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DOI:

[10.1016/j.cherd.2019.03.014](https://doi.org/10.1016/j.cherd.2019.03.014)

Document Version

Accepted author manuscript

[Link to publication record in Manchester Research Explorer](#)

Citation for published version (APA):

Muthia, R., van der Ham, A. G. J., Jobson, M., & Kiss, A. (2019). Effect of boiling point rankings and feed locations on the applicability of reactive distillation to quaternary systems. *Chemical Engineering Research & Design*. <https://doi.org/10.1016/j.cherd.2019.03.014>

Published in:

Chemical Engineering Research & Design

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Effect of boiling point rankings and feed locations on the applicability of reactive distillation to quaternary systems

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Keywords

Reactive distillation, mapping method, applicability graph, quaternary systems

Highlights

- Insights into reactive distillation (RD) design and techno-economic applicability
- Conceptual process design and preliminary economic ranking of RD processes
- Influence of feed locations on the performance of reactive distillation systems
- Effect of boiling point order on the applicability of RD to quaternary systems

Abstract

Reactive distillation (RD) offers major benefits such as costs reduction and energy saving, but the understanding and design of RD processes usually demand complex tasks that include extensive studies and rigorous simulations. To reduce this complexity and the time required, this study applies a novel mapping method to quickly provide insights into the RD applicability to reversible quaternary systems ($A + B \rightleftharpoons C + D$). Generic cases are used to produce applicability graphs (i.e. plots of reflux ratio vs number of theoretical stages) and multiple RD column configurations. The systems are assumed to have ideal properties and fixed key parameters (i.e. relative volatilities and chemical equilibrium constants). This study focuses on quaternary reactions considering different boiling point rankings and feed locations. Using the mapping method, quick results are achievable regarding the preliminary economic ranking of RD processes, and the optimal feed locations with reduced energy requirement (i.e. lower reflux ratio). Ultimately, this study provides a much better understanding of the effect of boiling point orders and feed locations on the RD applicability and conceptual design, being a valuable tool in early techno-economic evaluations.

1 **1. Introduction**

2 Reactive distillation (RD) is a process intensification technique that allows simultaneous
3 separation to take place when chemical reaction occurs. RD gives benefits to equilibrium-
4 limited reactions by pulling the chemical equilibrium towards complete conversion and
5 allowing high selectivity (e.g. avoiding potential consecutive reactions) due to the continuous
6 removal of products from the RD column (Baur et al., 2000). Among others, RD offers great
7 advantages in costs reduction by simplifying complex processes and integrating the reaction
8 and separation functions into a single unit with reduced requirements (Kiss, 2018). The use of
9 exothermic heat of reaction to drive the liquid vaporization reduces the energy requirement
10 (Kaur and Sangal, 2017). There are also health, safety and environmental improvements
11 mainly because of less emissions from plants, lower levels of reactive hold-up and decreased
12 risks of runaway reactions (Taylor and Krishna, 2000; Shah et al., 2012).

13 The commercialization of reactive distillation has expanded for over 30 years (Stankiewicz,
14 2003). The most well-known RD process is in the methyl acetate production via esterification,
15 which has been established since 1984 by Eastman Kodak Company (Agreda et al., 1990).
16 The syntheses of ethers, i.e. methyl tert-butyl ether, ethyl tert-butyl ether and tert-amyl methyl
17 ether, are other remarkable examples where RD technology is applied (Sharma and Mahajani,
18 2002). Furthermore, the implementation of RD is very appealing to other reactions involving
19 reversible quaternary systems ($A + B \rightleftharpoons C + D$) and ternary systems ($A + B \rightleftharpoons C$ and $A \rightleftharpoons B +$
20 C). Some examples include the hydration of cyclohexene to obtain cyclohexanol (Chen et al.,
21 2014), dehydration of glycerol to acetol (Chiu et al., 2006), isoamyl acetate production via
22 esterification of isoamyl alcohol and acetic acid (González et al., 2017) and diethyl carbonate
23 synthesis via trans-esterification of propylene carbonate and ethanol (Wang et al., 2014).

24 Studies related to RD technology provide various methods to design and control the column
25 operation, specify and modify its physical structures and/or evaluate the economic aspect. For
26 example, a set of equations can be used to calculate the minimum reflux ratio (RR_{\min}) for both
27 single- and double-feeds RD columns (Barbosa and Doherty, 1988a; Barbosa and Doherty,
28 1988b). The location of reactive zone in binary reactions can be visualized by using the
29 McCabe-Thiele and Ponchon Savarit methods (Lee et al., 2000). Other studies determined
30 heuristic approaches, by considering basic knowledge and industrial experience, to specify the
31 operating conditions and the physical aspects of RD (Subawalla and Fair, 1999). Other RD
32 design methods have been also reported in literature (Buzad and Doherty, 1994; Ciric and Gu,
33 1994; Almeida-Rivera et al., 2004; Groemping et al., 2004; They et al., 2005; Jantharasuk et
34 al., 2011). However, in contrast to much information available, the complexity of designing

1 and understanding RD processes remained a strong barrier that has hindered the rapid
2 commercialization of RD for over 15 years (Chen et al., 2000; Malone and Doherty, 2000;
3 Harmsen, 2007; Segovia-Hernández et al., 2015; Li et al., 2016). In other words, simplicity is
4 strongly needed in the assessment of RD design at early stages of industrial R&D.

5 The present study aims to provide insights into RD processes by using a mapping method that
6 was initially developed to quickly assess the applicability of RD based on pre-defined maps
7 (i.e. applicability graphs) obtained from generic cases (Muthia et al., 2018). In that initial
8 work, we validated the approach using two case studies, i.e. hydrolysis of methyl lactate and
9 transesterification of methyl benzoate and benzyl alcohol, and showed that one can
10 successfully use the generic cases to predict the applicability of RD to real systems. The
11 following assumptions for the generic cases are applied in this study: ideal thermodynamics
12 (neither azeotropes nor liquid split); fixed values of key parameters, i.e. relative volatility (α)
13 and chemical equilibrium constant (K_{eq}); and vapor-liquid and chemical equilibria on each
14 stage. The key feature of the mapping method is RD applicability graphs produced from the
15 generic cases, which are the plots of reflux ratio (RR) vs number of theoretical stages (NTS).

16 This study focuses on quaternary systems, as they are the most encountered reactions for the
17 operation of RD in chemical industry. Beyond the scope of this paper, the mapping method is
18 promising for applications to ternary systems, but further studies are required to further
19 develop the method for those systems. The application of RD to quaternary systems
20 investigated in this study considers different boiling point (T_b) rankings and feed locations.
21 Insights into the RD applicability related to economics and conceptual design are provided
22 early on. This is in contrast to the conventional way, where knowledge about these aspects
23 usually requires rigorous simulations and/or detailed calculations and at such level that
24 understanding can be obtained only at the final stage of conceptual design studies (Seider et
25 al., 2003; Towler and Sinnott, 2012). Firstly, the mapping method is used to obtain a
26 preliminary economic ranking, thus providing an overview on the most beneficial RD
27 configurations in chemical processes. Secondly, the mapping method is used to investigate the
28 possibilities of reducing the energy requirement (i.e. operating RD with lower reflux ratios)
29 by finding the optimal feed locations. A recent review revealed that extensive studies about
30 the design of RD are available in literature, but there is still a lack of understanding in terms
31 of process optimization (Segovia-Hernández et al., 2015).

