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**Kinetic model assessment for the synthesis of γ -valerolactone from *n*-butyl levulinate and levulinic acid hydrogenation over the synergy effect of dual catalysts Ru/C and Amberlite
IR-120.**

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Abstract

The production of platform molecules from the valorization of lignocellulosic biomass is increasing. Among these platform molecules, γ -valerolactone (GVL) is a promising one and could be used for different industrial applications. This molecule is synthesized from levulinic acid (LA) or alkyl levulinates (AL) through a tandem hydrogenation/cyclization (lactonization) cascade. A lot of investigations have been carried out to develop the best catalyst for the hydrogenation step by using solely LA or AL. However, one should keep in mind that in the AL production via fructose alcoholysis, there is also LA production, and both are present in the product mixture during the further conversion. To the best of our knowledge, no article exists describing the hydrogenation of LA and AL simultaneously in one-pot. Also, the literature reporting the use of solid catalyst for the second cyclization step is rare. To fill this gap, the hydrogenation of levulinic acid and butyl levulinate was studied over Ru/C and Amberlite IR-120. Several kinetic models were evaluated via Bayesian inference and K-fold approach. The kinetic assessment showed that a non-competitive Langmuir-Hinshelwood with no dissociation of hydrogen where LA, BL and H_2 are adsorbed on different sites (NCLH1.2) and non-competitive Langmuir-Hinshelwood with dissociation of hydrogen where LA, BL (butyl levulinate) and H_2 are adsorbed on different sites (NCLH2.2) are the best model to describe this system. The presence of LA and Amberlite IR-120 allows to increase the kinetics of cyclization steps, and *in fine* to accelerate the production of GVL.

Keywords: γ -valerolactone, kinetic modeling, Bayesian statistics, cross-validation, levulinic acid.

1. INTRODUCTION

The valorization of biomass to chemicals, fuels, or materials is essential to decrease the use of fossil raw materials, reduce CO₂ emission, and favor circular economies [1,2]. To make biorefineries efficient in production and energy consumption, knowledge of kinetics, catalysis, and thermodynamics is vital [3]. Currently, academia and the private sectors are focusing their efforts on the valorization of lignocellulosic biomass (LCB) to avoid the dilemma of “food versus fuels” [4].

There are several platform molecules that can be derived from the valorization of LCB [5–7] such as 1,4-diacid, 5-HMF and 2,5-FDCA, 3-HPA, aspartic acid, glutamic acid, glucaric acid, itaconic acid, sorbitol, ABE or levulinic acid.

Research interest at the academy and industrial levels for the utilization of levulinic acid platform chemicals is increasing. Levulinic acid (LA) or alkyl levulinate (AL) are produced from the solvolysis or alcoholysis of cellulose-hemicellulose [8][9]. The market for these molecules (LA or AL) is growing [10,11] due to the versatile use and valorization of these molecules in different industrial sectors.

The hydrogenation of LA or AL leads to the production of γ -valerolactone (GVL), also considered as a platform molecule [12–14]. GVL is regarded as an excellent green aprotic polar solvent [15][16][17][18] due to its low vapor pressure and high flash point [14,19,20]. GVL is stable, renewable and non-toxic [21]. This platform molecule can be used as an intermediate in the production of many value-added chemicals [1]. From an energetic viewpoint, GVL is a good intermediate for bio-jet fuel production via its decomposition into butene [22][23].

There are several routes for the production of GVL [24,25]; among those involving the hydrogenation of AL or LA, one can distinguish:

- use of molecular hydrogen [26–38];
- in situ* decomposition of formic acid into hydrogen [39–47];
- the Meerwein–Ponndorf–Verley reaction, i.e., the use of alcohols for catalytic transfer hydrogenation [48–50].

Hence, one can notice that the literature concerning the production of GVL from the hydrogenation of AL or LA is quite vast. Researchers have put a lot of effort into developing catalysts, evaluating reaction pathway, and using different feedstock.

The most common method for the production of GVL is the hydrogenation of AL or LA with molecular hydrogen over Ru/C, since it provides the best atom economy. This reaction comprises of two steps [38] (Fig. 1) : the hydrogenation of the substrate to obtain an intermediate, and the cyclization of the intermediate to GVL. During the first step, the carbonyl group of AL or LA is hydrogenated. During the second step, the hydroxyl group of the intermediate reacts with the ester group leading to a cyclization.

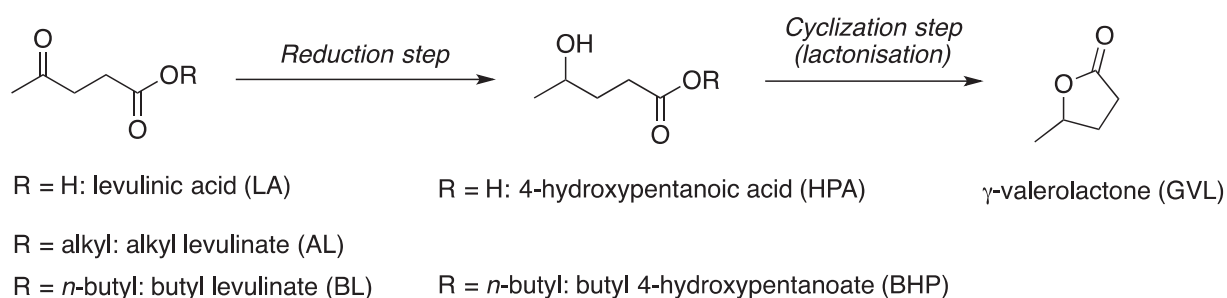


Fig. 1. Hydrogenation of LA or AL to GVL over Ru/C.

One should remember that the alcoholysis of fructose leads to LA and the corresponding AL [51]. Thus, in the case of GVL production from fructose via alcoholysis and consecutive hydrogenation, LA is also present in this chemical system. Piskun et al. [38] observed that protons from LA dissociation can catalyze the cyclization step. Thus, it could be beneficial for the production of GVL to start from the products of fructose alcoholysis, namely the presence of LA and AL.

Highly concentrated LA solutions present a corrosion risk, which is why, the use of AL has gained interest, but its reactivity is lower compared to LA. A good compromise could be to use a mixture of both reactants. Methyl, ethyl and n-butyl levulinates are the three most studied AL. A thermal risk assessment of AL hydrogenation [36] showed that the risk of thermal runaway for the hydrogenation of methyl levulinate is higher than for butyl levulinate (BL), and the use of butanol for the alcoholysis of carbohydrates into BL is a promising route [52].

The choice of solvent is also important. A previous study showed that the solubility of hydrogen is higher in GVL solvent [28]. Besides, the hydrogenation of BL in GVL solvent allows to work at higher temperature due to the high vapor pressure of the different chemicals [52].

For that reason, the current study focuses on:

- 1) the hydrogenation of LA and BL reactants into GVL.
- 2) the use of cation exchange resins under H form to catalyze the cyclization reaction. The cyclization step has been shown to be slower for alkyl levulinates than for LA [38]. The use of such acid resin catalyst can significantly increase the reaction kinetics. To the best of our knowledge, there are few studies in this field [53,54]. In the first study, Moreno-Marrodan and Barbaro [53] demonstrated that the use of heterogeneous catalyst based on sulfonated cation exchange resin and embedded Ru nanoparticles leads to the complete conversion of LA to GVL with remarkable selectivity at low temperatures and H₂ pressure, as well as excellent catalyst durability and no need

for additives. The second study [54] analysed the hydrogenation of LA to GVL catalysed by a commercial Ru supported catalyst in combination with Amberlyst A70, showing a high selectivity to GVL also at mild processing conditions.

This article aims to assess different plausible kinetic models for the hydrogenation of BL and LA synergically catalyzed by Ru/C and Amberlite IR-120 via Bayesian inference. The Bayesian approach to developing a kinetic model for catalytic systems has become increasingly popular [55–59].

Traditionally, kinetic models are evaluated mainly by the fit to experimental data and the coefficient of determination. Some researchers incorporate in the assessment: the residual analysis, the credible intervals of the estimated parameters, the number of parameters via the Akaike information criterion (AIC) [60] and the correlation matrix. All these analyses are done solely on the training dataset, and few studies use a validation dataset to evaluate prediction quality of the kinetic model. The validation stage is rarely done because some experimental data are not used in the regression stage, so the model accuracy is lower. To overcome this issue, cross-validation, and more particularly the K-fold approach, is used in this investigation [61]. Cross-validation is also a way to determine the best model.

2. MATERIALS AND METHODS

2.1 Chemicals

Chemicals used in this work, without further purification, are described in this section. Hydrogen gas (H_2 purity > 99.999 vol %) from Linde. *n*-Butyl levulinate (BL purity = 98 wt%), CAS: 2052-15-5, was purchased from Alfa-Aesar. Levulinic acid (LA purity > 98 wt%), CAS: 123-76-2, was purchased from Acros Organics. γ -Valerolactone (GVL purity $\geq$ 99 wt%), CAS: 108-29-2, was purchased from Sigma Aldrich. Acetone (analytical grade), CAS: 67-64-1, was bought from VWR. Ruthenium, 5% on activated carbon powder, reduced, nominally 50% water, CAS: 7440-18-8, was purchased from Alfa-Aesar. Amberlite IR120, H-Form, ion-exchange resin, CAS: 78922-04-0, was purchased from Acros Organics.

2.2 Analytical methods

For the quantitative analysis of the samples, gas chromatography coupled with flame ionization detection technique was used. The equipment (GC) used is from supplier Scion Instruments, equipped with a GC capillary column ZB-5, a versatile, low polarity column. This column is composed of 95% of dimethylpolysiloxane and 5% of phenyl groups. The column has 30m of length, 0.32 mm of diameter and 0.25 μ m of film internal coating.

Helium (99.99%) was used as the carrier gas at a constant flow rate of 1.2 mL.min⁻¹ to transfer the sample from the injector, through the column, and into the FID-detector. The temperature of the injector and the detector were set at 250°C. The oven temperature ramp was set to 50°C (1 min) - 20°C min⁻¹ - 200°C (1 min). Samples were diluted in acetone and injected into the GC. The injection volume was 1 μ L, and the split ratio was 20:1.

2.3 Kinetic experiments

To develop kinetic models for GVL synthesis from the hydrogenation of *n*-BL with LA, a total of 14 experiments were carried out varying the initial operating conditions such as pressure, temperature, initial concentrations of the reactants, and catalyst loadings (Table 1). GVL was used as a solvent. Experiments carried out with Amberlite IR 120 were done at a reaction temperature lower than 120°C to avoid the leaching of sulfonic groups.

Table 1. Experimental matrix for the kinetic study of the reaction scheme shown in Fig. 1.

Run	Pressure bar	Temp K	mcat_Ru kg (50% weight moisture)	mcat_Ambk g (dried)	m0GVL kg	m0BL kg	m0LA kg	BL ₀ mol/m ³	GVL ₀ mol/m ³	LA ₀ mol/m ³
1	20.6	404.15	0.0005	0	0.083	0.027	0.015	1272	7703	1105
2	20	403.15	0.0005	0	0.083	0.042	0	1912	6311	0
3	20.6	404.15	0.0005	0	0.083	0.027	0.015	1249	6606	1029
4	22	383.15	0.0005	0	0.083	0.027	0.015	1249	6606	1029
5	10.9	403.15	0.0005	0	0.083	0.027	0.015	1351	7169	1029
6	21.5	402.15	0.0005	0	0.083	0.022	0.02	1055	6831	1372
7	21.4	374.15	0.0005	0	0.083	0.027	0.015	1450	6606	1300
8	21.6	373.15	0.001	0	0.083	0.027	0.015	1313	6841	1077
9	21.7	383.15	0.0015	0	0.083	0.022	0.02	879	5950	719
10	21.3	385.15	0.0015	0.010479	0.083	0.022	0.02	1093	6819	1390
11	9.7	375.15	0.0012	0.010	0.083	0.022	0.025	1117	6769	1597
12	9.5	414.15	0.0010	0.000	0.083	0.027	0.010	1415	7431	582
13	10.5	391.15	0.0012	0.010	0.083	0.022	0.025	1058	6586	1707
14	15.4	394.15	0.0012	0.006	0.083	0.03	0.015	1337	6515	1154

The autoclave (stirred tank reactor) used for these experiments is a stainless- steel laboratory-scale vessel with a capacity of 300 mL, which is equipped with a stirrer set at 1000 RPM, an electrically heating jacket, a cooling coil, a pressure sensor, and a temperature sensor (Fig. 2). Each experiment was carried out in isothermal and isobaric conditions. During these experiments, samples were taken at different times and then analyzed via GC-FID.

The concentration measurement uncertainty was evaluated via the standard deviation of replicate measurements. Each sample was analyzed thrice.

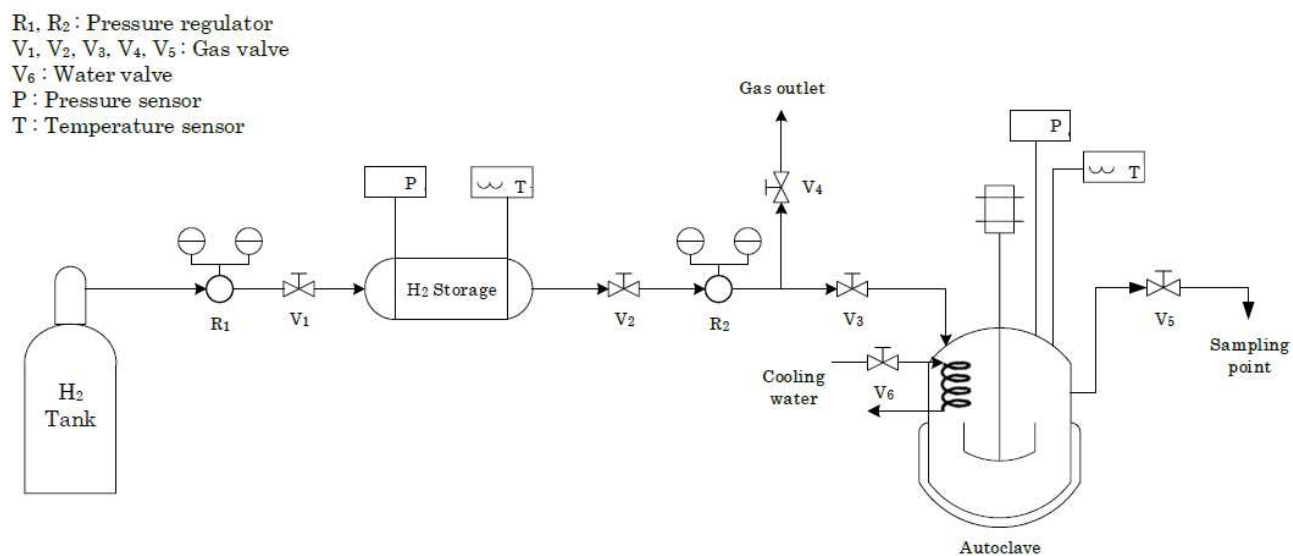


Fig. 2. Experimental setup of the autoclave.

3. RESULTS

This section describes the effects of experimental parameters such as temperature, pressure, and concentrations. on the hydrogenation and cyclization reactions from a phenomenological viewpoint. The hydrogenation of butyl levulinate or levulinic acid over Ru/C to GVL is a two-step reaction illustrated by Fig. 1[28,29,36,38]. In the first step, the carbonyl group of BL or LA is hydrogenated. In the second step, the hydroxyl group of the intermediate attacks the ester group leading to a cyclization.

The hydrogenation step was evaluated by the ratios $[BL]/[BL]_0$ and $[LA]/[LA]_0$. The cyclization step was assessed by the ratios $[BHP]/[BL]_0$ and $[HPA]/[LA]_0$. The standard deviations for BL, LA, BHP and HPA concentrations were found to be in average 1.76%, 3.64%, 2.41% and 3.64%, respectively. All the samples were analyzed three times. In Figs 3-8, the replicated points are also displayed.

3.1 Repeatability

In addition, to replicate three times the sample analysis, two similar experiments were reproduced to evaluate the repeatability of the results. Experiments 1 and 3 (Table 1) were carried in similar operating conditions in the experimental matrix. Fig. 3 shows that the protocol used in this study is repeatable. Experiments 1 and 3 resulted in practice in identical results.

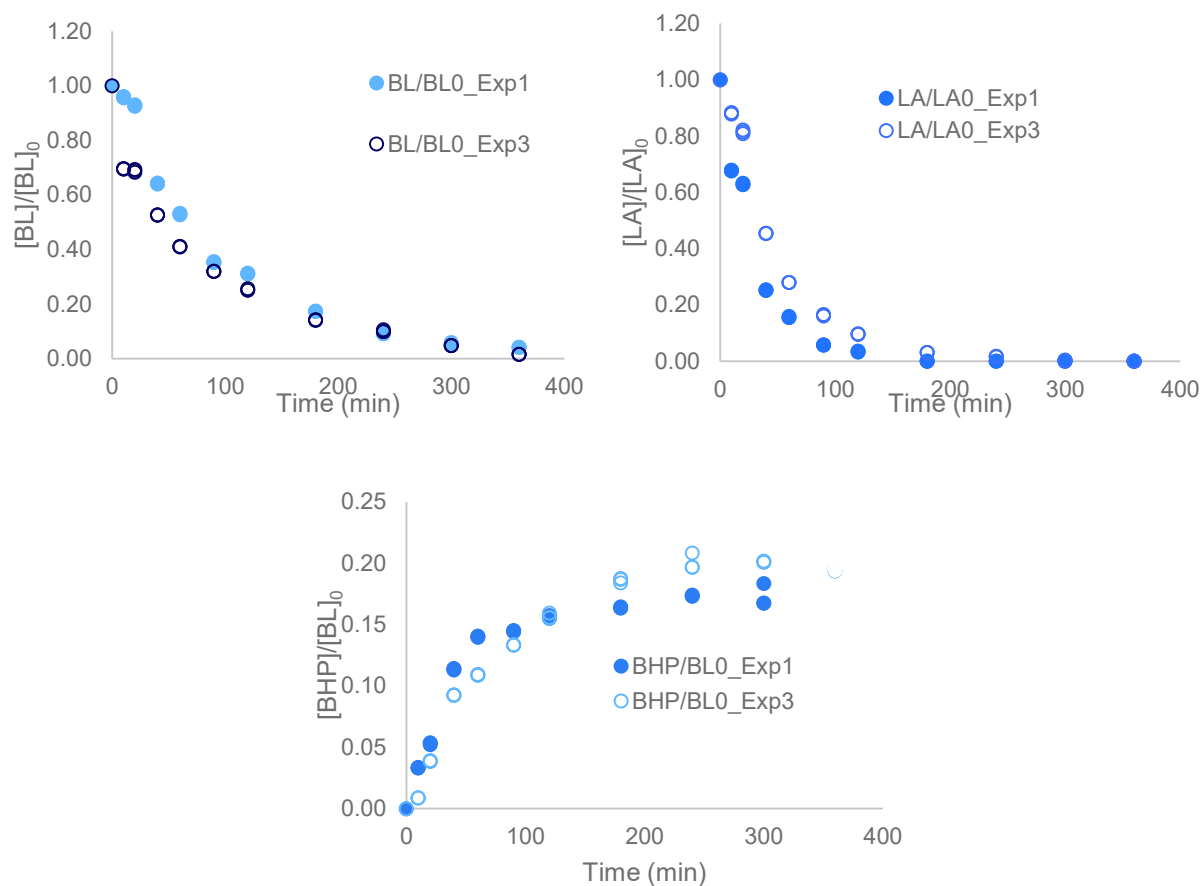


Fig. 3. Evolution of $[BL]/[BL]_0$, $[LA]/[LA]_0$ and $[BHP]/[BL]_0$ for experiments 1&3.

