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# Accelerating the identification of the rate controlling steps by conducting microkinetic modeling on surrogate networks†

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Identifying the rate-controlling steps in an unknown reaction network can be time-consuming due to its inherent complexity. Here we present a strategy to simplify this process by focusing expensive barrier calculations on significant elementary steps. The strategy is constructed by determining significant elementary steps using the degree of rate control data, which is derived from microkinetic modeling calculations performed on surrogate networks, which are a series of networks generated by assigning fictitious values to unknown barriers while all the reaction energies are computed using density functional theory. The barriers for significant elementary steps are then calculated iteratively to refine the network. We demonstrate this strategy for the reaction of Fischer–Tropsch synthesis, which has already been extensively studied in our previous work. Applying the strategy, we identified the most rate-controlling step, achieving a 77% reduction in the number of transition state calculations compared to traditional methods. Additionally, a detailed analysis of the strategy reveals the correlation between the parameters in the strategy and its performance. We validate the practicability of the strategy by applying it onto testing networks and the potential limitations of the strategy are also discussed.

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## Introduction

A catalytic reaction network is typically complex; a system involving hundreds of intermediates can result in a network with numerous reactions and pathways. To extract useful information from such a network, a large number of calculations must be performed. Traditional calculations for mechanistic studies can be divided into three parts, as illustrated in Fig. 1: reaction energy calculations (structural optimizations of initial and final states of each elementary step), transition state (TS) calculations (TS search using NEB or constrained optimization or other methods), and microkinetic modeling (MKM). Conducting high-fidelity calculations for the

entire reaction network is both costly and time-consuming. Typically, TS calculations are the most time-consuming, often being several orders of magnitude more computationally expensive than other calculations. Therefore, reducing the number of TS calculations is the key to improving the efficiency of reaction kinetics studies.

This process can be significantly simplified by recognizing that only a small portion of reactions controls most of the properties of the entire network. The majority of the network properties, except for the kinetic parameters of the rate-controlling steps, have small impacts on the overall reaction network. The rate-controlling step can be simply interpreted as the slowest step of the favored pathway and it can strongly influence the overall reaction rate of the network. In order to identify rate-controlling steps, one popular method is the degree of rate control (DRC). This approach for analyzing multistep reaction mechanisms was introduced by Campbell,<sup>1–4</sup> in which the “degree of rate control for elementary step  $i$ ”,  $DRC^i$ , is defined as

$$DRC^i = \frac{k_i}{r} \left( \frac{\partial r}{\partial k_i} \right)_{k_{j \neq i}, K_i} = \left( \frac{\partial \ln r}{\partial \ln k_i} \right)_{k_{j \neq i}, K_i} \quad (1)$$

where  $r$  is the net reaction rate to the product of interest,  $k_i$  is the forward rate constant for step  $i$ ,  $k_{j \neq i}$  are the rate constants for steps other than  $i$ , and  $K_i$  is the equilibrium constant for step  $i$ . The larger the numeric value of  $DRC^i$  is for a given step, the greater the influence of its rate constant is on the overall reaction rate  $r$ . The elementary step with the largest DRC value is

