Neural Network Based Modeling and Control for a Batch Heating/Cooling Evaporative Crystallization Process

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Abstract. Crystallization processes have been widely used for separation in many fields to provide a high purity product. In this work, dynamic optimization and neural network (NN) have been applied to improve the quality of the product: citric acid. In the dynamic optimization, optimization problems maximizing both crystal yield and crystal size have been formulated. The neural networks have been developed to provide NN models to be used in the formulation of not only neural network inverse control (NNDIC) but also neural network model predictive control (NNMPC) strategies. The Levenberg Marquadt algorithm has been used to train the network and optimal neural network architectures have been determined by a mean squared error (MSE) minimization technique. In addition, a neural network model has been designed to provide estimates of the temperature and the concentration of the crystallizer. These estimates have been incorporated into the NNMPC controller. In the NNDIC controller, another neural network model has been applied to predict the set point of jacket temperature. The simulation results have shown that the obtained crystal size is increased by 19% and 30% compared to that by cooling and evaporation methods respectively and the obtained yield is increased more than 50%. The robustness of the proposed controller is investigated with respect to parameters mismatches. The results have shown that the NNMPC controller provides superior control performances in all case studies.

Keywords: Neural network, batch crystallization, optimization, heating/cooling evaporation.
1. Introduction

Batch crystallization, separation and purification behavior, is widely used in many fields such as food, chemical and pharmaceutical to produce high value-added specialty chemicals with high purity, desired Crystal Size Distribution (CSD) and shape. In general, batch cooling crystallization is one of the most method to apply in industrial processes owing to it operates easily. On the contrary, batch evaporative crystallization is economically more favorable than batch cooling crystallization if the solubility of solute depends on slightly temperature. With the object of enhancing the productivity of batch crystallization process, there are a number of researches focused on optimization and controlling both of batch crystallization by cooling and evaporation processes [1-6]. Paengjuntuek et al. [7] studied the optimization of batch crystallization processes. Mesbah et al. [8] presented a model-based control approach for optimal operation of a seeded fed-batch evaporative crystallization that was operated at isothermal and vacuum pressure.

Moreover, combining heating/cooling and evaporation in batch crystallization is the main topic of interest due to it can produce higher supersaturation, higher crystal yield and larger crystal size than all conventional crystallization. This method is only suitable for the system where the solubility varies significantly with temperature [9]. The combined evaporative crystallization with other process presents several advantages such as reduce equipment requirements, time cycle and saving cost of separation and recovery in some case. Choong and Smith [9] optimized and compared batch heating/cooling evaporative crystallization with the other crystallizations such as batch evaporative, semi-batch evaporative and batch cooling crystallizations. The results showed that the heating/cooling evaporative crystallization provided larger average crystal size 35%, 12% and 8% than crystallization by batch cooling and unseeded constant evaporation, batch seeded evaporative and semi-batch evaporative, respectively. In addition, the heating/cooling evaporative crystallization provided higher crystal yield than other modes of crystallization about 47%.

The batch crystallization process is a highly complex and nonlinear system. Therefore, an accurate mathematical model of the process is rarely available. Therefore, an artificial neural network approach, one of several techniques widely successfully applied to provide the modeling of nonlinear systems, is needed. An advantage of the implementation of the neural network is that it is easy to design and use and can provide the estimates of the relationship of input and output to explain the process. Kittisupakorn et al. [10] has applied the neural network to predict the concentration of a hydrochloric acid in hydrochloric acid recovery process. Arpornwichanop and Shomchoam [11] has applied neural network to estimate the substrate concentration in a fed-batch bioreactor. Moreover, there have been many researches applied the neural network to modeling nonlinear processes [12-19].

Normally, a production’s goal is to achieve product properties such as crystal size, purity and crystal size distributions that influence downstream processing operations such as filtering, drying and storage. However, since the batch crystallization process is a high complexity and nonlinear system, the control of this process based on the application of neural network is a significant challenge. For instance, Damour et al [20] presented the implementation of Neural Network Model Predictive Control (NNMPC) for an industrial crystallization process using model-based observers. A neural network model based on the estimates of crystal mass was used as internal model to predict the process output. Besides, their work compared the performances between NNMPC and industrial data from a PID-controlled process. Kittisupakorn et al. [21, 34-36] used neural network as a model in the model predictive control (MPC) algorithm for a steel pickling process. Furthermore, the neural network is incorporated into a model based predictive control technique in many researches [22-24].