3.2 Effect of temperature

The effect of temperature on the kinetics was evaluated through Experiments 3 and 4 (Table 1) because only temperature change in these two experiments. Fig. 4 shows that hydrogenation steps are faster with increasing temperature. The temperature increase leads to a rise in the cyclization kinetics, and hence the decrease of the intermediate concentrations (BHP and HPA) is faster.

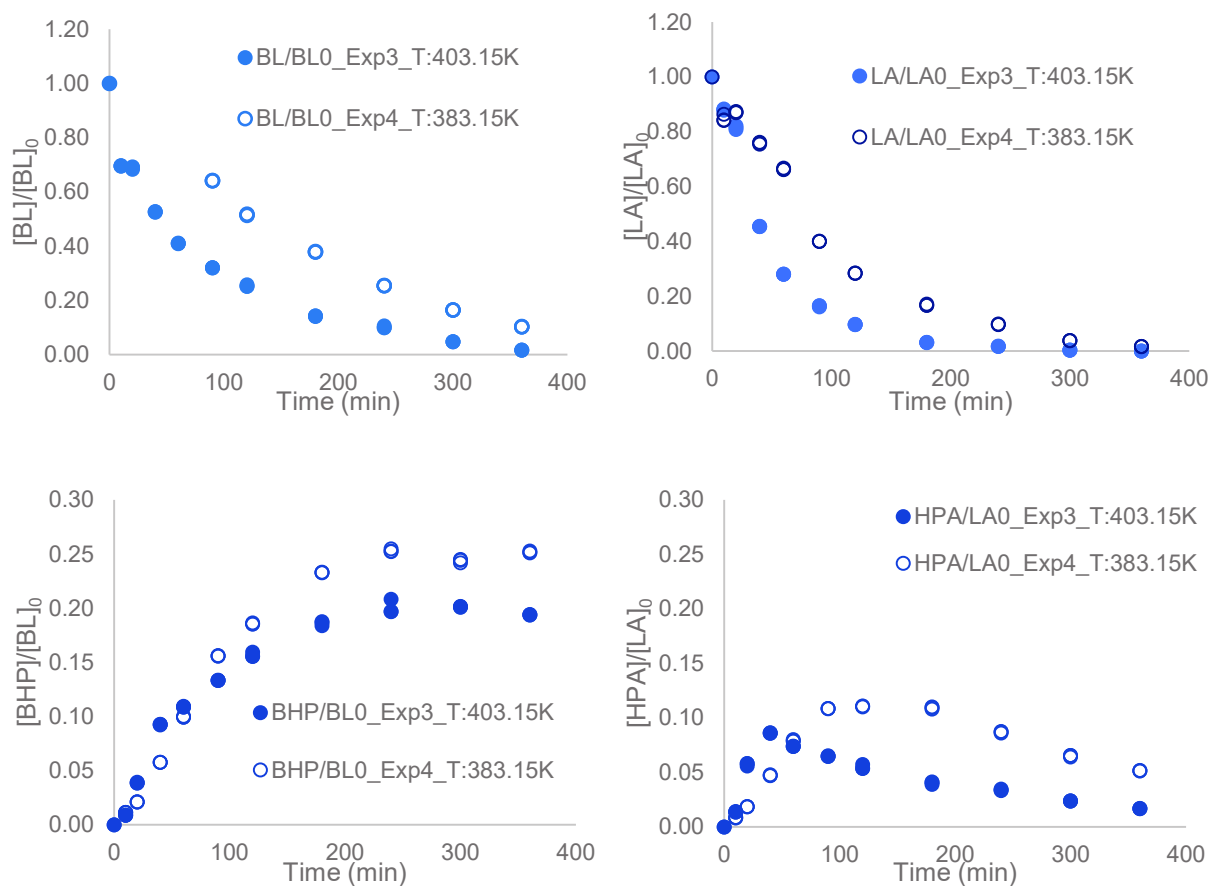


Fig. 4. Effect of temperature on hydrogenation and cyclization reactions for experiments 3&4.

3.3 Effect of hydrogen hydrostatic pressure

Fig. 5 shows the H_2 pressure effect by comparing Experiments 3 and 5 (Table 1). One can observe that the increase of hydrostatic pressure leads to accelerate the hydrogenation steps. The consequence of this acceleration is an increase of intermediate concentrations (BHP and HPA). The pressure has an indirect effect on the cyclization step.

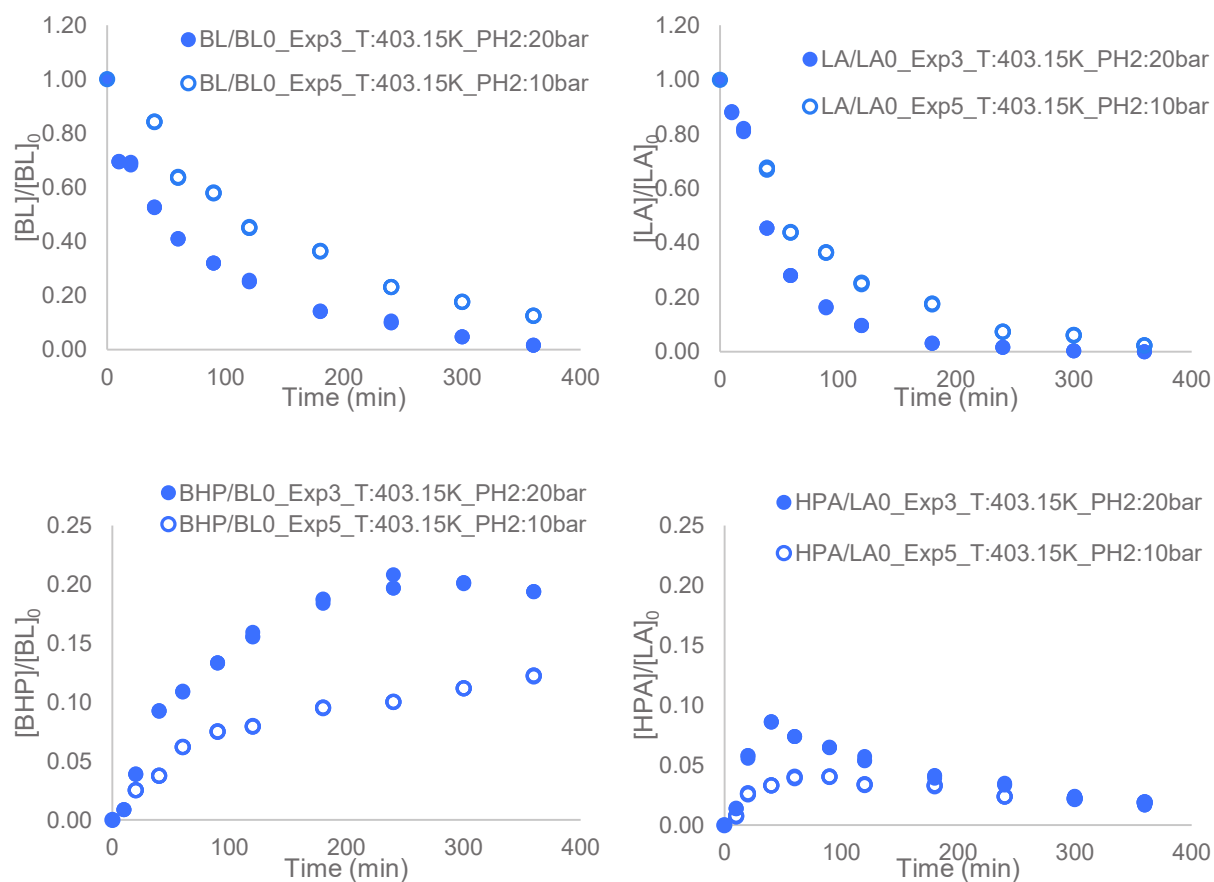


Fig. 5. Effect of pressure on the kinetics of hydrogenation and cyclization steps for experiments 3&5.

3.4 Effect of catalyst Ru/C

The effect of Ru/C loading on the kinetics were evaluated by comparing Experiments 7 and 8. The increase of Ru/C augments hydrogenation kinetics, leading to the rise of intermediate concentrations (Fig. 6).

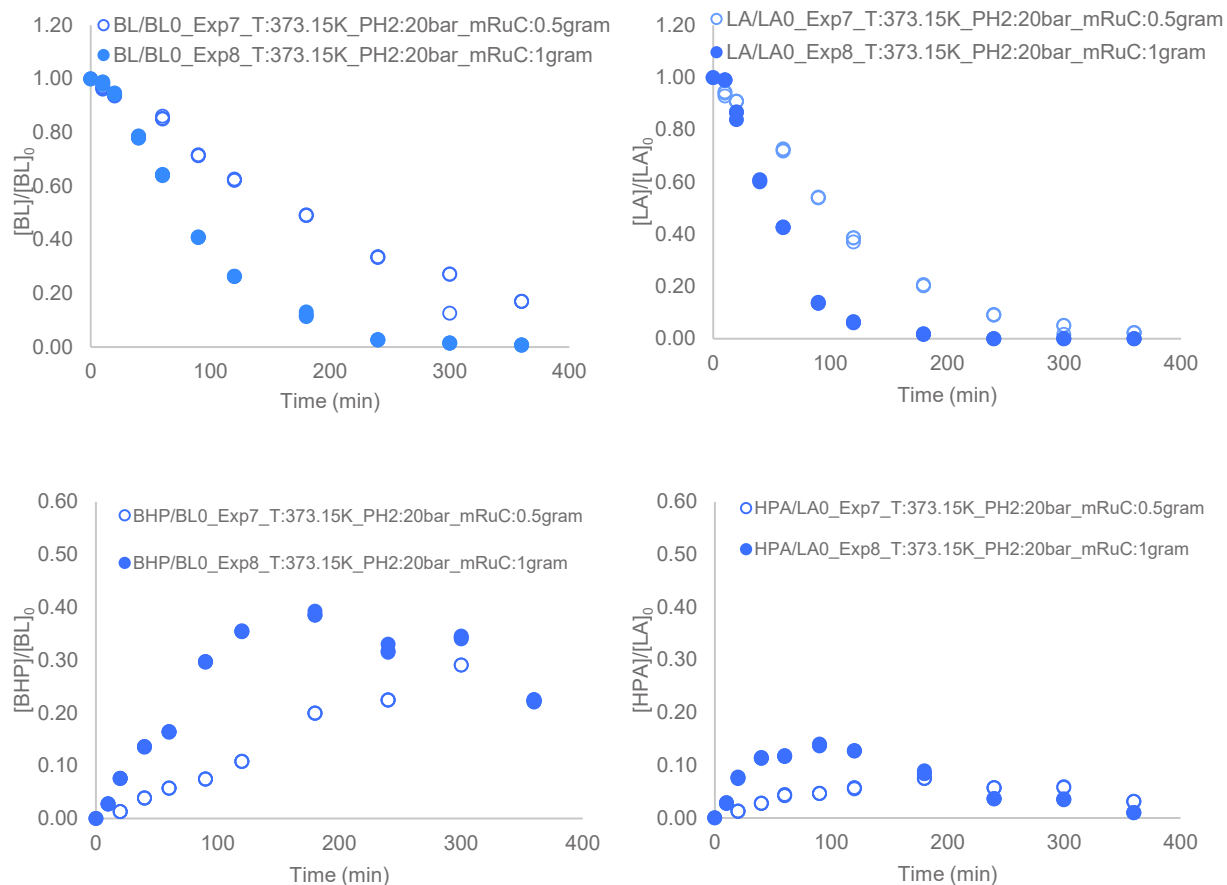


Fig. 6. Effect of pressure on Ru/C loadings on the kinetics for Experiments 7&8.

3.5 Effect of catalyst Amberlite IR-120

Experiments 9 and 10 (Table 1) give information on the catalytic effect of Amberlite IR-120. Fig. 7 shows that this catalyst does not have an impact on the first reaction step as expected. The HPA concentrations were too low to be detected, but one can notice that Amberlite IR-120 strongly affects the consumption kinetics of the intermediate (BHP).

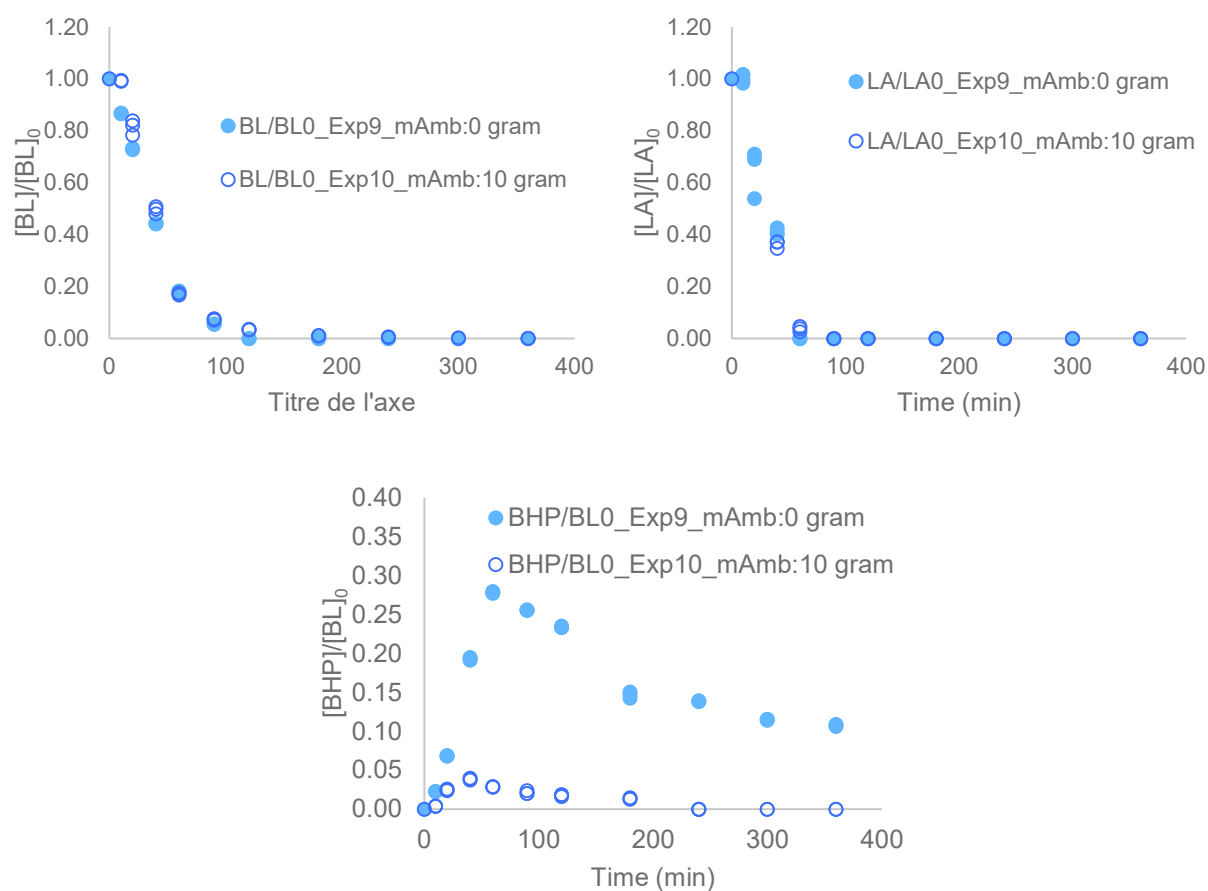


Fig. 7. Effect of Amberlite IR120 loadings on the kinetics of hydrogenation and cyclization for Experiments 9&10.

3.6 Effect of LA amount

Levulinic acid dissociates in the reaction mixture producing protons, which can catalyze the second reaction step. To verify this assumption, the results from Experiments 3 and 6 were compared. The kinetics of BL hydrogenation are similar (Fig. 8). The kinetics of LA hydrogenation is to be slightly faster when the concentration of LA is higher. One can notice that the increase of LA concentration accelerates the cyclization of HPA and BHP steps (Fig. 8).

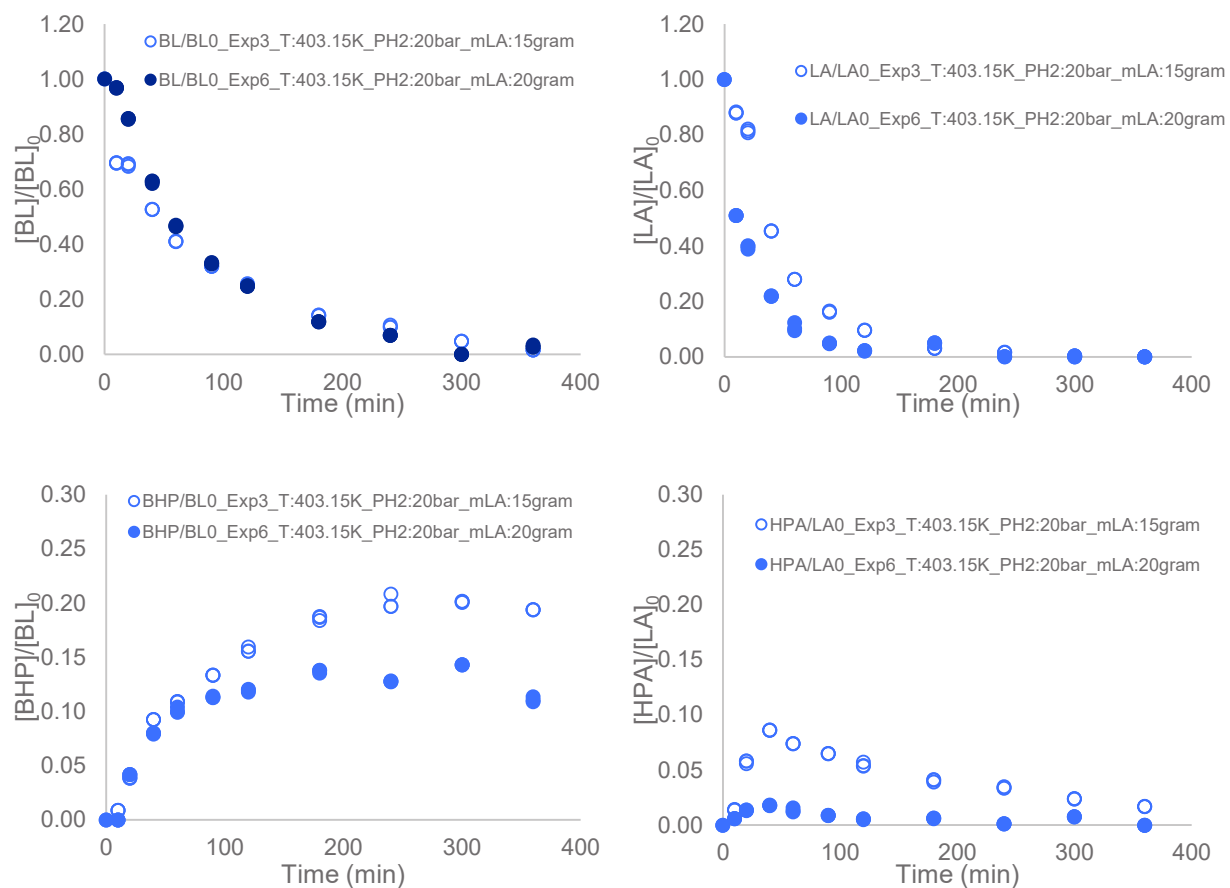


Fig. 8. Effect of LA concentration on the kinetics of hydrogenation and cyclization steps for Experiments 3&6.

