

MATHEMATICAL IDENTIFICATION OF HOMOGENISATION PROCESSES IN ARGON STIRRED LADLE

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Mathematical models processed results of experimental investigation obtained during ladle gas argon bubbling realised by stir elements situated in the ladle bottom. Exact theoretical description of processes occurring at argon bubbling into steel would be very complex and it would lead to a system of non-linear partial differential equations describing transfer of momentum, heat, components, and with excitation function in the form of equation of so called deterministic chaos (argon bubbling). On the basis of pouring ladle model diagram and concentrations courses, the simplified linear physically adequate model was proposed, which described behaviour of steel concentration in pouring ladle during its bubbling. The analysed process was understood in the form of a cybernetic model.

Key words: steel, ladle, gas argon bubbling, stir elements, physical modelling, mathematical identification

Matematička analiza procesa homogenizacije u loncu s argonskim miješanjem. Matematičkim modelima obrađeni su rezultati eksperimentalnog istraživanja homogenizacije u loncu s argonskim mješačima. Egzaktan teorijski opis procesa uvođenja mjehurića argona u čeličnu talinu bio bi vrlo kompleksan i vodio bi ka sustavu nelinearnih diferencijalnih jednadžbi za opis prijenosa momenta gibanja, topline i elemenata, uz dodatnu funkciju takozvanog determinističkog kaosa (gibanje mjehurića argona). Temeljem dijagramskog modela i koncentracijskih krugova predložen je pojednostavnjen, fizikalno odgovarajući linearni model, koji prikazuje promjenu koncentracije čelične taline u ljevačkom loncu tijekom upuštanja mjehurića. Analizirani proces smatran je pritom kibernetičkim modelom.

Ključne riječi: čelik, lonac, upuštanje argonskih mjehurića, mješači, fizikalno modeliranje, matematička identifikacija

INTRODUCTION

Previously presented paper “Physical Modelling of Bath Homogenisation in Argon Stirred Ladle“ was devoted to results of bath behaviour in the ladle model investigation during its gas argon bubbling realised by one or two stir elements situated in the ladle bottom. Study was performed with use of physical modelling method on a scale model 1 to 10. Development of homogenisation processes after start of bubbling was evaluated on the basis of electrical conductivity and temperature change, which were measured at three points of the ladle volume by conductivity and temperature sensors. The executed works were realised for conditions of 180 tons steel ladles.

It was found that the rate of homogenisation in the steel ladle is significantly dependent on the volume flow of argon. Mathematical method for the identification process was used for determining the optimum flow of argon.

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MATHEMATICAL DESCRIPTION OF PROCESSES AT ARGON BUBBLING

It was appropriate to create a physical-mathematical (physically adequate) model for the measured time courses of tracing substance concentrations in physical model of pouring ladle (PL) occurred during argon bubbling into the steel.

Description of situation

Schematic representation of situation at argon bubbling into bath in the model of steel ladle (mLP) is shown in Figure 1.

Argon bubbles flow from eccentrically situated stir element (P) in the bottom of mLP at constant volumetric flow (q). They „disrupt“ the layer of concentration enriched and coloured water with thickness (h_k) and there occurs gradual mixing of enriched and clear liquid (steel, water). Two (pressure) forces II and I act basically against each other on molecules of water in proximity of the sensors.

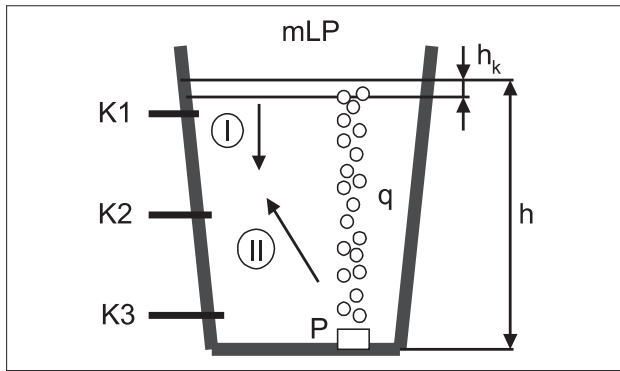


Figure 1. Argon bubbling into ladle model

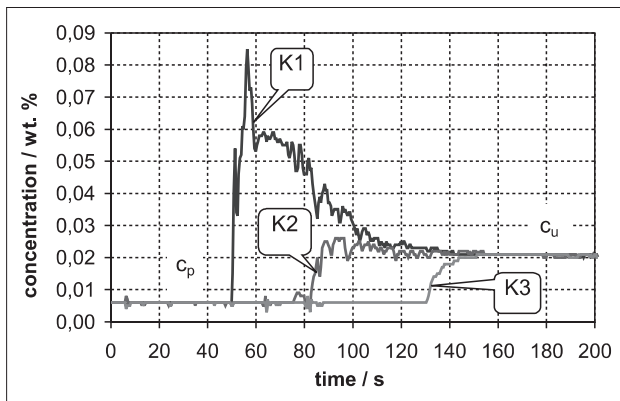


Figure 2. The time response of concentrations on the sensors K1, K2 and K3

Measured data

Analysis and synthesis of mathematical models was realised with use of the data measured on physical model, where courses of the measured concentrations, (with period of sampling $\Delta\tau \approx 0,5$ s), had on the sensors K1, K2 and K3 are shown in Figure 2.