The aim of this study is to improve the product quality of a citric acid-water system [9], regarding to a batch heating/cooling evaporative crystallization, by controlling the system at an optimal trajectory by Neural Network Model Predictive Control. A neural network model is developed to represent the process and the obtained model is used in the MPC strategy. To select proper neural network models, the Lenvenberg-Marquaudt algorithm is used to train the neural network and the structure of neural network is chosen based on minimum the mean squared error (MSE) of training and testing data.
2. Mathematical Model of Batch Heating/Cooling and Evaporative Crystallization

A mathematical model of batch heating/cooling and evaporative crystallization consists of two mass balances of solute and solvent, energy balance and crystallization model [9]. The assumptions made in the development of the process are: (1) agglomeration and breakage of crystals are negligible, (2) total nucleation rate is the sum of primary and secondary nucleation rates, (3) crystal size is independent growth, (4) the solution is well mixed and (5) crystal nuclei produced from primary and secondary nucleation have negligible size [25]. By performing mass balances, the following equations are obtained:

The mass balance of solute:

\[
\frac{dC_s}{dt} = -k_v \frac{d}{M} \left( m_3 M \right) - C_s \frac{d}{dt} \ln M
\]  
(1)

The mass balance of solvent:

\[
\frac{dM}{dt} = -Q_{evap}
\]  
(2)

where \( C_s \) is the concentration of solute, \( k_v \) is the volumetric shape factor, \( \rho_c \) is the density of crystals, \( m_3 \) is the third moment of crystal size, \( M \) is the mass of solvent and \( Q_{evap} \) is the evaporation rate.

The evaporation rate is a function of temperature and can be calculated by the following equation:

\[
Q_{evap} = \frac{a(P_v - P_a)A_s}{\sqrt{2\pi mRT}}
\]  
(3)

where \( P_v \) is the saturation vapor pressure, \( P_a \) is the partial pressure in the crystallizer, \( A_s \) is the evaporation surface area, \( a \) is the evaporation coefficient, \( m \) is the molecular weight and \( R \) is the gas constant.

The energy balance around the crystallizer is shown as follows:

\[
\frac{dT}{dt} = \frac{UA_j(T_j - T) - H_{evap}Q_{evap} + H_{crys}k_v\rho_t \frac{dm_3}{dt} M}{C_pM_{tot}}
\]  
(5)

\[
\frac{dT_j}{dt} = \frac{F_j \rho_j C_{pj}(T_{jp} - T_j) - UA_j(T_j - T)}{\rho_j C_{pj}V_j}
\]  
(6)

and

\[
H_{evap} = Ar \times abs(1 - T / T_j)^\nu
\]  
(7)

where \( T \) is the temperature in the crystallizer, \( U \) is the overall heat transfer coefficient, \( A_j \) is the total heat transfer surface area, \( T_j \) is the jacket temperature, \( H_{evap} \) is the heat of vaporization, \( H_{crys} \) is the heat of crystallization, \( C_p \) is the heat capacity of the solution, \( M_{tot} \) is the total mass of solution, \( F_j \) is the water flow rate in jacket, \( \rho_t \) is the density of water in jacket, \( C_{pj} \) is the heat capacity of the water in jacket, \( T_{jp} \) is the set point of jacket temperature and \( V_j \) is the jacket volume.

For the crystallization model can be described by following equations:

The crystal growth rate:

\[
G = k_g(T)(C_s - C^*)^\delta
\]  
(8)
where $G$ is the crystal growth rate, $k_g$ is the crystal growth rate constant, $C^*$ is the solubility of the solute and $g$ is the crystal growth rate exponent.

The Population Balance Equation with the assumption of crystal growth rate, variable volume and well mixed batch crystallizer can be given as:

$$\frac{\partial n}{\partial t} + \frac{\partial (Gn)}{\partial L} + n \frac{d (\ln M)}{dt} = 0$$

(9)

where $n$ is the number density of the crystals, $L$ is the characteristic crystal length and $M$ is the total mass of solvent at any instant of time.