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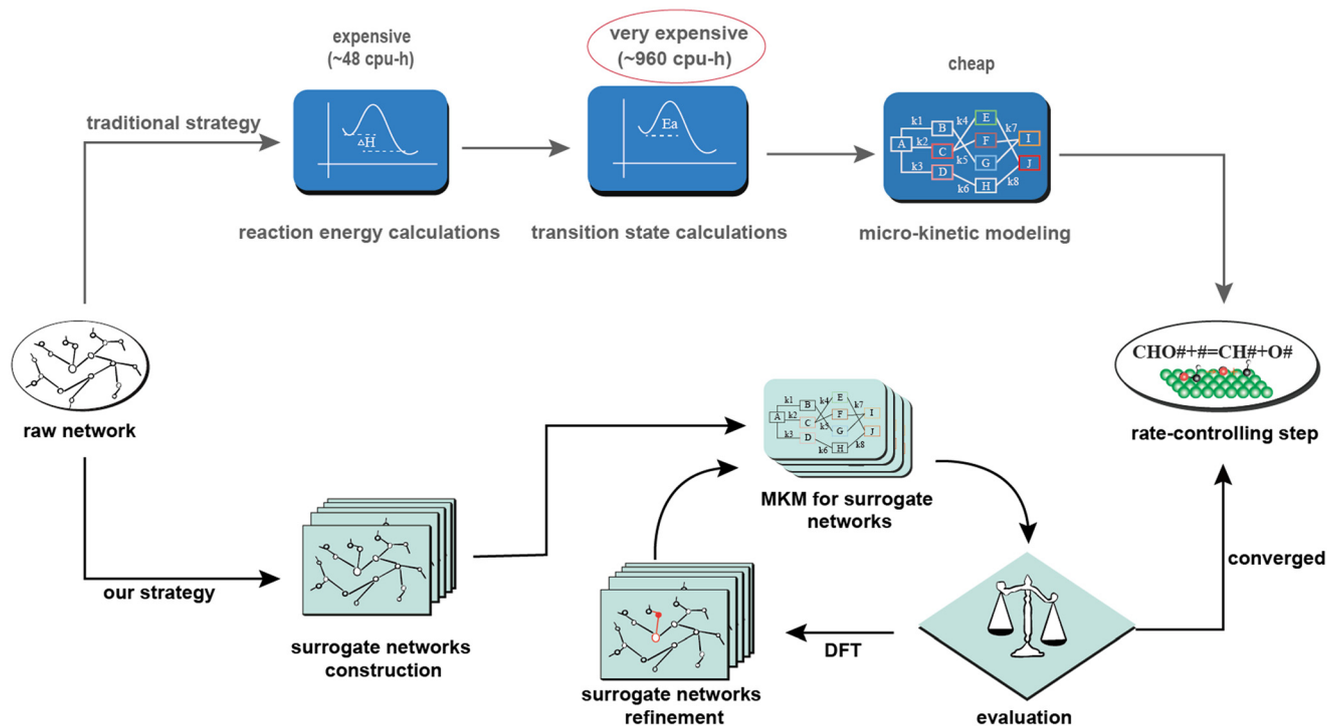
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**Fig. 1** The traditional method (upper) and our strategy (lower) of determining rate-controlling steps. Our strategy efficiently identifies the rate-controlling steps in the reaction network by minimizing the number of TS calculations. First, surrogate networks with accurate reaction energies and fictitious energy barriers are constructed. MKM on the surrogate networks provides kinetic analysis to identify significant elementary steps. The activation energies of the highest-priority steps are calculated iteratively using DFT, with the network being updated to improve accuracy. This process continues until convergence is achieved.

identified as the most rate-controlling step, which is crucial for understanding the mechanism of the reaction network and valuable for designing more effective catalysts. The challenge is that obtaining the DRC data requires MKM of the network, which involves extensive calculations of reactions and activation energies. Therefore, a key question arises: how quickly can the rate-controlling step of an unknown network be identified with as little calculation as possible?

To address this, researchers have proposed various solutions,<sup>5–14</sup> primarily relying on approximations like the BEP relation<sup>15–20</sup> or machine learning models to minimize computational resources and time. As a representative example, by using the prediction of machine-learning and approximations, such as scaling relations, Ulissi *et al.* developed an approach to determine the significant reactions on the favored pathway: by focusing only on these reactions with full density functional theory (DFT) calculations at each iteration, they achieved remarkable reduction of time and resource cost on studying the network of syngas.<sup>9</sup> Despite all the progress in this field, two main improvements could still be made. One is to achieve a more optimal trade-off between accuracy and efficiency. We need to determine when to use an approximate method that saves time but is less accurate, and when to choose a high-fidelity DFT calculation, which is more precise but also more time-consuming. The second improvement is to develop a better strategy to identify significant elementary steps.

In this work, we aim to address these issues. We developed a workflow to identify the rate-controlling steps of a reaction network, significantly reducing computational effort by performing a limited number of TS calculations. The strategy used DFT calculations to obtain reaction energies, and the MKM was conducted using CATKINAS, a software developed by our group and widely used.<sup>21–30</sup> We saved time and computational resources by reducing barrier calculations, as shown in Fig. 1. Our workflow uses DRC data as an indicator to find significant elementary steps and calculate the corresponding TS energy at each iteration. The characteristics of the reaction network are refined at each iteration, and the rate-controlling steps of the network are determined at the end of the refinement. We demonstrated this workflow for the Fischer-Tropsch (FT) reaction,<sup>31–46</sup> a process that utilizes synthesis gas to convert into hydrocarbons and oxygenated hydrocarbons. Using the strategy, we identified the most rate-controlling step with reducing the number of TS calculations by 77%.

