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FLUX EFFECT ONTO PERITECTIC PHASES GROWTH IN THE ZINC COATING

Wpływ topnika na wzrost faz perytektycznych w powłoce cynkowej

A model of the (Zn) – coating formation on the iron/steel substrate is proposed. The model assumes the phases’ sub-layers creation in a sequence. This sequence is referred to the Fe-Zn phase diagram. However, this sequence of phases’ appearance is perturbed by the flux presence in the zinc bath. The flux effect on the coating morphology and appearance/disappearing of some sub-layers is analysed. The phases’ formation is treated as the result of the peritectic reaction accompanying the coating solidification. A comparison of the coating formations before and after flux decay is delivered. Thus, a function which describes the flux decay is also analysed. Additionally, a ternary Fe-Zn-F(flux) phase diagram is considered. The varying zinc concentration across the phases sub-layers is described with the use of the function which determines the flux decay. The behaviour of the solidification path before and after flux decay is discussed due to the adequate equations formulated in frame of the current model.

Keywords: Fe-Zn-flux phase diagram; flux disappearing; coating thickening

1. Introduction

Technology of the hot-dip galvanizing requires to apply a flux which is able to improve the adhesion of the (Zn) – coating settled on the steel substrate, [1-11]. As the (Zn) – coating consists of two layers: first layer being directly in the contact with substrate and second layer being the result of the substrate/coating system emerging from the zinc bath, the adhesion between substrate and first layer is of the main significance for the technology, [12-15].

Therefore, the $\Gamma_1$ and $\delta$ – phases appearance is the subject of the critical analysis in the current approach. However, other phases formation, like $\zeta$ or $\zeta' \equiv \zeta + \eta$ will also be considered, especially in the agreement with the solidification sequence resulting from the phase diagram, as discussed in literature, [16,17].

2. Model for the zinc coating growth

A schematic model for the sub-layers formation in the zinc coating is shown in Fig. 1. It results from this scheme that the sub-layers appear in the coating sequentially. Each sub-layer phase is the product of the peritectic transformation under non-equilibrium condition, that is due to undercooled peritectic reaction.

The solidification accompanied by peritectic reaction is preceded by the substrate dissolution within the zone denoted $dx$, to produce the $N_0^F$ – liquid solution, Fig. 1. It occurs according to the reaction: $\text{liquid}(N_0^F) + \text{Fe} \rightarrow \text{liquid}(N_0^F)$. 

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The solidification is isothermal one due to the imposed real temperature and therefore, the Number of the Degree of Freedom is equal to zero. In fact, $f = c + 1 = 0$ with $c = \text{Fe, Zn, F (flux)}$; $p = \text{liquid} (N^F_0)$, $Γ_1, δ, ζ$, (where flux consists of Zn and Cl).

It is assumed, in the first approximation, that the intersection of the $T_R$ – real temperature with the liquidus line defines the end of solidification path denoted as $N^F$, Fig. 3. Strictly analyzing, the $N^F$ equilibrium solution of the iron in zinc results from the intersection of the $T_{Zn}^m$ – isotherm with the liquidus line, [19,20], $(T_{Zn}^m$ – the melting point of zinc). So, the length of the solidification path is $N^F_0 + N^F$ within the period of time when the flux is not yet evaporated, Fig. 3.

It seems that the formation of channels used by this system for the boundary diffusion is a natural phenomenon accompanying the peritectic reaction. Various issues related with the peritectic solidification were described in detail in literature, [21]. An important case of the peritectic solidification is the situation when the primary $α$ phase is not a substrate for the nucleation of $β$ – phase, Fig. 4a. Then, the $β$ – peritectic phase nucleates in the liquid surrounding the $α$ – phase. The concentration of the liquid surrounding the $α$ – phase changes during the