Mostly, the moment transformation method is applied to convert Eq. (9). The moment of the crystal size distribution are given in the following expression:

$$\frac{dm_0}{dt} = (B_p + B_s) \cdot m_0 \frac{d (\ln M)}{dt}$$

(10a)

and

$$\frac{dm_i}{dt} = iGm_{i-1} + B l_{i} - m_i \frac{d (\ln M)}{dt}$$

(10b)

where $B_p$ and $B_s$ are the primary and secondary nucleation rate, respectively, $L_0$ is the characteristic crystal length of a newly formed crystal, and $m_0$ and $m_i$ are the zero and $i$ moment of crystal size ($i = 0, 1, 2, 3, 4, \ldots$), respectively.

For nucleation, both primary and secondary nucleation is used to describe in the empirical power law form:

Primary nucleation:

$$B_p = k_p(T)(C_s - C^*)^p$$

(11)

Secondary nucleation:

$$B_s = k_s(T)(C_s - C^*)^{p/\nu} M_{crys}^b$$

(12)

where $k_p$ and $k_s$ are the primary and secondary nucleation rate constant, respectively, $p$ and $s$ are primary and secondary nucleation rate exponent, respectively, $M_{crys}$ is the magma density of crystal and $b$ is the magma density exponent.

The solubility of the solute can be expressed by a polynomial equation.

$$C^* = C_0 + C_1 T + C_2 T^2 + C_3 T^3$$

(13)

In this work, the objective functions of optimization are to maximize the average crystal size (weight mean crystal size), $L_{aw}$ and the crystal yield. The expression for average crystal size and crystal yield are shown as follows:

$$L_{aw} = \frac{m_3}{m_3}$$

(14)

$$\text{crystal yield} = C_s(t_0) M(t_0) - C_s(t_f) M(t_f)$$

(15)

The crystallization kinetics and physical properties of citric acid-water system are shown in Table 1 and the initial values as shown in Table 2 which were excerpted from Bohlin and Rasmuson [28], Choong and Smith [9] as well as Alexander Apelblat [26].
3. Dynamic Optimization

The dynamic optimization computes the optimal operating temperature policy to control the crystallizer. In batch crystallization process, the quality of crystal effect to downstream process such as filtration, drying and storage [29]. In this study, the crystal quality which is crystal yield and crystal size is focused [9]. The aims of a dynamic optimization are to maximize the crystal yield and the crystal size. In this work, two optimization problems are considered.

Problem 1 (OPT1):

\[
\max \ m_d(t_f)/m_d(t_f) \quad (16)
\]

T(t) subject to the crystallizer model equations:

\[
T_{min} \leq T \leq T_{max} \quad (17)
\]

\[
L_{aw, min} \leq L_{aw} \leq L_{aw, max} \quad (18)
\]

\[
M_{min} \leq M \quad (19)
\]

Problem 2 (OPT2):

\[
\min C_s(t_f) \cdot M(t_f) \quad (20)
\]

T(t) subject to the crystallizer model equations:

\[
T_{min} \leq T \leq T_{max} \quad (21)
\]

\[
L_{aw, min} \leq L_{aw} \leq L_{aw, max} \quad (22)
\]

\[
M_{min} \leq M \quad (23)
\]

where \( T_{min} \) and \( T_{max} \) are chosen as 40 and 70 °C, respectively, \( L_{aw, min} \) and \( L_{aw, max} \) are chosen as 315 and 550 µm, respectively and the minimum total mass of solvent, \( M_{min} \), is 0.5 kg. The final batch time, \( t_f \), is 6500 s.

4. Neural Network

4.1. Neural Network Forward Model

The neural networks consist of an input layer for receiving data from external source, one or some hidden layer(s) for processing input data and an output layer for displaying the output values. There are some neurons in each layer that are connected with some connections to previous and next layers. A neuron includes input and output values, weight factors as well as bias and transfer function. The neural networks are trained using a training algorithm and a training data set in order to adjust the connection weights and biases.

For this neural network, the weight factors and biases are the coefficients that determine the relationship of the network inputs by randomly. The neural network uses tan-sigmoid as activation function of the nodes in hidden layer and linear function is used as the activation function in its output layer. The Levenberg-Marquardt algorithm is applied for training the network. The objective of neural network training is to minimize the error function which is mean square error (MSE) between the predicted neural network values and actual targeted value. Eq. (24) is shown the MSE calculation.

\[
MSE = \frac{1}{n} \sum_{i=1}^{n} (T_{net} - T_{pt})^2 \quad (24)
\]
where \( n \) is the number of data, \( T_{ac} \) is the actual targeted temperature value and \( T_p \) is the predicted neural network value.