Several facts are obvious from the development of concentration in the sensors:

- start and progress of gas argon bubbling into steel can be approximately considered in the form of the Heaviside unit step function and it is therefore possible to consider the development of concentration as a unit step response,
- the sensors reacted only after elapsing of certain “dead time”, which is proportional to the distance of individual sensors from the liquid level in the mLP,
- the overshoot of courses (apparently proportional to the magnitude of the force I or rather to the difference of the forces I and II) also descends with the distance from the liquid level in the mLP,
- steady-state (final) value of concentration is proportional to the proportion of volumes of pure water and water with enriched concentration.

Physical-mathematical model

Exact theoretical description of processes occurring at argon bubbling into steel would be very complex and it would lead to a system of non-linear partial differen-

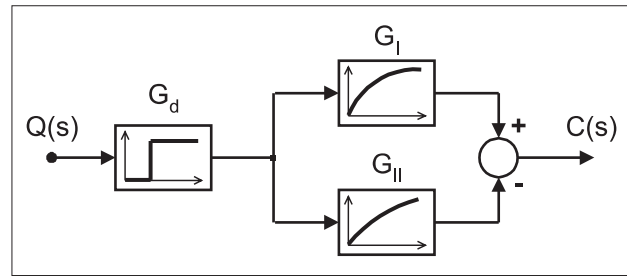


Figure 3. Block diagram of processes in ladle model (mLP)

tial equations describing transfer of momentum, heat, components, and with excitation function in the form of equation of so called deterministic chaos (argon bubbling). On the basis of the mLP diagram and concentrations courses, the simplified linear physically adequate model was proposed, which described behaviour of steel elements concentration in PL during its bubbling. Analysed process was understood in the form of a cybernetic model, which can be transparently visualised by so called block diagram shown in Figure 3.

It is series-parallel connection of three components, namely component of time delay and two parallel proportional (inertial) components acting against each other.

Two parallel and antagonistically connected simplest proportional systems with inertia of the 1st order (with transfers G_I a G_{II}) are assumed for the part of the model without time delay [1]:

$$G_I(s) = \frac{k_1}{\tau_1 s + 1}, G_{II}(s) = \frac{k_2}{\tau_2 s + 1} \tag{1}$$

where:

- k_1, k_2 - coefficients of transfer (amplification) of systems / $\% \times s / m^3$,
- τ_1, τ_2 - time constants of systems / s,
- s - complex variable in the Laplace transform / s^{-1} .

For this part of the model it is then possible to compose on the algebra basis of transfer the following continuous L-transfers (for zero initial conditions):

$$G(s) = \frac{k_1}{\tau_1 s + 1} - \frac{k_2}{\tau_2 s + 1} = \frac{(k_1 - k_2) + (k_1 \tau_2 - k_2 \tau_1) s}{(\tau_1 s + 1)(\tau_2 s + 1)} \tag{2}$$

Wherefrom for the L-image and original of transfer function $H(s)$ and $h(t)$ we get the final expression for standard concentration transfer function, which can be used at the same time as non-linear regression model F1 with three parameters k_1, τ_1 and τ_2 :

$$h(t) = c_n(t) = 1 - k_1 \cdot \exp(-t / \tau_1) + (k_1 - 1) \cdot \exp(-t / \tau_2) \tag{3}$$

Regression model for all concentrations

Figure 4 shows courses of the regression function based on the model F1, for standard concentration values in all three sensors K1, K2 and K3.

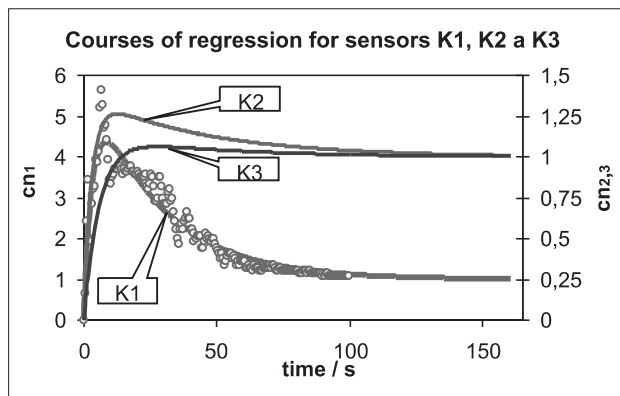


Figure 4. Courses of regression of F1 model for data

It is obvious from the results that the model F1 is suitable and usable for description of concentration course for all three sensors. It is also obvious that the transfer coefficient (gain coefficient) k_1 decreases with the distance of the sensor from the liquid level in the mLP, while the time constant τ_2 increases in this dependence.