At the beginning of the zinc coating formation the neighboring bath contains: Zn + F (flux), Fig. 2a. Next, the $dx$ – dissolution zone is created by the reaction: \textit{liquid (Zn)} + Fe $\rightarrow$ \textit{liquid} $(N^F_0)$, Fig. 2b. The liquid from the $dx$ – dissolution zone diffuses towards the bath to promote two first sub-layers formation due to peritectic reaction: $α + \text{liquid} (N_1) \rightarrow Γ_1$ which is followed by the subsequent in sequence peritectic reaction: $Γ_1 + \text{liquid} (N_2) \rightarrow δ$, ($δ \equiv δ_c$, in fact), Fig. 2c. At the same time the remaining liquid enters into the reaction with the zinc to transform it into its equilibrium solution: \textit{liquid}$(N_2)$ + \textit{liquid}$(\text{Zn})$ $\rightarrow$ \textit{liquid}$(N^F)$.

Now, the $N^F$ - equilibrium zinc solution forms the liquid in the $dx$ – dissolution zone according to the reaction: \textit{liquid} $(N^F) + \text{Fe} \rightarrow \text{liquid} (N^F_0)$, as mentioned above, Fig. 2d. Simultaneously, the birth/nucleation of the $δ_p$ – phase occurs on the previously formed $δ_c$ – phase sub-layer, when solidification time is: $t = t_c$, Fig. 2d. Nothing opposites to the $ζ$ – phase appearing which forms two sub-layers: $ζ$ – itself and $ζ_2 \equiv ζ + η$, Fig. 2d. At the $t_c$ – time the flux disappears/decays and the growth of the $δ_c$– phase sub-layer is arrested. Instead, the $δ_p$ – phase sub-layer is formed, exclusively, Fig. 2e. Now, the Zn – solute concentration in the dissolution zone attains a new value: $N^F_0 \rightarrow N^F_0$, which occurred at time $t = t_c$, Fig. 2d, Fig. 2e.

All the phenomena discussed above, Fig. 2, can be related to the Fe-Zn phase diagram. Recently, a new phase diagram has been published, [18]. This phase diagram is exclusively dedicated to the hot dip galvanizing technology, Fig. 3. The $T_R$ – real temperature of the hot dip galvanizing is superimposed on the diagram. Also, the $N^F_0$ – solute concentration, typical for the beginning of the galvanizing, (described in Fig. 1) is marked in the phase diagram, Fig. 3. As the peritectic transformations occur at the $T_R$ – real temperature they become the undercooled peritectic reactions.

The solidification is isothermal one due to the imposed real temperature and therefore, the Number of the Degree of Freedom is equal to zero. In fact, $f = c + 1 = 0$ with $c = \text{Fe, Zn, F (flux)}$; $p = \text{liquid} (N^F_0), Γ_1, δ, ζ$, (where flux consists of Zn and Cl).

It is assumed, in the first approximation, that the intersection of the $T_R$ – real temperature with the liquidus line defines the end of solidification path denoted as $N^F$, Fig. 3. Strictly analyzing, the $N^F$ equilibrium solution of the iron in zinc results from the intersection of the $T_{Zn}^m$ – isotherm with the liquidus line, [19,20], $(T_{Zn}^m$ – the melting point of zinc). So, the length of the solidification path is $N^F_0 + N^F$ within the period of time when the flux is not yet evaporated, Fig. 3.

It seems that the formation of channels used by this system for the boundary diffusion is a natural phenomenon accompanying the peritectic reaction. Various issues related with the peritectic solidification were described in detail in literature, [21]. An important case of the peritectic solidification is the situation when the primary $α$ phase is not a substrate for the nucleation of the $β$ – phase, Fig. 4a. Then, the $β$ – peritectic phase nucleates in the liquid surrounding the $α$ – phase. The concentration of the liquid surrounding the $α$ – phase changes during the
growth of the $\beta$–phase. Finally, the adequate thermodynamic conditions are created to promote melting of the $\alpha$–phase and further growth of the $\beta$–phase, [20]. This process of peritectic phase formation is called the peritectic reaction (as opposed to peritectic transformation shown in Fig. 4b).