The steps of neural network structure designing are summarized in Fig. 1. First step is preparing input and output data for training, validation and testing which are obtained by solving Eq. (1) – (13) and varies the value of manipulated variable (the set point of jacket temperature, \( T_{jsp} \)) in step changes. Second step is to design the network structure and choose the error function. Next are to initialize the weight factors, train the network, test and validate the network and then examine the error. If the error is unacceptable, the weight factors or structure will be reset to new values but if the error is acceptable, the neural network model will be obtained.

Figure 2 shows the input and output chosen to represent the forward model which is 6 input nodes that consists of \( C_S(k-1), C_3(k), T(k-1), T(k), T_{ip}(k-1) \) and \( T_{jp}(k) \) as well as 2 output nodes that compose of \( C_r(k+1) \) and \( T(k+1) \).

The input and output data sets which consist of training, testing and validating data sets are obtained from mathematical model to train the neural network by varied the set point of jacket temperature. In the generated data for network training consist of plant certainty case called nominal case and plant uncertainty cases that compose of -30% of \( U \), +30% of \( H_{crys} \), -30% of \( H_{evap} \), +30% of \( k_g \) and +30% of \( k_p \). The input and output data sets which are trained in neural network forward model are 18 sets. The data sets which are generated from mathematical model need to be nominalized in order to overcome the significant minimum and maximum values used in the training process. The raw process data generated are scaled down to between 0.05-0.95 using the following equations:

\[
value_{sd} = \frac{\left( value_{ac} - value_{min} \right) \left( 0.95 - 0.05 \right)}{\left( value_{max} - value_{min} \right)} + 0.05
\]

and the actual value (scaled up) is given by

\[
value_{ac} = \frac{\left( value_{sd} - 0.05 \right) \left( value_{max} - value_{min} \right)}{0.95 - 0.05} + value_{min}
\]

where \( value_{sd}, value_{ac}, value_{min}, value_{max} \) are the scaled down, actual, minimum and maximum values of the data, respectively.

The input and output data sets which are nominalized are integrated and randomized to train test and validate the neural network. The training, testing and validating data set are 60%, 30% and 10% of the all data, respectively.

### 4.2. Neural Network Inverse Model

In this part, the neural network direct inverse control is used for control the crystallization process by training neural network inverse model. These steps of neural network structure designing are same as the neural network model (Fig. 1). The neural network inverse model is applied to predict the future value of the manipulated variable (the set point of jacket temperature, \( T_{ip} \)). The input and output data pattern of neural network inverse model is given in Fig. 3.

The input and output of inverse model is shown in Fig. 3 which is 9 input nodes that consists of \( \text{error}(k-1), \text{error}(k), C_s(k-1), C_s(k), T(k-1), T(k), T(k+1), T_{ip}(k-1), T_{jp}(k) \) as well as 1 output nodes that is \( T_{ip}(k+1) \).

The input and output data sets are generated from closed-loop control that is feedback controller (PID) to train the neural network [30-33]. In this section, the generated data for network training, which are the same as the forward model, consist of nominal and plant uncertainty cases (-30% of \( U \), +30% of \( H_{crys} \), -30% of \( H_{evap} \), +30% of \( k_g \) and +30% of \( k_p \)). In addition, the data sets of 18 sets, used to train the neural network inverse model generated from mathematical models, are normalized based on the equation 25. Then the data sets are combined and randomized. The data sets are divided into 3 groups which consist of 60% training data set, 30% testing data set and 10% validating data set.

### 5. Neural Network Model Based Predictive Controller
Model predictive control using a neural network has been focused by some researchers [21-24]. The neural network model based predictive control strategy in this work is shown in Fig. 4. The neural network forward model is used to predict the future process outputs over the prediction horizon (p). The predictions are passed to the optimization which minimizes a specified objective function. Sequential quadratic programming (SQP) algorithm is used to solve the optimization problem of minimizing as given in Eq. (27). The objective function of the model predictive control strategy is described as follows:

$$\min \sum_{i=1}^{p} W_1 \left[ T \left( k + i \right) - T_{sp} \left( k + i \right) \right]^2 + W_2 \left[ \Delta T_{sp} \right]^2$$  \hspace{1cm} (27)

Subject to

$$25 \ C \leq T_{sp} \left( k + 1 \right) \leq 100 \ C, \quad i = 1, 2, 3, \ldots, p$$ \hspace{1cm} (28)

$$T \left( k + p \right) = T_{sp} \left( k + p \right)$$ \hspace{1cm} (29)

where \( p \) is a parameter specifying the prediction horizon, \( W_1 \) is weighting parameter used to give different weights to different squared tracking error, \( W_2 \) is weighting parameter of different squared of manipulated variable and \( T_{sp} \) is the set point of crystallizer temperature as obtained from optimization.

