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Integrated Process Design and Control of Reactive Distillation Processes

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Abstract: In this work, integrated design and control of reactive distillation processes is presented. Simple graphical design methods that are similar in concept to non-reactive distillation processes are used, such as reactive McCabe-Thiele method and driving force approach. The methods are based on the element concept, which is used to translate a system of compounds into elements. The operation of the reactive distillation column at the highest driving force and other candidate points is analyzed through analytical solution as well as rigorous open-loop and closed-loop simulations. By application of this approach, it is shown that designing the reactive distillation process at the maximum driving force results in an optimal design in terms of controllability and operability. It is verified that the reactive distillation design option is less sensitive to the disturbances in the feed at the highest driving force and has the inherent ability to reject disturbances.

Keywords: Process design, Process control, Driving force, Reactive distillation, Element-based method

1. INTRODUCTION

Traditionally, process design and process control are considered as independent problems, that is, a sequential approach is used where the process is designed first, followed by the control design. The limitations with the sequential approach are related to dynamic constraint violations, for example, infeasible operating points, process overdesign or under performance. Therefore, this approach does not guarantee robust performance (Seferlis and Georgiadis, 2004). Furthermore, process design decisions can influence process control and operation. To overcome the limitations associated with the sequential approach, operability and controllability are considered simultaneously with process design, in order to assure that design decisions give the optimum operational and economic performance. In control design, operability addresses stability and reliability of the process using a priori operational conditions and controllability addresses maintaining desired operating points of the process subject to disturbances.

A number of methodologies and tools have been proposed for addressing the interactions between process design and control, and they range from optimization-based approaches to model-based methods (Luyben and Floudas, 1994; Nikacevic et al., 2012).

In this work, integrated design and control of reactive distillation processes is considered, since process design decisions will influence process operability and controllability. Numerous design algorithms for multi-component separation systems with reactions have accompanied the increasing interest in reactive distillation processes. In design, the input and (selected) output variables are specified and the task is to determine the optimal reactive distillation process configuration (for example, minimum number of stages), and the optimal design parameters (for example, optimum reflux ratio, optimal feed location) that achieve the given product specification. It is intended to achieve the optimal design in such way that it is also an operable process at pre-defined conditions under presence of disturbances.

Pérez-Cisneros et al. (1997) have proposed an element mass balance approach to design the reactive distillation processes, which employs the traditional graphical tools similar in concept to design of non-reactive distillation columns, such as McCabe-Thiele method and driving force approach by Bek-Pedersen and Gani (2004). Moreover, Hamid et al. (2010) have proposed an integrated process design and controller design methodology. However, their methodology covers the aspects related to design and control of non-reactive binary distillation processes. In this work, the method of Hamid et al. (2010) is extended to also cover a ternary compound reactive distillation process (using element-based approach) and criteria of selecting the optimal design and the controller structure selection will be presented. In order to demonstrate the application of the aforementioned approach, production of methyl-tert-butyl-ether (MTBE) from methanol and isobutene using a reactive distillation column is considered.

2. REACTIVE DISTILLATION COLUMN DESIGN

The computation of simultaneous chemical and physical equilibrium plays an important role in the prediction of the
limits for conversion and separation of a specific reactive separation process, particularly for the reactive distillation systems. Using the Gibbs free energy minimisation approach, Pérez-Cisneros et al. (1997) proposed solution procedures where the multicomponent chemical and physical equilibrium is posed as an “element phase” equilibrium problem. This transformation is based on the concept of chemical model as proposed by Michelsen (1989). This concept is derived from chemical model theory, where, the equations of chemical equilibrium together with any appropriate physical model yielding the chemical potentials are incorporated into an element-based model (called the chemical model). The main difference between the chemical model algorithm and those developed earlier, is the use of the chemical models in a way that renders the chemical and physical equilibrium problem formally identical to the physical equilibrium problem for a mixture of element (representing the system). Further details can be found in (Pérez-Cisneros, 1997; Daza et al., 2003). The reaction for MTBE synthesis is given as follows:

\[
\text{Isobutene} (C_4H_8) + \text{Methanol} (CH_3O) \rightarrow (C_7H_{12}O)
\]