3. Flux effect onto the (Zn) – coating formation

The contribution of the third component which is flux designated as $F$, to the hot dip galvanizing technology, is to be examined in order to describe the solidification process under investigation. It is the presence of the $F$ – flux in the galvanizing that makes the $\Gamma_1$ – phase growth and, as mentioned above, also promotes the appearance of the $\delta_c$ – phase of the compact morphology, [22].

The model also assumed that the $\zeta$ – phase sub-layer stops growing at the $t_C$ – time, that is, at an instant when the decay of flux and products of its combustion/decomposition which evaporate is completed, Fig. 5. Further extension of the model, to better its adaptation to the needs of the galvanizing technology, requires the development of a hypothesis regarding the flux life in the process. It has been assumed with good result, after comparing the simulation with measurement, that 90% of the flux fades immediately upon being introduced to the zinc bath. The remaining 10% (or 0.1 in mass fraction) is actively involved in the process of solidification, Fig. 6.

Hypothetical function which describes the flux behavior in the process of the (Zn) – coating growth on iron, Fig. 6 has three characteristic points (indicated by dots): the first – for the time, $t_{B\Gamma_1}$ (birth of the $\Gamma_1$ – phase), the second – for the time, $t_{B\zeta}$ (birth of the $\zeta$ – phase) and the third – for the $t_C$ – time. It allows for defining the $F(t)$ – function which describes the flux evaporation, Eq. (1).

The process of flux decay was divided into two phenomena: burning out of the flux and the effect of combustion products on the coating solidification. Both phenomena are juxtaposed, and the resultant hyperbole, $F(t)$, could be plotted hypothetically, Fig. 6.
of the Fe-Zn binary phase diagram, exactly, when the ternary system to reach a constant value on the transformation at a time

- the variability of zinc concentration in the sub-layers of the intermetallic phases under the influence of the \( F \) – flux, for \( t \leq t_c \), and after the flux decay, \( t > t_c \).

The variability of zinc concentration in the \( \zeta \) – peritectic phase emerging in the sequence as a third one and designated here with the symbol P3;

\[
N_{P3}(t) = \frac{k_4 N_3}{k_4 N_3 + k_4 N_1} \left( 1 - F(t) \right) \left( t_c - t \right) \leq t \leq t_c, \quad (4)
\]

A consequence of this behavior of the Zn – solute concentration in individual sub-layers of the peritectic phases is the variability in the \( N_0(t) \) – function, Fig. 7. This function makes the hypothetical solidification path, Fig. 6, defined as well as possible in frame of the present model.

The problem of the length of the solidification path is not fully solved, but it is known that the solidification process combined with diffusion is proceeding in such a way that remaining liquid phase of the \( N^F \) – concentration goes into the zinc bath which is of the same concentration. On the other hand, the \( N^F \) – equilibrium solution (zinc bath) still serves for the substrate dissolution. The proposed function which describes the kinetics of flux decay, Eq. (1), is used to determine the variability of the peritectic phases concentration during the period of flux existence in the bath. It is shown by the following equations written successively for:

- the variability of zinc concentration in the sequence as a first one and designated here with the symbol P1;

\[
N_{P1}(t) = \begin{cases} 
0 & \text{for } t \leq t^{\Gamma^1}_B, \\
k_2 N_1 & \text{for } t^{^\Gamma^1}_B \leq t \leq t_c, \\
k_2 N_1 + k_3 N_2 + k_4 N_3 & \text{for } t_c \leq t,
\end{cases} \quad (2)
\]

- the variability of zinc concentration in the sequence as a second one and designated here with the symbol P2;

\[
N_{P2}(t) = \begin{cases} 
0 & \text{for } t \leq t^{\delta}_B, \\
k_3 N_2 & \text{for } t^{\delta}_B \leq t \leq t_c, \\
k_3 N_2 + k_3 N_2 + k_4 N_3 & \text{for } t_c \leq t,
\end{cases} \quad (3)
\]

\[
F(t) = \frac{t - t_c}{a t + b}, \quad t^{\Gamma^1}_B \leq t
\]

\[
a = \frac{t^{\Gamma^1}_B F^{\zeta} - t^{\Gamma^1}_B F^{\Gamma^1}}{t^{\Gamma^1}_B - t^{\Gamma^1}_B} = -60.2;
\]

\[
b = \frac{t^{\Gamma^1}_B (F^{\Gamma^1} - F^{\zeta}) + t_c (t^{\Gamma^1}_B F^{\zeta} - t^{\Gamma^1}_B F^{\Gamma^1})}{F^{\Gamma^1} F^{\zeta}} = -1500
\]