The simulations have been carried out using the neural network model to find a set of suitable control parameter \( W_1, W_2, p \) and \( m \). The prediction horizon \( p \) corresponds to the future time interval used to compute prediction with the neural network model. The control horizon is equal or less than the prediction horizon (\( m \leq p \)). In the same IAE performance case, the choice of \( W_1 \) and \( W_2 \) are chosen 1 and 2, respectively. The prediction horizon and control horizon are set at 6 and 6, respectively. In the best performance case, \( W_1 \) and \( W_2 \) are varied and then fixed at 1 and the prediction horizon and control horizon are set at 10 and 1, respectively.

6. Results and Discussion

6.1. Dynamic Optimization

The dynamic optimization calculates the optimal temperature profile by optimizing the objective function. Figure 5 shows crystallizer temperature profiles that provide maximum average crystal size and Fig. 6 shows crystallizer temperature profiles that provide maximum crystal yield. In the optimization order, it classified into 4 sets which consist of 1, 5, 8 and 10 intervals. In the optimization problems 1 and 2 get the maximum average crystal size and crystal yield at 10 intervals. The dynamic optimization is applied to be a set point of crystallizer temperature in PID control, neural network direct inverse control (NNDIC) and neural network model based predictive control. Moreover, it is applied in nominal case and mismatch case for controlling.

The final product qualities in terms of average crystal size \( (L_{aw}) \) and crystal yield are shown in Table 3. It can be seen that OPT1 provides larger average crystal size than OPT2 about 6.46% but OPT2 gives more crystal yield than OPT1 about 3.39%. Moreover, the average crystal size and crystal yield which are obtained from the heating/cooling evaporative crystallization method are more than cooling and evaporation method. From Table 4, it can be seen that the heating/cooling evaporative crystallization method produce the average crystal size of 19% and 30% larger than cooling and evaporation method, respectively as well as the obtained crystal yield from heating/cooling evaporative crystallization method is higher than cooling and evaporation method about 50%.

6.2. Neural Network Forward Model

The simulation demonstrates that the well-trained neural network model can be applied to predict the concentration and crystallizer temperature profiles in this process. The optimal neural network structure consists of 6 nodes in input layer, 14 nodes in first hidden layer, 12 nodes in second hidden layer, and 2 nodes in output layer and can be written in the form of [6-14-12-2]. It can be seen that the neural network forward model gives good accuracy for the prediction of the system. The mean squared error index for the data test and validate of neural network forward model are shown in Table 5.
6.3. Neural Network Inverse Model

In this part, the optimal structure of the neural network inverse model consists of 9 nodes in the input layer, 2 nodes in the first hidden layer, 12 nodes in the second hidden layer, and 1 node in the output layer and can be written in the form of [9-2-12-1]. The neural network inverse model can provide sufficiently good jacket temperature set point, \( T_{jsp} \). The mean squared error index of both testing and validation are given in Table 6.

6.4. Comparisons among PID, NNDIC and NNMPC Controllers

Here, the optimal temperature profiles obtained from the optimization with the objective function 1 and 2 (OPT1 and OPT 2) are computed and the PID, NNDIC and NNMPC controllers have been formulated to provide tracking of the profile by manipulating of the jacket temperature set point. The optimal crystallizer temperature profiles are shown in Figs. 5-6. In this case, the optimization 1 (OPT1) is chosen to study. The simulations are divided into 2 cases: a nominal case and parameter mismatches case.

6.4.1. Nominal case

In this case, the PID and NNMPC controllers are tuned to achieve their best control performances. Table 7 shows the best performances of the PID and NNMPC controllers. Figures 7-8 show the response profiles of crystallizer temperatures and the set point of jacket temperature. It can be seen that both controllers can bring the temperature closely to set point with small overshoot and oscillation. However, the PID controller provides more overshoot and oscillation than the NNMPC controller does.