It is evident that the selection of the elements has an important role in the present formulation. They are traditionally chosen as the “natural” chemical elements present in the reaction mixture, but, indeed, one is free to select any reaction invariant fragment of the reactants. The element matrix is constructed based on the rules provided by Pérez-Cisneros et al. (1997) and it is as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>C₄H₈ (1)</td>
</tr>
<tr>
<td>B</td>
<td>1</td>
</tr>
</tbody>
</table>

Therefore, the ternary system of compounds can be reduced into a binary system of elements A and B and the reaction can be rewritten as: \(A + B \rightarrow \) . The first component (element \(A\)) and the second component (element \(B\)) form the third element (element \(AB\)). Having the ternary system of compounds represented in form of a binary element system, similar graphical design methods, that are applied to non-reactive binary distillation column design, such as McCabe-Thiele method can be used. However, in order to use the McCabe-Thiele method, a reactive equilibrium curve is required. The reactive equilibrium curve is constructed through sequential computation of reactive bubble points (Pérez-Cisneros, 1997). In order to generate the reactive data set, Wilson thermodynamic model for prediction of the liquid phase behaviour and SRK equation of state for prediction of vapour phase behaviour were used. Note that the calculation of reactive vapour-liquid equilibrium (VLE) data set is in terms of compounds. Therefore, a ternary compound data set is obtained. To convert this data set to be represented in form of a binary element system the following expressions are used where mole fractions of elements \(A\) and \(B\) are calculated in the liquid phase:

\[
W'_\text{v} = \frac{x_1 + x_2}{x_1 + x_2 + 2 \cdot x_3}
\]

In the above equations \(W'_\text{v}\) and \(W'_\text{b}\) are the liquid mole fractions of elements \(A\) and \(B\), respectively. For calculation of the element mole fractions in vapour phase (\(W'_\text{v}^A\) and \(W'_\text{v}^B\)), the equations used are the same as (1) and (2) where instead of liquid molar fraction (\(x_i\)), the vapour molar fraction (\(y_i\)) is used. Fig. 1 depicts the temperature (\(T\))-\(W'_\text{d}\) diagram for MTBE reactive system.

![Fig. 1. T-W'_d diagram for MTBE reactive system (P = 1 atm).](image)

The design task is to separate a binary element mixture that is 70 mole percent element \(A\) (\(x_{W'_d} = 0.7\), \(x_{W'_d} = 0.3\)) into 50 element mole percent bottoms product (\(W'_\text{d} = 0.50\)) and 99 element mole percent distillate (\(W'_\text{d} = 0.99\)) product. Note that based on the binary element reaction matrix, element \(A\) and \(B\) correspond to isobutene and methanol, respectively. The element feed flow rate is 100 Kg-mole element/hr at 300K and 1 atm. The operating pressure of the reactive distillation column is 1 atm and pressure drop across the column is assumed to be negligible. The reflux element ratio (RR) is 2. The physical and chemical equilibrium curve is constructed using the data set presented in Fig. 1. Theoretical reactive stages are calculated from the reactive McCabe-Thiele method. A partial reboiler, total condenser and chemically saturated liquid reflux are set for the column. In order to design the described reactive distillation column for MTBE synthesis, McCabe-Thiele method is used. Fig. 2 depicts the reactive distillation column design using reactive McCabe-Thiele method. As it is shown in Fig. 2, the reactive distillation column has five reactive stages.