4. DISCUSSION

4.1 Kinetics

From the Results section, several reaction mechanisms are possible for the hydrogenation and cyclization steps. The work of Capecci et al.[29] described that the surface reaction for the hydrogenation step of BL followss a non-competitive Langmuir-Hinshelwood model with no dissociation of molecular hydrogen. It means that there are two kinds of Ru sites and that the carbonyl group and hydrogen adsorb on each of them without competing. It was also found that the cyclization step can be catalyzed by Ru/C catalyst.

In this reaction system, BL and LA underwent similar reaction pathways. We neglected the esterification reaction of levulinic acid by butanol, because from the experimental data levulinic acid consumption is faster than the BHP cyclization.

Therefore, in the present work, seven kinetic models were evaluated for the hydrogenation step:

- competitive Langmuir-Hinshelwood with no dissociation of hydrogen (LH1),
- competitive Langmuir-Hinshelwood with dissociation of hydrogen (LH2),
- non-competitive Langmuir-Hinshelwood with no dissociation of hydrogen where LA and BL are in competitive adsorption on the same site but not hydrogen (NCLH1.1),
- non-competitive Langmuir-Hinshelwood with no dissociation of hydrogen where LA, BL and H₂ are adsorbed on different sites (NCLH1.2),
- Eley-Rideal with no adsorption of hydrogen on the active sites (ER),
- non-competitive Langmuir-Hinshelwood with dissociation of hydrogen where LA and BL are in competitive adsorption on the same site but not hydrogen (NCLH2.1),
- non-competitive Langmuir-Hinshelwood with dissociation of hydrogen where LA, BL and H₂ are adsorbed on different sites (NCLH2.2).

For the sake of clarity, the derivation of the seven kinetic models are described in Supplementary Material (S1). Table 2 displays the hydrogenation rate expression for each model.

Table 2. Rate expression for the hydrogenation steps.

MODELS	Rate expression for BL, $R_{BL,Hyd}$	Rate expression for LA, $R_{LA,Hyd}$
(LH1)	$\frac{k_1 * [H_2] * [BL] * \omega_{Cat.}}{\left(\begin{array}{l} K_{H_2} * [H_2] + K_{BL} * [BL] + K_{BHP} * [BHP] \\ + K_{LA} * [LA] + K_{HPA} * [HPA] + 1 \end{array} \right)^2}$	$\frac{k_1 * [H_2] * [LA] * \omega_{Cat.}}{\left(\begin{array}{l} K_{H_2} * [H_2] + K_{BL} * [BL] + K_{BHP} * [BHP] \\ + K_{LA} * [LA] + K_{HPA} * [HPA] + 1 \end{array} \right)^2}$
(LH2)	$\frac{k_1 * [H_2] * [BL] * \omega_{Cat.}}{\left(\begin{array}{l} K_H * \sqrt{[H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_i * [BL] * \sqrt{[H_2]} \\ + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{i2} * [LA] * \sqrt{[H_2]} + 1 \end{array} \right)^2}$	$\frac{k_1 * [H_2] * [LA] * \omega_{Cat.}}{\left(\begin{array}{l} \sqrt{K_H * [H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_i * [BL] * \sqrt{[H_2]} \\ + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{i2} * [LA] * \sqrt{[H_2]} + 1 \end{array} \right)^2}$
(ER1)	$\frac{k_1 * [H_2] * [BL] * \omega_{Cat.}}{\left(\begin{array}{l} K_{BL} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] \\ + HPA * [HPA] + 1 \end{array} \right)}$	$\frac{k_1 * [H_2] * [LA] * \omega_{Cat.}}{\left(\begin{array}{l} K_{BL} * [BL] + K_{BHP} * [BHP] \\ + K_{LA} * [LA] + HPA * [HPA] + 1 \end{array} \right)}$
(NCLH1.1)	$\frac{k_1 * [H_2]}{(1 + K_{H_2} * [H_2])} * \frac{[BL] * \omega_{Cat.}}{\left(\begin{array}{l} 1 + K_{BHP} * [BHP] + K_{BL} * [BL] \\ + K_{HPA} * [HPA] + K_{LA} * [LA] \end{array} \right)}$	$\frac{k_1 * [H_2]}{(1 + K_{H_2} * [H_2])} * \frac{[LA] * \omega_{Cat.}}{\left(\begin{array}{l} 1 + K_{BHP} * [BHP] + K_{BL} * [BL] \\ + K_{HPA} * [HPA] + K_{LA} * [LA] \end{array} \right)}$
(NCLH1.2)	$\frac{k_1 * [H_2]}{(1 + K_{H_2} * [H_2])} * \frac{[BL] * \omega_{Cat.}}{(1 + K_{BHP} * [BHP] + K_{BL} * [BL])}$	$\frac{k_1 * [H_2]}{(1 + K_{H_2} * [H_2])} * \frac{[LA] * \omega_{Cat.}}{(1 + K_{HPA} * [HPA] + K_{LA} * [LA])}$
(NCLH2.1)	$\frac{k_1 * [H_2]}{K_H * \sqrt{[H_2]} + 1} * \frac{[BL] * \omega_{Cat.}}{\left(\begin{array}{l} K_{BL} * [BL] + K_C * \sqrt{[H_2]} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] \\ + K_{C2} * \sqrt{[H_2]} * [LA] + K_{HPA} * [HPA] + 1 \end{array} \right)}$	$\frac{k_1 * [H_2]}{\sqrt{K_H * [H_2]} + 1} * \frac{[LA] * \omega_{Cat.}}{\left(\begin{array}{l} K_{BL} * [BL] + K_C * \sqrt{[H_2]} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] \\ + K_{C2} * \sqrt{[H_2]} * [LA] + K_{HPA} * [HPA] + 1 \end{array} \right)}$
(NCLH2.2)	$\frac{k_1 * [H_2]}{K_H * \sqrt{[H_2]} + 1} * \frac{[BL] * \omega_{Cat.}}{(K_{BL} * [BL] + K_C * \sqrt{[H_2]} * [BL] + K_{BHP} * [BHP] + 1)}$	$\frac{k_1 * [H_2]}{\sqrt{K_H * [H_2]} + 1} * \frac{[LA] * \omega_{Cat.}}{(K_{LA} * [LA] + K_{C2} * \sqrt{[H_2]} * [LA] + K_{HPA} * [HPA] + 1)}$

For the second cyclization reaction, four different types of reaction were considered: non-catalytic cyclization of BHP and HPA, the catalytic routes due to the presence of acid sites on Ru/C, the acidic groups on Amberlite IR-120 and the protons from the dissociation of levulinic acid. Due to space limitation, the derivation for these rate equations are explained in Supplementary Material (S3).

Rate equations for the cyclization steps are:

$$R_{BHP_noncat} = k_{BHP_noncat} \cdot [BHP] \quad (1)$$

$$R_{HPA_noncat} = k_{HPA_noncat} \cdot [HPA] \quad (2)$$

$$R_{BHP_RuC} = k_{BHP_RuC} \cdot [BHP] \cdot \frac{1}{K_{\blacksquare HPA} \cdot [HPA] + K_{\blacksquare BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.RuC} \quad (3)$$

$$R_{HPA_RuC} = k_{HPA_RuC} \cdot [HPA] \cdot \frac{1}{K_{\blacksquare HPA} \cdot [HPA] + K_{\blacksquare BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.RuC} \quad (4)$$

$$R_{BHP_SO3H} = k_{BHP_SO3H} \cdot [BHP] \cdot \frac{1}{K_{SO3H-HPA} \cdot [HPA] + K_{SO3H-BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.Amb} \quad (5)$$

$$R_{HPA_SO3H} = k_{HPA_SO3H} \cdot [HPA] \cdot \frac{1}{K_{SO3H-HPA} \cdot [HPA] + K_{SO3H-BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.RuC} \quad (6)$$

$$R_{BHP_diss} = k_{BHP_diss} \cdot [BHP] \cdot [LA] \quad (7)$$

$$R_{HPA_diss} = k_{HPA_diss} \cdot [HPA] \cdot [LA] \quad (8)$$

4.2 Material balances

Kinetic experiments were carried out under isobaric and isothermal conditions. Our previous study [46] found that external and internal mass transfer can be assumed to be negligible by using the operating conditions described in Section 2.

Material balances for different compounds in the liquid phase can be expressed as:

$$\frac{dC_{BL}}{dt} = -R_{BL_hyd} \quad (9)$$

$$\frac{d[H_2]_{liq}}{dt} = k_L \cdot a \cdot ([H_2]_{liq}^* - [H_2]_{liq}) - R_{BL_hyd} - R_{LA_hyd} \quad (10)$$

$$\frac{dC_{BHP}}{dt} = R_{BL_hyd} - R_{BHP_noncat} - R_{BHP_RuC} - R_{BHP_SO_3H} - R_{BHP_diss} \quad (11)$$

$$\frac{dC_{BuOH}}{dt} = R_{BHP_noncat} + R_{BHP_RuC} + R_{BHP_SO_3H} + R_{BHP_diss} \quad (12)$$

$$\frac{dC_{LA}}{dt} = -R_{LA_hyd} \quad (13)$$

$$\frac{dC_{HPA}}{dt} = R_{LA_hyd} - R_{HPA_noncat} - R_{HPA_RuC} - R_{HPA_SO_3H} - R_{HPA_diss} \quad (14)$$

$$\frac{dC_{Water}}{dt} = R_{HPA_noncat} + R_{HPA_RuC} + R_{HPA_SO_3H} + R_{HPA_diss} \quad (15)$$

$$\begin{aligned} \frac{dC_{GVL}}{dt} = & R_{BHP_noncat} + R_{BHP_RuC} + R_{BHP_SO_3H} + R_{BHP_diss} + R_{HPA_noncat} + R_{HPA_RuC} + R_{HPA_SO_3H} + \\ & R_{HPA_diss} \end{aligned} \quad (16)$$

The term $[H_2]_{liq}^*$ is the concentration of hydrogen at the gas-liquid interface. The values of this term

were determined through Henry's constant in GVL solvent $He(T) = \frac{[H_2]_{liq}^*}{P_{H_2,Reactor}}$ [38]. In this study

the values of $k_L \cdot a$, i.e., volumetric gas to liquid mass transfer coefficient for hydrogen was expressed as a function of density, viscosity and temperature [19].

4.3 Modeling

The commercial software Athena Visual Studio, using Bayesian statistics, was used for the simulation and estimation stages [62,63]. A Bayesian framework is considered to be more suitable for multi-response parameter estimation than the classical method of nonlinear least squares [55,64]. For the parameter estimation in Bayesian, one needs to determine the determinant criterion [65].

The concentrations of BL, BHP, LA and HPA were used as observables during the parameter estimation stage. The ODEs (9-16) were integrated by the DDAPLUS solver, included in Athena Visual Studio software. This solver is a modified Newton algorithm with a fixed leading coefficient backward difference formula to approximate the first-order derivative [66].

The subroutine package named GREGPLUS, was used to minimize the objective function $S(\partial)$, to calculate the credible intervals for each estimated parameter and to produce the normalized parameter covariance matrix.

To minimize the objective function $S(\partial)$, GREGPLUS uses successive quadratic programming starting from the user's initial guesses [62,64].

$$S(\partial) = (a + b + 1) \cdot \ln|v(\partial)| \quad (17)$$

where, a is the number of events in response, b is the number of responses and $|v(\partial)|$ is the determinant of the covariance matrix of the responses. Each element of this matrix is defined as

$$v_{ij}(\partial) = \sum_{u=1}^n [Y_{iu} - f_{iu}(\partial)] \cdot [Y_{ju} - f_{ju}(\partial)] \quad (18)$$

With Y_{iu} the experimental concentration and $f_{iu}(\partial)$ the estimated value for response i and event u ; Y_{ju} the experimental concentration and $f_{ju}(\partial)$ the estimated value for response j and event u .

The interval estimates for each estimated parameter are calculated from the final quadratic expansion of the objective function. The precision of the estimated parameters was evaluated by the marginal highest posterior density (HPD). The 95% marginal HPD was calculated by GREGPLUS package.

The modified Arrhenius equation is used to decrease the correlation between the pre-exponential factor and the activation energy.

$$k(T_R) = k(T_{ref}) \cdot \exp\left(-\frac{E_a}{R\chi} \left(\frac{1}{T_R} - \frac{1}{T_{ref}}\right)\right) \quad (19)$$

where T_{ref} is the reference temperature chosen in the considered experimental temperature range.

During the modeling stage of the seven models, the following constants tended to approach zero during the preliminary iteration:

- the adsorption constants for BHP and HPA on Ru/C for the hydrogenation steps,
- the activation energy for the cyclization of BHP and HPA over Amberlite IR-120,
- the adsorption constants for BHP and HPA over Amberlite IR-120 for the cyclization steps,
- the activation energy for the cyclization of BHP and HPA due to the acid sites from Ru/C,
- the adsorption constant for BHP and HPA over the acid sites from Ru/C,
- The equilibrium constant K_{C2} was also found to be low in model NCLH2.2.

For that reason, it was deemed reasonable to fix the values of these estimated parameters to zero.

Table 3 shows the regression parameters for each model:

-Sum of Squared Residuals (SSR) $\left(Y_{ju} - f_{ju}(\theta)\right)^2$

-Objective function defined by Equation 17 $S(\theta)$

-Number of estimated parameters

- Akaike information criterion (AIC) calculated by

$$AIC: \text{number of independent event} \cdot \ln \left(\frac{[y_{ju} - f_{ju}(\theta)]^2}{\text{number of independent event}} \right) + 2 \cdot \text{Number of estimated parameters} \quad (20).$$

The AIC value [60] allows including the number of estimated parameters in the model discrimination. Models with low AIC are the most reliable.

Table 3. Regression parameters for each model

	SSR	Objective function	Number of estimated parameters	AIC
LH1	11693900	24125	19	13224.9
LH2	11665200	24127	19	13221.3
NCLH1.1	11697000	24154	19	13225.3
NCLH1.2	10757800	23987	19	13102.4
ER1	11685300	24153	19	13223.8
NCLH2.1	11773600	24142	17	13230.9
NCLH2.2	10443600	23940	18	13056.9

Table 3 shows that NCLH1.2 and NCLH2.2 yield to low values of SSR, objective function and AIC. Due to space limitation, the estimated parameters, normalized covariance matrix and fit of these two models hereby reported, whereas the results obtained for the rest of the models analysed are included in the Supplementary Materials (S3). The results obtained in the present work is in agreement with with our previous work [29], where we found that NCLH1 was the most probable model for the hydrogenation of BL.

Table 4 shows the estimated values and the associated credible intervals (HPD%). The credible intervals for K_{H_2} and for $k_{HPA_noncat}(T_{Ref})$ are large due to the difficulty of estimating these values. The difficulty in estimating K_{H_2} is linked to the fact that the hydrogen pressure does not significantly affect the kinetics. The high uncertainty for $k_{HPA_noncat}(T_{Ref})$ is linked to the difficulty to tract HPA due to its high reactivity. The HPD intervals can be considered as low or medium for the other estimated parameters showing that the variation of the operating conditions was significant. Table 4 shows that the rate constant of LA hydrogenation is higher than the one of BL hydrogenation. The rate constants of HPA cyclization are higher than the ones of BHP cyclization (Table 4).

Table 4. Estimated Values at $T_{ref} = 392.72$ K and statistical Data for NCLH1.2