6.4.2. Parameter mismatches case

The robustness of the controller is studied according to parameter mismatches. The parameter mismatches consist of 5 cases: decreasing 30% of \( U \), increasing 30% of \( H_{crys} \), decreasing 30% of \( H_{evap} \), increasing 30% of \( k_p \) and increasing 30% of \( k_g \) from its nominal value. The control performances of both controllers regarding to parameter mismatches are divided into 6 scenarios shown in Table 7. Figures 9-11 show the response profiles of crystallizer temperature and the set point of jacket temperature in parameter mismatch case (-30%U). Table 7 summarizes the control performance of the PID, NNDIC and NNMPC controllers in terms of the absolute error (IAE). It can be seen that the NNMPC controller provides superior control performance in all cases.

7. Conclusions

In this study, the implementation of a dynamic optimization and neural network of a batch heating/cooling evaporative crystallizer for production of citric acid is presented. This neural networks use tan-sigmoid and linear as activation functions. The Levenberg-Marquardt algorithm is applied for training the network. The optimal structure of neural network forward model consists of 6 nodes in input layer, 14 nodes in first hidden layer, 12 nodes in second hidden layer, and 2 nodes in output layer. The optimal structure of neural network inverse model consists of 9 nodes in the input layer, 2 nodes in first the hidden layer, 12 nodes in the second hidden layer, and 1 node in the output layer. In addition, the optimization problems 1 and 2 get the maximum average crystal size and crystal yield, respectively. The optimal product qualities based on two objective functions are carried out to obtain average crystal sizes and crystal yields. Furthermore, the neural network model predictive control (NNMPC) has been formulated and tested under plant/model mismatches. It was found that the NNMPC can provide superior control performance than the PID in all cases.

Acknowledgment
The authors would like to acknowledge the support from the Faculty of Engineering, Burapha University under the contract # 36/2553 and The Institutional Research Grant (The Thailand Research Fund)(IRG 5780014) and Chulalongkorn University (Contract No. RES_57_411_21_076).

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Appendix I: List of Tables:

Table 1. Crystallization and physical properties of citric acid–water system.

| Symbol | Unit | Value |
|--------|------|-------|
| $A_j$  | m²   | 0.07  |
| $A_r$  | kJ/kg | 2891.83 |
| $A_s$  | m²   | 0.019 |
| $B$    |      | 0.84  |
| $C_0$  | kg/kgH$_2$O | 0.91176 |
| $C_1$  | kg/kgH$_2$O °C | 0.034857 |
| $C_2$  | kg/kgH$_2$O °C$^2$ | $-2.8785 \times 10^{-4}$ |
| $C_3$  | kg/kgH$_2$O °C$^3$ | $3.7228 \times 10^{-6}$ |
| $C_p$  | kJ/kg °C | 2.1745 |
| $C_{pj}$ | kJ/kg °C | 4.184 |
| $F_j$  | kg/s | 0.00003 |
| $G$    |      | 0.65  |
| $H_{crys}$ | kJ/kg | 117 |
| $k_g$  | m/(s(kg/kg)$^g$) | $0.02652 \exp(-3584/T)$ |
| $k_p$  | no./(kg/kg)(kgH$_2$O s) | $1.0 \times 10^{-7}$ |
| $k_v$  | no./(kg/kg)(kg/kg)(kgH$_2$O s) | $0.88774 \exp(4781/T)$ |
| $L_0$  | m    | $10 \times 10^{-6}$ |
| $M$    | kg/kmol | 18 |
| $M_{crys}$ | kg/kgH$_2$O | 0.024 |
| $N_r$  |      | 0.321 |
| $P$    |      | 3.54  |
| $P_a$  | kPa  | 10.1325 |
| $R$    | m³ Pa/K mol | 8.314 |
| $S$    |      | 0.543 |
| $T_c$  | K    | 647.13 |
| $U$    | W/m² °C | 0.5 |
| $V_j$  | m³   | 0.0014 |
| $A$    |      | 0.5   |
| $q_c$  | kg/m³ | 1540 |
| $q_l$  | kg/m³ | 1000 |

Table 2. Initial value in this crystallization process.

| Symbol | Value |
|--------|-------|
| $T_m$  | °C    |
| $M$    | kg    |
| $M_{crys}$ | kg/kgH$_2$O | 2.871 |

Table 3. The simulation results obtained from two dynamic optimization problems.