32 Prior studies investigating the effect of volatilities on the RD performance are available in
33 literature with different focuses and approaches. Luyben and Yu (2008) ranked quaternary
34 reactions with various boiling point orders by using detailed economic calculations but

1 considering only a fixed equilibrium constant. In this work, we use a range of chemical
2 equilibrium constants and rank the quaternary reactions with a simpler approach, based on
3 number of theoretical stages and reflux ratio that provide an indication of the capital
4 investment and energy requirement. Chen and Yu (2008) used the same approach as that in
5 Luyben and Yu (2008) to study the effect of relative volatility ranking on RD configurations,
6 but to ternary decomposition reactions only. In this study, we focus on quaternary reactions,
7 which are more complex by the presence of more components in the systems. Our previous
8 work (2018) focused on a single set of quaternary systems with both reactants as mid-boiling
9 components and products as lightest and heaviest components ($T_{b,C} < T_{b,A} < T_{b,B} < T_{b,D}$), and
10 fixed feed locations at both end sides of the reactive section. That work provided valuable
11 insights into RD performance considering low or high relative volatilities between product-
12 reactant (α_{CA} and α_{BD}) and both reactants (α_{AB}). In our current study, we include all
13 quaternary systems with different boiling point rankings and obtain the optimal feed locations
14 considering varied feed stages.

15 Summing it up, this work gives useful knowledge regarding industrial RD processes, covering
16 two aspects related to essential assessment in conceptual design studies: economic ranking of
17 process alternatives, and optimal process configurations. Furthermore, this study provides a
18 valuable understanding of the effect of boiling point rankings on the RD processes.

19

20 **2. Approach and methodology**

21 A novel mapping method – described in our previous work – has been employed to predict
22 the applicability of RD based on the applicability graphs of generic cases and to study the
23 effect of relative volatilities, chemical equilibrium and kinetics (represented by Damköhler
24 number) on RD processes (Muthia et al., 2018). This method was also used to determine the
25 optimal RD configurations that can operate at lowest costs (Muthia et al., 2018a).

26 The key feature of the mapping method is the RD applicability graph, illustrated in Figure 1.
27 A boundary line in the applicability graph splits the plot into applicable and not-applicable
28 areas. On that line, each NTS has a RD configuration with the lowest reflux ratio possible. On
29 the boundary line and inside the applicability area at any NTS, the operation of reactive
30 distillation is conceivable. Multiple RD configurations are available inside the applicability
31 area which give flat-optimum solutions (Muthia et al., 2018). For the sake of clarity, the
32 applicability graphs shown in this work have a maximum scale of 100 for both x- and y-axes.
33 Each RD applicability graph is coupled with a representation of column configurations within
34 the applicable area. The selection of that representation can be based on various consideration,

1 e.g. at a fixed NTS, at $NTS=2 \cdot NTS_{min}$, at NTS with $RR=1.2 \cdot RR_{min}$, or at any other points. In
2 this study, each representation of RD configurations is at $NTS=2 \cdot NTS_{min}$ which is only based
3 on the well-known rule of thumb in conventional distillation regarding the prediction of the
4 optimum configuration. That representation is selected by considering the availability of
5 multiple RD configurations with reflux ratios up to 10% larger than the lowest RR. This
6 consideration is logically accepted as that marginal reflux ratios difference is commonly
7 negligible in the RD application. Considering the flat-optimum solutions in the applicability
8 graphs, various trends of RD configurations can be obtained when different applicability
9 graphs are studied and compared. The representations of RD columns are selected based on
10 the decrease of number of reactive stages for a higher K_{eq} when applicability graphs of
11 various equilibrium constants are compared. This selection criterion is logical since a better
12 reaction performance is achieved for a higher K_{eq} .

13 The schematic procedure to generate an applicability graph is given in Figure 2. It might be
14 preferred to specify relative volatilities of components following the boiling point rankings of
15 the quaternary systems. For instance, with the order of $T_{b,C} < T_{b,A} < T_{b,D} < T_{b,B}$, specifying
16 α_{CA} , α_{AD} and α_{DB} might be preferred. This study only focuses on equilibrium-limited reactions
17 as similar knowledge is expected to be the outcome for the study of their kinetically
18 controlled reactions, as proven in our previous work (2018). Since vapor-liquid and chemical
19 equilibria are reached on each stage of the column, specifying liquid hold-up or residence
20 time (resulting in the Damköhler number) is not required as an input for the simulation in this
21 work.

22 The quaternary reactions are classified into six groups based on the boiling point rankings, as
23 shown in Table 1. For consistency reasons, the naming convention of the groups follows the
24 rule defined by Luyben and Yu (2008). To deliver comparable results for different groups,
25 several key relative volatilities have to be specified uniformly. In this study, α_{AB} and α_{CD} are
26 specified at 2 and 6, respectively (see Table 1). Only group I_r has different specified α_{AB} and
27 α_{CD} since its boiling point order does not allow defining those relative volatilities at 2 and 6,
28 respectively. Both groups II_p and II_r are disregarded from the investigation, since from a
29 thermodynamic point of view, it is unfeasible to obtain two heaviest / lightest products from
30 two lightest / heaviest reactants (in stoichiometric systems).

31 Using the mapping method, a preliminary economic ranking of RD is obtained here by
32 considering the number of theoretical stages and reflux ratio as two main variables, which
33 give the first economic assessment. NTS indicates the height of column which is proportional
34 to the capital investment and RR affects the column diameter and is directly proportional to

1 the amount of hot utility required (energy usage), respectively (Kiss, 2013). To perform this
2 assessment comprehensively, the chemical equilibrium constants are varied at values of: 0.01,
3 0.1, 0.2, 1 and 10. These values cover the practical range of reactions in terms of the RD
4 application.

5 All simulations are performed in Aspen Plus v8.6. The RD scheme is presented in Figure 3
6 (a). The RD column operates at atmospheric pressure, assuming negligible pressure drop. The
7 reactants are fed as saturated liquid, in an equimolar ratio. The lighter reactant is fed on the
8 bottom of reactive zone and the heavier reactant is introduced to the top of reactive zone,
9 therefore a counter current flow occurs which allows reaction to take place along the reactive
10 stages. Sensitivity analysis is carried out by varying the position and the length of rectifying,
11 reactive and stripping sections. The optimization tool is used to minimize reflux ratio for any
12 converged solutions by considering product purity at top and bottom (min. 99 mol%) as a
13 hard constraint. Next, the optimal feed locations are assessed for the quaternary systems as
14 given in the study of preliminary economic ranking. Any configurations in RD applicability
15 graphs can be selected for this investigation. The resulting representations of column
16 configurations in the previous section are used as base cases, in terms of numbers of
17 rectifying, reactive and stripping stages. Sensitivity analysis is carried out by varying both
18 positions of feed stages, as shown in Figure 3 (b). The optimization tool is used to minimize
19 the reflux ratio by considering the same constraint as in the study of preliminary economic
20 ranking.

21

22 **3. Results and discussion**

23 *3.1. Preliminary economic ranking of RD processes*

24 Simulations in Aspen Plus v8.6 were performed for groups I_p, I_r, III_p and III_r to obtain their
25 applicability areas accounting for various equilibrium constants (from 0.01 to 10). There is no
26 applicability graph available in group I_r for any specified K_{eq} which indicates that the required
27 product purity cannot be achieved, hence a single RD column is not applicable. An advanced
28 RD configuration (involving two columns) might be used for this group. The RD setup might
29 be adjusted by adding side-draw product stream(s) to the column as both desired products are
30 mid-boiling compounds (Luyben and Yu, 2008). At least an additional conventional
31 distillation is needed to obtain the products at high purity. Besides, the application of reactive
32 dividing wall column (R-DWC) could be considered as another option. As the application of a
33 single column is aimed in this study, group I_r is disregarded from our further consideration.

1 Figure 4 (a), (c) and (e) depict the RD applicability graphs for groups I_p , III_p and III_r ,
2 respectively. In group III_r , the applicability area of the system with $K_{eq}=0.01$ could not be
3 obtained for both NTS and RR up to 100. For these groups, the applicability area becomes
4 larger when K_{eq} is higher due to a higher conversion which gives the possibilities of having
5 RD configurations with lower capital investments and reduced energy requirements (i.e. lower
6 NTS and RR), which is as expected.