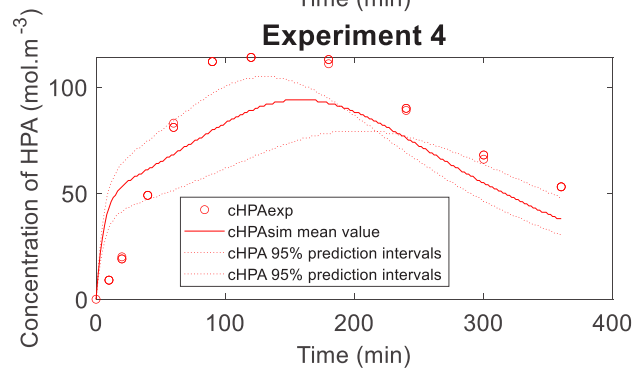
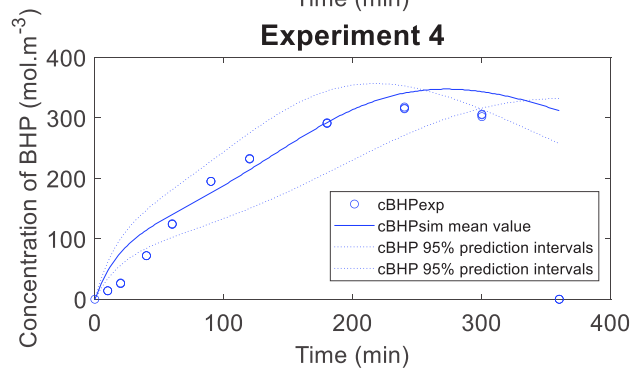
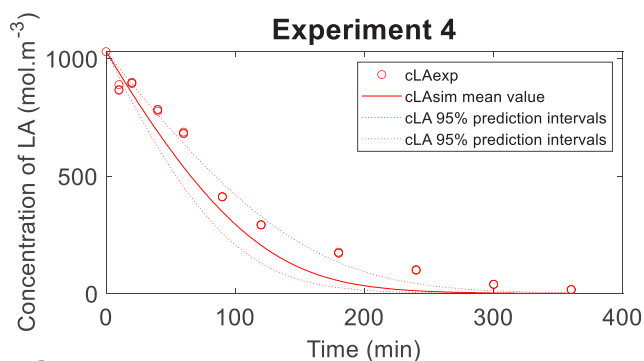
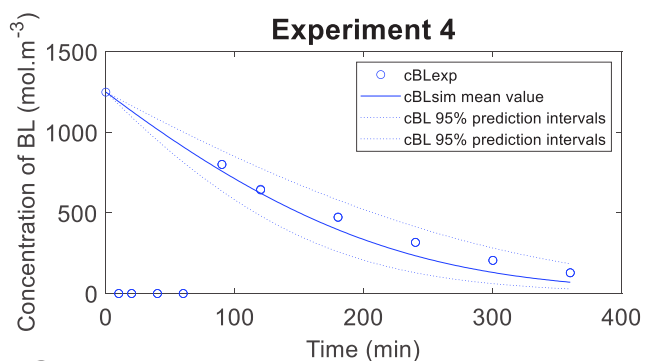
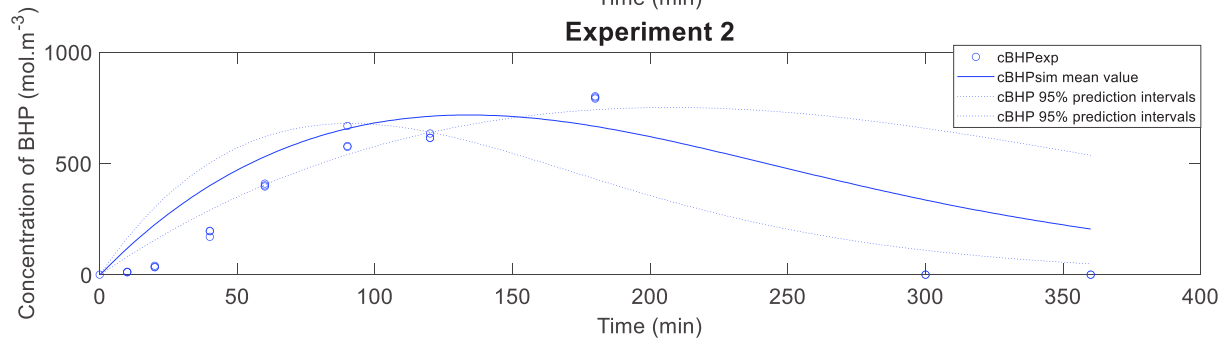
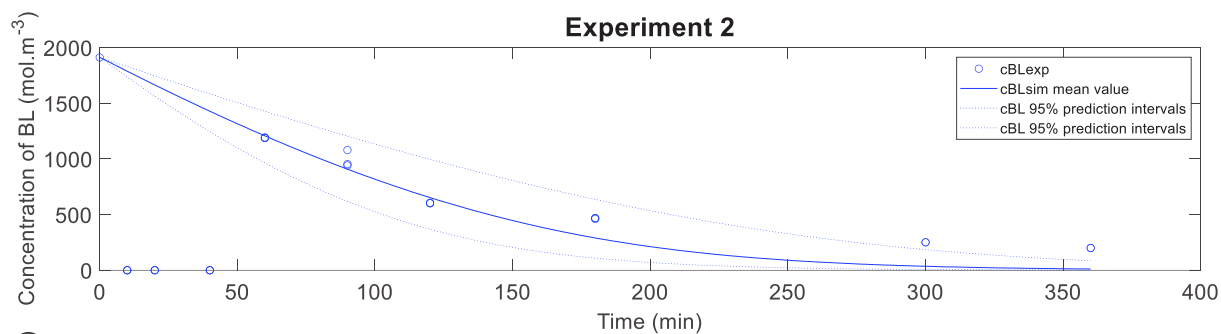
Parameters		Units	Estimates	HPD%
Par1	$k_{BL_hyd}(T_{Ref})$	$m^3 \cdot mol^{-1} \cdot s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	3.11E-06	14.57
Par2	Ea_{BL_hyd}	$J \cdot mol^{-1}$	3.62E+04	7.43
Par3	K_{H2}	$m^3 \cdot mol^{-1}$	7.36E-04	>100%
Par4	$K_{BL}^{\wedge}$	$m^3 \cdot mol^{-1}$	9.14E-04	29.02
Par5	$K_{BHP}^{\wedge}$	$m^3 \cdot mol^{-1}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ Amb^{-1}$	4.17E-05	49.21
Par7	$Ea'_{BHP_cat_Amb}$	$J \cdot mol^{-1}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	s^{-1}	5.78E-05	30.35
Par9	Ea_{BHP_noncat}	$J \cdot mol^{-1}$	8.67E+04	32.38
Par10	$k_{LA_hyd}(T_{Ref})$	$m^3 \cdot mol^{-1} \cdot s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	8.08E-06	12.03
Par11	Ea_{LA_hyd}	$J \cdot mol^{-1}$	4.65E+04	6.47
Par12	$K_{LA}^{\wedge}$	$m^3 \cdot mol^{-1}$	1.75E-03	15.80
Par13	$K_{HPA}^{\wedge}$	$m^3 \cdot mol^{-1}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ Amb^{-1}$	4.84E-04	49.37
Par15	$Ea'_{HPA_cat_Amb}$	$J \cdot mol^{-1}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	s^{-1}	1.12E-06	>100%
Par17	Ea_{HPA_noncat}	$J \cdot mol^{-1}$	4.22E+05	23.87
Par18	$K_{BHP-SO3H}$	$m^3 \cdot mol^{-1}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	$m^3 \cdot mol^{-1}$	Fixed to zero	-
Par20	$k_{BHP_RuC}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	2.43E-05	16.96
Par21	Ea_{BHP_RuC}	$J \cdot mol^{-1}$	Fixed to zero	-
Par22	$k_{HPA_RuC}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	5.80E-05	7.69
Par23	Ea_{HPA_RuC}	$J \cdot mol^{-1}$	Fixed to zero	-
Par24	$K_{\blacksquare BHP}$	$m^3 \cdot mol^{-1}$	Fixed to zero	-
Par25	$K_{\blacksquare HPA}$	$m^3 \cdot mol^{-1}$	Fixed to zero	-
Par26	$k_{BHP_diss}(T_{Ref})$	$m^3 \cdot mol^{-1} \cdot s^{-1}$	1.70E-06	16.83
Par27	Ea_{BHP_diss}	$J \cdot mol^{-1}$	1.06E+05	13.31
Par28	$k_{HPA_diss}(T_{Ref})$	$m^3 \cdot mol^{-1} \cdot s^{-1}$	4.73E-06	6.99
Par29	Ea_{HPA_diss}	$J \cdot mol^{-1}$	6.78E+04	7.96

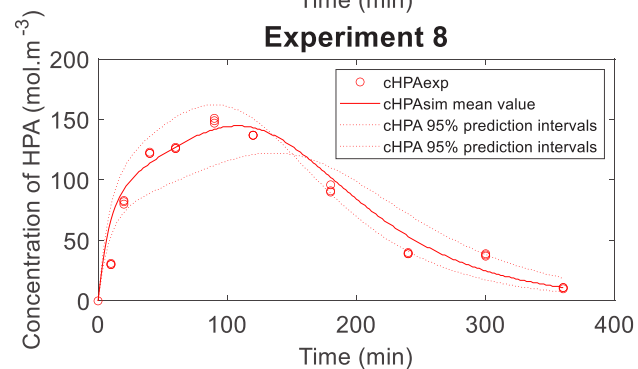
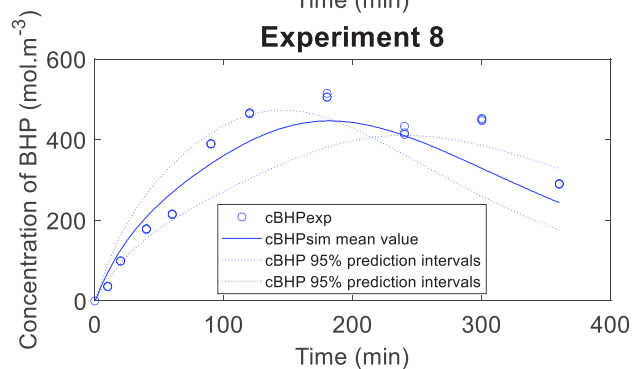
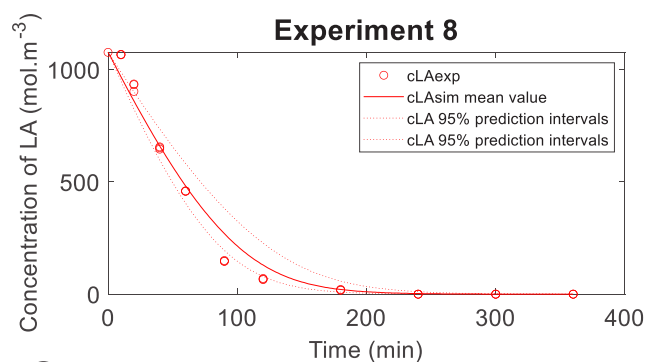
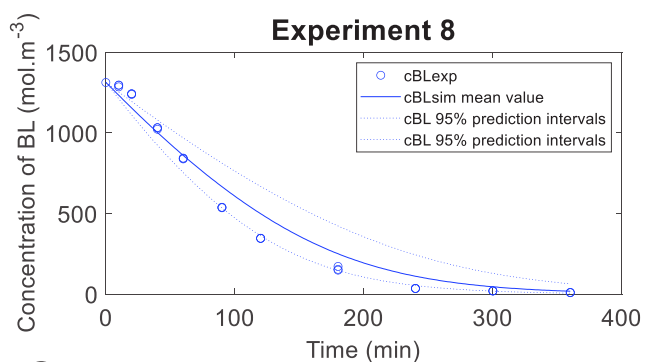
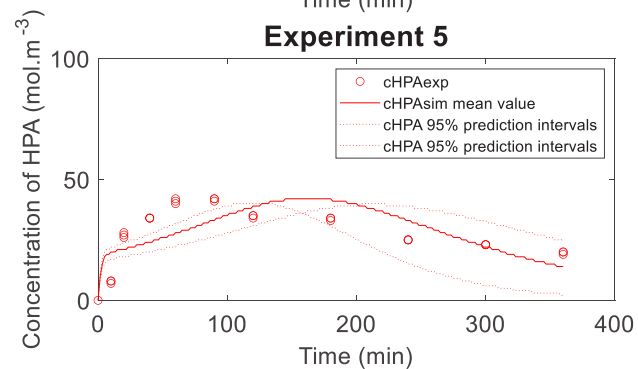
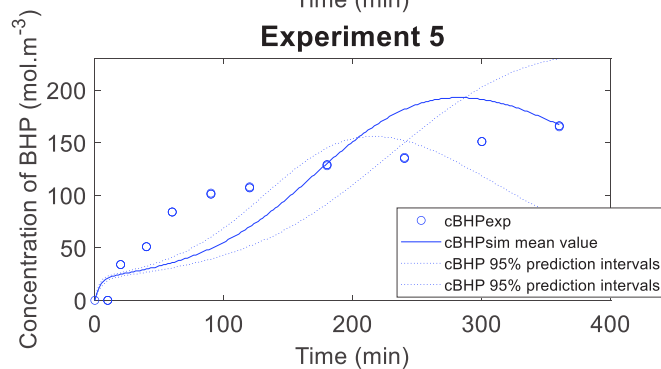
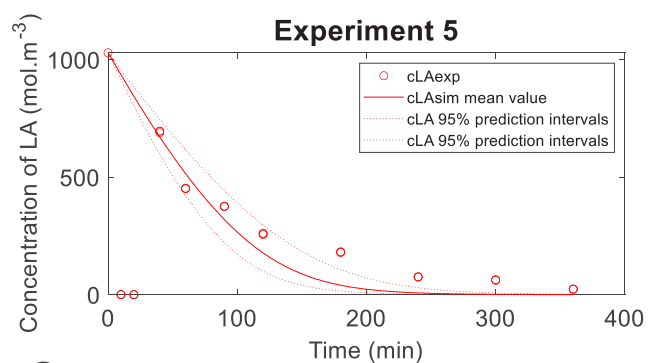
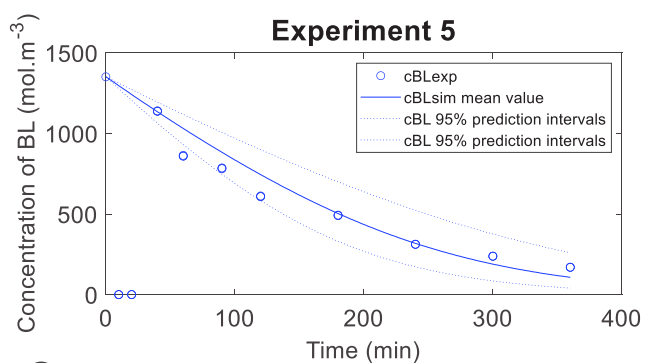
Table S3.11 shows the correlation between the estimated parameters. In general, the correlations are low. The significant correlation between $k_{BL_hyd}(T_{Ref})$ and $K_{BL}^{\wedge}$ are due to the difficulty to

estimate the adsorption constant, and the strong correlation $k_{BHP_noncat}(T_{Ref})$ and Ea_{BHP_noncat} is because the non-catalytic cyclization of BHP is relatively slow.

Fig. S4.1 displays the parity plots for BL, BHP, LA and HPA. NCLH2.1 model can predict BL and LA concentrations very well. The prediction of BHP and HPA concentrations is slightly lower due to the difficulty of tracking these intermediates.

Fig. 9 shows the fit of the model to the experimental concentrations with the 95% Prediction Intervals and the mean estimated values. From these graphs, one can notice that the model fits the experiments, and most of the experimental concentrations lie between the intervals. The intermediate concentrations for Experiment 10 are low due to the presence of Amberlite IR-120.





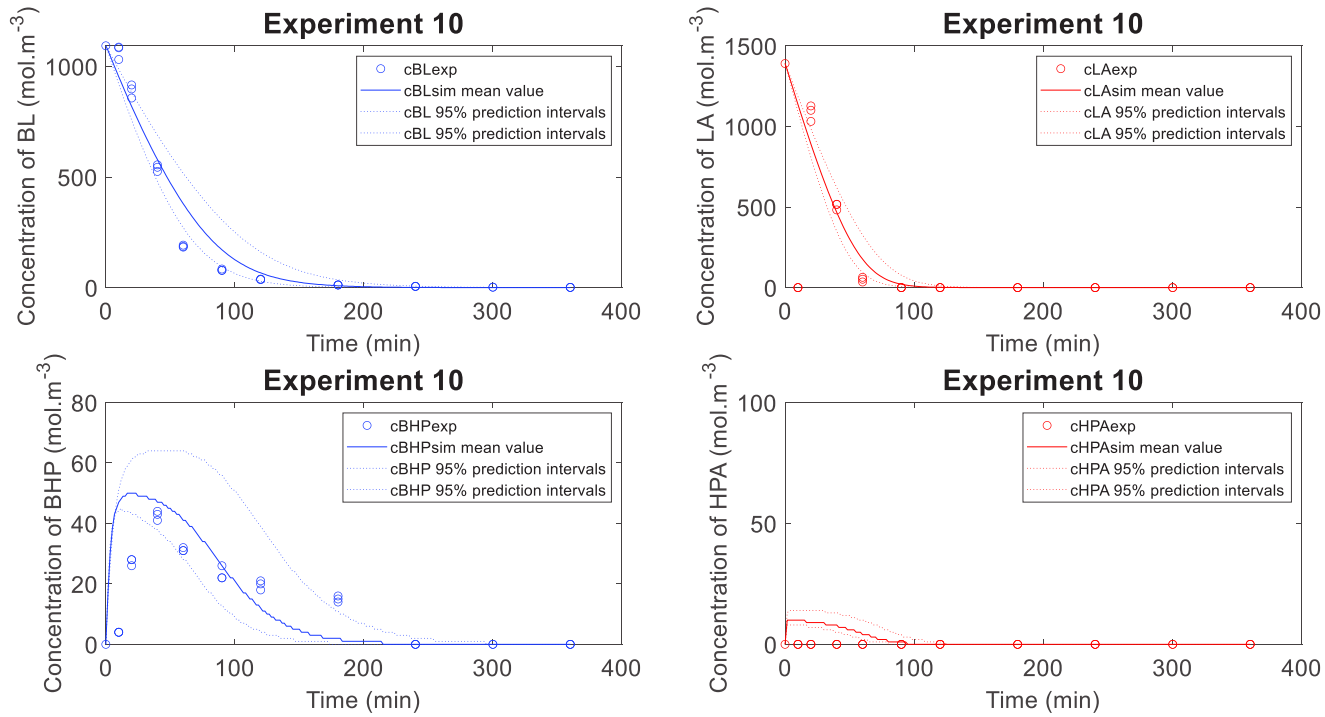


Fig. 9. Fit of Model NCLH1.2 to the experimental concentrations with prediction intervals.

NCLH22

Table 5 displays the estimated values and their credible intervals. During the modeling for this model, it was not possible to estimate K_{H2} and K_{BL} for that reason, their values were fixed to zero. The 95% HPD for $k_{HPA_{noncat}}(T_{Ref})$ is higher due to the high reactivity of HPA. The HPD intervals for the other parameters can be assumed to be medium or low. From Table 5, one can notice that the rate constant of LA hydrogenation is higher than the one of BL. The rate constants of HPA cyclization (from LA dissociation, Ru/C and Amerlite IR-120) are higher than the ones of BHP cyclization (Table 5).

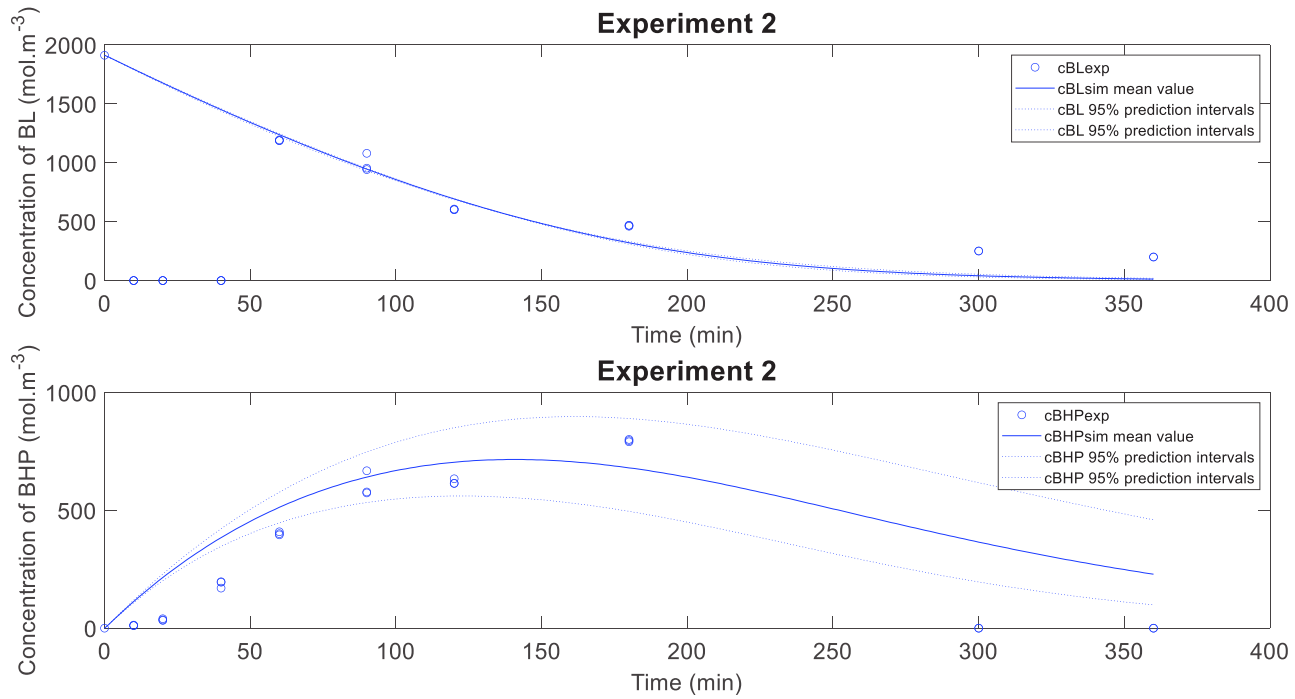
Table 5. Estimated Values at $T_{ref} = 392.72$ K and Statistical Data for NCLH2.2