| Problem | $L_{aw}(\mu m)$ | Crystal yield (kg) |
|---------|-----------------|--------------------|
| OPT1    | 451             | 1.585              |
| OPT2    | 420             | 1.641              |
Table 4. Comparison of the product quality among different crystallization method.

| Method                          | L_{90} (µm) | Crystal yield (kg) |
|---------------------------------|-------------|--------------------|
| Unseeded cooling\(^a\)          | 296         | 0.787              |
| Evaporation\(^a\)               | 315         | 0.787              |
| Heating/cooling evaporative     | 451         | 1.585              |

\(^a\) Obtained from Choong and Smith [18].

Table 5. Mean squared error value of testing and validation in the neural network forward model.

| Data sets | No. of samples | Mean squared error |
|-----------|----------------|--------------------|
| Testing data 1 | 583         | 5.3528x10^{-5}    |
| Testing data 2 | 583         | 8.0095x10^{-5}    |
| Validation data | 389         | 4.2852x10^{-5}    |

Table 6. Mean squared error value of testing and validation in the neural network inverse model.

| Data sets | No. of samples | Mean squared error |
|-----------|----------------|--------------------|
| Testing data 1 | 583         | 9.4699x10^{-6}    |
| Testing data 2 | 583         | 8.3586x10^{-6}    |
| Validation data | 388         | 1.1091x10^{-5}    |

Table 7. Performance of controller for nominal and parameters mismatch cases.

| Cases                                           | IAE values |
|-------------------------------------------------|------------|
|                                                 | NNMPC | PID | NNDIC |
| Nominal                                         | 41.21  | 67.90 | 79.85 |
| -30%U                                           | 44.96  | 81.34 | 93.76 |
| +30%k\(_p\)                                     | 41.43  | 67.91 | 80.29 |
| +30%k\(_g\)                                     | 41.40  | 68.00 | 81.12 |
| +30%H\(_crys\)                                  | 42.13  | 68.75 | 80.87 |
| -30%H\(_{evap}\), +30%H\(_{crys}\)              | 41.49  | 69.00 | 80.01 |
| -30%U, +30%H\(_{crys}\), -30%H\(_{evap}\), +30%k\(_g\), +30%k\(_p\) | 45.42  | 84.70 | 98.13 |
Appendix II: List of Figures

Prepare input and output data for training, validation and testing from mathematical model

Design the structure of network (the number of hidden layers and nodes in hidden Layer, Activation function) and choose the error function

Initialize the weight factors

Train the network with training data sets

Test and validate the network

Examine the error

Obtain the neural network model

Set the new network structure

Change the weight factors

Fig. 1. Steps of neural network structure designing.

Neural network forward model

| \(C_i(k-1)\) | \(C_i(k)\) | \(C_i(k+1)\) |
|---|---|---|
| \(T(k-1)\) | \(T(k)\) | \(T(k+1)\) |
| \(T_{\text{inf}}(k-1)\) | \(T_{\text{inf}}(k)\) |

Output

Input

Fig. 2. Input and output data pattern for forward model.

Neural network inverse model

| error(k-1) | error(k) | error(k+1) |
|---|---|---|
| \(T(k-1)\) | \(T(k)\) | \(T(k+1)\) |
| \(C_i(k-1)\) | \(C_i(k)\) |
| \(T_{\text{inf}}(k-1)\) | \(T_{\text{inf}}(k)\) |

Output

Input

Fig. 3. Input and output data pattern for inverse model.
Fig. 4. The neural network based model predictive control strategy.

Fig. 5. Temperature profile for the optimization problem 1.
Fig. 6. Temperature profile for the optimization problem 2.

Fig. 7. The crystallizer temperature control and the manipulated variable (T_{jsp}) in nominal case of PID controller.
Fig. 8. The crystallizer temperature control and the manipulated variable ($T_{jsp}$) in nominal case of NNMPC controller.

Fig. 9. The crystallizer temperature control and the manipulated variable ($T_{jsp}$) in parameter mismatch case (-30% U) of PID controller.
Fig. 10. The crystallizer temperature control and the manipulated variable ($T_{jsp}$) in parameter mismatch case (-30%U) of NNDIC controller.

Fig. 11. The crystallizer temperature control and the manipulated variable ($T_{jsp}$) in parameter mismatch case (-30%U) of NNMPC controller.