heavy liquid metal coolant technology (Robertson et al. 1988; Fazio et al. 2001; Aiello et al. 2004; Ivanov et al. 2005, 2017; Zhang et al. 2005, 2007; Lim et al. 2010; Martinelli et al. 2008a; Weisenburger 2008; Schroer and Konys 2009; Steiner 2009; Hwang and Lim 2010; Askhadullin et al. 2011; Schroer et al. 2012; Zhang 2013; Nuclear Energy Agency 2015; Tsisar et al. 2017; Salaev et al. 2018).

The MASKA-LM software complex implements basic models and approximations with the following features:

• modeling of the distribution of dissolved impurities along the circulation path of lead coolant in various operating modes of a reactor unit, taking into account the condition of the surfaces of structural steels;
• simulation of a transient distribution of impurities when changing operating modes;
• predicting the deviation of the quality indicators of lead coolant from normal operating conditions for various initiating events;
• simulation of the distribution of the gas phase in lead coolant;
• simulation of the transfer of solid-phase impurities along the circulation circuit.

The use of commercial CFD codes in this area has not yet been widely applied because of the need to build a complex apparatus of user functions for coordinated calculations of thermal hydraulics and processes of interaction and formation of impurities. For example, the works (Marino 2015; Marino et al. 2018) on modeling physicochemical processes in lead coolant show the possibility of using this approach. However, even for the simplest cases (individual sections, areas, etc.), the dimension of computational grids reaches excessively large values at which the use of only a CFD code will require significant machine resources. At the same time, the general accuracy of the calculations is mainly determined by the accuracy of the equations of physicochemical processes. “Massive” calculations of three-dimensional non-stationary distributions of impurity components in the primary circuit in lead coolant at “life cycle” mode using a CFD code with fine grids are problematic due to unreasonable calculation time, and the use of coarse grids eliminates the “hydrodynamic” advantages of CFD.

Mathematical model of the processes of mass transfer and physico-chemical interaction of impurities in the volume of the coolant and on the surfaces of structural steels

A system of three-dimensional equations of an incompressible multicomponent medium in a Cartesian coordinate system was used to describe the processes of formation, transformation, and transfer of impurities in the primary circuit of a reactor with HLMC (Alekseev et al. 2010a). The system includes hydrodynamic equations in the Boussinesq approximation, equations of conservation and transfer of thermal energy, equations of physicochemical kinetics of impurities (Kumaev et al. 2002), equations of conservation and transfer of components of a multicomponent medium. The system of equations is solved by finite-difference methods.

The medium is assumed to be incompressible with temperature-dependent thermophysical properties. The presence of dissolved and dispersed particulate impurities does not affect the hydrodynamics, heat transfer, and thermophysical properties of lead. Concentration distributions of dissolved impurities and particles are affected by molecular, Brownian, and turbulent diffusion. On the surfaces of structural materials, oxide films are formed that impede the diffusion exchange of oxygen and iron between the coolant and structural materials.

The system of equations describing the non-stationary thermohydrodynamics of the components of a multicomponent medium is presented in the form of equations of motion, conservation of mass and thermal energy of an inhomogeneous continuous medium (Kumaev et al. 2002, 2005; Alekseev et al. 2010a, b, c). The components of the medium are fuel, structural and other materials that are in a solid state and do not move in space, liquid media that can move, as well as components of various impurities that undergo transformations in physicochemical reactions. The viscosity and thermal conductivity coefficients of a liquid are presented as the sum of molecular and turbulent components.

The equations of motion for the characteristics of a multicomponent medium averaged over components that are not impurities were obtained in (Kumaev et al. 2002, 2005) and in the Cartesian coordinate system in the Boussinesq approximation and have the following form:
\[
\frac{\partial U_i}{\partial \tau} + \frac{\partial U_i}{\partial x_i} = -\frac{1}{\rho} \frac{\partial \rho}{\partial x_i} - \frac{1}{\rho} \left( \frac{\partial}{\partial x_i} \mu \frac{\partial U_i}{\partial x_i} \right) + g_i \rho \frac{\partial t}{\partial x_i} - \Lambda \frac{\partial U_i}{\partial x_i}, \quad (1)
\]

where: \( U_i \) – projections (components) of the medium velocity vector in the Cartesian coordinate system; \( \rho \) – density of medium averaged over temperature and components; \( \mu \) – dynamic effective viscosity coefficient; \( \rho t \) – averaged only over components and temperature-dependent density of the medium; \( g_i \) – projections of the gravitational acceleration vector on the coordinate axes; \( \Lambda \) – tensor of drag coefficients.

The drag coefficients of the medium are also used to simulate the solid and liquid states of the medium. Zero values of the resistance coefficients correspond to the liquid state of the components; large values (tending to infinity) correspond to the solid state of the components.

Environmental properties are obtained by averaging the initial properties of the components and are determined by the following dependencies:

\[
\rho = \sum \rho_i \varepsilon^*_n; \quad \rho_i = \text{density of a component;}
\]

\[
\mu = \sum \mu_i \varepsilon^*_n; \quad \mu_i = \text{n component dynamic viscosity coefficient;}
\]

\[
U_i = \sum u_i^* \varepsilon^*_n \quad \text{– projections of the velocity vector of the averaged motion of the medium;}
\]

\[
\varepsilon^*_n = \frac{U_i}{u_i} \quad \text{– volume fraction of component with number n;}
\]

\[
u_i^* = U_i + \Delta u_i^* \quad \text{– projections of the velocity vector of the medium component with number n.}
\]

The drag coefficients of the medium \( \Lambda \) are used to simulate the solid and liquid states of the medium. For components in the liquid state, \( \Lambda \) is set equal to zero, for components in the solid state it is set to a larger value, which ensures that the velocity of the medium is equal to zero.

The equations of transport of non-impurity components of the medium, taking into account stratification and taking into account the sources of components that can occur during chemical reactions, have the following form:

\[
\frac{\partial \varepsilon^*_n}{\partial \tau} + \frac{\partial (\varepsilon^*_n U_i)}{\partial x_i} = -\frac{\partial}{\partial x_i} (\varepsilon^*_n \Delta u_i^*) + \sum_{j=1}^{N} J^m, \quad (2)
\]

For components that are impurities, diffusion processes are additionally taken into account in the transport equations:

\[
\frac{\partial C^m}{\partial \tau} + \frac{\partial}{\partial x_i} (C^m U_i) = \frac{\partial}{\partial x_i} (C^m \Delta u_i^*) + \frac{\partial}{\partial x_i} D^m \frac{\partial C^m}{\partial x_i} + \sum_{j=1}^{N} J^m, \quad (3)
\]

where \( \Delta u_i^* \) – components of the velocity vectors of the relative motion of the components of the medium (stratification velocity); \( \varepsilon^m \) – volume fraction of component \( n \), that is in a liquid state; \( J^m \) – source of component \( n \) due to loss of component \( p \); \( \hat{N}^m \) – mass concentration of the impurity component \( n \), that is in the liquid; \( G^m \) – mass source of the impurity component \( n \) due to a decrease in the impurity component \( p \); \( D^m \) – component \( n \) diffusion coefficient.

The left-hand sides of equations (2.3) describe the transfer of medium components due to averaging motion. The right-hand sides of equations (2) describe the process of separation of a mixture of components in a liquid state, sources of components due to chemical reactions or other processes, as well as diffusion processes of component transfer. Solid components do not delaminate. The initial and boundary conditions that describe the flows of impurities onto solid surfaces, as well as the values of the concentrations of impurities at the input, are added to the transport equation.

Modeling of heat and mass transfer and heat exchange processes in a multicomponent medium taking into account stratification is carried out by solving the energy equation in the following form:

\[
\frac{\partial (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda)}{\partial \tau} + \frac{\partial}{\partial x_i} (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda U_i) = \frac{\partial}{\partial x_i} (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda U_i) + \frac{\partial}{\partial x_i} (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda U_i) + \sum_{j=1}^{N} \frac{\partial}{\partial x_i} (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda \Delta u_i^*) + \frac{\partial}{\partial x_i} (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda \Delta u_i^*) + \sum_{j=1}^{N} \frac{\partial}{\partial x_i} (\varepsilon^*_n \rho \mu \tau \rho \varepsilon^*_n \lambda J^m), \quad (4)
\]

where: \( c \) – the specific heat of the medium averaged over the components; \( \lambda \) – thermal conductivity coefficient averaged over the components of the medium; \( U_i \) – medium volume velocity; \( \Delta u_i^* \) – projections of the component stratification velocity vector \( n \); \( \varepsilon^m \) – volume fraction of a component \( n \) in a liquid state.

The first two terms on the left side of equation (4) describe the convective transfer of thermal energy. The third term is the transfer of thermal energy due to the separation of components.

**Description of the processes of physicochemical interaction of impurities in the coolant volume and on the surfaces of structural steels**

Mass transfer processes in nonisothermal circuits with HLMC are closely related to the interaction of the coolant with structural steels. Structural steels are the main sources of metallic impurities entering the coolant (iron, chromium, nickel, etc.). In turn, the coolant is a supplier of dissolved oxygen for structural steels, which is necessary for the formation of protective oxide films on steel surfaces that prevent the development of corrosion processes.
Dissolved oxygen in the coolant is both in free form, and in the form of various compounds with a coolant and metal impurities. Moreover, in quantitative terms, free oxygen in the solution is negligible relative to the bound forms of dissolved oxygen.