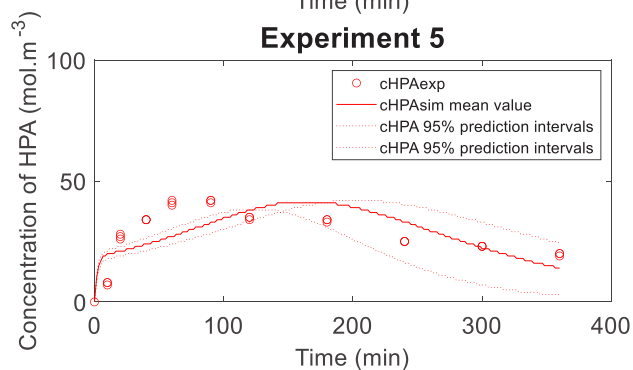
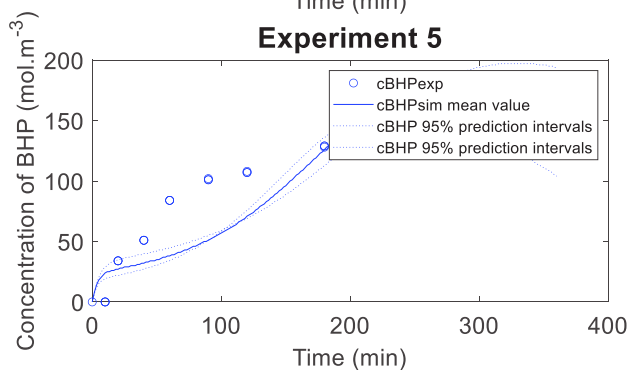
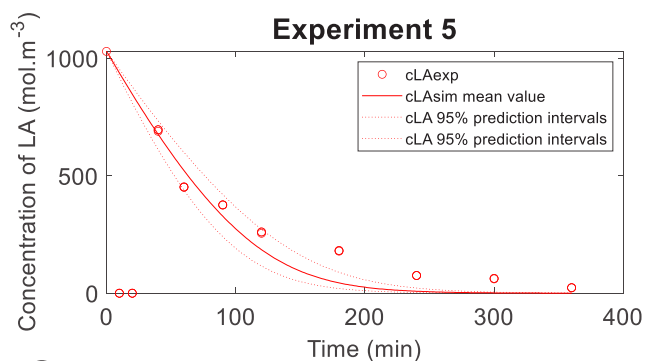
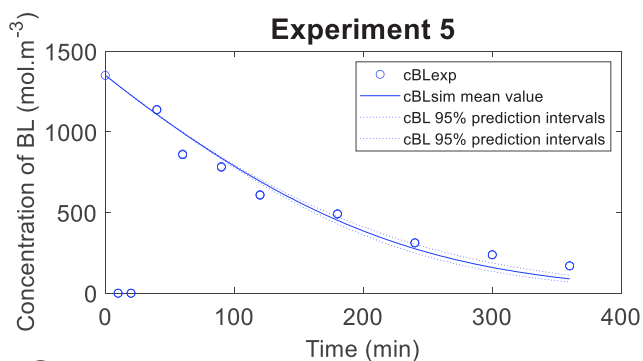
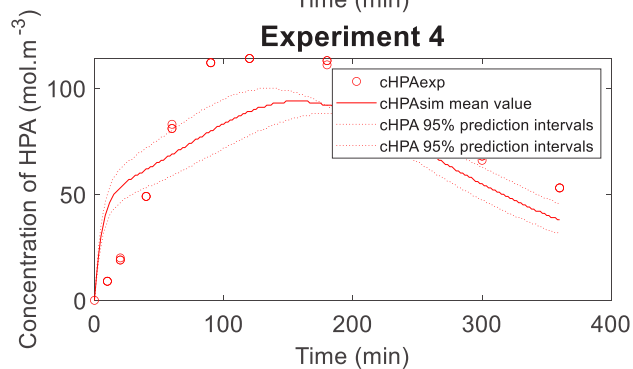
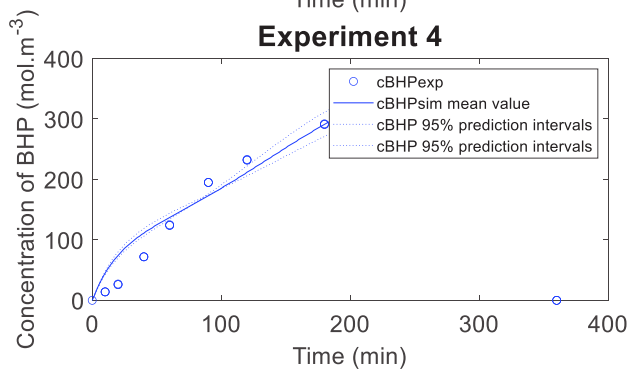
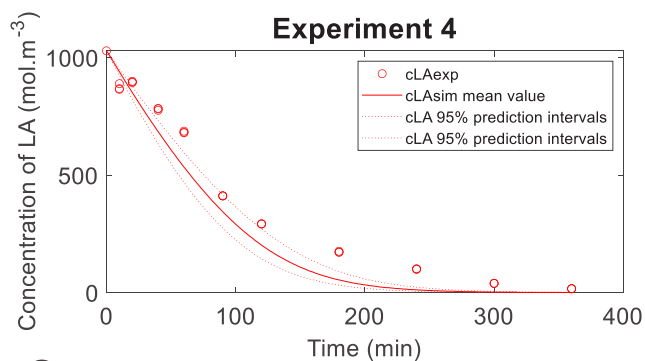
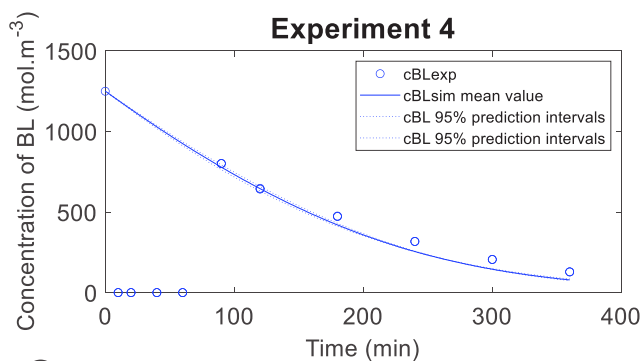
Parameters		Estimates	HPD%
Par1	$k_{BL_hyd}(T_{Ref})$	3.02E-06	11.42
Par2	Ea_{BL_hyd}	3.69E+04	7.17
Par3	$K_{H2.}$	Fixed to zero	-
Par4	$K_{BL}^{\wedge}$	Fixed to zero	-
Par5	$K_{BHP}^{\wedge}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	4.36E-05	48.29
Par7	$Ea'_{BHP_cat_Amb}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	5.93E-05	30.40
Par9	Ea_{BHP_noncat}	7.78E+04	35.92
Par10	$k_{LA_hyd}(T_{Ref})$	7.75E-06	9.17
Par11	Ea_{LA_hyd}	4.61E+04	6.51
Par12	$K_{LA}^{\wedge}$	1.69E-03	15.86
Par13	$K_{HPA}^{\wedge}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	4.79E-04	49.34
Par15	$Ea'_{HPA_cat_Amb}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	1.25E-06	>100%
Par17	Ea_{HPA_noncat}	4.15E+05	24.08
Par18	$K_{BHP-SO3H}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	Fixed to zero	-
Par20	$k_{BHP_RuC}(T_{Ref})$	2.41E-05	17.61
Par21	Ea_{BHP_RuC}	Fixed to zero	-
Par22	$k_{HPA_RuC}(T_{Ref})$	5.74E-05	7.61
Par23	Ea_{HPA_RuC}	Fixed to zero	-
Par24	$K_{\blacksquare BHP}$	Fixed to zero	-
Par25	$K_{\blacksquare HPA}$	Fixed to zero	-
Par26	Kc	1.59E-04	25.39
Par27	Kc2	Fixed to zero	-
Par28	$k_{BHP_diss}(T_{Ref})$	1.69E-06	16.19
Par29	Ea_{BHP_diss}	1.09E+05	12.83
Par30	$k_{HPA_diss}(T_{Ref})$	4.73E-06	6.71
Par31	Ea_{HPA_diss}	6.70E+04	8.01

Table S3.12 shows the correlation between the estimated parameters. One can notice a significant correlation between the following parameters: $k_{BL_hyd}(T_{Ref})$ and Kc ; $k_{BHP_noncat}(T_{Ref})$ and Ea_{BHP_noncat} ; $k_{LA_hyd}(T_{Ref})$ and K_{LA} and $k_{HPA_noncat}(T_{Ref})$ and Ea_{HPA_noncat} .

Fig. S4.2 shows the parity plot for BL, LA, BHP and HPA. Similar to Model NCLH1.2, the prediction for BL and LA is better than for the intermediates.

Fig. 10 shows the fit of the model to the experimental concentrations with the 95% prediction. The fitting is similar to the previous model.





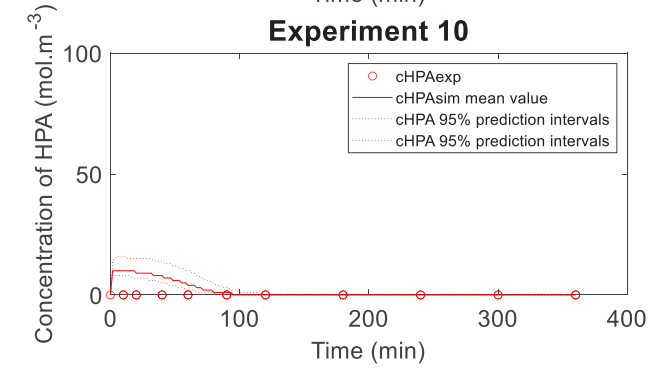
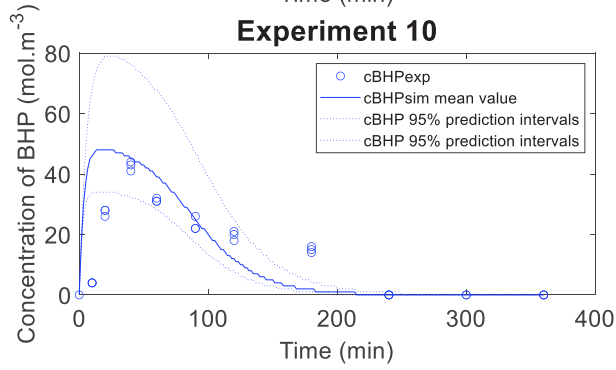
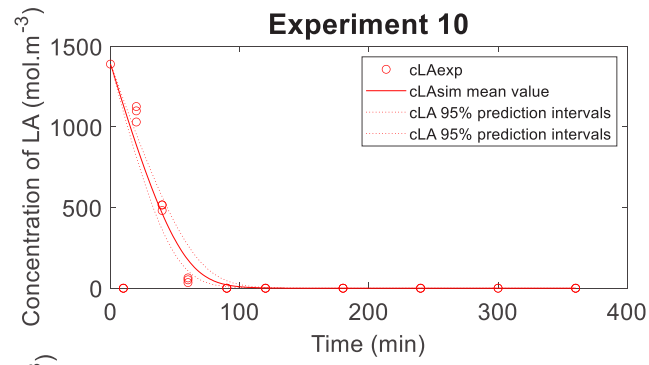
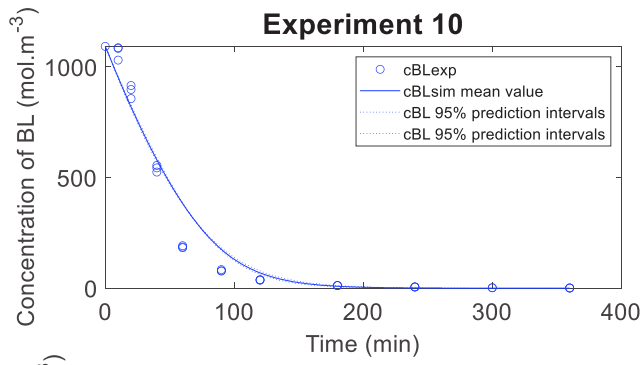
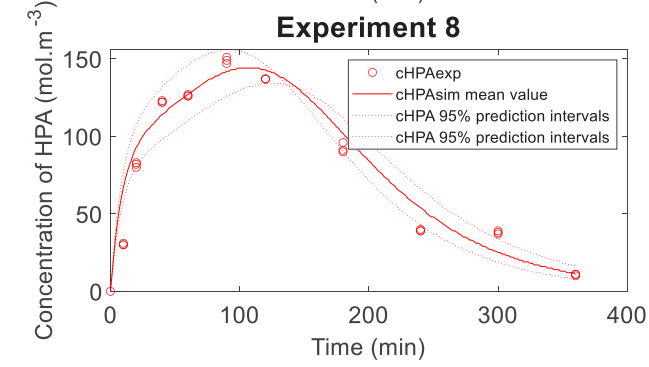
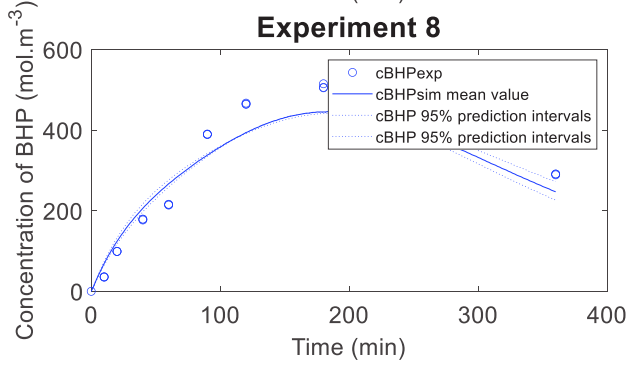
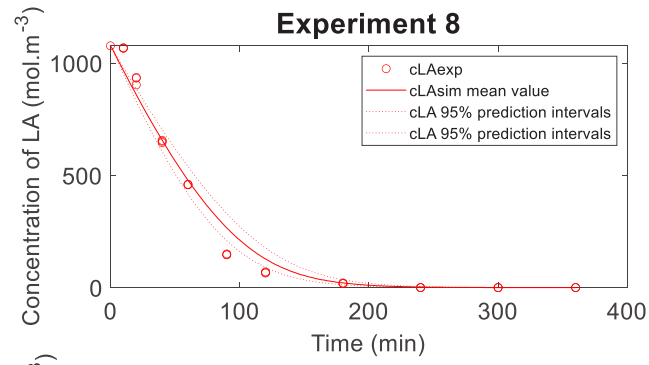
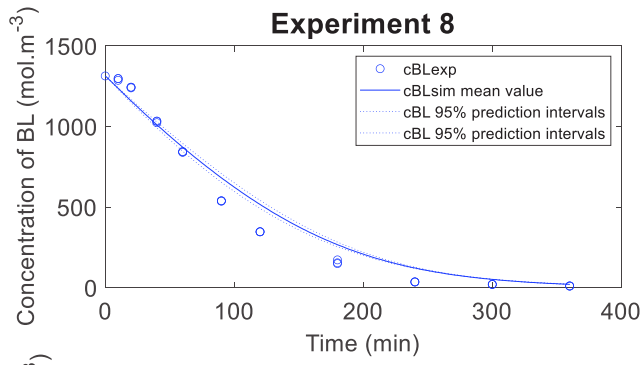


Fig. 10. Fit of Model NCLH2.2 with prediction intervals to the experimental concentrations.

4.4 Cross-validation: K-fold

The final stage of the assessment was the cross-validation one. Cross-validation was used to evaluate the predictability of the models and to determine the most probable one. The K-fold method was used [61]. The 14 experiments were divided randomly into 7 folds (Table 6). The regressions (a.k.a training) were made on 6 folds and validation (a.k.a testing) on the remaining fold as illustrated by Table 7.

Table 6. Distribution of the 14 experiments in the 7 folds.

FOLD	EXPERIMENTS
Fold 1	10 11
Fold 2	14 9
Fold 3	8 1
Fold 4	13 7
Fold 5	12 2
Fold 6	6 4
Fold 7	5 3

Table 7. Different Sets for regression and validation.

Set	Regression/Train	Validation/Test
Set 1	Folds 1-2-3-4-5-6	Fold 7
Set 2	Folds 7-1-2-3-4-5	Fold 6
Set 3	Folds 6-7-1-2-3-4	Fold 5
Set 4	Folds 5-6-7-1-2-3	Fold 4
Set 5	Folds 4-5-6-7-1-2	Fold 3
Set 6	Folds 3-4-5-6-7-1	Fold 2
Set 7	Folds 2-3-4-5-6-7	Fold 1

The kinetic constants are estimated from each regression, and these estimated constants are used for the validation. To evaluate the prediction capacity of a model the $CV_{(K)}$ number is calculated.

$$CV_{(K)} = \frac{1}{7} \cdot \sum_{K=1}^7 (Y_{i,experimental} - Y_{i,simulated})_K^2 \quad (21)$$

The lower the $CV_{(K)}$, the better the model is predictable. Table 8 shows that $CV_{(K)}$ number is lower for NCLH1.2 and NCLH2.2. The standard deviation of $CV_{(K)}$ was calculated for each model, and it was found that the standard deviation was lower for NCLH1.2 and NCLH2.2. This means that the validation step was similar for each set.

Table 8. $CV_{(K)}$ and standard deviation for each model

	$CV_{(K)}$	$SD(CV_{(K)})/\%$
LH1	3 586 050	99.88
LH2	3 656 893	93.19
NCLH1.1	3 635 269	94.19
NCLH1.2	2 168 016	28.66
ER1	3 595 498	95.09
NCLH2.1	3 690 655	92.00
NCLH2.2	1 978 294	27.47

In Supplementary Materials (S5), the estimated values for each regression set were displayed for NCLH1.2 and NCLH2.2. One can notice that the estimation was similar for each set compared to the estimation with the whole experimental data, i.e., the ones displayed in Tables 4 and 6.

To validate the models by cross-validation, Fig. 11 displays the coefficient of determination for the training step, test set and all data (i.e., Figs S4.1 and S4.2). One can notice that these values are similar, meaning that both models are validated.

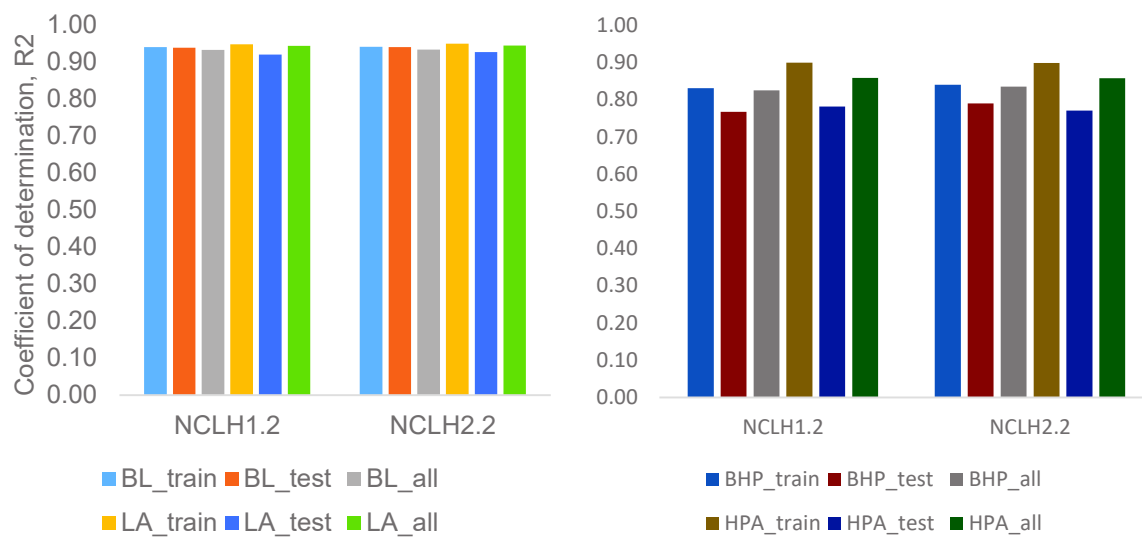


Fig. 11. Coefficient of determinations for training, test and all for the different models.