USE OF MODEL FOR EVALUATION OF EXPERIMENTS

Proposed physically adequate model F1 was used for evaluation of the physical modelling results at argon bubbling in the ladle model. Its parameters were used for determination of other aggregated parameters, and all of them were used for definition of an optimum range of blown argon flow. The paper documents results of one stir element for position „A“ in the ladle bottom.

Approaches to solution

Three groups of parameters were chosen for characterisation of homogeneity „intensity“ measure in respect to blown argon flow q : basic physical parameters of model (2 parameters – see the model in Figure 3 and time delay, i.e. k_1 , k_2 , and τ_d), aggregated parameter defining derivative time constant (1 parameter, i.e. τ_D), and aggregated parameters defining maximum of model transfer function (2 parameters, i.e. t_m , respectively t_{max} , and h_m). Aggregated parameters were chosen in order to achieve simplicity, clarity and appropriate interpretability of homogenisation process.

Parameters of physically adequate model

Graphical dependencies of foregoing mathematical model parameters on blown argon flow for the sensor K2 are shown in Figure 5:

It is obvious from the Figure 5 that physically adequate model, in conformity with the assumed behaviour of real system, has evaluated a trend of dependence of the transport delay τ_d on the argon flow rate. At higher flow rate homogenisation starts sooner and runs more rapidly.

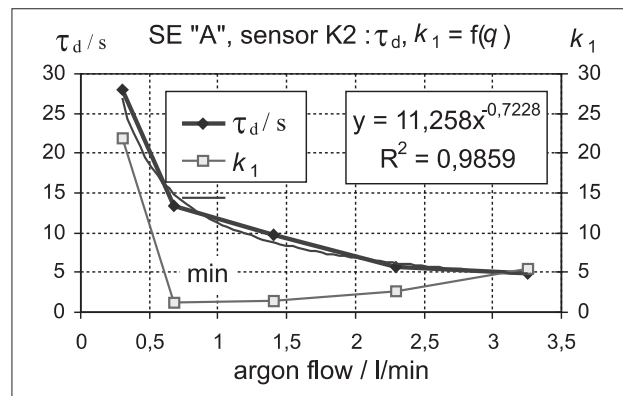


Figure 5. Dependencies of parameters τ_d , k_1 on argon flow for the sensor K2

Derivative time constant

Overall L-transfer of the defined model has character of so called real derivative member with a delay of the 2nd order [2], and derivative time constant τ_D is function of all three parameters. The constant τ_D can be expressed from the equation (2) with use of the relation:

$$\tau_D = k_1(\tau_2 - \tau_1) + \tau_1 \quad (4)$$

Lower values of this derivative time constant τ_D express faster reaction of the model on change of the input value, which is in this case represented by the volume flow of argon.

Graphical dependence of this time constant on the argon flow (for measurement at the sensor K2) is shown in Figure 6.

It is obvious from the Figure 6 that increase of the volume flow of argon to 1,4 l/min causes a significant step change of the value τ_D and therefore also more rapid changes in chemical composition of the model liquid (and development of homogenisation). The value of τ_D does not change significantly with further increase of the argon flow rate.

Transfer function overshoot

This approach uses analytical relations for determination of transfer function position and transfer function maximum values. Graphical dependencies of the as-

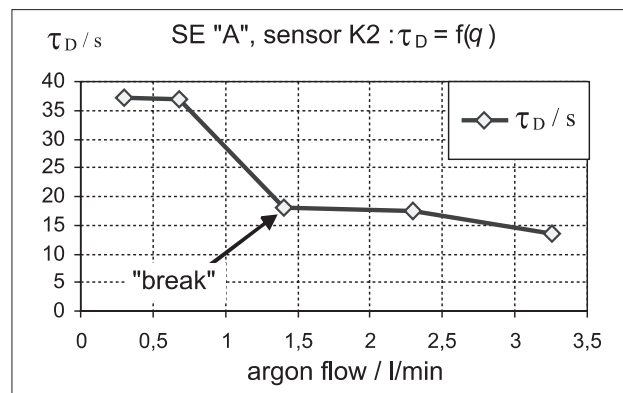


Figure 6. Dependence of derivative time constant τ_D on argon flow for the sensor K2