The main metal impurity, which should be taken into account when calculating the simulation of mass transfer processes in the circulation circuit with HLMC, is an iron impurity. In lead melt, iron can be both free and in oxygen-bound form. The relationship between these forms of existence of dissolved iron in the coolant depends on the oxygen content and can vary widely.

In the model of interaction of dissolved impurities in the coolant, only those forms of existence of oxide compounds that play a decisive role should be taken into account: PbO, Fe₃O₄.

The most important in modeling physicochemical interaction of impurities are the processes of formation and growth of protective oxide films on the surfaces of structural steels. The composition of oxide film, as a rule, varies from chrome spinel near the boundary with the coolant to chromium spinel at the border with steel. In this work, only the process of magnetization film at the “coolant – oxide film” interface was taken into account, and the formation of chromium spinel is not modeled separately, but is completely determined by the model of magnetite film formation. This approach can be considered as simplified, but it gives quite adequate description of the evolution of the oxide film.

**Oxide film growth kinetics model**

Oxidative kinetics leading to the parabolic law of growth of an oxide film is usually used when the effective diffusion of reagents is independent of time and size of the film. The oxidation mechanism of ferritic-martensitic steels is determined by the diffusion of Fe ions in the direction of the coolant and variations in the oxygen potential. In this approach, oxidation of the surface of steels is described by the parabolic velocity equation with the rate constant predicted by Wagner’s theory if the resulting oxide film has an “ideal” crystal lattice, and ion diffusion is dominant.

**General Provisions**

For simplicity, we consider the growth of a single-layer film. In the developed (Kumaev et al. 2002, 2005; Alekseev et al. 2010c) model of the growth of a single-layer film, at the first stage, the iron flux “remaining” in the wall layer is determined as the difference between the fluxes through the oxide film and the flux that has gone into the coolant. The “oncoming” oxygen flow from the coolant is determined according to the stoichiometry of the resulting magnetite.

In this approximation, it is unambiguously assumed that the growth of the film (magnetite) occurs due to the reaction on the surface of the oxide layer:

\[ 3\text{Fe} + 4\text{PbO} \rightleftharpoons \text{Fe}_3\text{O}_4 + 4\text{Pb}, \quad (5) \]

and at the same time, all four components of the reaction are in chemical equilibrium. Since the rate of chemical reactions is much higher than the characteristic rates of all other physical processes, the chemical equilibrium can be considered steady in the volume of the coolant at each moment of time and the distribution of components substantially depends on thermo-hydraulic processes.

Let us consider a model assuming the presence of a diffusion “reaction” layer at the interface between the oxide film and the coolant. The main components interacting with oxygen are iron and lead. We consider the oxide film as a single layer and consisting of magnetite. At the steel-coolant boundary, we distinguish four subregions: steel (conventionally Fe), oxide, an intermediate diffusion layer, and HLMC (Fig. 1).

Intermediate layer is a virtual layer where the above-described surface reaction is carried out (5). For flows of components and sources of impurities in the media presented in Fig. 1, we use the notation: \( J^\beta_{\partial} \) – O (PbO) oxygen flow in the coolant; \( J^\beta_{\partial} \) – Fe oxygen flow in the coolant; \( J^\beta_{\partial} \) – oxygen flow in oxide film; \( J^\beta_{\partial} \) – iron flow in oxide film; \( \varepsilon^\beta_{\partial} \) – oxygen flow (source) in the intermediate layer; \( \varepsilon^\beta_{\partial} \) – iron flow (source) in the intermediate layer; \( \gamma^\beta_{\partial} \) – the amount of oxygen and iron in the intermediate layer, for impurity concentrations \( c_{\partial} \), where \( n = O, \text{Pb}, \text{Fe}, \text{Fe}_3\text{O}_4 \) and activity \( \gamma_{\partial} \), \( C_{\partial} \) – saturation concentration.

The chemical equilibrium of the oxygen-lead system with the formation of the [PbO] complex is described in the standard way:

\[
K^\beta = \frac{a_{\text{PbO}}}{a_{\text{Fe}}} \exp \left( \frac{- \Delta G_{\text{PbO}}}{RT} \right) \quad (6)
\]

Lead activity is equal to one; therefore, this ratio shows the ratio of oxygen concentrations and the dynamic complex of PbO. For the liquid phase, the existence of a separate oxygen phase [O] does not have deep physical meaning, and its concentration is much lower than the concentration of [PbO]. The concentration of [O] can be considered as the concentration of “instantly free” oxygen.

Using the data (Nuclear Energy Agency 2015) for the Gibbs energy parameters for the formation of lead oxide and relation (6), it is easy to estimate that \( a_{\text{PbO}} / a_{\text{O}} \sim 10^4 \), so for oxygen activity it is enough to understand the activity of the complex [PbO].

For HLMC medium with impurities (Pb, PbO, Fe, \text{Fe}_3\text{O}_4) we introduce the notation \( \beta \). In the presence of iron in the coolant, three reactions are formally possible:
\[ Pb + O = PbO \]
\[ 3Fe + 4O = Fe_3O_4 \]
\[ 3Fe + 4PbO = Fe_3O_4 + 4Pb \]  \( (7) \)

Chemical equilibrium constant for the second reaction:
\[ k^y_p = \frac{1}{(a^y_3)^2} \exp\left(\frac{-\Delta G_{PbO}}{RT}\right) \]  \( (8) \)

The chemical equilibrium constant for the third reaction, provided that the activity of magnetite is equal to unity:
\[ k^\gamma_p = (a^\gamma_3)^3 \exp\left(\frac{-4\Delta G_{PbO} - \Delta G_{\gamma}^{\text{m}}}{RT}\right) \]  \( (9) \)

where \( a^\gamma_j \) – activities corresponding to the coolant component.

The equilibrium constants \((6, 8, 9)\) in the dynamic and chemical equilibrium of all components are obviously related:
\[ k^\gamma_p = \left(\frac{k^\gamma_3}{k^\gamma_4}\right)^4 \frac{k^\gamma_5}{k^\gamma_6} \]  \( (10) \)

Obviously, in conditions of dynamic equilibrium, it is sufficient to consider any two of the three equations. From the point of view of the thermodynamics of the lead-iron-oxygen system, it is more convenient to consider the first two equations. And the area of existence of iron oxides of various compositions in terms of oxygen potentials overlaps with the region of existence of oxygen solutions in the lead melt.

Let’s consider the surface layer at the interface of oxide film-coolant (medium \( \gamma \)). As we discussed above, if we formally approach this region as well as the coolant region, then there will be no differences between them and, moreover, there will not even be a need to introduce this region for consideration. Since a possible film growth is assumed in this medium, that is, a kind of phase transition:
\[ Fe_3O_4\text{(solid phase)} + Pb\text{(liquid phase)} \rightarrow \text{dissolved(Fe, PbO) in Pb\text{(liquid phase)}} \]  \( (11) \)
then in this environment we assume the possibility of only the third reaction from \( (11) \).

Equilibrium constant in this environment:
\[ k_j = (a^\gamma_3)^3 \exp\left(-\frac{4\Delta G_{PbO} - \Delta G_{\text{etch}}}{RT}\right) \]  \( (12) \)

Once again, we note that in general case, knowledge of the equilibrium constant is not enough for this medium, since we will assume that the existence of equilibrium is not a prerequisite. In this case, it is necessary to know the parameters of forward and reverse reactions \((7)\). In addition, the third reaction \((7)\), which occurs on the surface of magnetite, is in reality not “one-step” as it is written, but contains steps for adsorption and desorption of the elements involved. Therefore, the reaction constants do not have to be identical with the constants in a homogeneous medium. The oxygen-coolant system is assumed in chemical equilibrium, similarly to formula \((6)\). It should also be noted that the heterogeneous reaction rate is directly proportional to the area of contacting reagents. Therefore, “rough” surface, obviously, will provide a more intensive reaction process. Therefore, oxygen consumption by the steel surface will depend not only on its type, but also on the method of its manufacture. The latter is almost impossible to take into account when modeling, which can significantly reduce the physical justification for the direct application of the Wagner model (Robertson et al. 1988).

Let’s consider the layer of the film itself, which at this stage is assumed to be composed only of magnetite \( Fe_3O_4 \) (medium \( \delta \)). In general, film growth also occurs on the metal-magnetite interface, but since the oxygen diffusion rate is much lower than the rate of the chemical reaction of incorporation of oxygen into the magnetite lattice, it is sufficient to know the concentration at the metal-magnetite interface to calculate the oxygen flux.

The constant of chemical equilibrium in this medium, taking into account the fact that the activity of magnetite and iron are equal to unity:
\[ k^\delta_0 = \frac{1}{(a^\delta_3)^4} \exp\left(-\frac{\Delta G_{PbO} - \Delta G_{\text{etch}}}{RT^3}\right) \]  \( (13) \)

where \( a^\delta_j \) – oxygen activity in the medium \( \delta \), \( T^4 \) – film temperature.