5. Conclusions

One of the most common routes for the production of GVL is the hydrogenation of levulinic acid or alkyl levulinates. When alkyl levulinates produced from the alcoholysis of fructose, there is the presence of levulinic acid is also produced in the system. This paper proposed to investigate the kinetics of the hydrogenation of butyl levulinate and levulinic acid in GVL solvent over Ru/C. To increase the kinetics of the cyclization step, Amberlite IR-120 catalyst was added in the reaction mixture.

In the first analysis, it was found that the presence of LA can increase the kinetics of cyclization and Amberlite IR-120 has a significant catalytic effect on this reaction.

In the second step, several kinetic models, via Bayesian inference, were evaluated for the hydrogenation steps throughout K-fold approach. Seven kinetic models were evaluated: competitive Langmuir-Hinshelwood with no dissociation of hydrogen (LH1), competitive Langmuir-Hinshelwood with dissociation of hydrogen (LH2), non-competitive Langmuir-Hinshelwood with no dissociation of hydrogen where LA and BL are in competitive adsorption on the same site but not hydrogen (NCLH1.1), non-competitive Langmuir-Hinshelwood with no dissociation of hydrogen where LA, BL and H₂ are adsorbed on different sites (NCLH1.2), Eley-Rideal with no adsorption of hydrogen on the active sites (ER), non-competitive Langmuir-Hinshelwood with dissociation of hydrogen where LA and BL are in competitive adsorption on the same site but not hydrogen (NCLH2.1), and non-competitive Langmuir-Hinshelwood with dissociation of hydrogen where LA, BL and H₂ are adsorbed on different sites (NCLH2.2). It was found that NCLH1.2 and NCLH2.2 were the most probable models, and the prediction capacity of these models was higher compared to the other.

This paper showed the benefit to using such reaction mixture system for the production of GVL. From an industrial viewpoint, the hydrogenation of levulinic acid and alkyl levulinate can be consecutive to the alcoholysis process. Quantum mechanics calculation could give more information concerning the adsorption mechanism.

Notation

E_{a_i}	activation energy of reaction i [J.mol^{-1}]
$f(\partial)$	estimated concentration
H_e	Henry's coefficient [$\text{mol.m}^{-3}.\text{bar}^{-1}$]
k_i	rate constant of reaction i
K_i	adsorption rate of specie i
$k_{L,a}$	volumetric mass transfer coefficient [s^{-1}]
P	pressure [bar]
R_i	reaction rate i [$\text{mol.m}^{-3}.\text{s}^{-1}$]
R	gas constant [$\text{J.K}^{-1}.\text{mol}^{-1}$]
$S(\partial)$	objective function
T	temperature [K]
$ \nu(\partial) $	determinant of the covariance matrix of responses
Y_i	experimental concentration of specie i

Greek letters

Θ	catalyst active sites
θ_i	active sites occupied by specie i
$\omega_{cat.}$	catalyst loading [kg.m^{-3}]

Subscripts and superscripts

Ref	reference
*	interfacial value

Abbreviations

AIC	Akaike information criterion
AL	Alkyl levulinate
Amb	Amberlite IR-120
BL	Butyl levulinate
BHP	Butyl 4-hydroxypentanoate
ER	Eley-Rideal kinetic model without hydrogen adsorption
GC	Gas chromatography
GVL	γ -valerolactone
HPA	4-hydroxypentanoic acid
HDP	Highest Posterior Density
LA	Levulinic acid
LCB	Lignocellulosic biomass
LH1	Langmuir-Hinshelwood kinetic model without hydrogen dissociation
LH2	Langmuir-Hinshelwood kinetic model with hydrogen dissociation
NCLH1.1	Non-competitive Langmuir-Hinshelwood kinetic model without hydrogen dissociation, where BL and LA are in competitive adsorption on the same site but not hydrogen
NCLH1.2	Non-competitive Langmuir-Hinshelwood kinetic model without hydrogen dissociation, where BL, LA and hydrogen are adsorbed on different sites

NCLH2.1	Non-competitive Langmuir-Hinshelwood kinetic model with hydrogen dissociation, where BL and LA are in competitive adsorption on the same site but not hydrogen
NCLH2.2	Non-competitive Langmuir-Hinshelwood kinetic model with hydrogen dissociation, where BL, LA and hydrogen are adsorbed on different sites
ODEs	Ordinary differential equation system
ROH	Co-product of the cyclyzation step(water or butanol)
Ru/C	Ruthenium on activated carbon
SSR	Sum of squared residuals

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Supporting information

Synergy effect of dual catalysts for the synthesis of γ -valerolactone from *n*-butyl levulinate hydrogenation over Ru/C and Amberlite IR-120.

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S1. Derivation of hydrogenation rate expression

Langmuir Hinshelwood with molecular adsorption of H₂ (LH1)



The adsorption and desorption steps are considered to be very fast reaching quasi-equilibria. Thus,

$$K_{BL*} = \frac{\theta_{*BL}}{[BL] \cdot \theta_*} \quad (\text{S9})$$

$$K_{LA*} = \frac{\theta_{*LA}}{[LA] \cdot \theta_*} \quad (\text{S10})$$

$$K_{H_2*} = \frac{\theta_{*H_2}}{[H_2] \cdot \theta_*} \quad (\text{S11})$$

$$K_{BHP*} = \frac{\theta_{*BHP}}{[BHP] \cdot \theta_*} \quad (\text{S12})$$

$$K_{HPA*} = \frac{\theta_{*HPA}}{[HPA] \cdot \theta_*} \quad (\text{S13})$$

K_{BL*} , K_{LA*} , K_{H_2*} , K_{BL} , K_{BHP*} and K_{HPA*} are adsorption constants.

Reactions (S4) and (S5) are considered to be the rate-determining steps for the hydrogenation steps.

All the reaction determining steps will be assumed to be irreversible ones.

$$R_{BL_hyd} = R_4 = k_4 * \theta_{H_2} * \theta_{BL} * \omega_{Cat.} = k_4 * K_{H_2*} * [H_2] * \theta_* * K_{BL*} * [BL] * \theta_* * \omega_{Cat.} \quad (\text{S14})$$

$$R_{LA_hyd} = R_5 = k_5 * \theta_{H_2} * \theta_{LA} * \omega_{Cat.} = k_5 * K_{H_2} * [H_2] * \theta_* * K_{LA} * [LA] * \theta_* * \omega_{Cat.} \quad (S15)$$

The mass balance for active sites leads to

$$1 = \theta_{H_2} + \theta_{BL} + \theta_{BHP} + \theta_{LA} + \theta_{HPA} + \theta_* \quad (S16)$$

By introducing eqs (S9)-(S13) in eq (S16), one gets

$$1 = K_{H_2} * [H_2] * \theta_* + K_{BL} * [BL] * \theta_* + K_{BHP} * [BHP] * \theta_* + K_{LA} * [LA] * \theta_* + K_{HPA} * [HPA] * \theta_* + \theta_* \quad (S17)$$

Resulting in

$$\theta_* = \frac{1}{K_{H_2} * [H_2] + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] + K_{HPA} * [HPA] + 1} \quad (S18)$$

Hence, the rate of hydrogenation can be expressed as

$$R_{BL_hyd} = k_{BL_hyd} * \frac{[H_2] * [BL]}{(1 + K_{H_2} * [H_2] + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] + K_{HPA} * [HPA])^2} * \omega_{Cat.} \quad (S19)$$

$$\text{with, } k_{BL_hyd} = k_4 * K_{H_2} * K_{BL}$$

$$R_{LA_hyd} = k_{LA_hyd} * \frac{[H_2] * [LA]}{(1 + K_{H_2} * [H_2] + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] + K_{HPA} * [HPA])^2} * \omega_{Cat.} \quad (S20)$$

$$\text{with, } k_{LA_hyd} = k_5 * K_{H_2} * K_{LA}$$

Langmuir Hinshelwood with hydrogen dissociation (LH2)





$$K_{BL} = \frac{\theta_{BL}}{[BL] \cdot \theta_*} \quad (S32)$$

$$K_H = \frac{\theta_H^2}{[H_2] \cdot \theta_*^2} \quad (S33)$$

$$K_{BHP} = \frac{\theta_{BHP}}{[BHP] \cdot \theta_*} \quad (S34)$$

If reactions (S26) and (S27) are the rate-determining steps and (S24)-(S25) are very rapid, thus

$$R_{BL_hyd} = k_{26} * \theta_H * \theta_{BLH} * \omega_{cat.} = k_{26} * \sqrt{K_H * [H_2]} * \theta_* * \theta_{BLH} * \omega_{cat.} \quad (S35)$$

$$K_{24} = \frac{\theta_{BLH} \cdot \theta_*}{\theta_{BL} \cdot \theta_H} \text{ equivalent to } \theta_{BLH} = K_{24} * \frac{\theta_{BL} \cdot \theta_H}{\theta_*} = K_{24} * K_{BL} * [BL] \cdot \theta_* * \sqrt{K_H * [H_2]} \quad (S36)$$

$$R_{LA_hyd} = k_{27} * \theta_H * \theta_{LAH} * \omega_{cat.} = k_{27} * \sqrt{K_H * [H_2]} * \theta_* * \theta_{LAH} * \omega_{cat.} \quad (S37)$$

$$K_{25} = \frac{\theta_{LAH} \cdot \theta_*}{\theta_{LA} \cdot \theta_H} \text{ equivalent to } \theta_{LAH} = K_{25} * \frac{\theta_{LA} \cdot \theta_H}{\theta_*} = K_{25} * K_{LA} * [LA] \cdot \theta_* * \sqrt{K_H * [H_2]} \quad (S38)$$

Material balance on active site leads to

$$1 = \theta_H + \theta_{BL} + \theta_{BHP} + \theta_{BLH} + \theta_{LA} + \theta_{HPA} + \theta_{LAH} + \theta_* \quad (S39)$$

By introducing eqs (S32)-(S34) and (S36)-(S38) in eq (S39), we get

$$1 = \sqrt{K_H * [H_2]} * \theta_* + K_{BL} * [BL] * \theta_* + K_{BHP} * [BHP] * \theta_* + K_{24} * K_{BL} * [BL] * \sqrt{K_H * [H_2]} * \theta_* + K_{LA} * [LA] * \theta_* + K_{HPA} * [HPA] * \theta_* + K_{25} * K_{LA} * [LA] * \sqrt{K_H * [H_2]} * \theta_* + \theta_* \text{ equivalent to}$$

$$\theta_* = \frac{1}{\sqrt{K_H * [H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{24} * K_{BL} * [BL] * \sqrt{K_H * [H_2]} + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{25} * K_{LA} * [LA] * \sqrt{K_H * [H_2]} + 1} \quad (S40)$$

Hence, the rates of hydrogenation can be expressed as

$$R_{BL_hyd} = k_{26} \cdot K_H \cdot [H_2] \cdot K_{24} \cdot K_{BL} \cdot [BL].$$

$$\omega_{Cat.} = \frac{1}{\left(1 + \sqrt{K_H * [H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{24} * K_{BL} * [BL] * \sqrt{K_H * [H_2]} + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{25} * K_{LA} * [LA] * \sqrt{K_H * [H_2]}\right)^2} * \quad (S41)$$

$$R_{LA_hyd} = k_{27} \cdot K_H \cdot [H_2] \cdot K_{25} \cdot K_{LA} \cdot [LA].$$

$$\omega_{Cat.} = \frac{1}{\left(1 + \sqrt{K_H * [H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{24} * K_{BL} * [BL] * \sqrt{K_H * [H_2]} + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{25} * K_{LA} * [LA] * \sqrt{K_H * [H_2]}\right)^2} * \quad (S42)$$

The following notations are introduced: $k_{BL_hyd} = k_{26} \cdot K_H \cdot K_{24} \cdot K_{BL}$, $k_{LA_hyd} = k_{27} \cdot K_H \cdot K_{25} \cdot K_{LA}$,

$K_H = \sqrt{K_H}$, $K_i = K_{24} * K_{BL} * \sqrt{K_H}$ and $K_{i2} = K_{25} * K_{LA} * \sqrt{K_H}$. Hence, Equations (S41) and (S42)

become

$$R_{BL_hyd} = k_{BL_hyd} \cdot [H_2] \cdot [BL].$$

$$\omega_{Cat.} = \frac{1}{\left(1 + K_H * \sqrt{[H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_i * [BL] * \sqrt{[H_2]} + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{i2} * [LA] * \sqrt{[H_2]}\right)^2} * \quad (S43)$$

$$R_{LA_hyd} = k_{LA_hyd} \cdot [H_2] \cdot [LA].$$

$$\omega_{Cat.} = \frac{1}{\left(1 + K_H * \sqrt{[H_2]} + K_{BL} * [BL] + K_{BHP} * [BHP] + K_i * [BL] * \sqrt{[H_2]} + K_{LA} * [LA] + K_{HPA} * [HPA] + K_{i2} * [LA] * \sqrt{[H_2]}\right)^2} * \quad (S44)$$

Eley-Rideal with no adsorption of hydrogen (ER1)



$$K_{BL} = \frac{\theta_{BL}}{[BL] \cdot \theta_*} \quad (\text{S53})$$

$$K_{BHP} = \frac{\theta_{BHP}}{[BHP] \cdot \theta_*} \quad (\text{S54})$$

$$K_{LA} = \frac{\theta_{LA}}{[LA] \cdot \theta_*} \quad (\text{S55})$$

$$K_{HPA} = \frac{\theta_{HPA}}{[HPA] \cdot \theta_*} \quad (\text{S56})$$

K_{BL} , K_{BHP} , K_{LA} and K_{HPA} , are adsorption constants.

Reactions (S47)-(S48) are rate-determining steps

$$R_{BL_hyd} = R_{47} = k_{47} * [H_2] * \theta_{BL} * \omega_{Cat.} = k_{47} * [H_2] * K_{BL} * [BL] * \theta_* * \omega_{Cat.} \quad (\text{S57})$$

$$R_{LA_hyd} = R_{48} = k_{48} * [H_2] * \theta_{LA} * \omega_{Cat.} = k_{48} * [H_2] * K_{LA} * [LA] * \theta_* * \omega_{Cat.} \quad (\text{S58})$$

Material balance on active site leads to

$$1 = \theta_{BL} + \theta_{BHP} + \theta_{LA} + \theta_{HPA} + \theta_* \quad (S59)$$

By introducing eqs (S53)-(S56) in eq (S59), we get

$$1 = K_{BL} * [BL] * \theta_* + K_{BHP} * [BHP] * \theta_* + K_{LA} * [LA] * \theta_* + K_{HPA} * [HPA] * \theta_* + \theta_* \text{ equivalent to}$$

$$\theta_* = \frac{1}{K_{BL}*[BL]+K_{BHP}*[BHP]+K_{LA}*[LA]+K_{HPA}*[HPA]+1} \quad (S60)$$

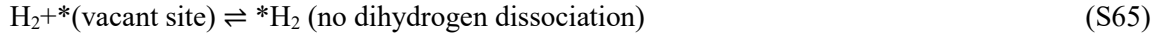
Hence, the rates of hydrogenation can be expressed as

$$R_{BL_{hyd}} = k_{47} * [H_2] * K_{BL} * [BL] * \frac{1}{(1+K_{BL}*[BL]+K_{BHP}*[BHP]+K_{LA}*[LA]+K_{HPA}*[HPA])} * \omega_{Cat}. \quad (S61)$$

$$R_{LA_{hyd}} = k_{48} * [H_2] * K_{LA} * [LA] * \frac{1}{(1+K_{BL}*[BL]+K_{BHP}*[BHP]+K_{LA}*[LA]+K_{HPA}*[HPA])} * \omega_{Cat}. \quad (S62)$$

Non-competitive Langmuir Hinshelwood with no dissociation of hydrogen (NCLH1.1)

Here we consider that BL and LA adsorb on a same site $\wedge$, and H_2 adsorb on a different site $*$



The adsorption and desorption steps are considered to be very fast. Thus,

$$K_{BL\wedge} = \frac{\theta_{\wedge BL}}{[BL] \cdot \theta_{\wedge}} \quad (S72)$$

$$K_{LA\wedge} = \frac{\theta_{\wedge LA}}{[LA] \cdot \theta_{\wedge}} \quad (S73)$$

$$K_{H2} = \frac{\theta_{H2}}{[H2] \cdot \theta_*} \quad (S74)$$

$$K_{BHP\wedge} = \frac{\theta_{\wedge BHP}}{[BHP] \cdot \theta_{\wedge}} \quad (S75)$$

$$K_{HPA\wedge} = \frac{\theta_{\wedge HPA}}{[HPA] \cdot \theta_{\wedge}} \quad (S76)$$

$K_{BL\wedge}$, $K_{LA\wedge}$, K_{H2} , $K_{BHP\wedge}$ and $K_{HPA\wedge}$ are adsorption constants.

Reactions (S66)-(S67) are considered to be the rate-determining steps for the hydrogenation steps.

From the experiments, the system is not reversible, so all the reaction determining steps will be assumed to be direct reactions.

$$R_{BL_hyd} = R_{66} = k_{66} * \theta_{H2} * \theta_{BL} * \omega_{Cat.} = k_{66} * K_{H2} * [H_2] * \theta_* * K_{BL} * [BL] * \theta_{\wedge} * \omega_{Cat.}$$

(S77)

$$R_{LA_hyd} = R_{67} = k_{67} * \theta_{H2} * \theta_{LA} * \omega_{Cat.} = k_{67} * K_{H2} * [H_2] * \theta_* * K_{LA} * [LA] * \theta_{\wedge} * \omega_{Cat.}$$

(S78)

Material balance on active site $\wedge$ leads to

$$1 = \theta_{BL} + \theta_{BHP} + \theta_{LA} + \theta_{HPA} + \theta_{\wedge}$$

(S79)

Material balance on active site $*$ leads to

$$1 = \theta_{H2} + \theta_*$$

(S80)

By introducing eq (S74) in eq (S80), we get

$$1 = K_{H2} * [H_2] * \theta_* + \theta_* \text{ equivalent to}$$

$$\theta_* = \frac{1}{K_{H2} * [H_2] + 1}$$

(S81)

By introducing eqs (S72), (S73), (S75) and (S76) in eq (S80), we get

$$1 = K_{BL} * [BL] * \theta_{\wedge} + K_{BHP} * [BHP] * \theta_{\wedge} + K_{LA} * [LA] * \theta_{\wedge} + K_{HPA} * [HPA] * \theta_{\wedge} + \theta_{\wedge}$$

equivalent to

$$\theta_{\wedge} = \frac{1}{K_{BL} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] + K_{HPA} * [HPA] + 1}$$

(S82)

Hence, the rate of hydrogenation can be expressed as

$$R_{BL_hyd} = R_{66} = k_{66} * K_{H2} * [H_2] * \frac{1}{(1 + K_{H2} * [H_2])} * K_{BL} * [BL] * \frac{1}{(1 + K_{BL} * [BL] + K_{BHP} * [BHP] + K_{LA} * [LA] + K_{HPA} * [HPA])} * \omega_{Cat.}$$

$$R_{BL_hyd} = k_{BL_hyd} * [H_2] * \frac{1}{(1+K_{H2}*[H_2])} * [BL] * \frac{1}{(1+K_{BL^\wedge}*[BL]+K_{BHP^\wedge}*[BHP]+K_{LA^\wedge}*[LA]+K_{HPA^\wedge}*[HPA])} * \omega_{Cat.} \quad (S83)$$

$$\text{with } k_{BL_hyd} = k_{66} * K_{H2} * K_{BL}$$

$$R_{LA_hyd} = R_{67} = k_{67} * K_{H2} * [H_2] * \frac{1}{(1+K_{H2}*[H_2])} * K_{LA^\wedge} * [LA] * \frac{1}{(1+K_{BL^\wedge}*[BL]+K_{BHP^\wedge}*[BHP]+K_{LA^\wedge}*[LA]+K_{HPA^\wedge}*[HPA])} * \omega_{Cat.}$$

Equivalent to

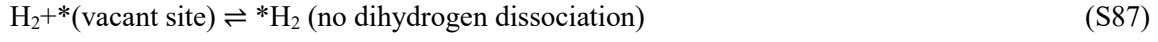
$$R_{LA_hyd} = k_{LA_hyd} * [H_2] * \frac{1}{(1 + K_{H2} * [H_2])} * [LA] * \frac{1}{(1 + K_{BL^\wedge} * [BL] + K_{BHP^\wedge} * [BHP] + K_{LA^\wedge} * [LA] + K_{HPA^\wedge} * [HPA])} * \omega_{Cat.}$$

(S84)

$$\text{With } k_{LA_hyd} = k_{67} * K_{H2} * K_{LA}$$

Non-competitive Langmuir Hinshelwood with no dissociation of hydrogen (NCLH1.2)

Here we consider that BL, LA and H₂ adsorb on different sites $\wedge$, 0 and *, respectively



The adsorption and desorption steps are considered to be very fast. Thus,

$$K_{BL\wedge} = \frac{\theta_{\wedge BL}}{[BL] \cdot \theta_{\wedge}} \quad (\text{S94})$$

$$K_{LA0} = \frac{\theta_{0LA}}{[LA] \cdot \theta_0} \quad (\text{S95})$$

$$K_{H_2} = \frac{\theta_{H_2}}{[H_2] \cdot \theta_*} \quad (\text{S96})$$

$$K_{BHP\wedge} = \frac{\theta_{\wedge BHP}}{[BHP] \cdot \theta_{\wedge}} \quad (\text{S97})$$

$$K_{HPA0} = \frac{\theta_{0HPA}}{[HPA] \cdot \theta_0} \quad (\text{S98})$$

$K_{BL\wedge}$, K_{LA0} , K_{H_2} , $K_{BHP\wedge}$ and K_{HPA0} are adsorption constants.