7 The representations of RD configurations, at various equilibrium constants, is shown in
8 Figure 4 (b), (d) and (f) for groups I_p , III_p , III_r , respectively. As observed earlier, the increase
9 of K_{eq} leads to a better reaction performance, therefore the NTS and RR decrease for each
10 group. In comparison between different groups, at a fixed K_{eq} , the number of theoretical
11 stages and reflux ratio increase from group I_p , III_p to III_r , respectively. This trend indicates the
12 cost ranking of these three groups – I_p , III_p , III_r – from the lowest to the highest cost,
13 respectively. Having obtained the RD configurations for all groups in Figure 4 (b), (d) and (f),
14 the column schemes are shown in Figure 5.

15 Considering the volatility order of group III_p ($T_{b,C} < T_{b,A} < T_{b,D} < T_{b,B}$), it is very important to
16 fully convert component B before it reaches the stripping section, otherwise its presence
17 becomes a hindrance to a high purity of product D at the bottom stream. Therefore, if group
18 III_p is compared to group I_p , a larger number of reactive stages is required. For any fixed
19 equilibrium constant - see Figure 4 (b) and (d) - the rectifying section of group III_p is smaller
20 than that of group I_p because of the larger relative volatilities of compounds ($\alpha_{CA}=4$ and
21 $\alpha_{CB}=8.4$ in group III_p , $\alpha_{CA}=2$ and $\alpha_{CB}=4$ in group I_p) which lead to an easier separation.
22 Besides, the stripping section of the RD column for group III_p is larger than that of group I_p as
23 the separation between reactant A and product D is more challenging ($\alpha_{AD}=3$ in group I_p and
24 $\alpha_{AD}=1.5$ in group III_p).

25 Regarding the relative volatility sets listed in Table 1, the configurations of group III_r are
26 expected to be mirror images of the RD columns of group III_p , shown in Figure 5 (b) and (c).
27 However, it is observed that a higher number of theoretical stages and a larger reflux ratio are
28 required for group III_r at various K_{eq} s, see Figure 4 (d) and (f). In contrast to group III_p , group
29 III_r needs a larger reactive section to fully convert the reactant A before it reaches the
30 rectifying section. This task is more difficult than that in group III_p because the reactant A is
31 the lightest, therefore it is easily vaporized and enter the upper level of the column. A higher
32 reflux ratio is required mainly because of more difficult separation in rectifying section
33 ($\alpha_{CB}=1.5$ in group III_r , $\alpha_{CA}=4$ and $\alpha_{CB}=8.4$ in group III_p).

1 To gain more understanding about the effect of chemical equilibrium constants on the reaction
2 and separation processes in the different groups, the liquid composition profiles are provided
3 in Figure 6. A low and a high K_{eq} s (0.1 and 10) are evaluated for each group. For all groups
4 with $K_{eq}=0.1$ - see Figure 6 (a), (c) and (e) - there is always an accumulation of a reactant
5 along the reactive zone because it helps to push the conversion of reaction to the products
6 side. Commonly, reactant A is the accumulated compound because it is more easily vaporized
7 than reactant B and the condensation process at the upper part of the column changes its phase
8 back to liquid. However, in group III_r having abundant reactant A will inhibit the desired RD
9 process as it interferes with the high purity of product C. Therefore, in this group at a low K_{eq} ,
10 B is the reactant that is accumulated.

11 If the systems with an equilibrium constant $K_{eq}=10$ are compared to those with $K_{eq}=0.1$, the
12 accumulation in the reactive zone of any reactant is then replaced by the product(s) due to a
13 better reaction performance. In group I_p - see Figure 6 (b) - the accumulation of products C
14 and D in the reactive zone can be observed, with a higher mol fraction of D present. Less
15 accumulated C is detected, because overall the separation of product C from any reactants is
16 easier ($\alpha_{CA}=2$, $\alpha_{CB}=4$, $\alpha_{AD}=3$, $\alpha_{BD}=1.5$). In group III_p - see Figure 6 (d) - component D is the
17 accumulated product because its separation from reactant A is more challenging than that of
18 product C considering their relative volatilities ($\alpha_{AD}=1.5$, $\alpha_{CA}=4$). The opposite difficulty
19 occurs in group III_r, see Figure 6 (f), in which the separation between reactant B and product
20 C is more challenging than the separation between reactant B and product D ($\alpha_{CB}=1.5$,
21 $\alpha_{BD}=4$), therefore C is the accumulated product in the column.

22

23 3.2. Optimal feed locations for RD units

24 The optimization of feed locations has been carried out by taking the RD configurations as
25 given in Figure 4 (b), (d) and (f) with $K_{eq}=1$ as the base cases. For all combinations of feed
26 locations, the numbers of rectifying, reactive and stripping stages remain the same as those in
27 the base cases, which limit the varied design parameters and give clarity to the presented
28 results. Figure 7 (a), (b) and (c) show reflux ratios as a contour plot for varied feed locations
29 of both reactants A and B for groups I_p, III_p and III_r, respectively, at $K_{eq}=1$. Cross and square
30 marks in each plot indicate the base case and a RD configuration with the most optimal feed
31 locations (i.e. with the lowest reflux ratio reducing energy requirements), respectively.
32 Moving from the crosses to the squares, the reflux ratios of the base cases decrease from 2.1
33 to 1.7 for group I_p (19% reduction), from 4.6 to 2.9 for group III_p (37% reduction), and from
34 8.6 to 6.6 for group III_r (23% reduction), by adjusting the feed locations.

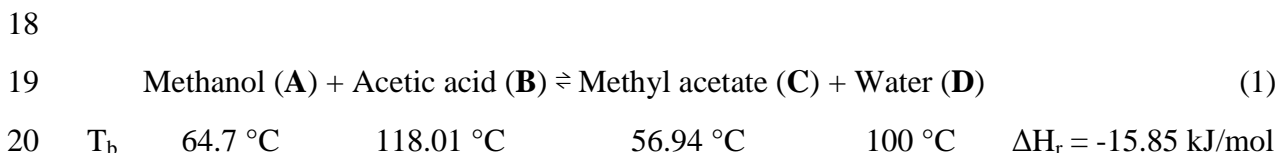
1 To clearly distinguish the RR changes, each plot in Figure 7 is partitioned based on certain
2 RD ranges and marked with distributed colors. The optimal region (the darkest zone) in each
3 plot is achieved by RD configurations with reflux ratios up to 3% higher than the lowest RR.
4 For all groups, it is always beneficial to have a shorter distance between the two feed stages,
5 therefore an immediate contact between two reactants occur before separation takes place.
6 Besides, it is essential to set the feed stage of reactant B (as the higher boiling reactant) above
7 that of reactant A in order to enhance the direct contact of both reactants in liquid phase,
8 therefore enhancing the reaction performance. Placing the feed inlets on a reverse order
9 causes a significant increase in reflux ratios. For instance, in group I_p, the feed stages of
10 reactants A and B at 24 and 11 require RR=2.1, while the reverse order gives RR=8.1. In
11 group III_p, RR=3.8 is needed when the feed inlets of reactants A and B are at stages 12 and 4,
12 while the reverse order requires RR=10.9. In group III_r, RR=6.9 is needed when the feed
13 inlets of reactants A and B are at stages 56 and 44, while the reverse order requires RR=44.3.
14 Moving from the crosses to the squares in Figure 7, the optimization pattern for different
15 groups are identified. In group I_p, the RD process is optimized by positioning the two feed
16 stages close to the centre part of the reactive zone. In group III_p, bringing the inlet of reactant
17 A up within the reactive zone reduces the energy requirements (lower RR). On the contrary,
18 lowering the feed stage of reactant B closer to the bottom section of reactive stages decreases
19 the energy requirements (lower RR) in group III_r. For a better insight, the representations of
20 RD configurations with lowest reflux ratios inside the darkest zone in Figure 8 (b), (d) and (f)
21 are coupled with the corresponding column configurations of the base cases, as shown in
22 Figure 8 (a), (c) and (e).