The condition of chemical equilibrium in the intermediate layer during the formation of magnetite:
\[ \frac{V^\delta_j}{V^\gamma_j} = \frac{\partial V^\gamma_j}{\partial V^\gamma_j} = 4 \frac{J^\delta_j}{J^\gamma_j} \]  \( (14) \)

The formulas for the sources of oxygen and iron in the surface layer \( \gamma \), \( J^\gamma_j \) and \( J^\gamma_s \), are proportional to the growth rate (or degradation) of this “reaction” layer, which in turn is proportional to the amount of oxygen and iron substance in it. Since the system in question is assumed in chemical equilibrium, the velocities of direct, \( R^- = K^- (C^\gamma_j)^4 (C^\gamma_s)^4 \) and reverse, \( R^+ = K^+ (C^\gamma_j)^4 (C^\gamma_s)^4 \) reactions are the same and from the chemical equilibrium condition \((12)\) it is possible to determine only the ratio of concentrations and “fluxes” of iron and oxygen. That is, the condition of chemical equilibrium and the assumption of the instantaneous establishment of this chemical equilibrium in comparison with the slow diffusion process does not make it possible to determine the slow growth of the film and additional assumptions about the mechanism of film growth are needed, namely, that the molar fluxes of oxygen and iron are treated as 4/3, in accordance with the kinetics of
reaction (7), with instant subsequent formation of magnetite on the surface.

The source of oxygen in the system is the coolant (oxygen content is regulated through external sources). Therefore, “diffusion” of oxygen from the coolant to the steel-coolant surface section and the formation of a conventional oxide film (magnetite in our model) are initially assumed. That is, oxygen is “consumed” on the growth of the inner and outer parts of the film, respectively:

\[ J_{\delta}^c |_{x \rightarrow \gamma} = J_{\gamma}^c + J_{\delta}^c \]  

Similarly for iron that diffuses from steel:

\[ J_{\delta}^f |_{x \rightarrow \gamma} = J_{\gamma}^f + J_{\delta}^f \]  

Note that the model considered in (Alekseev et al. 2010b, c) is a special case of the proposed model. According to (Alekseev et al. 2010b, c), iron fluxes at the interface can be calculated as follows:

\[ J_{\delta}^f |_{x \rightarrow \gamma} = \chi_{\delta} \left( C_{\delta}^f - C_{\delta}^f \right) \]  

where \( \chi_{\delta} \) - mass transfer coefficient of iron at the border.

Similarly, oxygen flows in media can be represented by \( \delta \) (Alekseev et al. 2010b, c) and \( \beta \) respectively:

\[ J_{\delta}^o |_{x \rightarrow \gamma} = 2D_{\delta}^o \frac{C_{\delta}^o - C_{\delta}^o}{\delta} \]  

where \( \chi_{\delta} \) - oxygen mass transfer coefficient.

Using ratios (16) and (17) we get:

\[ \theta \left( J_{\delta}^c - J_{\delta}^f \right) = J_{\gamma}^c - J_{\delta}^c \]  

where \( \theta = \frac{4M_{\text{Fe}}}{3M_{\text{Fe}}} \) - for molar (mass) flows, respectively, or

\[ \frac{4M_{\text{Fe}}}{3M_{\text{Fe}}} \left( D_{\delta}^o \frac{C_{\delta}^o - C_{\delta}^o}{\delta} - \chi_{\delta} \left( C_{\delta}^f - C_{\delta}^f \right) \right) = \chi_{\delta} \left( C_{\delta}^f - C_{\delta}^f \right) - 2D_{\delta}^o \frac{C_{\delta}^o - C_{\delta}^o}{\delta} \]  

Using relations (17–20) and the relationship between the activities of oxygen and iron in the wall layer \( \gamma \), we can obtain a nonlinear equation for the concentration (or activity) of oxygen in the wall layer in which the concentrations of iron and oxygen, \( C_{\delta}^o \) and \( C_{\delta}^f \), are calculated from equations (3–5). Concentrations \( C_{\delta}^o \) and \( C_{\delta}^f \) are calculated based on the properties of magnetite, namely, the activity of iron in magnetite is assumed equal to one. The activity of oxygen in magnetite is found from the condition of chemical equilibrium (13). Thus, the equation obtained from relation (20) will contain one unknown parameter, namely, the film thickness \( \delta \). Note that the obtained equation for the concentration \( C_{\delta}^o \) (or the activity corresponding to it) parametrically depends only on the concentrations in other regions (\( \beta \) and \( \delta \)), which in turn depend on time. The thickness of the film cannot be expressed only through these parameters (concentration or fluxes), and the film growth will directly depend on time.

**Film growth model within the Wagner approach**

As was determined above, the diffusion processes of iron and oxygen in the oxide film are slow, and therefore the determining processes of its growth. Therefore, formulas (17) and (19) are a rather crude approximation.

According to Wagner’s theory, the flow of iron through the oxide layer as a function of coordinates can be represented (Zhang 2005):

\[ J_{\delta}^f = \frac{C_{\delta}^f D_{\delta}^f}{RT} \frac{d \mu_{\text{Fe}}}{dx} = \frac{1}{2} \mu_{\text{Fe}} D_{\delta}^f \frac{d \ln P_{\text{Fe}}}{dx} = \frac{2}{3} D_{\delta}^f C_{\delta}^f \frac{d \ln P_{\text{Fe}}}{dx} , \]  

where \( \mu_{\text{Fe}} \) - chemical potential of iron, \( P_{\text{Fe}} \) - partial pressure of oxygen, \( x \) - coordinate, for FeO.

The fulfillment of relation (23) gives a constant flow of iron [10]:

\[ J_{\delta}^f = \frac{2}{3} C_{\delta}^f \int_{x_{(\delta)}}^{x_{(0)}} D_{\delta}^f d \ln a_{\delta_{\text{Fe}}} \]  

where \( a_{\text{Fe}} \) - oxygen activity in oxide at a point with a coordinate \( x \), determined as \( a_{\text{Fe}} = \frac{P_{\text{Fe}}}{P_{\text{Fe}}^0} \), where \( P_{\text{Fe}}^0 = 1 \text{ atm.} \), \( c_{\text{Fe}} = \frac{c_{\text{Fe}^0}}{a_{\text{Fe}}} \) - the average density of iron in magnetite, where \( M_{\text{Fe}} = 3M_{\text{Fe}^0} + M_{\text{Fe}} \), \( M_{\text{Fe}^0} \) - molar masses of iron and oxygen, respectively, \( \rho_{\text{Fe}} \) - magnetite density, \( C_{\delta}^f (0) \) - oxygen saturation concentration in magnetite.

Using relation (24) and the dependence of the diffusion coefficient of iron cations, which takes into account the transport of iron ions over vacancies and along the grain boundaries of the metal in the framework of the theory of point defects (Topfer et al. 1995; Zhang 2005; Martinelli et al. 2008a, 2008b; Mikityuk 2010):

\[ D_{\delta} = D_{\text{Fe}} \exp \left( -E_{\text{Fe}} / RT \right) P_{\delta}^{1/3} + D_{\text{Fe}} \exp \left( -E_{\text{Fe}} / RT \right) P_{\delta}^{2/3} , \]  

where \( D_{\text{Fe}} = a_{\text{Fe}} \) - oxygen activity in oxide at a point with a coordinate \( x \), determined as \( a_{\text{Fe}} = \frac{P_{\text{Fe}}}{P_{\text{Fe}}^0} \), where \( P_{\text{Fe}}^0 = 1 \text{ atm.} \), \( c_{\text{Fe}} = \frac{c_{\text{Fe}^0}}{a_{\text{Fe}}} \) - the average density of iron in magnetite, where \( M_{\text{Fe}} = 3M_{\text{Fe}^0} + M_{\text{Fe}} \), \( M_{\text{Fe}^0} \) - molar masses of iron and oxygen, respectively, \( \rho_{\text{Fe}} \) - magnetite density, \( C_{\delta}^f (0) \) - oxygen saturation concentration in magnetite.
$D_c^\delta$ and $D_i^\delta$ – constants characterizing the transport of vacancies and between grains, respectively, we can obtain the value of the iron flux from the layer $\delta$ as a function of oxygen activity at the layer boundary ($x = \delta$):

$$J_c^\delta = \frac{C_c}{\delta} \left[ A \left[ (a_o(\delta))^{3/2} - (a_o(0))^{3/2} \right] - B \left[ (a_o(\delta))^{4/3} - (a_o(0))^{4/3} \right] \right]^{1/3} \left( \frac{(\delta)}{\delta_o} \right)^{2/3},$$

where $A = D_c^\delta \exp(-E_c/RT)$ and $B = D_c^\delta \exp(-E_i/RT)$.

Expression (35) shows that the film thickness at $t \rightarrow \infty$ reaches its limit $\delta_{\infty}$.

Neglecting the oxygen flux in the film, we can obtain the equation of film growth in the Zhang approximation (Zhang and Li 2005):

$$J_r^\delta = \frac{4M_r^o}{3M_r} (J_r^\delta - J_r^\delta) = \frac{4M_r^o}{3M_r} (J_r^\delta - X_r (C_i^\delta - C_r^\delta))$$

Then the film growth equation takes the following form:

$$\frac{d\delta}{dt} = \frac{K_p}{2\delta} \frac{X_r (C_i^\delta - C_r^\delta)}{C_r^\delta}$$

Denote the value $\chi_r \frac{(C_i^\delta - C_r^\delta)}{C_r^\delta}$, which reflects the dissolution rate (corrosion) of the film. Then the equation for changing the film takes an even simpler form:

$$\frac{d\delta}{dt} = \frac{K_p}{2\delta} - \chi_r$$

This equation corresponds to the Tedmon model (Martinelli et al. 2008b). If the parameters of the equation are considered as constants, then the equation has an analytical solution:

$$t = \frac{\delta - \delta_0}{\chi_r t} = \frac{K_p}{2\chi_r} \left[ \ln \left( \frac{2\chi_r t}{K_p} \right) - \ln \left( \frac{2\chi_r \delta}{K_p} \right) \right]$$

where $\delta_0$ – initial film thickness.