Daza et al. (2003) have extended the driving force (DF) method for non-reactive systems (Bek-Pedersen and Gani, 2004) to include reactive systems. Similar in concept to non-reactive systems, the driving force is defined as the difference in composition between two coexisting phases. The driving-force design method for reactive as well as non-reactive distillation systems is based on the availability of data for the vapour-liquid behaviour. In the case of reactive systems, the vapour-liquid equilibrium data must be based on the elements (see Fig. 1).
The driving-force diagram can only exploit binary interaction between compounds or elements in two coexisting phases, or two compounds on a solvent-free basis. Note that the element-based driving-force diagram fully incorporates the extent of reaction on an element basis, and, therefore, it can be applied in the design of reactive distillation columns. Provided the element vapour-liquid behaviour data exist, or can be computed (which is the case in this study, see Fig. 1), the reactive driving-force diagram can be obtained using (3) with respect to the elements.

\[
DF_i = W^i_x - W^i_y = \frac{W^i_x \alpha_y}{1 + W^i_x (\alpha_y - 1)} - W^i_y
\]

(3)

The driving force concept is used to find the optimal (design target) values of the process variables for separation systems. Based on identification of the largest driving force (see Fig. 3), defined as the difference in composition of a component \(i\) between the vapour phase and the liquid phase, which is caused by the difference in the volatilities of component \(i\) and all other components in the system. Fig. 3 shows the driving force diagram for MTBE reactive system at 1 atm. As the driving force decreases, separation becomes difficult and may become infeasible when the driving force approaches zero. On the other hand, as the driving force approaches its maximum value, the separation becomes easier. Therefore, from a process design point of view, a separation process should be designed/selected at the highest possible driving force, which will naturally lead to the optimal design with respect to the energy consumption (Bek-Pedersen and Gani, 2004).

In this work, the optimal feed location of the reactive distillation column is determined using the driving force diagram. Reactive McCabe-Thiele method has been only used to determine the number of stages. The feed and product specifications are already known since they were used in the reactive McCabe-Thiele method. The optimal feed location at the maximum driving force can be found using (4).

\[
N_F = N(1 - D_s)
\]

(4)

In (4), \(N\) is the number of stages which was obtained from the reactive McCabe-Thiele method (was found to be 5); \(D_s\) is the value corresponding to the maximum driving force on the \(x\)-axis \((D_s = 0.61)\). The optimal feed location is identified using the additional rules for driving force (Bek-Pedersen and Gani, 2004) and therefore it is stage 1 from the top of the column.
\[
\frac{dy}{dd} = \left( \frac{dy}{d\theta} \right) \left( \frac{d\theta}{dx} \right) \left( \frac{dx}{dd} \right)
\]  (5)

The reactive element operating lines for the rectifying section and stripping sections are given in (10) and (11). RR is the element reflux ratio, and RB is the element reboil ratio.

\[
W' = \frac{RR + 1}{RR} W'_d + \frac{1}{RR + 1} W'_d
\]  (10)

\[
W'_d = \frac{RB + 1}{RB} W'_d - \frac{1}{RB} W'_d
\]  (11)

Substituting (10) and (11) in (3) for \( W'_d \) gives the top and bottom element product composition with respect to the driving force in (12) and (13) which is:

\[
W'_d = DF \left( RR + 1 \right) + W'_d
\]  (12)

\[
W'_d = W'_d - DF; RB
\]  (13)

Equations (12) and (13) are differentiated with respect to \( DF_i \) and result in the following expressions:

\[
\frac{dW'_d}{dDF_i} = \left( RR + 1 \right) + \left( \frac{dW'_d}{dDF_i} \right)^{-1} W'_d
\]  (14)

\[
\frac{dW'_b}{dDF_i} = W'_d - RB \left( \frac{dDF_i}{dW'_d} \right)^{-1} W'_d
\]  (15)

The total element \( A \) mass balance is written as follows:

\[
F_i z_{W}\ = W'_d b^d + W'_b b^b
\]  (16)

Where, \( b^d \) and \( b^b \) are element \( A \) mass flows in top and bottom of the column, respectively. Substituting (12) and (13), one at the time, into (16) for \( W'_d \) and \( W'_b \), the total element \( A \) mass balance in terms of driving force is expressed as:

\[
F_i z_{W}\ = DF_i \left( RR + 1 \right) b^d + W'_d b^d + W'_b b^b
\]  (17)

or

\[
F_i z_{W}\ = W'_d b^d + W'_d b^d - b^b F_i RB
\]  (18)