23 Further investigation is performed by observing the liquid composition profiles of different
24 groups, as given in Figure 9. In comparison to the base case of group I_p, see Figure 9 (a) and
25 (b), the immediate contact of reactants for the optimized feed stages has reduced their
26 accumulation along the RD column. As consequence, there are less unreacted compounds
27 found in the rectifying and stripping sections which results in less difficulty in the separation.
28 In group III_p - see Figure 9 (c) and (d) - the improved reaction condition in the reactive zone
29 with modified feed locations significantly helps to eliminate reactant B (i.e. the heaviest
30 compound in this group). The average concentration of A in the reaction zone has increased
31 with at the same time a lower concentration of A at the bottom part of the reactive zone
32 compare Figure 9 (c) and (d) - which reduces the concentration of B even further and
33 improves the separation performance (lower reflux ratio) for A/D. In the base case of group
34 III_r, the accumulation of reactant B - see Figure 9 (e) - is needed to enhance the reaction

1 performance and to prevent reactant A from reaching the upper part of column. In the
 2 improved RD process, the distribution of accumulated reactant B in the reactive zone has been
 3 optimized which is implied by the shift of the peak of its mol fraction and the increase in
 4 average concentration of B in the reactive zone, as shown in Figure 9 (f). This shift reduces
 5 the amount of B going up the rectifying section, therefore the separation of product C is
 6 easier.

7 Although a closer distance between the two feed stages is preferred in each group, introducing
 8 reactants A and B at the same stage potentially increases the energy requirements (higher RR)
 9 because reactant A is partially vaporized at the same time when reactant B moves down to the
 10 lower level of the column prior to reaching the reaction conditions. This observation indicates
 11 the competition between reaction and separation on a reactive stage.

12
 13 For the validation of our findings, we refer to a RD configuration with the optimal feed
 14 locations for the methyl acetate production, as provided by Tang et al. (2005) by performing
 15 rigorous simulations and detailed economic calculations. The synthesis of methyl acetate via
 16 esterification is given in Eq. (1). Based on its boiling points ranking, this reaction is classified
 17 into group III_p ($T_{b,C} < T_{b,A} < T_{b,D} < T_{b,B}$).



21
 22 The optimal RD configuration reported by Tang et al. (2005) consists of 1 rectifying, 34
 23 reactive and 4 stripping stages including reboiler. The feed locations of acetic acid (the
 24 heavier reactant) and methanol (the lighter reactant) are at stage 3 and 26, respectively,
 25 counted from the top down, which matches the predicted region of the optimal feed locations
 26 of group III_p as suggested by this work, see Figure 7 (b) and Figure 8 (c), (d). This example
 27 confirms the promising application of the mapping method. Besides the methyl acetate
 28 production, Tang et al. (2005) assessed the optimal RD designs for other acetic acid
 29 esterification reactions producing ethyl acetate, isopropyl acetate, n-butyl acetate and amyl
 30 acetate. Due to the higher level of complexity in those systems (i.e. the presence of ternary
 31 and heterogeneous azeotropes), it was reported that the operation of a single RD column is not
 32 feasible and therefore RD columns with decanter and/or additional stripper are required. For
 33 these different RD setups, the approach provided by this work is not applicable.

34

1 **4. Conclusions**

2 This work successfully demonstrates the use of the mapping method for obtaining insightful
3 knowledge of RD processes by using generic cases. Two essential aspects in conceptual
4 design (i.e. economic ranking of configurations and process optimization) can be investigated
5 at an early stage, instead of being assessed at the final stage of conventional studies.

6 The boiling point rankings of components have significant effects on RD configurations. In
7 the case of comparable chemical equilibrium constants and relative volatilities, the
8 preliminary cost ranking of quaternary systems for the application of a single RD column is:
9 group $I_p < III_p < III_r$ (from the lowest to the highest cost). A single RD column setup cannot
10 be applied to group I_r considering the challenging separation of both mid-boiling products
11 from the lightest and the heaviest boiling reactants. The presented approach can be used to
12 assess other equilibrium constants and relative volatility sets, significantly improving the
13 understanding of RD processes for different cases.

14 The mapping method is also useful to determine the regions where the optimal feed locations
15 are obtained. In group I_p , it is more beneficial to position both feed inlets closer to the centre
16 of the reactive zone. In group III_p , lower energy usage (reflux ratios) can be obtained by
17 putting the inlet of reactant A within the reactive zone, while the inlet of reactant B is kept at
18 the top of the reactive zone. In contrast, for group III_r , it is more beneficial to feed reactant B
19 within the reactive zone, while keeping the inlet of reactant A at the bottom of the reactive
20 zone. For all groups, a shorter distance between two feed inlets gives benefits, with the inlet
21 of the heavier reactant always above the inlet of the lighter reactant. However, feeding both
22 reactants at the same stage is detrimental for the energy usage since the competition between
23 reaction and separation phenomena hinders the RD performance.

24

25 **Acknowledgment**

26 The contribution of full financial fund from the LPDP (Indonesia Endowment Fund for
27 Education) for R. Muthia is greatly acknowledged. A. A. Kiss gratefully acknowledges the
28 Royal Society Wolfson Research Merit Award. The author also thanks all the participants of
29 the *Distillation & Absorption* conference 2018 (Florence, Italy) for the useful discussions, and
30 the reviewers for their insightful comments and suggestions.

31

32

33

1 **Nomenclature**

2	K_{eq}	chemical equilibrium constant [-]
3	NTS_{min}	minimum number of theoretical stages [-]
4	NTS	number of theoretical stages [-]
5	RR	reflux ratio [mol/mol]
6	RR_{min}	minimum reflux ratio [mol/mol]
7	T_b	boiling point temperature [°C]
8	α_{ij}	relative volatility between component i and j [-]
9	ΔH_r	heat of reaction [kJ/mol]

10

11

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1 **Table**

2 **Table 1.** Groups of quaternary systems ($A + B \rightleftharpoons C + D$) based on boiling point (T_b) orders.

Group	Boiling point ranking	Specified variables		Set variables		α set following T_b order
		α_{AB}	α_{CD}	α_{CA}	α_{BD}	
I_p	$C < A < B < D$	2	6	2	1.5	$\alpha_{CA} = 2, \alpha_{AB} = 2, \alpha_{BD} = 1.5$
I_r	$A < C < D < B$	6	2	0.6	0.6	$\alpha_{AC} = 1.7, \alpha_{CD} = 2, \alpha_{DB} = 1.7$
II_p	$C < D < A < B$	disregarded from consideration, for thermodynamic reasons				
II_r	$A < B < C < D$					
III_p	$C < A < D < B$	2.1	6	4	0.7	$\alpha_{CA} = 4, \alpha_{AD} = 1.5, \alpha_{DB} = 1.4$
III_r	$A < C < B < D$	2.1	6	0.7	4	$\alpha_{AC} = 1.4, \alpha_{CB} = 1.5, \alpha_{BD} = 4$

3

1 **Figure captions** (auto-updated)

2 Figure 1. An illustrative applicability graph for reactive distillation (Muthia et al., 2018).

3 Figure 2. Schematic procedure to generate RD applicability graph (Muthia et al., 2018).

4 Figure 3. RD columns with (a) fixed feed inlets on the top and the bottom parts of reactive
5 zone and (b) varied feed inlets along the RD column.

6 Figure 4. RD applicability graphs and their configurations at $NTS=2 \cdot NTS_{min}$ for: (a) group I_p ,
7 (b) group III_p , (c) group III_r . All the relative volatilities are according to Table 1 (Muthia et al.,
8 2018b).