Expression (35) shows that the film thickness at $t \rightarrow \infty$ reaches its limit $\delta_{\infty}$.

At $\delta_{\infty} \ll \delta$ and $\frac{(C_i^\delta - C_r^\delta)}{C_r^\delta}$, expression (35) can be approximately written as

$$\delta - \delta_0 = \left( \frac{K_p t}{2\chi_r} \right)^{1/2} - \left( \frac{2\chi_r t}{K_p} \right)^{1/2} - \left( \frac{2\chi_r \delta}{K_p} \right)^{1/2}$$

In this case, we can assume that the parabolic film growth is quite well performed and the film corrosion can be neglected at times $t \ll \frac{2\chi_r \delta}{K_p}$.

**Approximation of a semi-empirical account of diffusion through an oxide film**

The derivation of the film growth rate constant of the parabolic law (31) is based on the model (Topfer et al. 1995), where, when modeling the diffusion of iron cations, the assumption was made that the concentration of vacancies and interstitial ion concentrations are independent of porosity and grain size of magnetite, that is, ion diffusion within the framework of the vacancy model for ideal “magnetite” (Hallstrom et al. 2011). Such models are quite fundamental in nature and allow one to calculate the diffusion coefficient of iron ions, which is necessary for calculating the film growth rate constant in the
framework of the Wagner model. The main result of the model (Topfer et al. 1995) and similar ones (Diekmann 1998) is that in the region of low partial oxygen pressures \( K_p \sim P_{O_2}^{0.16} \). EXPERIMENTAL STUDIES OF FILM GROWTH ON VARIOUS ALLOYS SHOW SLIGHTLY DIFFERENT DEPENDENCES OF THE FILM GROWTH CONSTANT ON THE PARTIAL PRESSURE OF OXYGEN THAN WAS OBTAINED FOR PURE MAGNETITE USING MODELS OF THE TYPE (Topfer 2005; Hallstrom et al. 2011).

Since in reality the film for the overwhelming majority of materials under consideration is not pure magnetite, but is at least two-layer, the models developed in (Topfer 2005; Hallstrom et al. 2011) it’s not necessarily to describe the dependence of the film growth rate constant (conditional name) on the partial oxygen pressure like \( K_p \sim P_{O_2}^{0.16} \). Since in the works (Surman 1973; Smith 1982; Saito et al. 1985; Sato et al. 2002) the dependencies \( P_{O_2}^{0.14} \), \( P_{O_2}^{0.15} \), \( P_{O_2}^{0.14} \) and \( P_{O_2}^{0.27,0.31} \) have been experimentally found for different types of steel correspondingly. These empirical estimates imply the fulfillment of the parabolic law of film growth (31), i.e., without taking into account possible mass transfer processes leading to its dissolution (corrosion). The latter was taken into account in (33) and for large times provides a nonparametric change in the growth of the oxide film. Such a different dependence of the film velocity constant is determined by both the composition of the metal so that by the properties of the grains, which leads to the failure of the theoretical dependence \( K_p \sim P_{O_2}^{0.16} \), characteristic to the “ideal” magnetite (Diekmann 1998). In (Sato et al. 2002), it was assumed that the film growth rate constant is generally determined by the diffusion of chromium ions in chromium oxide, the theoretical dependence of which \( K_p \sim P_{O_2}^{0.16} \), which, in principle, is somewhat closer to the experimental values.

Note that in (Surman 1973; Smith 1982; Saito et al. 1988; Diekmann 1998; Sato et al. 2002) the oxidation of steel surface was carried out at low partial oxygen pressures typical for partial pressures in liquid lead, but in a gaseous medium. Therefore, the processes of film dissolution and erosion, in principle, could not play a significant role. The results of experiments in a stationary coolant (Steiner 2009, Ashkadullin et al. 2011) led to an empirical dependence \( P_{O_2}^{0.125} \) for steels EP823-(Sh) and E302-(Sh). For T91 steel, the available experimental data shows a dependence of the type \( P_{O_2}^{0.11} \).

The parabolic dependence of the film growth (31), in principle, is quite well confirmed experimentally (Aziello et al. 2004), if corrosion-erosion processes have low intensity (for example, in a gaseous medium), and the Wagner approach is currently the only reasonably substantiated approximation. Deviations from the parabolic law can be explained within the framework of the approach (Zhang and Li 2005; Steiner et al. 2008), in which the film growth is described by equation (34).

In the general case, the film growth constant can be written as:

\[
K_p = A \exp \left( -\frac{Q}{RT} \right) \left( P_{O_2}^0 \right)^\gamma \left( P_{O_2}^0 \right)^\delta
\]

\[37a\]

\( P_{O_2}^0, P_{O_2}^0 \) – partial oxygen pressure at the interface between the coolant-oxide and steel-oxide media, respectively, while unknown parameters are determined empirically.

The partial pressure at the coolant-oxide interface, using relation (10), is defined as:

\[
P_{O_2}^0 = C_p \exp \left( \frac{2\Delta G_{M-\gamma}}{RT} \right)
\]

\[37b\]

The value of the partial pressure at the oxide-steel interface is much smaller than the value of the partial pressure of oxygen at the surface and can be neglected (Robertson and Manning 1988). In the case of lead – bismuth coolant, it can be estimated that since the binding energy of bismuth oxide is much lower than the binding energy of lead oxide, the formation of the former does not occur until a significant amount of lead oxide is formed. Therefore, in all calculations, the formation of bismuth oxide is neglected for its smallness.

When describing the film growth constant using formula (37), the ratio of fluxes at the boundary (22) takes the following form:

\[
\frac{4M_p C_p^0}{3M_p} \left( K_p \cdot C_p \left( C_p^0 - C_p^\delta \right) \right) = \delta \left( C_p^0 - C_p^\delta \right)
\]

\[38\]

Given the discussion of relation (31), this expression means that \( \left( \frac{K_p}{P_{O_2}^{3/16}} \right) \cdot \lambda \left( C_p^0 - C_p^\delta \right) \), where \( \lambda = \frac{K_p}{P_{O_2}^{3/16}} \).

That is, upon reaching the maximum value of the film \( \delta \), the oxygen concentrations in the film and in the volume are equal. That is, the global isoconcentration regime is established.

**OXIDE FILM GROWTH CRITERION**

Based on the foregoing, in the previous section, we rewrite equation (38) in the following form:

\[
\frac{4M_p}{3M_p} \left( \frac{\delta}{\delta^\infty} \right)^{1/2} = \frac{\lambda_\delta \left( C_p^0 - C_p^\delta \right)}{C_p \left( C_p^0 - C_p^\delta \right)}
\]

\[39\]

The expression for the maximum value of the film \( \delta \), the oxygen concentration in the film and in the volume is reached at its surface, takes the form:

\[
\delta = \frac{K_p \cdot 3M_p}{2 \lambda_\delta C_p^0 \alpha_p \rho_M} = 0.2K_p \frac{C_p^0}{\lambda_\delta C_p^0 \alpha_p}
\]

\[40\]

Here, the densities of lead and magnetite are taken for temperature 550 °C, \( C_p^0 \) – saturation concentration of iron is reached at its surface, provided that there is no iron in the coolant flow and the coolant-oxide and steel-oxide media, respectively, while unknown parameters are determined empirically.

The partial pressure at the coolant-oxide interface, using relation (10), is defined as:

\[
P_{O_2}^0 = C_p \exp \left( \frac{2\Delta G_{M-\gamma}}{RT} \right)
\]

\[37b\]

The value of the partial pressure at the oxide-steel interface is much smaller than the value of the partial pressure of oxygen at the surface and can be neglected (Robertson and Manning 1988). In the case of lead – bismuth coolant, it can be estimated that since the binding energy of bismuth oxide is much lower than the binding energy of lead oxide, the formation of the former does not occur until a significant amount of lead oxide is formed. Therefore, in all calculations, the formation of bismuth oxide is neglected for its smallness.

When describing the film growth constant using formula (37), the ratio of fluxes at the boundary (22) takes the following form:

\[
\frac{4M_p C_p^0}{3M_p} \left( K_p \cdot C_p \left( C_p^0 - C_p^\delta \right) \right) = \delta \left( C_p^0 - C_p^\delta \right)
\]

\[38\]

Given the discussion of relation (31), this expression means that \( \left( \frac{K_p}{P_{O_2}^{3/16}} \right) \cdot \lambda \left( C_p^0 - C_p^\delta \right) \), where \( \lambda = \frac{K_p}{P_{O_2}^{3/16}} \).

That is, upon reaching the maximum value of the film \( \delta \), the oxygen concentrations in the film and in the volume are equal. That is, the global isoconcentration regime is established.
versa for iron. Note that the parameter \( \delta_c = \delta (T, P_{O_2}, \nu, d_i) \) will characterize either the growth or dissolution of the film with respect to the initial thickness to an equilibrium value \( \delta_i \).