## Result

### General workflow

As mentioned before, traditional reaction mechanism calculations consist of three parts: calculating reaction energies, determining the TSs, and performing the microkinetic modeling (MKM). Comparing the computational cost of each calculation, it



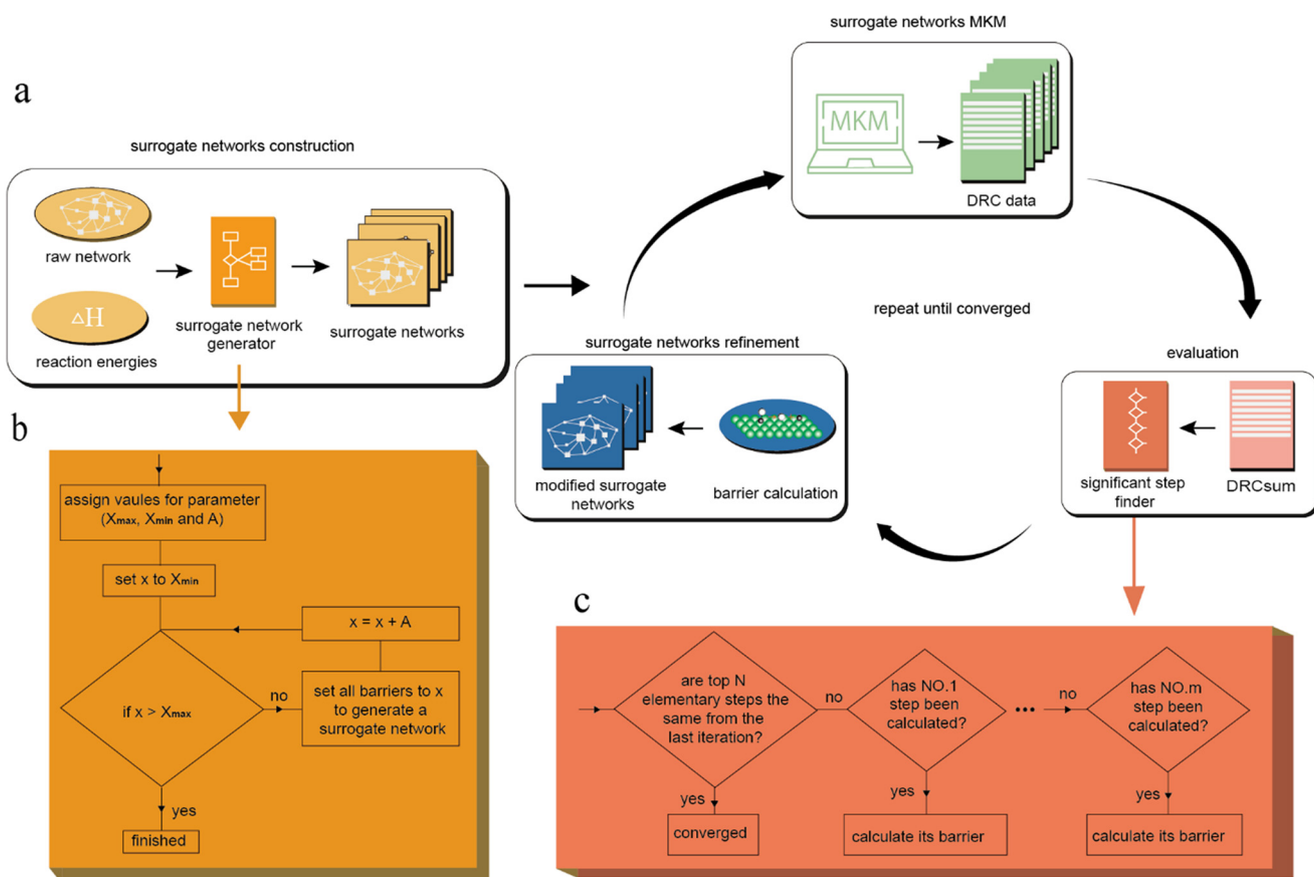
is well known that the TS calculations are much more expensive than the other calculations. This leads to the core of our strategy: minimizing the number of TS calculations while maintaining a high level of accuracy. We managed to achieve this goal by constructing a workflow to find the significant elementary steps and calculate their TS energies iteratively. As illustrated in Fig. 2a, the entire workflow is divided into four parts: surrogate network construction, surrogate network MKM, evaluation and surrogate network refinement.

In constructing the surrogate networks, the first step is to define all possible chemical species and elementary steps to be considered in the network. Then the adsorption energies of all intermediates are calculated to obtain the reaction energy for each elementary step. In this way, a raw reaction network with only the thermodynamic information is built. Next, a surrogate network is generated by assigning fictitious values to energy barriers of all the elementary steps in the network. To construct a series of surrogate networks, as shown in Fig. 2b, we began with assigning values to  $X_{\max}$  (the biggest fictitious barrier value),  $X_{\min}$  (the smallest fictitious barrier value) and  $A$  (the interval of fictitious barrier between surrogate networks) in our strategy. As the fictitious barrier  $x$

changes from  $X_{\max}$  to  $X_{\min}$ , a series of surrogate networks were constructed. For clarity, when we built the surrogate networks, the fictitious energy barrier  $x$  is only added to the exothermic forward/backward reactions in each step. For example, assuming the energies of the initial state and the final state for a specific elementary step are  $E_1$  and  $E_2$ , respectively, the energy of TS will be  $E_1 + x$  when  $E_1 < E_2$ ; otherwise, it will be  $E_2 + x$  if  $E_2 < E_1$ .