Differentiating (17) and (18) with respect to the \( F_i \) and \( z_{W}\) (assuming that the changes in composition, and top and bottom element flowrates \( b^d \) and \( b^b \) with respect to the feed flowrate is negligible), the expressions for \( dW'_d / dF_i \), \( dW'_b / dW_i \) are obtained. Having these derivatives, the solution to (9) is expressed by (19). Note that in (19), \( a_1, ..., a_8 \) are constants.

Values of \( dDF_i / dW'_d \) are calculated and shown in Fig. 4. Note that in Fig. 4, two other points (points II and III) which are not at the maximum are identified as candidate alternative designs for a distillation column, which will be used for verification purposes.

It must be noted that the expressions for \( dW'_d / dDF_i \) and \( dW'_b / dDF_i \) in (19) are equal to 1 at point (I) in Fig. 4 (maximum driving force) and greater than 1 in any other point.
Fig. 4. Driving force diagram for \( W_L-W_H \) separation (reactive zone only – top figure) and its corresponding derivative of \( FDI \) with respect to \( W_L \) (bottom figure).

Furthermore, at point A the value of \( \frac{dF_D}{dW_L} \) is equal to zero. Therefore, equation (19) at Point A (maximum driving force) can be expressed as:

\[
\frac{dy}{dd} = \begin{bmatrix} \frac{dW_d}{dF_f} & \frac{dW^*_d}{dF_f} \\ \frac{dW^*_d}{dz_{W^*_d}} & \frac{dW^*_d}{dz_{W^*_d}} \end{bmatrix} \begin{bmatrix} 0 \\ 0 \end{bmatrix} = \begin{bmatrix} 0 \\ 0 \end{bmatrix}
\]

Equation (20) reveals that the sensitivity of controlled variables to disturbances in the feed is minimum at the maximum driving force.

3.3. Selection of the Controller Structure

The controlled variables are defined as top and bottom element A composition \( W^*_d \) and \( W^*_d \). In this case, the potential manipulated variables are reflux ratio (RR) and reboil ratio (RB). (12) and (13) give the top and bottom product compositions with respect to the driving force.

Hence, they are differentiated with respect to RR and RB. Therefore, (6) can now be expressed as:

\[
\frac{dy}{du} = \begin{bmatrix} \frac{dW_d}{dR} & \frac{dW^*_d}{dR} \\ \frac{dW^*_d}{dR} & \frac{dW^*_d}{dR} \end{bmatrix} = \begin{bmatrix} \frac{dW_d}{dR} & \frac{dW^*_d}{dR} & \frac{dW^*_d}{dR} & \frac{dW^*_d}{dR} \end{bmatrix}
\]

From Fig. 4, it is known that \( \frac{dF_D}{dR} \) at the maximum driving force is equal to zero. Furthermore, assuming that \( \frac{dW^*_d}{dR} = \frac{dW^*_d}{dR} = 0 \), (22) is obtained. The best controller structure can easily be determined by looking at the value of \( \frac{dy}{du} \). It can be noted from (22) that since the values of \( \frac{dW^*_d}{dR} \) and \( \frac{dW^*_d}{dR} \) are bigger, controlling \( W^*_d \) by manipulating RR and controlling \( W^*_d \) by manipulating RB will require less control action.

\[
\frac{dy}{du} = \begin{bmatrix} \frac{dW_d}{dR} & \frac{dW^*_d}{dR} \\ \frac{dW^*_d}{dR} & \frac{dW^*_d}{dR} \end{bmatrix} = \begin{bmatrix} \frac{dW_d}{dR} & 0 \\ 0 & -\frac{dF_D}{dR} \end{bmatrix}
\]

This is because only small changes in RR and RB are required to move \( W^*_d \) and \( W^*_d \) in a bigger direction. This pairing between controlled-manipulated variables is also further verified by obtaining the transfer functions between the pairs using a reactive distillation dynamic model based on elements (Pérez Cisneros, 1997). Note that most of the modelling of dynamic reactive distillation operations has been done by introducing a rate of reaction expression in the component mass balances. However, when the chemical reactions occurring are fast enough to reach the equilibrium (for example, MTBE reactive system) chemical equilibrium condition is implicitly incorporated into the element mass balances through the functionality of the phase compositions on the element chemical potentials (Pérez-Cisneros, 1997).