**Parabolic law for the growth of an oxide film**

From relations (22) and (38) it follows that the condition of the parabolic law of film growth corresponds to the condition

\[
\frac{J_{\delta_c}}{J_{Fe}} = \frac{\delta}{\delta_i} \gg 1 \quad (41)
\]

Based on the smallness parameter for equation (39), parabolic behavior is expected for times \( \sim \frac{1}{\nu C_{Fe} \rho_{Fe} M_{Fe}} = 10^{3}-10^{5} \) from several thousand hours to several tens of thousands of hours. The latter means that parabolic approximation to this value will take place over a relatively short period of time, up to several hundreds of hours.

In order for the parabolic film growth to last for several thousand hours, the saturation parameter of the film growth \( \delta \) must be in the order of \( 10^{-2} \) m.

This corresponds to the fact that the film growth constant should be of the order of \( 10^{-3} \) m/s at \( T = 600 \) °C and an order of magnitude or two less at temperatures up to 400 °C. Such a film growth constant is very large and gives a huge value of the film thickness, which does not correspond to the known experimental data.

When the activity of iron at the surface of the film is of the order \( 10^{-3} \), parabolic film growth is provided for times from several thousand hours to several tens of thousands of hours depending on temperature.

**Double layer film**

An oxide film, as a rule, consists of two different layers and a transition region between the inner oxide layer and steel (Robertson and Manning 1988; Martinelli et al. 2008a; Alekseev et al. 2010a; Nuclear Energy Agency 2015). The inner layer is chromium-rich spinel, in which the concentration of chromium is approximately equal to its concentration in the metal, while the outer layer is a porous iron-rich oxide. A thin transition layer is observed in which the iron concentration gradually changes from the concentration in the oxide to the concentration in the alloy, the chromium concentration is practically unchanged, and the oxygen concentration gradually changes from the concentration in the oxide to zero.

Numerous experimental data show that the layers of magnetite and spinel are comparable in their thicknesses, which is explained in principle in the framework of the “accessible space” model (Robertson and Manning 1988; Martinelli et al. 2008a). The model assumes that the spinel layer Fe-Cr grows into an accessible space created by vacancies formed in steel due to diffusion of iron, while the mobility of iron ions is several orders of magnitude higher than the mobility of chromium ions. Vacancies can accumulate up to the formation of nanocavities and nanochannels, which facilitates the access of oxygen and provides the formation of spinel. But the rate of vacancy formation is completely determined by the diffusion of iron cations, and it is this process that determines the rate of both magnetite growth and the spinel formation rate. That is why the ratio of the thicknesses of magnetite and spinel Fe\(_{13}\)Cr\(_{6}\)O\(_{19}\) is approximately 1.16 (Robertson and Manning 1988; Martinelli et al. 2008a). In turn, the degree of diffusion of iron cations depends on the partial pressure at the interface between the media and determines the rate constant of the (parabolic) film growth \( K \) (40).

Based on the foregoing and following the ideas developed in (Robertson and Manning 1988; Martinelli et al. 2008a), it does not make much sense to complicate the engineering approach for calculating the changes not only in the magnetite film, but also in its spinel part, since the latter is completely determined by the diffusion (departure) of iron cations from the metal into magnetite. Let’s consider, in general terms, an equation similar to equation (28) for film growth, taking into account spinel formation and magnetite growth:

\[
\frac{1}{S} \frac{dv_{\text{Fe}}}{dt} = \frac{1}{S} \frac{dv_{\text{Fe,ox}}}{dt} + \frac{P_{\text{Fe}}}{M_{\text{Fe}}} \frac{\partial \delta_{\text{Fe,Mag}}}{dt} + \frac{P_{\text{Fe}}}{M_{\text{Fe}}} \frac{\partial \delta_{\text{Fe,Sp}}}{dt} = \frac{J_{\delta}}{4M_{\text{Fe}}},
\]

where \( v_{\text{Fe}}, v_{\text{Fe,ox}}, v_{\text{Sp}} \) — the amount of the substance of the oxide, which is the sum of the substance of magnetite and spinel. Based on the above accepted model, spinel growth is proportional to magnetite growth, \( \delta_{\text{Sp}} = a \delta_{\text{Mag}} \) and the film growth equation can be rewritten as:

\[
\frac{P_{\text{Fe}}}{M_{\text{Fe}}} \frac{\partial \delta_{\text{Sp}}}{dt} = \frac{P_{\text{Fe}}}{M_{\text{Fe}}} \frac{\partial \delta_{\text{Mag}}}{dt} + \frac{J_{\delta}}{4M_{\text{Fe}}},
\]

\[
\frac{\partial \delta}{dt} = \frac{J_{\delta}}{4M_{\text{Fe}}},
\]

\[
\frac{\rho_{\text{Fe}}}{M_{\text{Fe}}} \frac{\partial \delta_{\text{Fe,Mag}}}{dt} = \frac{\rho_{\text{Fe}}}{M_{\text{Fe}}} \frac{\partial \delta_{\text{Fe,Sp}}}{dt} + \frac{J_{\delta}}{4M_{\text{Fe}}},
\]

where \( \rho_{\text{Fe}} \) — density of iron. The film growth rate is determined by diffusion of iron cations from the metal into magnetite.
or

\[
\frac{1 + \rho_\text{ne}_M}{\rho_\text{ne}_M} = \frac{M_\text{ne}}{M_\text{ne}} \left( \frac{\partial \delta}{\partial t} + \frac{J}{4M_\text{ne}} \right), \quad \text{(43)}
\]

where \( \rho_\text{ne}_M \) and \( M_\text{ne} \) are densities and molar masses of magnetite and spinel \( Fe_3O_4 \), where \( x \) reflects the stoichiometry of the compound, respectively. Note that the densities of spinel and magnetite are fairly close to each other (Zhang 2013) and since \( \frac{\rho_\text{ne}_M}{\rho_\text{ne}_M} \approx 1 \), with accuracy of less than 3% at any time, equation (43) takes the form of (31) with high accuracy. Thus, within the framework of this model, a separate calculation of the growth of magnetite and spinel films is not made much sense, but only significantly complicates the calculations.

The interface between the outer and inner layers with sufficient accuracy coincides with the original metal surface (Robertson and Manning 1988; Schroer et al. 2012; Tsisar et al. 2017). Based on the mass balance and on the assumption that iron does not enter the coolant, but remains in the oxide outer and inner layers, and that all oxidized chromium is in the inner layer, we can write the ratio of the thicknesses of the outer layer and the inner (Robertson and Manning 1988):

\[
\frac{\delta_{o}}{\delta_{i}} = x \left( C_{Fe} / C_{Cr} \right)_{i}^{1/3} - 1 , \quad \text{(44)}
\]

where \( C_{Fe} \) and \( C_{Cr} \) determine the concentration of iron and chromium in the alloy (mol / m³).

It should be noted that the steel studied in sufficient detail for the structure of the oxide film does not give an unambiguous answer to the question of the presence or absence of an external layer of magnetite. But, in general, it can be argued that under nominal and similar modes, and in the case of dynamic experiments (coolant velocity 1–3 m/s) (Zhang and Li 2007; Weisenburger et al. 2008; Schroer et al. 2009; Tsisar et al. 2017), the magnetite layer is most likely absent due to erosion or dissolution of iron oxide films in the coolant. Although in some cases the same steels (for example, T91) show the presence of a layer of magnetite in dynamic experiments (Alekseev et al. 2010a; Schroer et al. 2012).

In static experiments, as a rule, a sufficiently bright two-layer structure of the film is observed and, as a rule, it is significantly thicker compared to dynamic experiments, in equal conditions.

**Model of volume sources in a coolant flow**

For an individual unit volume of coolant, the activities of oxygen, iron and magnetite in a state of chemical equilibrium are connected by the relation (the coolant index \( \beta \) in this section is omitted):

\[
k = \frac{a^{\text{Fe,Mg}}_\beta}{a^{\text{Fe}}_\beta}, \quad \text{(45)}
\]

Chemical equilibrium occurs due to the loss of oxygen and iron, the accumulation of magnetite:

\[
a^{\text{Fe,Mg}}_\beta = a^{\text{Fe}}_\beta + \Delta a^{\text{Fe}}_\beta; \quad a^{\text{Fe}}_\beta = a^{\text{Fe}}_\beta - \Delta a^{\text{Fe}}_\beta; \quad a^{\text{Fe,Mg}}_\beta = a^{\text{Fe,Mg}}_\beta - \Delta a^{\text{Fe,Mg}}_\beta \quad \text{(46)}
\]

The accumulation of magnetite is associated with a decrease in oxygen and iron ratio:

\[
\Delta a^{\text{Fe}}_\beta = \alpha \Delta a^{\text{Fe}}_\beta; \quad \Delta a^{\text{Fe}}_\beta = \beta \Delta a^{\text{Fe}}_\beta; \quad \alpha + \beta = 1. \quad \text{(47)}
\]

Considering the stoichiometric coefficients of the reaction of magnetite formation and the ratio of the atomic weights of oxygen and iron, we can obtain the coefficients \( \alpha = 0.726 \) and \( \beta = 0.274 \).