Reactions (S88)-(S89) are considered to be the rate-determining steps for the hydrogenation steps.

From the experiments, the system is not reversible, so all the reaction determining steps will be assumed to be direct reactions.

$$R_{BL_{hyd}} = R_{88} = k_{88} * \theta_{H_2} * \theta_{BL} * \omega_{Cat.} = k_{88} * K_{H_2} * [H_2] * \theta_* * K_{BL} * [BL] * \theta_{BL} * \omega_{Cat.}$$

(S99)

$$R_{LA_{hyd}} = R_{89} = k_{89} * \theta_{H_2} * \theta_{0LA} * \omega_{Cat.} = k_{89} * K_{H_2} * [H_2] * \theta_* * K_{LA0} * [LA] * \theta_0 * \omega_{Cat.}$$

(S100)

Material balance on active site $\wedge$ leads to

$$1 = \theta_{BL} + \theta_{BHP} + \theta_{\wedge}$$

(S101)

Material balance on active site 0 leads to

$$1 = \theta_{0LA} + \theta_{0HPA} + \theta_0$$

(S102)

Material balance on active site $*$ leads to

$$1 = \theta_{H_2} + \theta_*$$

(S103)

By introducing eq (S96) in eq (S103), we get

$$1 = K_{H_2} * [H_2] * \theta_* + \theta_* \text{ equivalent to}$$

$$\theta_* = \frac{1}{K_{H_2} * [H_2] + 1}$$

(S104)

By introducing eqs (S94) and (S97) in eq (S101), we get

$$1 = K_{BL} * [BL] * \theta_{\wedge} + K_{BHP} * [BHP] * \theta_{\wedge} + \theta_{\wedge} \text{ equivalent to}$$

$$\theta_{\wedge} = \frac{1}{K_{BL} * [BL] + K_{BHP} * [BHP] + 1}$$

(S105)

By introducing eqs (S95) and (S98) in eq (S102), we get

$$1 = K_{BL0} * [BL] * \theta_0 + K_{BHP0} * [BHP] * \theta_0 + \theta_0 \text{ equivalent to}$$

$$\theta_0 = \frac{1}{K_{BL0}*[BL] + K_{BHP0}*[BHP] + 1} \quad (S106)$$

Hence, the rates of hydrogenation can be expressed as

$$R_{BL_{hyd}} = R_{88} = k_{88} * K_{H2} * [H_2] * \frac{1}{(1 + K_{H2}*[H_2])} * K_{BL} * [BL] * \frac{1}{(1 + K_{BL}*[BL] + K_{BHP}*[BHP])} * \omega_{Cat}.$$

Equivalent to

$$R_{BL_{hyd}} = k_{BL_{hyd}} * [H_2] * \frac{1}{(1 + K_{H2}*[H_2])} * [BL] * \frac{1}{(1 + K_{BL}*[BL] + K_{BHP}*[BHP])} * \omega_{Cat}.$$

(S107)

With $k_{BL_{hyd}} = k_{88} * K_{H2} * K_{BL}$

$$R_{LA_{hyd}} = R_{89} = k_{89} * K_{H2} * [H_2] * \frac{1}{(1 + K_{H2}*[H_2])} * K_{0LA} * [LA] * \frac{1}{(1 + K_{LA}*[LA] + K_{HPA}*[HPA])} * \omega_{Cat}.$$

Equivalent to

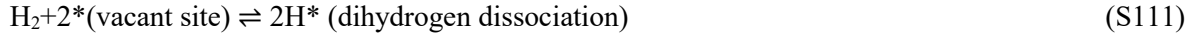
$$R_{LA_{hyd}} = k_{LA_{hyd}} * [H_2] * \frac{1}{(1 + K_{H2}*[H_2])} * [LA] * \frac{1}{(1 + K_{LA}*[LA] + K_{HPA}*[HPA])} * \omega_{Cat}.$$

(S108)

With $k_{LA_{hyd}} = k_{89} * K_{H2} * K_{0LA}$

Non-competitive Langmuir Hinshelwood with hydrogen dissociation (NCLH2.1)

Here we consider that BL-LA and H₂ adsorb on 2 different sites * and ^



The adsorption and desorption steps are considered to be very fast. Thus,

$$K_{\text{BL}^} = \frac{\theta_{\text{^BL}}}{[\text{BL}].\theta_{\text{^}}} \quad (\text{S120})$$

$$K_{\text{LA}^} = \frac{\theta_{\text{^LA}}}{[\text{LA}].\theta_{\text{^}}} \quad (\text{S121})$$

$$K_H = \frac{\theta_H^2}{[\text{H}_2].\theta_*^2} \quad (\text{S122})$$

$$K_{\text{BHP}^} = \frac{\theta_{\text{^BHP}}}{[\text{BHP}].\theta_{\text{^}}} \quad (\text{S123})$$

$$K_{\text{HPA}^} = \frac{\theta_{\text{^HPA}}}{[\text{HPA}].\theta_{\text{^}}} \quad (\text{S124})$$

$K_{BL}^{\wedge}$, $K_{LA}^{\wedge}$, $K_{HPA}^{\wedge}$, K_H and $K_{BHP}^{\wedge}$ are adsorption constants.

Steps (S112) and (S113) are supposed to be very fast and in equilibrium

$$K_{112} = \frac{\theta_{BLH} \cdot \theta_*}{\theta_{BL} \cdot \theta_H} = \frac{\theta_{BLH} \cdot \theta_*}{\theta_{BL} \cdot \sqrt{K_H * [H_2]} \cdot \theta_*} \quad (S125)$$

$$\theta_{BLH} = K_{112} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL}^{\wedge} \cdot [BL] \cdot \theta_{\wedge} \quad (S126)$$

$$K_{113} = \frac{\theta_{LAH} \cdot \theta_*}{\theta_{LA} \cdot \theta_H} = \frac{\theta_{LAH} \cdot \theta_*}{\theta_{LA} \cdot \sqrt{K_H * [H_2]} \cdot \theta_*} \quad (S127)$$

$$\theta_{LAH} = K_{113} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA}^{\wedge} \cdot [LA] \cdot \theta_{\wedge} \quad (S128)$$

Reactions (S114)-(S115) are considered to be rate-determining steps for hydrogenation steps. From the kinetic experiments, we did not observe that hydrogenation was a reversible step, so direct reaction was considered.

$$R_{BL_hyd} = R_{114} = k_{114} * \theta_H * \theta_{BLH} * \omega_{Cat.} = k_{114} * \sqrt{K_H * [H_2]} * \theta_* *$$

$$K_{112} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL}^{\wedge} \cdot [BL] \cdot \theta_{\wedge} * \omega_{Cat.} \quad (S129)$$

$$R_{LA_hyd} = R_{115} = k_{115} * \theta_H * \theta_{LAH} * \omega_{Cat.} = k_{115} * \sqrt{K_H * [H_2]} * \theta_* *$$

$$K_{113} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA}^{\wedge} \cdot [LA] \cdot \theta_{\wedge} * \omega_{Cat.} \quad (S130)$$

Material balance on active site $\wedge$ leads to

$$1 = \theta_{BL} + \theta_{BLH} + \theta_{BHP} + \theta_{LA} + \theta_{LAH} + \theta_{HPA} + \theta_{\wedge}$$

$$1 = K_{BL}^{\wedge} \cdot [BL] \cdot \theta_{\wedge} + K_{112} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL}^{\wedge} \cdot [BL] \cdot \theta_{\wedge} + K_{BHP}^{\wedge} \cdot [BHP] \cdot \theta_{\wedge} + K_{LA}^{\wedge} \cdot [LA] \cdot \theta_{\wedge} +$$

$$K_{113} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA}^{\wedge} \cdot [LA] \cdot \theta_{\wedge} + K_{HPA}^{\wedge} \cdot [HPA] \cdot \theta_{\wedge} + \theta_{\wedge}$$

$$\theta_{\wedge} = \frac{1}{K_{BL}^{\wedge} \cdot [BL] + K_{112} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL}^{\wedge} \cdot [BL] + K_{BHP}^{\wedge} \cdot [BHP] + K_{LA}^{\wedge} \cdot [LA] + K_{113} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA}^{\wedge} \cdot [LA] + K_{HPA}^{\wedge} \cdot [HPA] + 1}$$

(S131)

Material balance on active site $*$ leads to

$$1 = \theta_H + \theta_* \Leftrightarrow \theta_* = \frac{1}{\sqrt{K_H * [H_2]} + 1} \quad (S132)$$

Hence, the rates of hydrogenation can be expressed as

$$R_{BL_hyd} = k_{114} \cdot K_H \cdot K_{112} \cdot K_{BL^\wedge} \cdot [H_2] \frac{1}{\sqrt{K_H * [H_2]} + 1} [BL] \cdot \frac{1}{K_{BL^\wedge} \cdot [BL] + K_{112} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL^\wedge} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + K_{LA^\wedge} \cdot [LA] + K_{113} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA^\wedge} \cdot [LA] + K_{HPA^\wedge} \cdot [HPA] + 1} \omega_{Cat}.$$

Equivalent to

$$R_{BL_hyd} = k_{BL_hyd} \cdot [H_2] \cdot [BL] \frac{1}{\sqrt{K_H * [H_2]} + 1} \cdot \frac{1}{K_{BL^\wedge} \cdot [BL] + K_c \cdot \sqrt{[H_2]} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + K_{LA^\wedge} \cdot [LA] + K_{c2} \cdot \sqrt{[H_2]} \cdot [LA] + K_{HPA^\wedge} \cdot [HPA] + 1} \omega_{Cat}. \quad (S134)$$

With $k_{BL_hyd} = k_{114} \cdot K_H \cdot K_{112} \cdot K_{BL^\wedge}$, $K_c = K_{112} \cdot \sqrt{K_H} \cdot K_{BL^\wedge}$ and $K_{c2} = K_{113} \cdot \sqrt{K_H} \cdot K_{LA^\wedge}$

$$R_{LA_hyd} = k_{115} \cdot K_H \cdot K_{113} \cdot K_{LA^\wedge} \cdot [H_2] \frac{1}{\sqrt{K_H * [H_2]} + 1} [BL] \cdot \frac{1}{K_{BL^\wedge} \cdot [BL] + K_{112} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL^\wedge} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + K_{LA^\wedge} \cdot [LA] + K_{113} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA^\wedge} \cdot [LA] + K_{HPA^\wedge} \cdot [HPA] + 1} \omega_{Cat}.$$

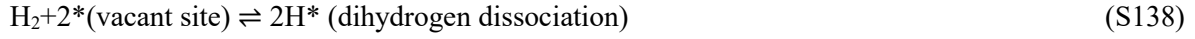
Equivalent to

$$R_{LA_hyd} = k_{LA_hyd} \cdot [H_2] \cdot [LA] \frac{1}{\sqrt{K_H * [H_2]} + 1} \cdot \frac{1}{K_{BL^\wedge} \cdot [BL] + K_c \cdot \sqrt{[H_2]} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + K_{LA^\wedge} \cdot [LA] + K_{c2} \cdot \sqrt{[H_2]} \cdot [LA] + K_{HPA^\wedge} \cdot [HPA] + 1} \omega_{Cat}. \quad (S135)$$

With $k_{LA_hyd} = k_{115} \cdot K_H \cdot K_{113} \cdot K_{LA^\wedge}$

Non-competitive Langmuir Hinshelwood with hydrogen dissociation (NCLH2.2)

Here we consider that BL, LA and H₂ adsorb on different sites $\wedge$, 0 and *



The adsorption and desorption steps are considered to be very fast. Thus,

$$K_{BL\wedge} = \frac{\theta_{\wedge BL}}{[BL].\theta_{\wedge}} \quad (\text{S147})$$

$$K_{LA0} = \frac{\theta_{0LA}}{[LA].\theta_0} \quad (\text{S148})$$

$$K_H = \frac{\theta_H^2}{[H_2].\theta_*^2} \quad (\text{S149})$$

$$K_{BHP\wedge} = \frac{\theta_{\wedge BHP}}{[BHP].\theta_{\wedge}} \quad (\text{S150})$$

$$K_{HPA0} = \frac{\theta_{0HPA}}{[HPA].\theta_0} \quad (\text{S151})$$

$K_{BL^\wedge}$, K_{LA0} , K_{HPA0} , K_H and $K_{BHP^\wedge}$ are adsorption constants.

Steps (S139) and (S140) are supposed to be very fast and in equilibrium

$$K_{139} = \frac{\theta_{BLH} \cdot \theta_*}{\theta_{BL} \cdot \theta_H} = \frac{\theta_{BLH} \cdot \theta_*}{\theta_{BL} \cdot \sqrt{K_H * [H_2]} \cdot \theta_*} \quad (S152)$$

$$\theta_{BLH} = K_{139} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL^\wedge} \cdot [BL] \cdot \theta^\wedge \quad (S153)$$

$$K_{140} = \frac{\theta_{0LAH} \cdot \theta_*}{\theta_{0LA} \cdot \theta_H} = \frac{\theta_{0LAH} \cdot \theta_*}{\theta_{0LA} \cdot \sqrt{K_H * [H_2]} \cdot \theta_*} \quad (S154)$$

$$\theta_{0LAH} = K_{140} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA0} \cdot [LA] \cdot \theta_0 \quad (S155)$$

Reactions (S141)-(S142) are considered to be rate-determining steps for hydrogenation steps. From the kinetic experiments, we did not observe that hydrogenation was a reversible step, so direct reaction was considered.

$$R_{BL_hyd} = R_{141} = k_{141} * \theta_H * \theta_{BLH} * \omega_{Cat.} = k_{141} * \sqrt{K_H * [H_2]} * \theta_* *$$

$$K_{139} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL^\wedge} \cdot [BL] \cdot \theta^\wedge * \omega_{Cat.} \quad (S156)$$

$$R_{LA_hyd} = R_{142} = k_{142} * \theta_H * \theta_{0LAH} * \omega_{Cat.} = k_{142} * \sqrt{K_H * [H_2]} * \theta_* *$$

$$K_{140} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA0} \cdot [LA] \cdot \theta_0 * \omega_{Cat.} \quad (S157)$$

Material balance on active site $^\wedge$ leads to

$$1 = \theta_{BL} + \theta_{BLH} + \theta_{BHP} + \theta^\wedge$$

$$1 = K_{BL^\wedge} \cdot [BL] \cdot \theta^\wedge + K_{139} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL^\wedge} \cdot [BL] \cdot \theta^\wedge + K_{BHP^\wedge} \cdot [BHP] \cdot \theta^\wedge + \theta^\wedge$$

$$\theta^\wedge = \frac{1}{K_{BL^\wedge} \cdot [BL] + K_{139} \cdot \sqrt{K_H * [H_2]} \cdot K_{BL^\wedge} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + 1} \quad (S158)$$

Material balance on active site 0 leads to

$$1 = \theta_{0LA} + \theta_{0LAH} + \theta_{0HPA} + \theta_0$$

$$1 = K_{LA0} \cdot [LA] \cdot \theta_0 + K_{140} \cdot \sqrt{K_H * [H_2]} \cdot K_{LA0} \cdot [LA] \cdot \theta_0 + K_{HPA0} \cdot [HPA] \cdot \theta_0 + \theta_0$$

$$\theta_0 = \frac{1}{K_{LA0} \cdot [LA] + K_{140} \cdot \sqrt{K_H \cdot [H_2]} \cdot K_{LA0} \cdot [LA] + K_{HPA0} \cdot [HPA] + 1} \quad (S159)$$

Material balance on active site * leads to

$$1 = \theta_H + \theta_* \Leftrightarrow \theta_* = \frac{1}{\sqrt{K_H \cdot [H_2]} + 1} \quad (S160)$$

Hence, the rates of hydrogenation can be expressed as

$$R_{BL_hyd} = k_{141} \cdot K_H \cdot K_{139} \cdot K_{BL^\wedge} \cdot [H_2] \frac{1}{\sqrt{K_H \cdot [H_2]} + 1} [BL]$$

$$\cdot \frac{1}{K_{BL^\wedge} \cdot [BL] + K_{139} \cdot \sqrt{K_H \cdot [H_2]} \cdot K_{BL^\wedge} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + 1} \omega_{Cat}.$$

Equivalent to

$$R_{BL_hyd} = k_{BL_hyd} \cdot [H_2] \cdot [BL] \frac{1}{\sqrt{K_H \cdot [H_2]} + 1} \cdot \frac{1}{K_{BL^\wedge} \cdot [BL] + K_c \cdot \sqrt{[H_2]} \cdot [BL] + K_{BHP^\wedge} \cdot [BHP] + 1} \omega_{Cat}. \quad (S161)$$

With $k_{BL_hyd} = k_{141} \cdot K_H \cdot K_{139} \cdot K_{BL^\wedge}$, $K_c = K_{139} \cdot \sqrt{K_H} \cdot K_{BL^\wedge}$

$$R_{LA_hyd} = k_{142} \cdot K_H \cdot K_{140} \cdot K_{LA0} \cdot [H_2] \frac{1}{\sqrt{K_H \cdot [H_2]} + 1} [LA] \cdot \frac{1}{K_{LA0} \cdot [LA] + K_{140} \cdot \sqrt{K_H \cdot [H_2]} \cdot K_{LA0} \cdot [LA] + K_{HPA0} \cdot [HPA] + 1} \omega_{Cat}.$$

Equivalent to

$$R_{LA_hyd} = k_{LA_hyd} \cdot [H_2] \cdot [LA] \frac{1}{\sqrt{K_H \cdot [H_2]} + 1} \cdot \frac{1}{K_{LA0} \cdot [LA] + K_{c2} \cdot \sqrt{[H_2]} \cdot [LA] + K_{HPA0} \cdot [HPA] + 1} \omega_{Cat}. \quad (S162)$$

With $k_{LA_hyd} = k_{142} \cdot K_H \cdot K_{140} \cdot K_{LA0}$ and $K_{c2} = K_{140} \cdot \sqrt{K_H} \cdot K_{LA0}$

S2. Derivation of cyclization rate expression

Non-catalytic effect

In this route, the cyclization of HPA or BHP to GVL is not catalyzed.