In the MKM phase of surrogate networks, we conducted MKM on all the surrogate networks to obtain their kinetic information, which contains DRC data. Since our goal in this work was to identify the key elementary steps within a highly complex reaction network, coverage dependence was not explicitly considered to streamline the calculation process. In the evaluation phase, we introduced an indicator,  $\text{DRC}_{(\text{sum})}$ , to assess the significance of each elementary step. For a specific elementary step  $i$ ,  $\text{DRC}_{(\text{sum})}^i$  is defined as the sum of the absolute value of the corresponding DRC in each surrogate network as follows:

$$\text{DRC}_{(\text{sum})}^i = \sum_{z=1}^n w_z \times |\text{DRC}_z^i| \quad (2)$$



**Fig. 2** Workflow used to generate surrogate networks to iteratively determine significant elementary steps. (a) Network refinement methodology. In each iteration, the significant elementary step is determined based on kinetic information obtained from MKM of surrogate networks and TS calculation will be conducted on the significant elementary step using DFT to refine surrogate networks. (b) Flow chart of surrogate network generator in the surrogate networks construction phase. (c) Flow chart of the significant step finder in the evaluation phase.



where  $n$  is the total number of surrogate networks,  $\text{DRC}_z^i$  is the DRC value of elementary step  $i$  for the  $z$ -th surrogate network, and  $w_z$  is the weight of  $\text{DRC}^i$  for the  $z$ -th surrogate network. To prevent positive and negative values of DRCs cancelling each other, the  $\text{DRC}_{(\text{sum})}$  is calculated as the sum of the absolute values of the DRCs across all the surrogate networks. We then assessed the significance of each elementary step by referring to the value of its  $\text{DRC}_{(\text{sum})}$ . This information is later used as a reference to identify the most significant elementary step and to determine which energy barrier needs to be calculated during the evaluation phase. To identify the most significant step, it is first necessary to assess whether the calculation has reached convergence. Convergence is deemed achieved when the top  $N$  elementary steps, characterized by the highest  $\text{DRC}_{(\text{sum})}$  values in the current iteration, remain consistent with those from the previous iteration, where  $N$  is a predefined parameter. If convergence is not attained, further evaluations are conducted to determine which elementary step's TS should be calculated using DFT in the subsequent phase.

In the refinement phase of surrogate networks, we used DFT calculations to obtain the barrier of the chosen elementary step and updated all the surrogate networks by replacing the fictitious barrier of the selected step with its actual barrier. This process is repeated until the convergence criterion is met.

### Identification of rate-controlling steps

In this work, we demonstrated this workflow for the FT reaction network on Co(0001), which has been thoroughly studied in our previous work.<sup>47</sup> Since the structure of the reaction network and the reaction energies and activation energies have already been calculated,<sup>47</sup> we skipped the network determination, reaction energy and barrier calculations in our current workflow, and focused on demonstrating how our workflow determines the significant elementary step iteratively and refines the network. We set parameters  $X_{\text{max}}$ ,  $X_{\text{min}}$  and  $A$  to 1.5 eV, 0.0 eV and 0.1 eV, respectively, for the construction of surrogate networks. Detailed explanation about parameters setting will be included in the next section. For simplicity, the weight  $w_z$  in the definition formula of  $\text{DRC}_{(\text{sum})}$  (eqn (1)) was uniformly set to 1 for every surrogate network.

The refinement procedure is illustrated in Fig. 3a, which shows the entire reaction network with nodes as reaction species and lines as elementary steps. For visual simplicity, hydrogen is omitted. Refinements were performed under various settings of parameter  $N$ . As  $N$  increased, a greater number of TSs were required to be calculated for the iteration to terminate. The additional elementary steps that needed to be calculated by DFT, as  $N$  was increased to  $N + 1$ , are indicated by colored lines in Fig. 3a. Since parameter  $N$  defines the convergence criterion in the evaluation phase, increasing  $N$  requires more rate-controlling steps to converge before the iteration stops. This would result in more TS

calculations and potentially more accurate outcomes. The accuracy of the final result and the number of iterations before convergence are both plotted against the value of  $N$  in Fig. 3b. In this work, 'accuracy' is defined as the ratio of the sum of DRC values of the calculated  $M$  elementary steps to the total DRC sum across all elementary steps, with all DRC values derived from a reference set based on the full reaction network from prior work,<sup>47</sup> which conducted calculations of all the TS energies through DFT calculations. This metric is designed specifically for internal assessment of the strategy's predictive accuracy within this study and is not intended for direct application in real-world determinations of rate-controlling steps.