The next natural step to verify the pairing in (22) is calculating the relative gain array (RGA) for the design at the optimal feed location (Design I – \( N_F = 1 \)) and two other alternative designs (Design II – \( N_F = 2 \); and Design III – \( N_F = 3 \)). Note that in calculating RGA, \( RB \) is represented by heat addition to the reboiler duty instead of reboil ratio; and, MTBE top and bottom compositions represent \( W^*_d \) and \( W^*_d \), respectively. The transfer functions have the form as equation (23) with one zero and two poles.

\[
G(s) = \frac{K}{(1+\tau_s)(1+\tau'_s)}
\]

Using the transfer functions, RGA matrix for design B (at the maximum driving force) and designs A and C is calculated and they are as follows:

\[
RGA_B = \begin{bmatrix} 0.93 & 0.07 \\ 0.07 & 0.93 \end{bmatrix}, RGA_A = \begin{bmatrix} 9.06 & -8.06 \\ -8.06 & 9.06 \end{bmatrix}, RGA_{B} = \begin{bmatrix} -0.28 & 1.28 \\ 1.28 & -0.28 \end{bmatrix}
\]

As it can be seen the pairing at the maximum driving force (design I, feed location 1) has the closest values on the diagonal to unity. Therefore, it has the least interaction between the loops. Furthermore, the suggested pairing by RGA for design I matches the pairing that was obtained from the driving force. Furthermore, singular value analysis
(SVA) was performed. However, very large condition numbers (CN) were obtained with no specific trend. Thus, no particular conclusion can be made based on them.

The open-loop and closed-loop performance of the system has been tested with the Proportional-Integral (PI) controller in a discrete-time manner. The rigorous dynamic reactive distillation model (Pérez Cisneros, 1997) was used. The controller implementation is visualized in Fig. 5. Note, however, any other control strategy can be applied to perform closed-loop simulations.

Fig. 5. Discrete-time controller structure implementation.

Fig. 6 and Fig. 7 show the open-loop and closed-loop performance of the system at maximum driving force (Design (I)), respectively. The disturbance scenario is that after 15 samples, the feed flowrate of element A (isobutene) is increased from 70 kg-mole to 85 kg-mole (~12% step change in composition of isobutene). This disturbance results in a change in total feed flowrate by +15% and also a change in the composition in the feed composition. As it can be seen in Fig. 6, as a result of an increased isobutene flowrate, its recovery in the top has increased which results in a lower MTBE composition in the top. Furthermore, because of excess isobutene in the system and thereby shifting the reaction equilibrium, MTBE composition has increased in the bottom.

Fig. 6. Open-loop performance of Design B to a disturbance in the feed.

It can be seen in Fig. 7 that disturbance has been rejected with least interaction between the loops and both top and bottom compositions are well controlled using the selected pairing obtained from the driving force. Note that the control of the MTBE top composition is achieved with very small changes in $RR$.

Fig. 7. Closed-loop performance of the control structure for Design B to a disturbance in the feed.

4. CONCLUSIONS

Integrated process design and control of a ternary compound reactive distillation process was investigated in this work. The optimal design-control solutions were obtained analytically and verified through rigorous dynamic simulations. It is verified that the reactive distillation design option is less sensitive to the disturbances in the feed at the highest driving force and has the inherent ability to reject disturbances. Furthermore, it is advantageous to employ the element-based method for designing multicomponent and complex reacting systems.

REFERENCES