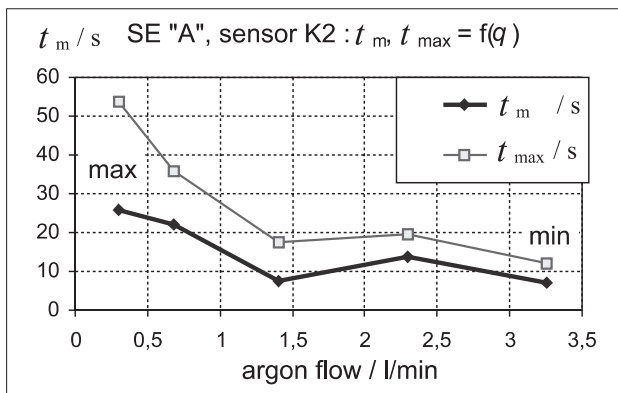


Figure 7. Dependencies of maximum times (t_m , t_{max}) of the flow of inert gas for the sensor K2

summed aggregated maximum parameters of the blown argon flow for the measurement at sensor K2 are shown in Figure 7 and Figure 8.

Diagram in Figure 7 shows development of times for achieving of maximum values in the impulse function in dependence on the argon flow rate (t_{mx} on the axis y is a general expression of time parameters t_m , which is a time of achieving the maximum regardless of transport delay τ_d , or τ_{max}). It is possible to obtain values of the parameter t_{max} by adding the parameter τ_m to the transport delay τ_d . Gradual convergence of t_m and t_{max} is given by decrease of the transport delay τ_d with increasing volume flow of argon. Faster circulation of volume of liquid results in faster achieving the maximum values of the impulse function. This speed, however, steeply falls at argon flow rates below 1,4 l/min.

The Figure 8 is devoted to a description of dependence of the maximum value of the impulse function (h_m – maximum value of dimensionless concentration) on the volume flow of argon. At its lowest flow rate (0,3 l/min) the injected substance gets to the sensor K2 in concentrated condition without being mixed. At increase of the argon flow rate to 0,68 l/min the faster dispersion of the marking substance at the expense of the speed of circulation of the volume of model liquid manifests itself the most from the viewpoint of this parameter (h_m). Further increase in argon flow rate brings about again a gradual enforcing of speed of circulation. At the highest argon flow rate (3,25 l/min) the rate of circulation is too high and injected substance gets to the sensor K2 without being able to disperse in the volume of the liquid.

CONCLUSION

Physically adequate mathematical model was proposed and verified for the measured time response of substance concentrations in the pouring ladle model bath.

This model was used for subsequent analysis of the input simulation parameters influence on its coefficients and aggregated parameters with consequences on appropriate (optimum) working mode setting for steel bubbling in the pouring ladle. Due to pre-set various

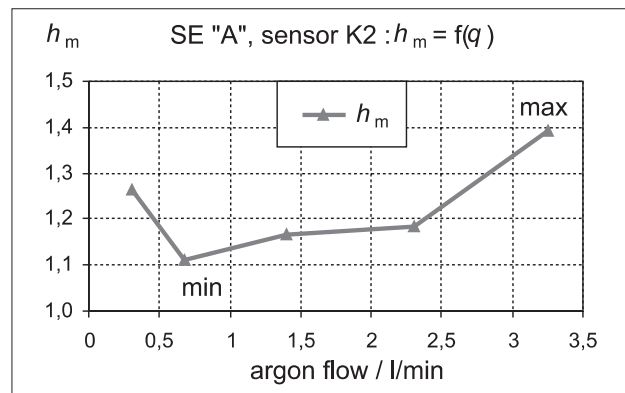


Figure 8. Dependencies of maximum values (h_m) on the flow of inert gas for the sensor K2

flows of argon blown into bath three groups of (aggregated) parameters for homogenisation intensity characterisation were determined and verified.

On the basis of the shape of their course, the argon flow optimum range was deduced – within limits from approx. 1,4 – 2,3 l/min (on the ladle model), which can be transformed with use of the appropriate volumetric flow scale factor to the realistic industrial equipment conditions, i.e. 243 - 400 l/min.

It was established in the previous paper entitled „Physical Modelling of Bath Homogenisation in Argon Stirred Ladle” that after achievement of a certain „breakpoint” volume flow its further increase brings low efficiency from the viewpoint of homogenisation times. Parameters assumed in this work enable moreover a more detailed investigation of the character of homogenisation process, correct setting of which forms an integral part of the advanced secondary metallurgy [3, 4]. For example magnitude of the maximum values of the impulse function (h_m) at individual argon flow rates gives evidence about behaviour e.g. of alloying additions injected into the volume of real pouring ladle at the initial stage of homogenisation.

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Note: The responsible translator for English language is B. Škandera, Czech Republic.