$$\text{Accuracy} = \frac{\sum_{i \in M} |\text{DRC}^i|}{\sum_{i \in Q} |\text{DRC}^i|} \quad (3)$$

where  $M$  is a collection of all the elementary steps that have been studied using DFT in our workflow, and  $Q$  is a collection of all the elementary steps in the network. As  $N$  increases, the accuracy of the final result and the number of iterations before convergence also increases (Fig. 3b). In addition, there are three significant increases on the accuracy curve, each representing the identification of one of the three most rate-controlling steps for the FT reaction network on Co(0001). In Fig. 3c, we show the top 3 most rate-controlling steps and their corresponding pathways determined by our strategy under different convergence criterion, in which the rate-controlling steps were highlighted in red. Only the top 3 rate-controlling steps are illustrated, because the DRC values of the top 3 rate-controlling steps, for the current studied network, were at least one magnitude higher than those of the rest elementary steps according to the reference set.

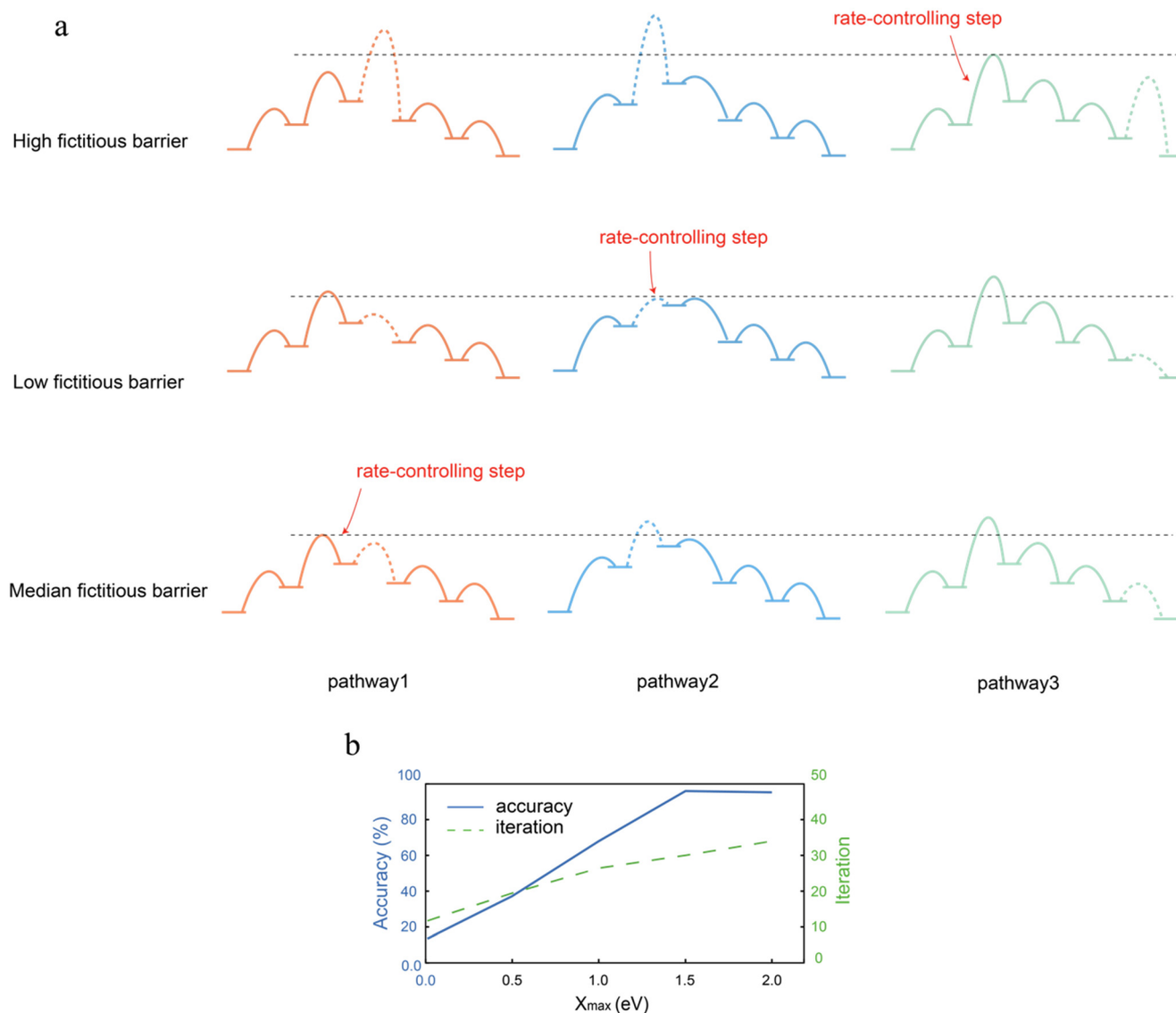
The first accuracy significant increase occurs when  $N$  was set to 1; with 11 iterations in total, the calculation converged. The most rate-controlling step was determined as  $\text{CH}_3\text{O} \rightleftharpoons \text{CH}_3 + \text{O}$ , which is the third most rate-controlling step according to the reference set. The second significant accuracy increase occurs when  $N$  was set to 2, with the convergence achieved after 16 iterations. In this case, the elementary step of  $\text{CHO} \rightleftharpoons \text{CH} + \text{O}$  was identified as the most rate-controlling step, which aligns with the result from the reference set. This means that using the method developed in this work, we only conducted 15 TS calculations to correctly predict the most rate-controlling step for the FT reaction network on Co(0001), reducing 77% of TS calculations. The third accuracy significant increase occurs when  $N$  was set to 5, with the convergence achieved after 29 iterations. This time we correctly identified the top 3 most rate-controlling steps and reached a value of accuracy of 0.97.