Combining relations (45–47), we can obtain an equation for determining the profit of magnetite in the process of establishing equilibrium:

\[
a^{\text{Fe,Mg}}_\beta = \frac{\left( \left( a^{\text{Fe}}_\beta + \Delta a^{\text{Fe}}_\beta \right) / a^{\text{Fe}}_\beta \right)^{1/3}}{k \left( \left( a^{\text{Fe}}_\beta - \beta \Delta a^{\text{Fe}}_\beta \right) / a^{\text{Fe}}_\beta \right)^{1/3}} \quad \text{(48)}
\]

Equation (48) is solved at each time step by the tangent method. According to the found values of the magnetite profit, the values of the loss of oxygen and iron are found at each step of solving the transport equations. Thus, the volume sources of impurities in the coolant are calculated by the relations:

\[
G^{\text{Fe}}_{i,0} = \Delta a^{\text{Fe}}_{i,0} C^{\text{Fe}}_{i} / \Delta t;
\]

\[
G^{\text{O}}_{i,0} = -\alpha \Delta a^{\text{Fe}}_{i,0} C^{\text{O}}_{i} / \Delta t; \quad \text{(49)}
\]

For the near-wall cells of the computational grid, surface flows of oxygen and iron are added to volume sources.

**Parabolic film growth constant**

As a rule, the parabolic film growth constant is determined experimentally in a gaseous medium or under the condition that the processes of corrosion (dissolution) of the film and erosion can be neglected. Obviously, these two effects will be quite small in a static experiment. Under the assumption that the yield of iron from steel to the coolant is small compared to its consumption by an oxide film, the Martinelli approximation (29) should work well, which ensures parabolic film growth when solving equation (28). In this case, the oxygen flux is inversely proportional to the thickness of the oxide film. Thus, to
determine the parabolic film growth constant, it is necessary to measure the oxidation rate of the surface at various temperatures and oxygen concentrations.

In the empirical determination of the parameters of the parabolic constant, the assumption of a parabolic growth of the oxide film in the static mode is fundamental. This assumption works fine provided that the diffusion yield of steel components in the coolant is small. In passivation mode in the initial period of formation of oxide films (Ivanov et al. 2005), the diffusion yield of metal components into the coolant is very intense, and its fraction can reach ~ 50% and the diffusion flux does not have to be proportional to the oxygen flux. According to the data of (Ivanov et al. 2005), for an oxidation duration of more than 400 hours, the proportion of steel components entering the coolant of the total amount of oxidized is ~ 1% and less, with an oxidation duration of 200 to 400 hours, this proportion is ~ 10%. Therefore, during long campaigns, at least more than several hundred hours, oxygen consumption occurs only on the surface and parabolic film growth in static experiments should be manifested to a greater extent than, for example, in dynamic ones. Thus, the empirical determination of the parabolic constant in the best way can be made by the thickness of the film after a sufficiently long oxidation (more than ~ 400 hours) and when using the static mode. Unfortunately, there are extremely few such experimental data.

Such experiments were carried out at JSC “SSC RFIPPE” (Askhadullin et al. 2011), where a technique was developed that made it possible to estimate the flows of oxygen consumed for the oxidation of the steel surface under variations in temperature and oxygen conditions of the coolant. There is a matrix of various experimental values of the flow of oxygen consumed for steel oxidation, and the temperature and partial pressure of oxygen in the coolant (Askhadullin et al. 2011; Ivanov et al. 2017).

Based on the available experimental data (Askhadullin et al. 2011; Ivanov et al. 2017), the following dependence was proposed for EP-832 steel:

\[
K_p = \exp(39.5) p_{O_2}^{0.125} \exp\left(-\frac{30000}{T}\right) \left[\text{m}^2 / \text{h}\right],
\]  
(50)

Similar experimental studies, by definition, were also carried out for EP-302 steel. The following dependence is proposed for a parabolic constant:

\[
K_p = \exp(31) p_{O_2}^{0.125} \exp\left(-\frac{20000}{T}\right) \left[\text{m}^2 / \text{h}\right],
\]  
(51)

For comparison, we present the dependences of parabolic constants for the basic oxygen regime, obtained in (Hwang and Lim 2010), based on an analysis of the available experimental data (Fazio et al. 2001; Barbier and Rusanov 2001; Aiello et al. 2004; Balbaud-Celerier et al. 2004; Zhang et al. 2005; Lim et al. 2010; Martinelli et al. 2008a; Hwang and Lim 2010; Nuclear Energy Agency 2015).

For steels of ferritic-martensitic class (9–12Cr) HT9, T91 the following dependence is obtained:

\[
K_p = 1.156 \times 10^{-2} \exp\left(-\frac{135693}{RT}\right) \left[\text{cm}^2 / \text{s}\right],
\]  
(52)

For austenitic steel AISI 316L:

\[
K_p = 3.027 \times 10^{-2} \exp\left(-\frac{161626}{RT}\right) \left[\text{cm}^2 / \text{s}\right],
\]  
(53)

In (Zhang 2013), the following dependences were also obtained for HT9 and T91 steels that take into account the partial pressure at the interface between media, namely, steel T91:

\[
K_p = 2.096 \times 10^{-4} c_o^{0.27} \exp\left(-\frac{157802}{RT}\right) \left[\text{m}^2 / \text{s}\right],
\]  
(54)

And for steel HT9:

\[
K_p = 9.735 \times 10^{-4} c_o^{0.27} \exp\left(-\frac{126802}{RT}\right) \left[\text{m}^2 / \text{s}\right],
\]  
(55)

where \(c_o\) (ppm, millionth share) – mass fraction of oxygen.

An analysis of the empirical dependences of parabolic constants (Fig. 2) shows that among ferritic-martensitic steels, oxygen consumption will be the smallest for EP-823 steel, which, under similar other conditions, will provide the smallest film growth. Austenitic steels (for example, the AISI 316L shown here) generally have lower oxygen consumption. Note that in (Barbier and Rusanov 2001; Balbaud-Celerier et al. 2004) the geometric data of the circulation loops are not presented in sufficient detail, on the basis of which the temperature dependences for the parabolic constant are obtained, which introduces additional calculated errors.

**Corrosion processes. Mass transfer coefficient**

It was also noted in (Zhang and Li 2007; Steiner 2009; Alekseev et al. 2010c) that the parabolic dependence does not necessarily describe the available experimental data. Further, when taking into account corrosion processes, we will adhere to the general classification of corrosion oxidation regimes adopted in (Zhang and Li 2007), namely, a comparison of \(K_p\): − parabolic constant) and \(Q_{c,o}\) – corrosion rates during corrosion oxidation:

- \(K_p < Q_{c,o}\) – corrosion oxidation process;
- \(Q_{c,o} < Q_{c,o}\) – metal corrosion without oxide formation;
- \(K_p > Q_{c,o}\) – corrosion oxidation affects corrosion;
- \(Q_{c,o} = Q_{c,o}\) – oxide corrosion, slight erosion;
- \(Q_{c,o} = Q_{c,o}\) – no new oxide formations.

The general concept of corrosion oxidation due to mass transfer effects is the assumption that the rate of corrosion oxidation is proportional to the corrosion of the metal itself (Zhang and Li 2007):

\[
Q_{c,o} \sim Q_c = \alpha(c_c - c_o) - \chi_0 (vD, d)P^n \exp(q(vD, d)/RT),
\]  
(56)
where \( Q \) – corrosion rate of the actual metal surface, \( V \) – coolant speed, \( \chi \) and \( q \) – constants depending on the properties of the coolant and flow geometry, \( n \) – determined by type of flow (Alekseev et al. 2010c).

At low coolant speeds, the corrosion rate of a metal surface is fully or partially controlled by the mass transfer effect (Chen et al. 1992; Balbaud-Célérier and Barbier 2001). The corrosion rate increases with increasing coolant velocity. At higher coolant velocities, the reactions of diffusion transfer of metal ions become equal to the diffusion rates of metal particles in the coolant or even become lower than them. As a result, the overall corrosion rate begins to be controlled by the reaction rates in the metal (activated controlled), and the corrosion rate ceases to depend on the speed of the coolant. At even higher speeds, mechanical erosion begins.

Thus, using the approach of (Zhang and Li 2007) for a pure metal, the corrosion rate is defined as

\[
Q_c \approx \chi \rho Fe C_{Fe}^{3},
\]

while

\[
\rho \approx \chi Fe C_{Fe}^{3}, \quad 3 ox c o Fe Fe \quad MQC M \gamma \rho \approx \gamma \rho \gamma \gamma \gamma (57)
\]

Thus, the activity of iron at the surface of the oxide film gives an understanding of how much the corrosion process on the film is slower compared to corrosion of the metal surface. As a rule, this value is about 100–1000. Based on the chemical equilibrium on the wall, the greater the oxygen activity in the coolant, the lower is the activity of iron and the less corrosion of the surface. Of course, it must be kept in mind that a large (close to saturation) oxygen activity can lead to excessive and undesirably large film growth, that is, the yield of iron can be minimized by an increase in oxygen in the coolant, but thereby its output will increase, in fact, in the oxide film, which also is not always desirable.

There are several expressions of mass transfer coefficients (Figs 2, 8) obtained on the basis of experimental data (Harriott and Hamilton 1965; Berger and Hau 1977; Silverman 1984):

\[
k_{b-h} = 0.0165 \nu^{-0.53} D^{0.67} V^{0.86} d^{-0.14}
\]

\[
k_{s} = 0.0177 \nu^{-0.579} D^{0.704} V^{0.875} d^{-0.125}
\]

\[
k_{h-h} = 0.0096 \nu^{-0.567} D^{0.654} V^{0.913} d^{-0.087}
\]

Using formulas (58), numerical values are obtained for the mass transfer coefficients, but these differences do not exceed 20% at a speed of 2 m/s and less at lower speeds. In further calculations, we use the expression \( k_{b-h} \). Fig. 3 shows typical values of the rates of dissolution (corrosion) of the film taking into account this mass transfer coefficient (58) depending on temperature.