The above function, Eq. (1), requires the selection of the parameters such as: \( F(t^{\Gamma^1}_B) = F^{\Gamma^1} \) and \( F(t^{\Gamma^1}_B) = F^{\zeta} \), Fig. 6, to make the model coherent in confrontation with the measurements of the \( N(x) \) – solute concentration. Additionally, the \( t_c \) – parameter is also to be known, and can be determined by the method shown in Fig. 5.

A hypothetical solidification path shown in a virtual ternary Fe-Zn-F system is plotted for the \( N_0 \) – nominal solute concentration treated as the starting point, Fig. 6. This solidification path is: \( N_0(t^{\zeta}) \equiv N^0_b \rightarrow N_0(t_c) \equiv N_1 \), Fig. 3.

The \( N_0 \) – nominal solute concentration varies smoothly in the ternary system to reach a constant value on the \emph{liquidus} line of the Fe-Zn binary phase diagram, exactly, when the \( t_c \) – time is attended. The flux as well as the product of its combustion do not exist more in the zinc bath. Now, the (Zn) – coating formation follows the solidification path in the Fe-Zn binary phase diagram: \( N_1 \rightarrow N^F \). It occurs until the appearance of the first solid-state transformation at a time \( t_M \sim 300 \text{ seconds of the galvanizing.} \)

The problem of the length of the solidification path is not fully solved, but it is known that the solidification process combined with diffusion is proceeding in such a way that remaining liquid phase of the \( N^F \) – concentration goes into the zinc bath which is of the same concentration. On the other hand, the \( N^F \) – equilibrium solution (zinc bath) still serves for the substrate dissolution. The proposed function which describes the kinetics of flux decay, Eq. (1), is used to determine the variability of the peritectic phases concentration during the period of flux existence in the bath. It is shown by the following equations written successively for:

- the variability of zinc concentration in the \( \Gamma^1 \) – peritectic phase emerging in the sequence as a first one and designated here with the symbol P1;

\[
N_{P1}(t) = \begin{cases} 
0 & \text{for } t \leq t^{\Gamma^1}_B, \\
k_2 N_1 & \text{for } t^{\Gamma^1}_B \leq t \leq t_c, \\
k_2 N_1 + k_3 N_2 + k_4 N_3 & \text{for } t_c \leq t,
\end{cases} \quad (2)
\]

- the variability of zinc concentration in the \( \bar{\delta} \) – peritectic phase emerging in the sequence as a second one and designated here with the symbol P2;

\[
N_{P2}(t) = \begin{cases} 
0 & \text{for } t \leq t^{\bar{\delta}}_B, \\
k_3 N_2 & \text{for } t^{\bar{\delta}}_B \leq t \leq t_c, \\
k_3 N_2 + k_3 N_2 + k_4 N_3 & \text{for } t_c \leq t,
\end{cases} \quad (3)
\]

- the variability of zinc concentration in the \( \zeta \) – peritectic phase emerging in the sequence as a third one and designated here with the symbol P3;

\[
N_{P3}(t) = \begin{cases} 
0 & \text{for } t \leq t^{\zeta}_B, \\
k_4 N_3 & \text{for } t^{\zeta}_B \leq t \leq t_c, \\
k_4 N_3 + k_4 N_1 & \text{for } t_c \leq t,
\end{cases} \quad (4)
\]

A consequence of this behavior of the Zn – solute concentration in individual sub-layers of the peritectic phases is the variability in the \( N_0(t) \) – function, Fig. 7. This function makes the hypothetical solidification path, Fig. 6, defined as well as possible in frame of the present model.