### Surrogate network assessment

We investigated the most important and innovative aspects of our workflow. Namely, we quantitatively assessed the construction and utilization of the surrogate networks. It is worth mentioning that the general benefit for incorporating surrogate networks into our workflow is that it prevents the







**Fig. 4** The construction of surrogate networks plays an important role in the identification of rate-controlling steps. (a) Schematic representation of rate-controlling step variations across different surrogate networks, assuming that the reaction network consists of three pathways. Solid lines indicate elementary steps with a calculated barrier, while dashed lines represent elementary steps with undetermined barriers. Different surrogate networks assign various fictitious barriers to these undetermined steps, leading to shifts in the rate-controlling step across networks. The figure demonstrates how varying fictitious barrier levels can shift the rate-controlling step within the reaction network, leading to calculations of more potentially significant elementary steps. (b) Illustration of accuracy and number of iterations before convergence as functions of  $X_{\max}$ . As  $X_{\max}$  increases, the number of iterations required for the calculation to converge increases. The highest accuracy was reached when  $X_{\max}$  was set to 1.5 eV.

in different surrogate networks are assigned to different fictitious values. For example, in a surrogate network with high fictitious barriers, all undetermined barriers are assigned to high values. In this case, pathway 3 becomes the favored pathway because the TS with the highest energy level in pathway 3 is still lower than the TSs with highest energy levels in the other two pathways. As a result, the elementary step containing this TS on pathway 3 is considered the rate-controlling step for the network. A similar process occurs in the networks with low and median fictitious barriers, leading to variations in the rate-controlling steps across different surrogate networks. This variation may result in additional

TS calculations for potentially important steps, thereby increasing the likelihood of identifying the actual rate-controlling step.

To investigate how the choice of surrogate networks influences the efficiency and accuracy of our strategy, we applied our workflow on the FT reaction network on Co(0001) with different range of fictitious barriers. We constructed 5 calculation groups, which sets  $X_{\max}$  to 0.0 eV, 0.5 eV, 1.0 eV, 1.5 eV and 2.0 eV, respectively, while  $X_{\min}$ ,  $A$  and  $N$  were set to 0.0 eV, 0.1 eV and 5 uniformly. As shown in Fig. 2b, as  $X_{\max}$  increases, the range of fictitious barriers become wider and more surrogate networks with higher