9 Figure 5. RD column schemes for (a) group I_p , (b) group III_p , (c) group III_r (Muthia et al.,
10 2018b).

11 Figure 6. The composition profiles of RD configurations at $NTS=2 \cdot NTS_{min}$ for group I_p
12 considering (a) $K_{eq}=0.1$ and (b) $K_{eq}=10$, group III_p considering (c) $K_{eq}=0.1$ and (d) $K_{eq}=10$,
13 group III_r considering (e) $K_{eq}=0.1$ and (f) $K_{eq}=10$. All relative volatilities are according to
14 Table 1.

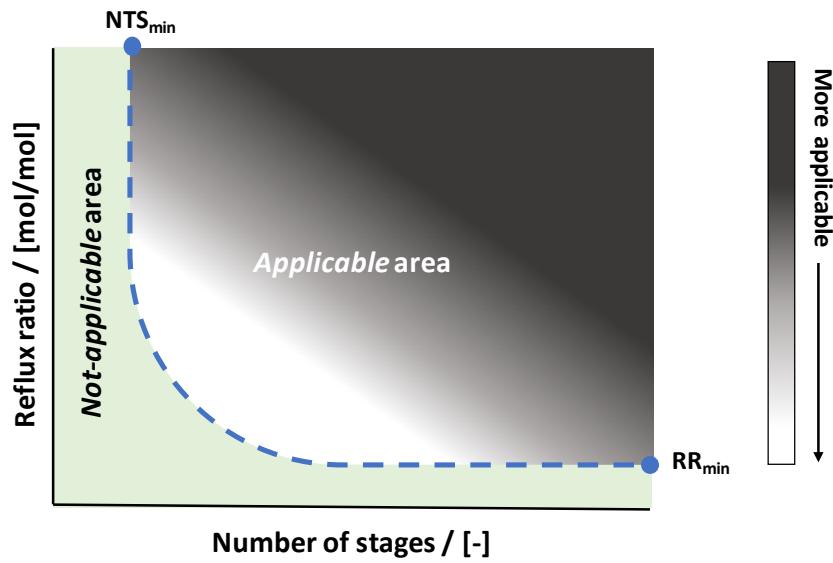
15 Figure 7. Varied feed locations of both reactants A and B and their corresponding RRs at
16 $K_{eq}=1$ for (a) group I_p , (b) group III_p and (c) group III_r . Cross and square indicate the base
17 cases and the RD configurations with the most optimal feed locations, respectively.

18 Figure 8. RD column schemes in case of fixed feed inlets at the top and the bottom parts of
19 reactive zone for (a) group I_p , (c) group III_p , (e) group III_r and in case of varied feed inlets to
20 obtain the lowest RR possible for (b) group I_p , (d) group III_p , (f) group III_r . The presented
21 numbers next to the column show the RD stages. All RD column configurations are at
22 $NTS=2 \cdot NTS_{min}$ in the applicability graph considering $K_{eq}=1$.

23 Figure 9. The composition profiles of RD column configurations in case of fixed feed inlets at
24 the top and the bottom parts of reactive zone for (a) group I_p , (c) group III_p , (e) group III_r and
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27 zone, and also the feed inlets in (a), (c), (e). The vertical dash lines in (b), (d), (f) present the
28 feed inlets. All RD column configurations are at $NTS=2 \cdot NTS_{min}$ in the applicability graph
29 considering $K_{eq}=1$, and relative volatilities according to Table 1.

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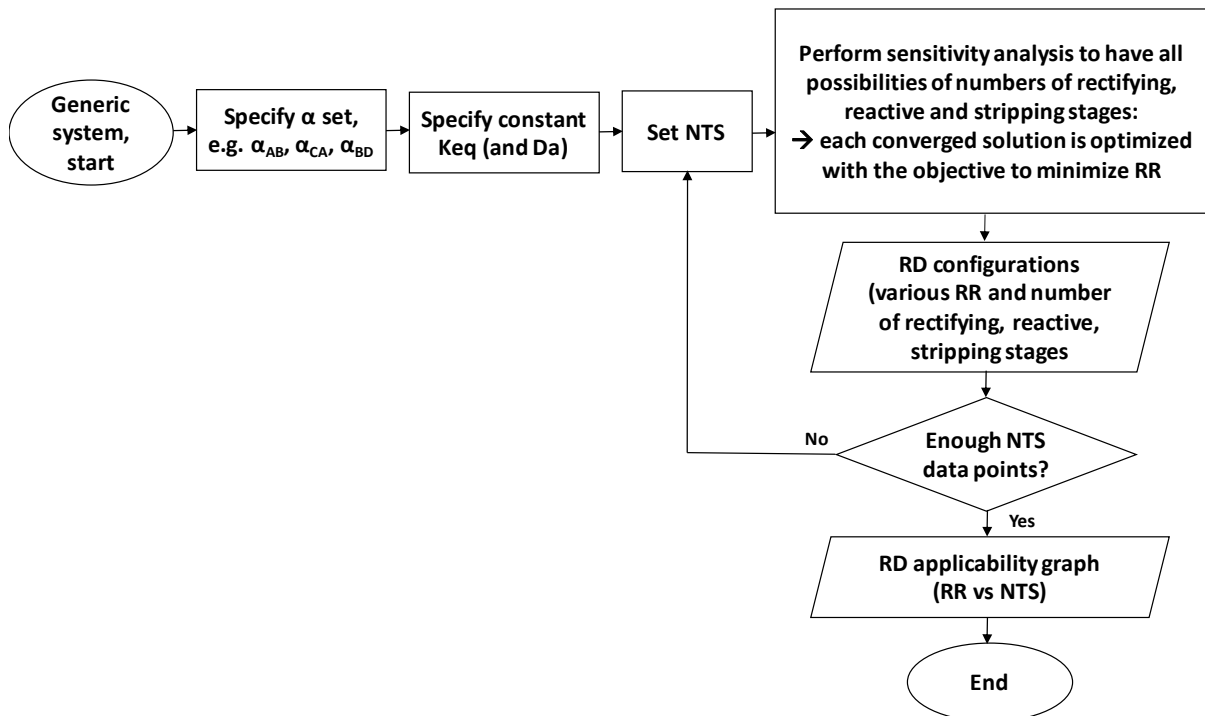
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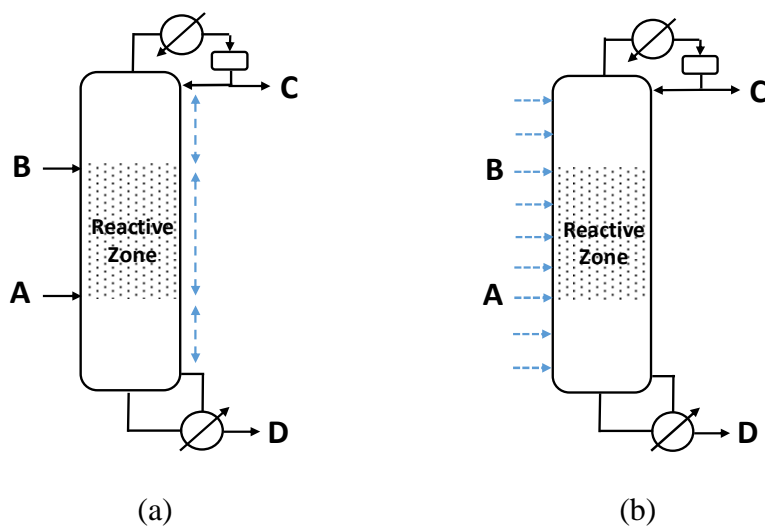
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10 **Figure 2.** Schematic procedure to generate RD applicability graph (Muthia et al., 2018).