When using experimental data to verify the calculation models, it is not always possible to extract data on the geometry of the working sections with sufficient accuracy, but from general considerations, the limits of the hydraulic diameters of these sections can be estimated. As a rule, hydraulic diameters are within the range of 10–100 mm. Corrosion rate parameter change \( \chi \) in (34) when changing the hydraulic diameter from 10 to 100 mm lies within 20%, as can be seen in Fig. 4.

**Justification of the calculation method**

The main goal of a computer program is to calculate the thickness of an oxide film and the associated concentrations of oxygen and iron. Thermohydraulic calculations are a necessary basis, but the accuracy of the calculated thermohydraulic characteristics (in our case, this is the field of temperatures and velocities) does not have to exceed the accuracy of the calculation of physicochemical processes (in our case, this is the thickness of the oxide film and the concentrations accompanying it). The main errors in the calculation are introduced by the uncertainty of empirical parameters.

The film growth in the static case is parabolic in nature. It is static experiments that serve as the basis for de-
terminating the parameters of the parabolic constant (37a), which is determined by the oxygen consumption of the film. Based on the fact that in the static case there is a parabolic dependence of the film thickness growth \( \delta^2 \sim K_p \cdot t \), then the relation between the relative errors of the film thickness and the uncertainty of the parabolic constant is written as \( \varepsilon_{K_p} = 2 \varepsilon_{\delta} \).

The experiments on oxygen consumption (Askhadullin et al. 2011; Ivanov et al. 2017; Salaev et al. 2018) make it possible to relatively accurately determine only the parameters of the dependence on temperature and partial pressure (37b). Therefore, the absolute values of the parabolic constant can be determined only by reference experiments. In our case, these are experiments (Barbier and Rusanov 2001; Balbaud-Celerier et al. 2004), since the experiments were performed for different types of steels on the same installation. Based on the data in Figs 8–10 (Balbaud-Celerier and Terlain 2004), we accept \( \varepsilon_{K_p} \approx 0.6 \).

Since the accepted empirical dependence (37a) is determined by three parameters, \( A, Q, \) and \( n \), the relative uncertainty of the parabolic constant is determined by the relative errors for these parameters. The dependence on the uncertainty of the degree of partial pressure can be neglected due to its smallness, then the uncertainty of the parameters \( A \) and \( Q \), \( \varepsilon_A \) and \( \varepsilon_Q \), contribute to the uncertainty of the parabolic constant as

\[
\varepsilon_{x_c} = \sqrt{(\varepsilon_A)^2 + (Q/T \cdot \varepsilon_Q)^2}
\]  

(60)

Since, based on the available data, it is impossible to unambiguously indicate which of the uncertainties makes the largest contribution, we accept this contribution to be the same, i.e.

\[
\varepsilon_A = |Q/T| \cdot \varepsilon_Q = 0.6 / \sqrt{2} = 0.43
\]  

(61)

That is, uncertainty \( \varepsilon_A \) may be high enough while uncertainty \( \varepsilon_Q \) should be small since \( |Q/T| \gg 1 \).

Thus, during variational calculations of the film thickness and impurity concentrations, the boundaries of the varied parameters are estimated \( A \) and \( Q \).

The errors of partial pressure and average flow rate, temperatures are taken from the available experimental data.

In dynamic experiments, the flow rate and hydraulic diameter were additionally varied. The speed varied based on the data of the error of the flow meter, this value, as a rule, lies within \( \pm 10\% \). The hydraulic diameter ranged from 10–100 mm (see previous section).

Based on the selected intervals, the average deviation of the calculated value from the experimental ones was calculated to get the relative error

\[
\varepsilon = \frac{1}{N} \sum_{i=1}^{N} \frac{\text{abs}(x_{\text{calc}} - x_{\text{exp}})}{x_{\text{exp}}}
\]  

(62)

where \( x_{\text{calc}} \) and \( x_{\text{exp}} \) calculated and experimental values of the simulated value, respectively.

Subsequent numerical calculations showed that the errors in the calculation of dissolved oxygen concentrations are not more than 15% when comparing the results obtained using the MASKA-LM and STAR-CCM + codes, and the oxide film thickness is not more than 30% for the static case (stationary coolant) and not more than 55% for the dynamic case (coolant with average speeds of 0.1–3 m/s) when comparing the calculated and experimental data.

When considering experimental errors for oxygen concentration, it should be noted that at the moment in the literature there is no described well-established measurement procedure with a proven error in a wide range of oxygen concentrations and it, in principle, can reach significant values. When considering experimental errors for oxide film thicknesses, we note that experimental results are very often presented with no errors at all, and those data presented with errors should be considered as

---

**Figure 4.** The value of the parameter of dissolution (corrosion) of the film (see equation (34)) depending on the hydraulic diameter.

**Figure 5.** Dependence of the saturation thickness of the oxide film on the coolant velocity at a basic oxygen concentration of \( 10^{-8} \) kg/kg.
a spread of experimental data without evaluating the experimental errors themselves (see, for example, Fig. 11).

**Simulation of film growth in the framework of the simplified 0D-model**

The joint solution of equations (34) and (38) within the framework of the point model, that is, without taking into account the exact geometry (Tadmon’s approach), allows a reasonable accuracy to estimate the film growth rate depending on temperature and coolant velocity. For this, we used the film growth parameters selected for EP-823 steel, since for this grade of steel there is a fairly large amount of experimental data.

The growth of an oxide film as a function of time is considered in detail in a subsequent chapter. The most indicative characteristics parameters of film growth are its maximum value (see equations (34–36)) and the characteristic film growth time according to the parabolic law \( t_{\chi} = \frac{\chi}{k^2} \). Based on (40), the saturation yield corresponds to the equality of the fluxes of iron through the film and in the coolant, \( J_{\nu}^f = J_{\nu}^c \), which corresponds to a zero flow of oxygen. This, of course, is performed provided that a constant concentration is maintained at the selected oxygen regime.

Figs 5–7 show the typical model dependences for the film saturation thickness on the oxygen velocity and concentration, as well as the characteristic time of transition to parabolic growth. The simulation parameters are taken for lead coolant and steel EP-823.

The growth of the oxide film upon saturation, depending on the coolant speed, is shown in Fig. 5. Obviously,
at low speeds, a substantial, almost exponential, increase in the thickness of the saturation film is observed. That is, the saturation thickness substantially depends on the velocity of the coolant.

Also, calculations showed that at temperatures of 450–650 °C, parabolic film growth is expected for times from less than an hour to several hundred hours (Fig. 6).

The calculations shown in Fig. 7 demonstrate that under the basic oxygen regime (oxygen concentration in the region of $10^{-8}$ kg/kg) and at standard coolant velocities (1–2 m/s), a very small film growth is expected (up to 1 μm). If you deviate, in the direction of increasing the oxygen content, from the basic mode, you can expect film growth to several tens of micrometers. At a low oxygen content (dash dotted line in Fig. 7), the formation of an oxide film is practically unrealistic, which can lead to corrosion.

Figs 8–10 show the calculations for steels EP-823, T91 and 316L for which both static and dynamic experiments were performed (Barbier and Rusanov 2001; Balbaud-Celerier and Terlain 2004), which allows a consistent analysis of these data, that is, the same parameterization of the parabolic constant is used for both types of tests. The coolant speed in the dynamic test is 1.9 m/s. Below is an analysis for $T = 600$ °C. Taking into account the numerical errors from the variation of the parameters, we obtain fairly good agreement with the experimental data. An interesting feature of the above calculations and experimental data is that although the steels EP-823 and T91 are quite close to each other in composition, the growth and thickness of the film for them is significantly different, which determines the adequacy of our engineering approach.

The authors of (Zhang and Li 2007; Weisenburger et al. 2011) analyzed the obtained experimental information for T91 steel. The experiments were carried out on the same installation of JSC “SSC RF-IPPE” as in (Balbaud-Celerier and Terlain 2004; Zhang et al. 2005), but with an average coolant speed of 1 m/s and under the basic oxygen regime. As discussed earlier, it is not possible to reconstruct the exact geometric parameters. Therefore, a certain spread in the values of the hydraulic diameter in the range of 0.005–0.1 m, which gives no more than 20% error in the value of the film thickness, is accepted. As an example, a calculation with an average coolant velocity of 0.3 m/s is presented to demonstrate sensitivity to this parameter. This speed was chosen intentionally to place most of the experimental data between the two calculated curves (Fig. 11). The calculated estimate demonstrates that having a spread of experimental data can be, inter alia, related to a “spread” in speed from 0.3 m/s to 1 m/s.

Numerical analysis of oxide film growth on surfaces of simple geometries using MASKA-LM and STAR-CCM+ codes

Distribution of oxygen at a given drain on the pipe wall

This section provides calculations of model oxygen distributions at various temperatures. Test calculations of the elementary model of HLMC flow in a pipe were carried out using the commercial StarCCM+ code and MASKA-LM code of own design.

The STAR CCM+ code interface allows you to use user-defined functions to model certain physical and chemical processes. The system of equations (33), (38), functions dependent on them, and semi-empirical dependencies of type (2, 20–25) were incorporated into the code for joint solution with the STAR CCM+ thermohydraulic models.