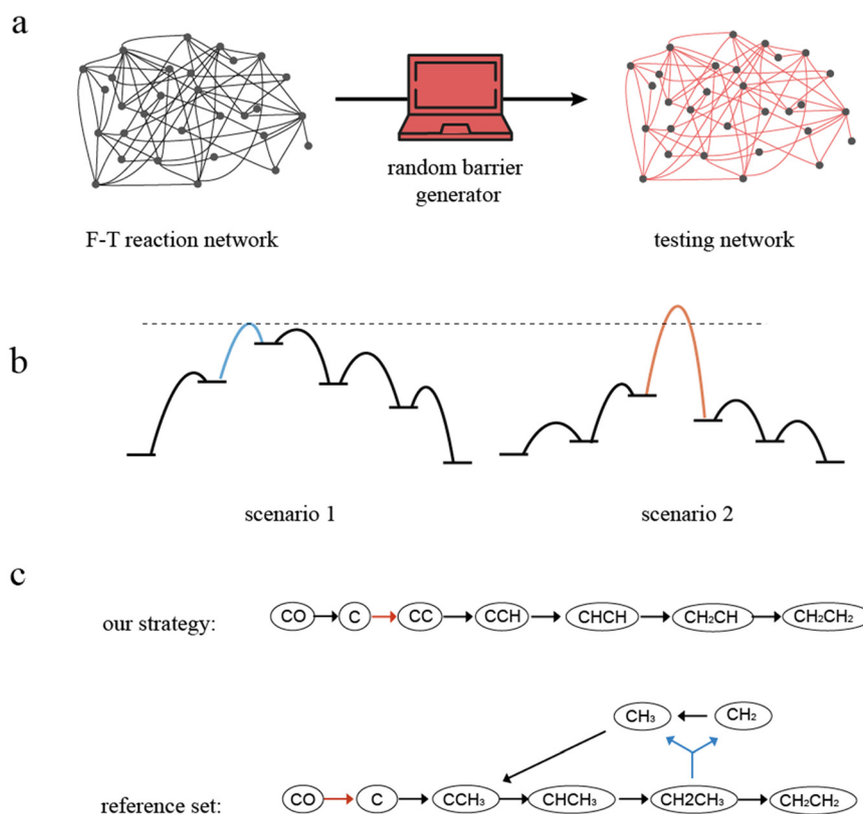


fictitious barriers will be generated for the corresponding group, leading to different results of refining processes. The accuracy of the final result and the number of iterations needed before convergence for each calculation are shown in Fig. 4b. It can be seen that as  $X_{\max}$  increases, the number of iterations increases. The calculation group with  $X_{\max}$  set to 1.5 eV reached the highest accuracy. The reason for this is that most of the barriers in the reaction network we studied lies in the range of 0.0–1.5 eV. Hence, the calculation group with  $X_{\min}$  and  $X_{\max}$  set to 0.0 eV and 1.5 eV is suitable for probing the rate-controlling steps of the network because its range of fictitious barriers matches the range of real barriers, ensuring that there are enough surrogate networks to investigate potentially significant elementary steps. In contrast, the groups with the range of fictitious barriers too narrow will not have enough surrogate networks to detect some rate-controlling steps and might miss some significant elementary steps. In addition, for the groups where the range of fictitious barriers is too wide, such as the calculation group with  $X_{\max}$  set to 2.0 eV, the surrogate networks with unusually high fictitious barriers will disproportionately influence the identification of the rate-controlling steps, thereby reducing the accuracy of the results.

### Examination with testing networks

We generated testing networks to demonstrate the generality of our rate-controlling step identification strategy and to assess its potential limitations. As shown in Fig. 5a, these testing networks were constructed based on the FT reaction network on Co(0001) previously studied.<sup>47</sup> By assigning random values to the barriers of all elementary steps while keeping the network structure and reaction energies unchanged, the testing networks were generated. Then we used the same strategy demonstrated above to determine the rate-controlling steps of those new testing networks. The parameters  $X_{\max}$ ,  $X_{\min}$ ,  $A$  and  $N$  were set to 1.5 eV, 0 eV, 0.1 eV and 5, respectively, as they achieved relatively high accuracy and efficiency for the original network, as shown in Fig. 4b.

Four testing networks were generated in total as the testing reference sets, and our strategy successfully predicted the most rate-controlling step for 3 of them as shown in Table 1. After analyzing the failed case, we find that there are mainly two types of scenarios where our strategy may encounter difficulties on the task of identifying the rate-controlling steps. As shown in Fig. 5b, the first scenario is that the rate-controlling step exists



**Fig. 5** Constructing new testing networks to further examine our current strategy. (a) Workflow of generating the testing networks. The testing networks are generated by assigning random values to the barriers of all elementary steps while keeping the network structure and reaction energies of previously studied FT reaction network unchanged. All the random barriers we generated were between 0.0 eV and 1.5 eV. (b) Two types of scenarios where our strategy may encounter difficulties discussed in the main text. (c) Favored pathways determined by the reference set and our strategy with rate-controlling steps highlighted in red. The reason for the disaccord is that there was an elementary step with an extremely low barrier in the favored pathway according to our generated reference set, which is highlighted in blue.



**Table 1** Results of applying our strategy to our generated networks to further testing our approach. The testing network, in which the barriers have been previously assigned, serves as the reference set

Testing networks	Amount of iterations	Top 3 reactions with biggest DRC(sum)	
		Our strategy (reaction #)	Reference set (reaction #)
T1	21	59, 3, 58	59, 3, 58
T2	15	43, 4, 6	3, 25, 58
T3	20	59, 58, 6	59, 6, 30
T4	20	3, 58, 59	3, 58, 59

on a thermodynamically unfavorable but kinetic favorable pathway, which means that the energies of intermediates are relatively high but energies of TSs are relatively low on this pathway. There is a possibility that our strategy would identify this pathway as an unfavored pathway and start searching for a possible rate-controlling step elsewhere. In the second scenario, if there is a pathway that is thermodynamically favorable but kinetically unfavorable and does not include the rate-controlling step, our strategy might mistakenly identify this pathway as favored and converge on an incorrect rate-controlling step. Thus, our strategy may encounter difficulties when certain types of elementary steps have unexpected high or low barriers.