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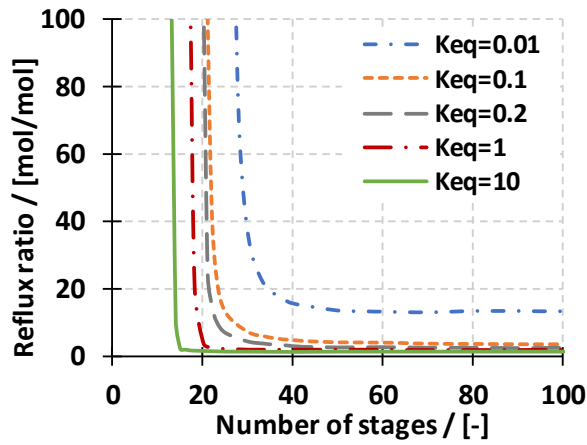
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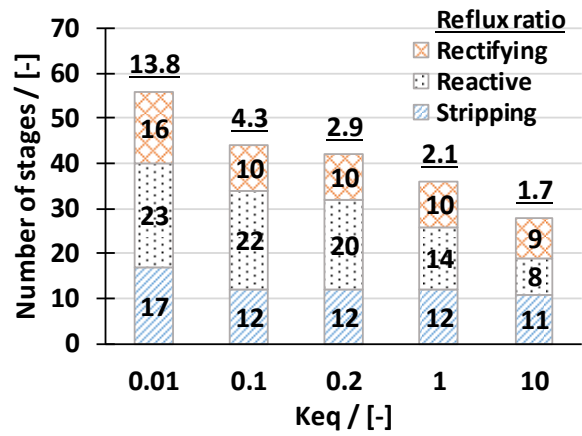
4 **Figure 3.** RD columns with (a) fixed feed inlets on the top and the bottom parts of reactive

5 zone and (b) varied feed inlets along the RD column.

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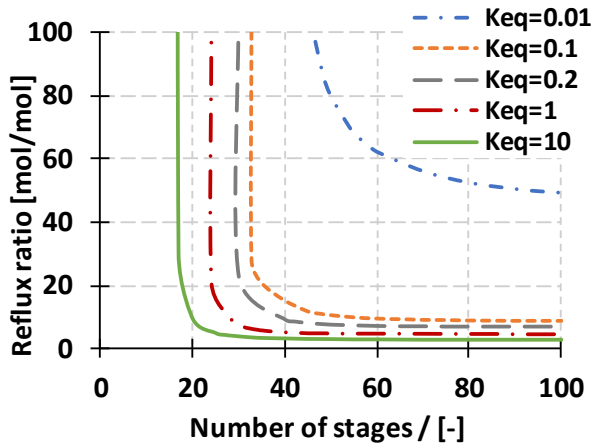
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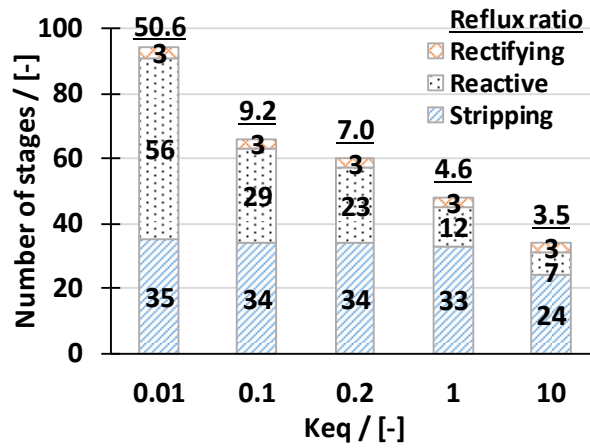
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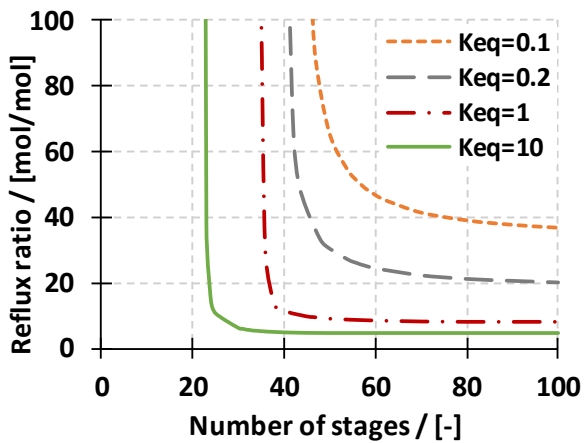
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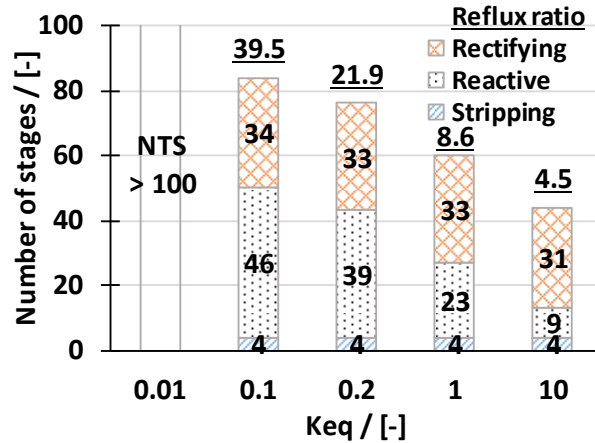
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8 **Figure 4.** RD applicability graphs and their configurations at $NTS=2 \cdot NTS_{min}$ for: (a) group I_p ,

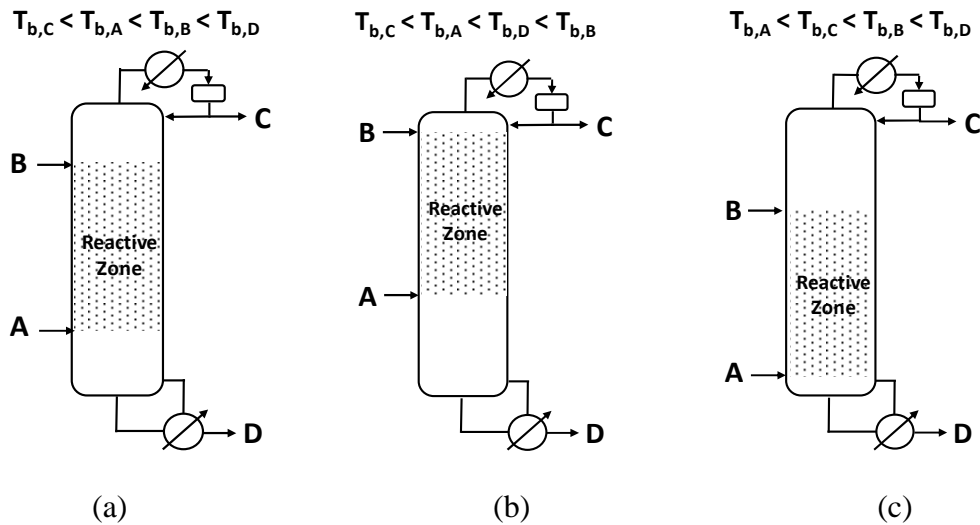
9 (b) group III_p , (c) group III_r . All the relative volatilities are according to Table 1 (Muthia et al.,

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2018b).

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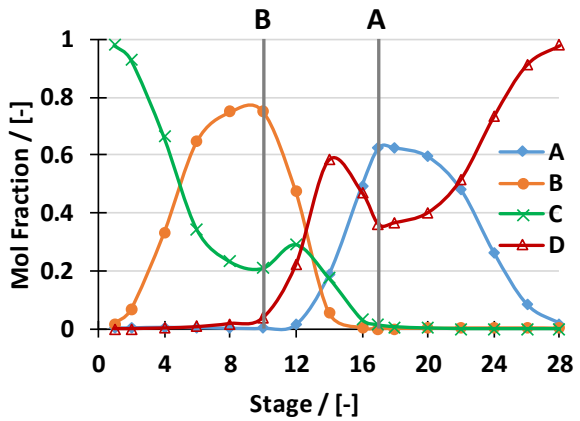
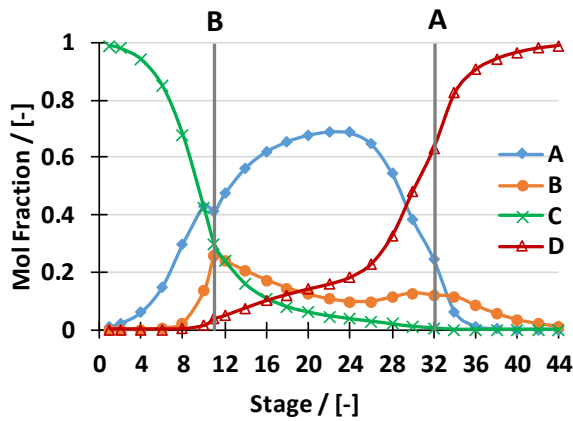
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Figure 5. RD column schemes for (a) group I_p, (b) group III_p, (c) group III_r (Muthia et al., 2018b).