The variability of the \( N_0 \) – parameter is shown more precisely to distinguish the periods of time when the individual nucleus (just born) join each to other to form a fully developed sub-layer. There are two periods: \( t^\delta_B \sim t^{\bar{\delta}}_B \) and \( t^{\zeta}_B \sim t^\zeta_B \). Both periods of time correspond with grey zones, respectively, Fig. 8. It is to be emphasized that the current model does not operate within the grey zones, Fig. 8.
The definition of the \( N_0(t) \) function is as follows:

\[
N_0(t) = \begin{cases} 
N_D(t) \frac{\tilde{\Gamma}^1(t) + N_{P2}(t) \tilde{\chi}^C(t)}{\tilde{\Gamma}^1(t) + \tilde{\chi}^C(t)}, & t \leq t_C, \\
N_D(t) \frac{\tilde{\Gamma}^1(t) + N_{P3}(t) \tilde{\chi}^C(t) + N_{P3}(t) \tilde{\chi}^C(t) + t_{\delta}^H - t_{\delta}^H}{\tilde{\Gamma}^1(t) + \tilde{\chi}^C(t) + \tilde{\chi}^C(t) + t_{\delta}^H - t_{\delta}^H}, & (5a) \\
N_0 = \frac{k_3 N_2 \tilde{\lambda}^D(t) + k_4 N_3 \tilde{\lambda}^C(t) + t_{\delta}^H - t_{\delta}^H}{\tilde{\lambda}^D(t) + \tilde{\lambda}^C(t) + \tilde{\lambda}^C(t) + t_{\delta}^H - t_{\delta}^H}, & (5b) \end{cases}
\]

A proper derivatives are present in the above definitions, Eq. (5). This formula corresponds well with the rate of the appropriate phases (\( \Gamma_1, \delta_C, \delta_P \) and \( \zeta \)) thickening, \( \lambda(t) \), which can be calculated on the basis of the laws, \( \lambda(t) \), for example, well determined in the industry condition, [23]. Both formulas, however, are generally associated with the data taken from the Fe-Zn phase diagram and additionally with the Lever Rule applied to the peritectic points of this phase diagram.

The compatibility of the solidification path and the results of the solute concentration measurement can be proved while calculating the \( \text{Zn} - \text{solute} \) concentrations of the \( \Gamma_1, P_2 \) and \( P_3 \) phases which is shown in Fig. 7.

The sum of different growth laws (determined previously for each phase sub-layer, separately, as shown in Fig. 5, for two sub-layers, only) is delivered in Fig. 9.

It also shows the specific sequence of growth, when the whole coating or some of its elements are examined, and not only each sublayer separately. The thick lines are referred to phases growing after the \( t_C \) threshold time, while phases growing in the period of the flux effect are indicated with thin lines, Fig. 9.

4. Concluding remarks

The presence of flux in the zinc bath significantly influences the occurrence of the (\( \text{Zn} \)) coating formation for the period of time, \( t \leq t_C \). First of all, the \( \delta = \delta_C - \peritectic \) phase has a different morphology in comparison with the \( \delta = \delta_P - \peritectic \) phase which appears after the flux disappearing. Additionally, the products of the flux decomposition/burning promote the \( \zeta - \peritectic \) phase nucleation and the \( \zeta_\eta = \zeta + \eta - \sublayer \) appearance. These products act as the substrates for the heterogeneous nucleation of the \( \zeta - \peritectic \) phase.

The function which describes the flux disappearing, Eq. (1), allows for presenting the Fe-Zn-F – virtual ternary phase diagram, Fig. 6. According to this ternary phase diagram the solidification path reduces to the path localized on the liquidus line for the Fe-Zn – binary phase diagram at the \( N_1 - \point \) well defined in Fig. 3. It proves that the \( \Gamma_1 - \phases \), according to the peritectic reaction, is not possible for the time \( t > t_C \), (when the solidification path reaches the \( N_1 - \point \)).

Even the sum of the growth laws, Fig. 9, allows for defining the \( t_C - \threshold \) time at which the flux effect onto the \( \text{(Zn)} - \coating \) formation is completed.

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