In the failed case of testing network T2, when the iteration stopped, our strategy identified the most rate-controlling step as  $C + C \rightleftharpoons CC$  as shown in Fig. 5c. However, the actual most rate-controlling step, according to the testing reference set, is  $C + O \rightleftharpoons CO$ . After thorough inspection, we found that the reason network T2 converged to an incorrect elementary step is that the actual favored pathway, according to the reference set, is a thermodynamically unfavorable pathway. This means that the energies of the species involved in this pathway are relatively high. However, the barrier of an elementary step ( $CH_2CH_3 \rightleftharpoons CH_2 + CH_3$ ) on this pathway, highlighted in blue in Fig. 5c, is unrealistically low (0.037 eV), which unexpectedly makes this pathway favorable. To identify the most rate-controlling step for the testing network T2, we increased the value of  $N$  to calculate more potential significant steps. After we increased  $N$  to 22, our strategy converged at the correct rate-controlling steps after 47 iterations.

## Discussion

The strategy presented in this paper could efficiently reduce the amount of TS calculations; it successfully identified the most rate-controlling step of the FT reaction network on Co(0001), which was calculated in our previous work,<sup>47</sup> with only 23% of TS calculations compared to the traditional method. The challenging aspect of this strategy lies in determining its parameters, including those for constructing surrogate networks and setting the convergence criterion. Based on our findings, for an

unknown reaction network, we recommend setting the fictitious barrier range to 0.0–2.0 eV and  $N$  to above 5% of the total number of the elementary steps of the reaction network. This setting achieved a great trade-off between accuracy and efficiency as shown in Fig. 3b and 4b.

For the reaction network tested in this work, which includes 26 species and 67 reactions, the good parameters  $X_{max}$ ,  $X_{min}$ ,  $A$ , and  $N$  were found to be 1.5 eV, 0.0 eV, 0.1 eV, and 5, respectively. This set choice has accurately predicted the top 3 rate-controlling steps while reducing the number of TS calculations by 55%, as shown in Fig. 4b. Compared to traditional calculations, our strategy focuses on identifying the rate-controlling steps as quickly as possible and performing barrier calculations only on those steps identified as significant. This approach saves substantial time and computational resources. Additionally, unlike methods that rely on machine learning or other approximations, the strategy developed in this work has greater generality. However, it is important to note that our strategy does not guarantee 100% accuracy in predicting rate-controlling steps if the parameters are not properly selected.

## Conclusion

In the current work, we developed a strategy to efficiently identify the rate-controlling steps of a given reaction network by leveraging DRC data to highlight significant elementary steps and allocate computational resources accordingly. By applying our strategy to the FT reaction network on Co(0001), we achieved a 77% reduction on TS calculations compared to the traditional method. This approach allows for significant reductions in time and computational costs when exploring unknown reaction networks.

The advancements of our method compared to previous approaches are summarized as follows:

1. Efficient rate-control predictions: by conducting MKM calculations on surrogate networks, our method enables the prediction of rate-controlling steps using DRC without the extensive barrier calculations.
2. Balanced accuracy and efficiency: by reducing the number of TS calculations while using high accuracy methods to conduct the calculations of reaction energies and MKM, our strategy reached excellent balance between accuracy and efficiency.
3. General applicability: unlike methods that rely on approximations such as BEP relations, our strategy does not depend on specific approximations, making it applicable to the study of any reaction network.

Finally, our method could be further advanced by integrating experimental data or machine learning models to predict transition state energies or adsorption energies. Combining these approaches has the potential to reduce computational costs and improve accuracy when exploring new reaction networks. Future work may focus on refining strategies to efficiently identify critical elementary steps and incorporating additional catalytic effects, such as coverage



effects, to enhance the practicality and effectiveness of the current method.

## Methods

All microkinetic modeling calculations are carried out *via* the CATKINAS package, which is a microkinetic simulation package developed by our group and widely used.<sup>21–30</sup>

## Data availability

The data that support the findings of this study are available from the corresponding author on reasonable request.

## Author contributions

P. H. and J. Z. conceived the project. H. Y. L. carried out all the calculations. H. Y. L. wrote the first draft of the manuscript and prepared figures and all the authors contributed to the revision of the manuscript.

## Conflicts of interest

The authors declare no competing financial interest.

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