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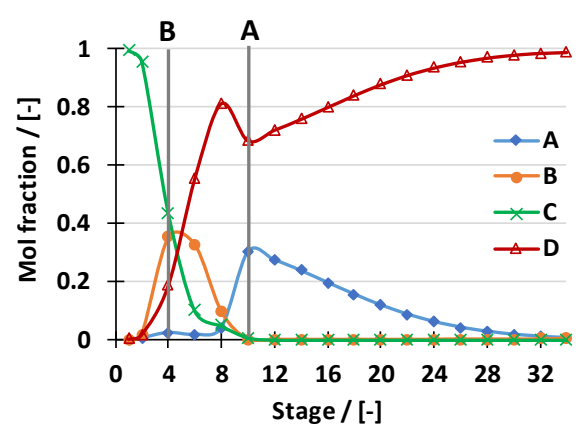
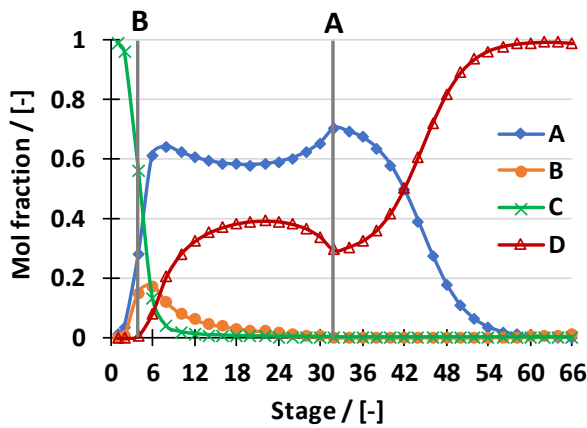


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(a)

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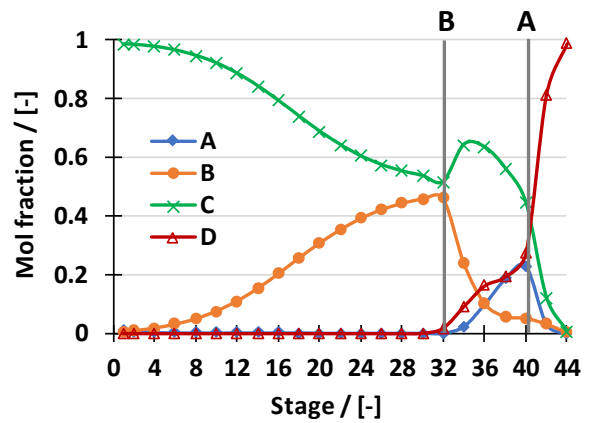
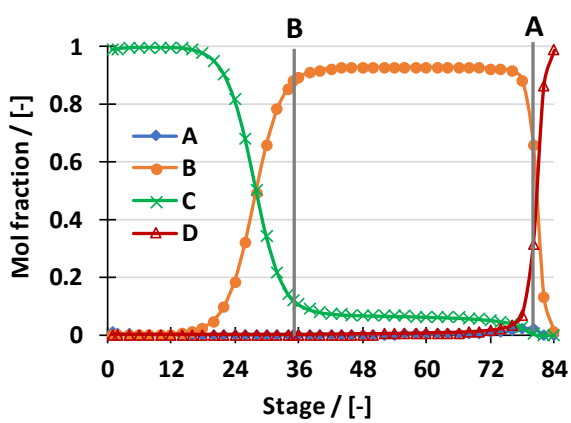


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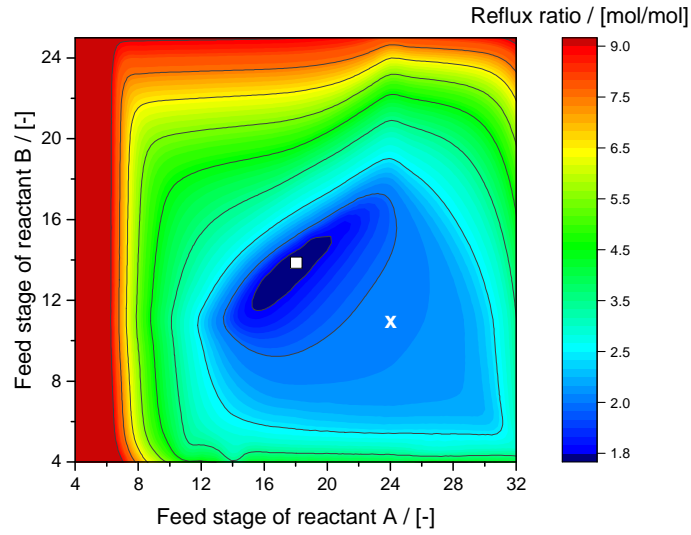
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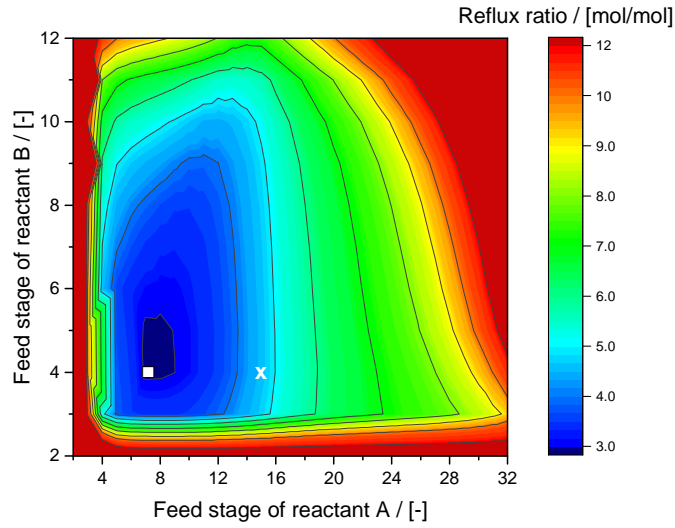
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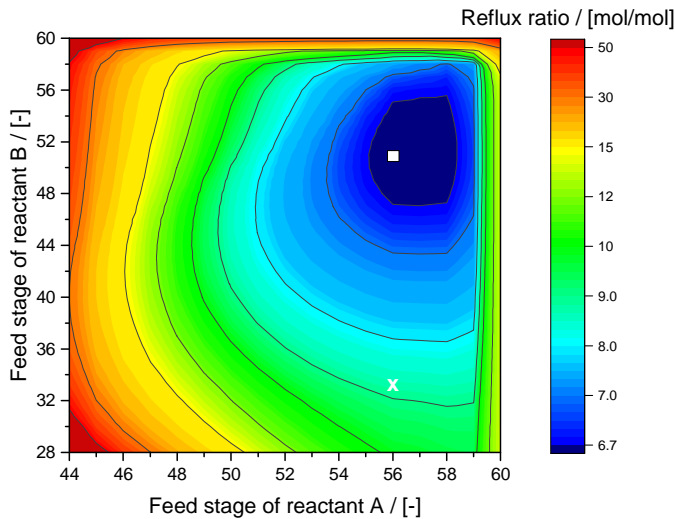
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(a)



(b)