For numerical analysis, a simple model was chosen for the flow of lead coolant of a given temperature in the pipe: the length of the computational domain is 1 m; diameter...
50 mm; design grid 1 million cells; Reynolds number is $10^5$; input mass concentration of oxygen $2 \times 10^{-8}$ kg/kg. To simulate oxygen corrosion, the following value was selected for the drain (consumption) of oxygen to the wall $4 \times 10^{-8}$ kg/m$^2$·s. The selected amount of oxygen consumption is very significant compared to that usually found in real calculations and was chosen only to demonstrate the distribution of oxygen.

When using the MASKA-LM code, a rectangular computational grid with constant spatial steps in the Cartesian coordinate system, consisting of 275,000 cells (50 × 50 × 110) (along the directions of the coordinate axes) was entered into the computational domain. That is, the number of cells and the quality of the grid is noticeably inferior to the StarCCM+ code.

The results of calculations using the StarCCM+ and MASKA-LM codes are shown in Figs 12, 13 (for coolant temperature of 400 °C and 500 °C, respectively, the differences are minimal). Calculations are given for various turbulence models (StarCCM+) and for various turbulent diffusion coefficients used in the MASKA-LM code. Changing the coefficient of turbulent diffusion by an order of magnitude gives the maximum error in calculating the oxygen concentration of less than 15%. It is worth noting that at the moment, the methodology for determining the measurement error of the oxygen activity sensor has not been developed, and according to qualitative estimates it can be at least 50%. In this case, the choice of a turbulence model or a simplified description of turbulent motion will not affect the description of available experimental results at all.
Thus, a turbulent diffusion coefficient of the order of \((1-3)\times10^{-5}\) gives a completely adequate description of the oxygen distribution with an error of no more than 15\% at the surface. With smaller oxygen sinks, the scatter of values is even smaller. The scatter in the values of the coefficient of turbulent diffusion gives a very small error in the thickness of the oxide film.

Numerical analysis of oxide film growth

This section presents model calculations of the growth of an oxide film in an EP-823 steel pipe and a beam of rods at various flow rates of HLMC and temperatures.

Figs. 14–16 show the calculations for different temperatures, oxygen concentrations and average speeds in the pipe using the MASKA-LM and STAR CCM+ codes.

Fig. 14 shows film growth versus time. All models, in accordance with Zhang’s approach, demonstrate the approximation of the film thickness to the maximum possible value. The calculation shown by the black solid line is made taking into account only the parabolic constant (there is no dependence on the coolant speed through taking into account mass transfer). As can be seen, for calculations at a speed of 1.8 m/s, the film growth rate close to the parabolic behavior is observed only from about 1 to 50 hours, while at a speed of 0.1 m/s it is observed up to several thousand hours. Although it is more correct to say that film growth is closer to the paralinear law in these areas. The Tedmon’s approach fairly well describes only areas close to saturation in film growth.

A similar behavior of the increase in the thickness of the oxide film is also observed at a temperature of \(T = 500^\circ\text{C}\). The main difference from the previous case is that the approximation to the asymptotic value of the film thickness occurs much later (Fig. 15). All models, including the simple 0D model, show very similar results. Iron concentration distributions also show fairly similar behavior, and the main difference, again, is observed near the surface.

Fig. 16 shows the calculations for an increased oxygen concentration; in general, the concentration of the basic oxygen regime is more than an order of magnitude higher. In this case, for a rather long time, a pronounced parabolic growth of the film is observed. Comparison of calculations using the codes in question was carried out only for temperatures \(T = 500^\circ\text{C}\) and \(T = 650^\circ\text{C}\). The maximum difference in the thickness of the oxide film is not more than 20\%.

For the following geometry, a bunch of rods was chosen with a diameter of 10 mm, a length of 0.97 m, in a triangular package, with a step of 14 mm; the simulated area is cut off along the planes of symmetry, and the grid contains \(3.1 \times 10^6\) cells when using the STAR CCM+ code; mass oxygen content of \(10^{-8}\) kg/kg; temperature \(500^\circ\text{C}, 650^\circ\text{C}\); coolant velocity 1.7 m/s (Re=\(\sim10^5\)), turbulence model K-\epsilon when using STAR CCM+ code.

Fig. 17 shows the calculation results for the thickness of the oxide film at various temperatures. As in the previous case, the difference in the obtained calculations does not exceed 20\%.

Note that in the above examples only isothermal calculations were considered.

Conclusion

An engineering model is presented for a self-consistent calculation of the growth of an oxide film in circulation loops with a heavy liquid metal coolant and concentrations of impurities (oxygen, iron, magnetite). The modeling of thermohydraulic and physicochemical processes is based on solving the associated three-dimensional equati-
ons of hydrodynamics, heat transfer, convective-diffusive transport, and the formation of chemically interacting impurity components in the coolant volume and on the surface of steels.

The change in the thickness of the oxide film on the metal surface and the corresponding exit of steel components into the coolant substantially depends on the steel grade. For a more adequate justification of the evolution of the oxide film, a semi-empirical model is proposed for using the empirical parameterization of the parabolic constant not only in the equation for changing the thickness of the oxide film, but also in the mass balance equation associated with it. The parabolic constant, which is determined by the degree of oxygen consumption by a steel, obviously significantly depends on the type of this steel and on the method of its preparation. Therefore, the direct application of the Wagner approach is unlikely to adequately describe such differences in steels, while the experimental parametric dependences on oxygen consumption by steel depending on temperature and oxygen partial pressure are obviously unique for each steel.

For an adequate experimental determination of the parametric dependences for a parabolic constant, at least two conditions are necessary: the experiment should be carried out for a static case or the average coolant velocity should be low in order to exclude the influence of corrosion-erosion processes; steel should have a sufficient initial oxide film to exclude the influence of the initial large flow of iron, and possibly corrosion of steel, since with a large yield of iron the film growth does not have to obey the parabolic law even in the static case.

The developed approach regarding accounting for the main physical and chemical processes is used not only in our MASKA-LM code, but is also implemented as user-defined functions in the STAR CCM+ code. The cross-verification calculations of the two test models showed a fairly good agreement between the results in terms of describing the increase in the thickness of the oxide film.

In the future, the developed approach will be applied to more complex systems in terms of geometry, expanded for non-isothermal loops, and applied to simultaneously account for the formation of deposits in loops.

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Appendix

Other Necessary Closing Relations

As the closing relations for the system of equations of motion and equations for the transport of impurities, we used dependences to estimate the values of the coefficients of turbulent thermal conductivity, turbulent viscosity, and turbulent diffusion (Reynolds 1979).

To calculate the coefficient of turbulent viscosity, we use the relations of the three-layer Karman model (Laptev 2007; FBU "STC NRS" 2012):

\[
\begin{align*}
\nu_T &= 0 \quad \text{for } y_\tau < 5 \\
\nu_T &= \frac{y_\tau}{5} - 1 \quad \text{for } 5 \leq y_\tau \leq 30 \\
\nu_T &= \frac{y_\tau}{2.5} - 1 \quad \text{for } y_\tau > 30
\end{align*}
\]

where \( \nu_T \) – turbulent viscosity, 
\( y_\tau \) – dimensionless coordinate in the boundary layer, 
\( v \) – kinematic viscosity, 
\( u^* \) – dynamic speed, \( \tau \) – wall shear stress, \( \rho \) – density. For pipes with a circular cross-section and the average shear stress along the perimeter of the bar in the assembly of smooth bars, the shear stress is calculated by the formula \( \tau = \frac{u^*}{2.5} \) (FBU "STC NRS" 2012), i.e. \( \nu_T = \frac{y_\tau}{2.5} - 1 \) – hydraulic resistance coefficient.

For pipes with circular cross-section at \( \text{Re} = 4 \times 10^3 \sim 10^6 \):

\[ \zeta = \frac{1}{(1.821 \cdot \lg \text{Re} \cdot 1.64)^3} \]

For assembling smooth rods in triangular grid with \( \text{Re} = 6 \times 10^3 \sim 10^6 \):

\[ \zeta = \zeta_0 (1 + (h/d)^{0.32}), \quad \zeta_0 = 0.21/\text{Re}^{0.25}, \] where \( h \) – grid period, \( d \) – rod diameter.

In (FBU "STC NRS" 2012) other correlations are also presented for calculating the hydraulic resistance of a rod assembly, which can be parametrically used in the code.

According to the Prandtl model, the average turbulent viscosity at \( y \geq \delta \) (that is, in the flow core) is calculated by the formula (Laptev 2007):

\[ \nu_T = 0.4 u^* \delta, \quad \text{where the thickness of the boundary layer is calculated in the formulation of the three-layer Karman model (derivation of the relation in [51]):} \]

\[ \delta = \frac{u}{u^*} 30 \exp \left[ 0.4 \left( \frac{\pi}{u^*} - 13.96 \right) \right] \]

Thus, the turbulent viscosity in the flow is defined as:

\[ \nu_T = 12 \exp \left[ 0.4 \left( \frac{\pi}{u^*} - 13.96 \right) \right] \]

The formulas used to determine the turbulent viscosity are not the only possible ones for use in calculations. In (Laptev 2007) alternative formulas are given for calculations both in the framework of the three-layer and in the framework of the two-layer model. In (Laptev 2007), a sufficient similarity of the results of using different models is demonstrated.

Estimates of the coefficient of turbulent diffusion of an admixture are based on the analogy of transport processes, the turbulent Schmidt number \( Sc_T = \nu_T / D_T \sim 1 \).