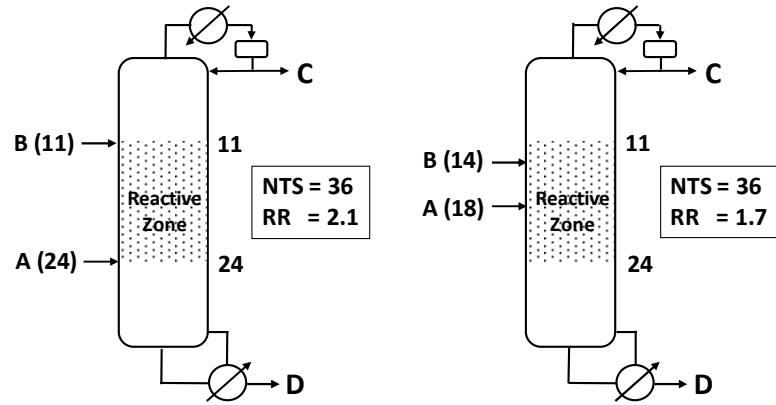


(c)

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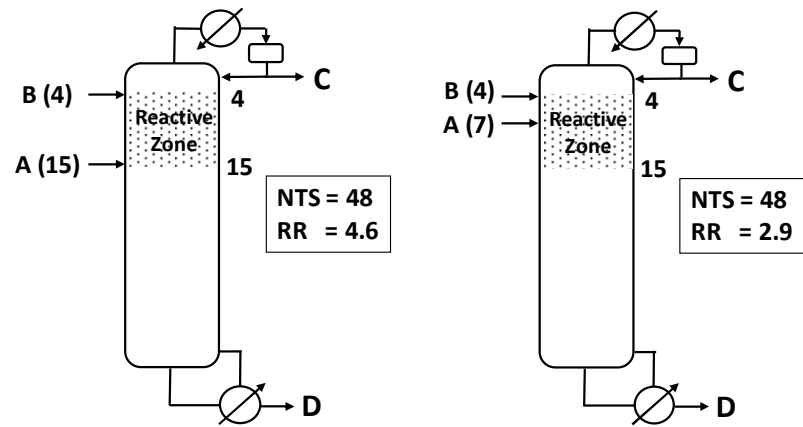
$$T_{b,C} < T_{b,A} < T_{b,B} < T_{b,D}$$



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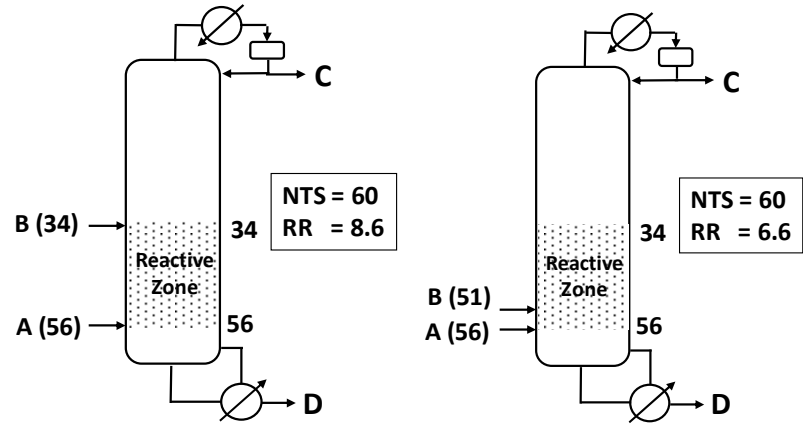
$$T_{b,C} < T_{b,A} < T_{b,D} < T_{b,B}$$



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$$T_{b,A} < T_{b,C} < T_{b,B} < T_{b,D}$$



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Figure 8. RD column schemes in case of fixed feed inlets at the top and the bottom parts of reactive zone for (a) group I_p , (c) group III_p , (e) group III_r and in case of varied feed inlets to obtain the lowest RR possible for (b) group I_p , (d) group III_p , (f) group III_r . The presented numbers next to the column show the RD stages. All RD column configurations are at $NTS=2 \cdot NTS_{min}$ in the applicability graph considering $K_{eq}=1$.

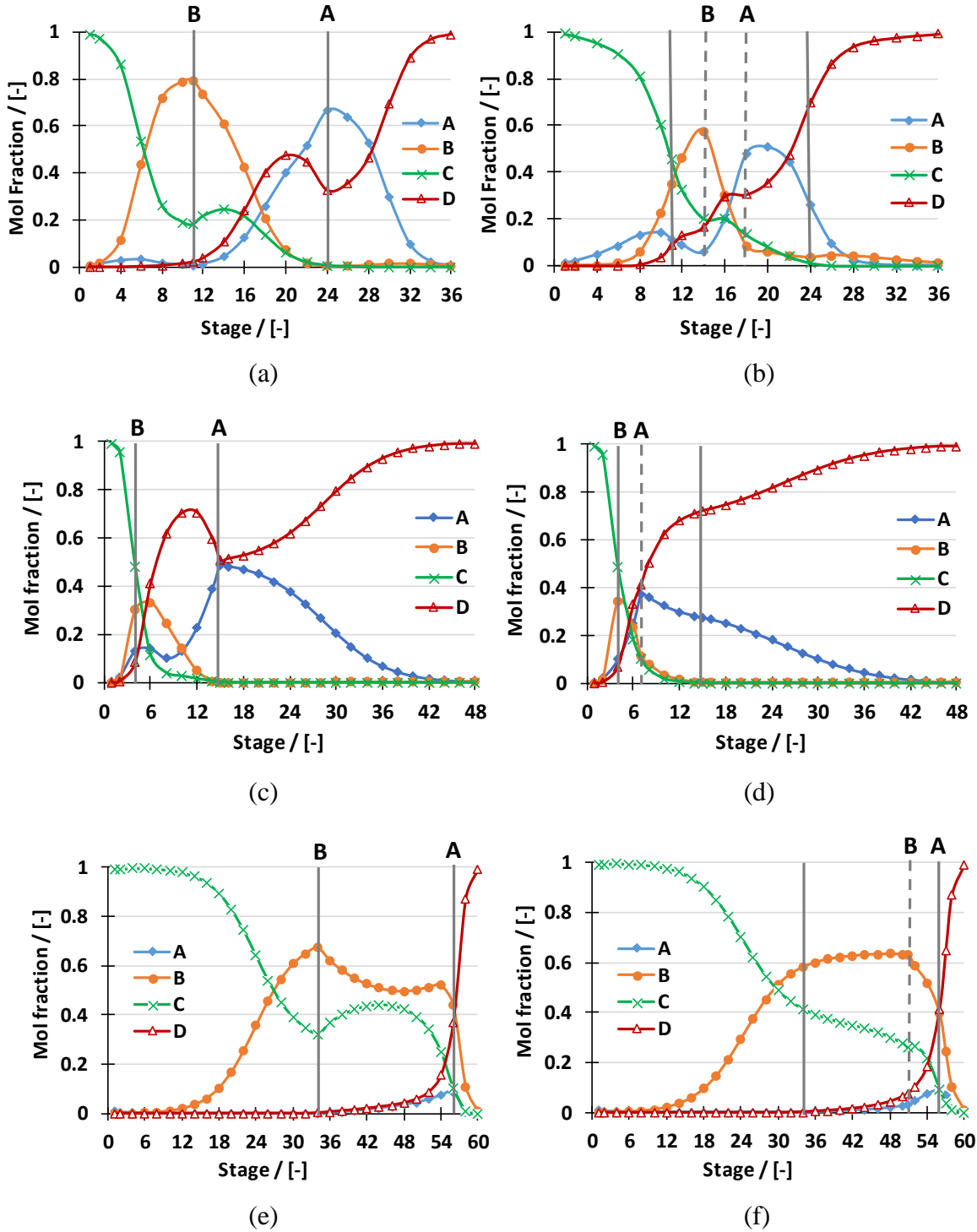


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