The rate equations can be expressed as

$$R_{\text{BHP}_{\text{noncat}}} = k_{\text{BHP}_{\text{noncat}}} \cdot [\text{BHP}] \quad (\text{S165})$$

$$R_{\text{HPA}_{\text{noncat}}} = k_{\text{HPA}_{\text{noncat}}} \cdot [\text{HPA}] \quad (\text{S166})$$

Catalyzed route by Ru/C

A Langmuir-Hinshelwood model was assumed and by considering that each site has the same affinity towards BL or LA. The following steps can be assumed



Reactions (S167), (S168) and (S171) are assumed to be fast and reversible, thus the adsorption can be expressed as

$$K_{\blacksquare\text{HPA}} = \frac{\theta_{\blacksquare\text{HPA}}}{[\text{HPA}] \cdot \theta_{\blacksquare}} \quad (\text{S172})$$

$$K_{\blacksquare\text{BHP}} = \frac{\theta_{\blacksquare\text{BHP}}}{[\text{BHP}] \cdot \theta_{\blacksquare}} \quad (\text{S173})$$

$$K_{\blacksquare GVL} = \frac{\theta_{\blacksquare GVL}}{[GVL] \cdot \theta_{\blacksquare}} \quad (S174)$$

Rate determining steps are (S169) and (S170), thus the rates

$$R_{BHP_RuC} = R_{169} = k_{169} \cdot \theta_{\blacksquare BHP} \cdot \omega_{Cat.RuC} \quad (S175)$$

$$R_{HPA_RuC} = R_{170} = k_{170} \cdot \theta_{\blacksquare HPA} \cdot \omega_{Cat.RuC} \quad (S176)$$

Mass balance on active site $\blacksquare$ leads to

$$1 = \theta_{\blacksquare HPA} + \theta_{\blacksquare BHP} + \theta_{\blacksquare} \quad (S177)$$

Thus,

$$1 = K_{\blacksquare HPA} \cdot [HPA] \cdot \theta_{\blacksquare} + K_{\blacksquare BHP} \cdot [BHP] \cdot \theta_{\blacksquare} + \theta_{\blacksquare}$$

$$\Leftrightarrow \theta_{\blacksquare} = \frac{1}{K_{\blacksquare HPA} \cdot [HPA] + K_{\blacksquare BHP} \cdot [BHP] + 1} \quad (S178)$$

The following notations are introduced $k_{BHP_RuC} = k_{169} \cdot K_{\blacksquare BHP}$ and $k_{HPA_RuC} = k_{170} \cdot K_{\blacksquare HPA}$. By combining (S178) with (S172), (S173), (S175) and (S176), the rates of cyclization catalyzed by Ru/C can be expressed as

$$R_{BHP_RuC} = k_{BHP_RuC} \cdot [BHP] \cdot \frac{1}{K_{\blacksquare HPA} \cdot [HPA] + K_{\blacksquare BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.RuC} \quad (S179)$$

$$R_{HPA_RuC} = k_{HPA_RuC} \cdot [HPA] \cdot \frac{1}{K_{\blacksquare HPA} \cdot [HPA] + K_{\blacksquare BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.RuC} \quad (S180)$$

Cyclization catalyzed route by Amberlite IR120

The active site of Amberlite IR120 is $-\text{SO}_3\text{H}$. The following mechanism was assumed





Reactions (S181), (S182) and (S185) are assumed to be fast and reversible, thus the adsorption can be expressed as

$$K_{\text{SO}_3\text{H}-\text{HPA}} = \frac{\theta_{\text{SO}_3\text{H}-\text{HPA}}}{[\text{HPA}] \cdot \theta_{\text{SO}_3\text{H}-}} \quad (\text{S186})$$

$$K_{\text{SO}_3\text{H}-\text{BHP}} = \frac{\theta_{\text{SO}_3\text{H}-\text{BHP}}}{[\text{BHP}] \cdot \theta_{\text{SO}_3\text{H}}} \quad (\text{S187})$$

$$K_{\text{SO}_3\text{H}-\text{GVL}} = \frac{\theta_{\text{SO}_3\text{H}-\text{GVL}}}{[\text{GVL}] \cdot \theta_{\text{SO}_3\text{H}}} \quad (\text{S188})$$

Rate determining steps are (S181) and (S182), thus the rates

$$R_{\text{BHP}_\text{SO}_3\text{H}} = R_{181} = k_{181} \cdot \theta_{\text{SO}_3\text{H}-\text{BHP}} \cdot \omega_{\text{Cat.Amberlite}} \quad (\text{S189})$$

$$R_{\text{HPA}_\text{SO}_3\text{H}} = R_{182} = k_{182} \cdot \theta_{\text{SO}_3\text{H}-\text{HPA}} \cdot \omega_{\text{Cat.Amberlite}} \quad (\text{S190})$$

Mass balance on active site $-\text{SO}_3\text{H}$ leads to

$$1 = \theta_{\text{SO}_3\text{H}-\text{HPA}} + \theta_{\text{SO}_3\text{H}-\text{BHP}} + \theta_{\text{SO}_3\text{H}-} \quad (\text{S191})$$

Thus,

$$1 = K_{\text{SO}_3\text{H}-\text{HPA}} \cdot [\text{HPA}] \cdot \theta_{\text{SO}_3\text{H}-} + K_{\text{SO}_3\text{H}-\text{BHP}} \cdot [\text{BHP}] \cdot \theta_{\text{SO}_3\text{H}-} + \theta_{\text{SO}_3\text{H}-}$$

$$\Leftrightarrow \theta_{\text{SO}_3\text{H}-} = \frac{1}{K_{\text{SO}_3\text{H}-\text{HPA}} \cdot [\text{HPA}] + K_{\text{SO}_3\text{H}-\text{BHP}} \cdot [\text{BHP}] + 1} \quad (\text{S192})$$

The following notations are introduced $k_{\text{BHP}_\text{SO}_3\text{H}} = k_{181} \cdot K_{\text{SO}_3\text{H}-\text{BHP}}$ and $k_{\text{HPA}_\text{SO}_3\text{H}} = k_{182} \cdot$

$K_{\text{SO}_3\text{H}-\text{HPA}}$. By combining (S192) with (S186), (S187), (S189) and (S190), the rates of cyclization

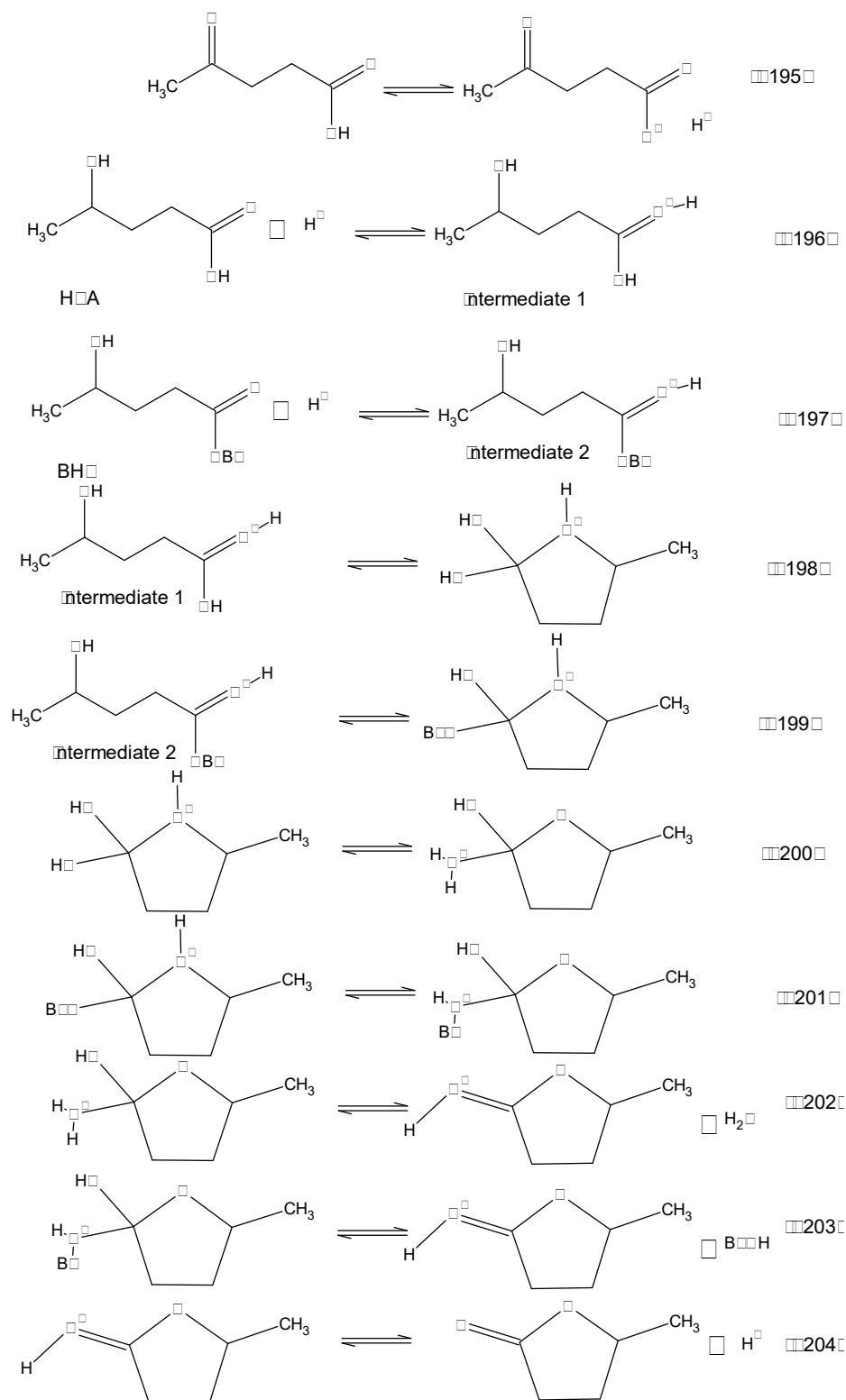
catalyzed by Ru/C can be expressed as

$$R_{\text{BHP}_\text{SO}_3\text{H}} = k_{\text{BHP}_\text{SO}_3\text{H}} \cdot [\text{BHP}] \cdot \frac{1}{K_{\text{SO}_3\text{H}-\text{HPA}} \cdot [\text{HPA}] + K_{\text{SO}_3\text{H}-\text{BHP}} \cdot [\text{BHP}] + 1} \cdot \omega_{\text{Cat.Amb}} \quad (\text{S193})$$

$$R_{HPA_SO3H} = k_{HPA_SO3H} \cdot [HPA] \cdot \frac{1}{K_{SO3H-HPA} \cdot [HPA] + K_{SO3H-BHP} \cdot [BHP] + 1} \cdot \omega_{Cat.RuC} \quad (S194)$$

Cyclization catalyzed route by LA dissociation

The dissociation of LA produces protons that can catalyze the cyclization reactions. The following mechanism is assumed



Reactions (S196)-(S197) are assumed to be fast and reversible, thus the equilibrium constants can be expressed as

$$K_{195} = \frac{[LA^-] \cdot [H^+]}{[LA]} \quad (S205)$$

$$K_{196} = \frac{[Intermediate1]}{[H^+].[HPA]} \text{ (S206)}$$

$$K_{197} = \frac{[Intermediate2]}{[H^+].[BHP]} \text{ (S207)}$$

The rate determining steps are (S198) and (S199), and can be expressed as

$$R_{BHP_diss} = k_{BHP_diss} \cdot [BHP] \cdot [LA]$$

$$R_{HPA_diss} = k_{HPA_diss} \cdot [HPA] \cdot [LA]$$

S3. Modeling results

NCLH1.1.

Table S3.1. Estimated values et $T_{ref}=392.15K$ and statistical data for NCLH1.1.

Parameters		Units	Estimates	HDP%
Par1	$k_{BL_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	3.57E-06	13.65
Par2	Ea_{BL_hyd}	$J.mol^{-1}$	3.20E+04	10.55
Par3	$K_{H2.}$	$m^3.mol^{-1}$	1.87E-03	151.37
Par4	$K_{BL}^{\wedge}$	$m^3.mol^{-1}$	9.79E-04	23.77
Par5	$K_{BHP}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	4.30E-05	48.27
Par7	$Ea'_{BHP_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	s^{-1}	6.35E-05	32.62
Par9	Ea_{BHP_noncat}	$J.mol^{-1}$	7.74E+04	40.02
Par10	$k_{LA_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	7.79E-06	13.97
Par11	Ea_{LA_hyd}	$J.mol^{-1}$	5.49E+04	7.01
Par12	$K_{LA}^{\wedge}$	$m^3.mol^{-1}$	3.54E-04	53.75
Par13	$K_{HPA}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	5.21E-04	54.44
Par15	$Ea'_{HPA_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	s^{-1}	5.39E-07	255.52
Par17	Ea_{HPA_noncat}	$J.mol^{-1}$	4.56E+05	39.09
Par18	$K_{BHP-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-

Par20	$k_{BHP_RuC}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	2.20E-05	20.80
Par21	Ea_{BHP_RuC}	$J.mol^{-1}$	Fixed to zero	-
Par22	$k_{HPA_RuC}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	6.96E-05	8.77
Par23	Ea_{HPA_RuC}	$J.mol^{-1}$	Fixed to zero	-
Par24	$K_{\blacksquare BHP}$	$m^3.mol^{-1}$	Fixed to zero	-
Par25	$K_{\blacksquare HPA}$	$m^3.mol^{-1}$	Fixed to zero	-
Par26	$k_{BHP_diss}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}$	1.36E-06	15.61
Par27	Ea_{BHP_diss}	$J.mol^{-1}$	1.04E+05	12.86
Par28	$k_{HPA_diss}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}$	3.71E-06	7.08
Par29	Ea_{HPA_diss}	$J.mol^{-1}$	7.41E+04	7.93

Table S3.2. Normalized parameter covariance matrix for NCLH1.1.

	Par1	Par2	Par3	Par4	Par6	Par8	Par9	Par10	Par11	Par12	Par14	Par16	Par17	Par20	Par22	Par26	Par27	Par28	Par29
Par1	1																		
Par2	0.082	1																	
Par3	0.662	-0.064	1																
Par4	0.618	0.369	-0.081	1															
Par6	0.027	0.007	0.022	0.015	1														
Par8	0.048	0.105	-0.029	0.066	0.006	1													
Par9	-0.073	-0.192	0.013	-0.115	-0.006	-0.938	1												
Par10	0.911	-0.028	0.62	0.531	0.015	0.07	-0.079	1											
Par11	-0.021	0.047	-0.019	0.015	-0.005	-0.08	0.112	0.083	1										
Par12	0.266	-0.547	0.262	-0.23	-0.01	0.078	-0.021	0.446	-0.135	1									
Par14	0.007	0.005	0.013	0	0	0	0	0.013	0.005	-0.006	1								
Par16	0.018	0.003	-0.007	0.029	0.003	0.026	-0.039	-0.039	-0.346	0.021	-0.002	1							
Par17	-0.008	-0.007	0.027	-0.034	-0.002	-0.023	0.034	0.036	0.279	-0.013	0.002	-0.967	1						
Par20	0.027	0.021	0.055	0.02	-0.002	-0.761	0.636	0.05	0.015	0.006	0.001	-0.006	0.006	1					
Par22	0.273	0.04	0.169	0.208	0.008	-0.022	0.016	0.187	0.211	0.034	-0.006	-0.347	0.306	0.017	1				
Par26	0.039	-0.2	-0.119	0.029	-0.036	-0.134	0.156	0.177	0.298	0.186	0.007	-0.121	0.093	-0.108	-0.059	1			
Par27	0.143	0.263	0.097	0.119	-0.006	0.233	-0.351	0.247	0.366	0.051	0.006	-0.133	0.107	0.06	0.087	0.212	1		

Par28	-0.176	-0.108	-0.311	-0.042	-0.015	0.049	-0.013	0.074	0.218	0.153	0.01	-0.202	0.145	-0.02	-0.44	0.403	0.139	1	
Par29	0.07	0.087	0.032	0.074	-0.002	-0.046	0.064	0.126	0.751	-0.052	0.001	-0.494	0.397	0.018	0.442	0.186	0.306	0.18	1

LH1Table S3.3. Estimated values et $T_{ref}=392.15K$ and statistical data for LH1.

Parameters		Units	Estimates	HDP%
Par1	$k_{BL_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	3.67E-06	14.96
Par2	Ea_{BL_hyd}	$J.mol^{-1}$	3.15E+04	10.58
Par3	$K_{H2.}$	$m^3.mol^{-1}$	1.73E-03	110.02
Par4	$K_{BL}^{\wedge}$	$m^3.mol^{-1}$	3.96E-04	17.53
Par5	$K_{BHP}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	4.30E-05	47.82
Par7	$Ea'_{BHP_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	s^{-1}	6.36E-05	32.33
Par9	Ea_{BHP_noncat}	$J.mol^{-1}$	7.75E+04	39.20
Par10	$k_{LA_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	8.12E-06	15.21
Par11	Ea_{LA_hyd}	$J.mol^{-1}$	5.47E+04	6.98
Par12	$K_{LA}^{\wedge}$	$m^3.mol^{-1}$	1.61E-04	41.45
Par13	$K_{HPA}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	5.20E-04	52.93
Par15	$Ea'_{HPA_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	s^{-1}	4.45E-07	263.18
Par17	Ea_{HPA_noncat}	$J.mol^{-1}$	4.70E+05	40.19
Par18	$K_{BHP-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par20	$k_{BHP_RuC}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ RuC^{-1}$	2.21E-05	20.53
Par21	Ea_{BHP_RuC}	$J.mol^{-1}$	Fixed to zero	-

Par14	0.009	0.005	0.012	0.005	0	0	0	0.014	0.006	- 0.004	1								
Par16	- 0.001	0.004	- 0.017	0.019	0.003	0.026	-0.039	- 0.055	- 0.359	0.008	- 0.002	1							
Par17	0.014	- 0.007	0.035	- 0.016	- 0.002	-0.023	0.035	0.058	0.299	0	0.002	-0.972	1						
Par20	0.034	0.019	0.062	0.033	- 0.002	-0.764	0.64	0.056	0.017	0.013	0.001	-0.008	0.008	1					
Par22	0.237	0.042	0.164	0.238	0.007	-0.028	0.022	0.157	0.218	0.03	- 0.007	-0.36	0.325	0.018	1				
Par26	0.003	- 0.209	- 0.111	- 0.023	- 0.036	-0.136	0.157	0.135	0.308	0.167	0.008	-0.127	0.101	-0.105	-0.064	1			
Par27	0.123	0.261	0.095	0.135	- 0.007	0.229	-0.347	0.225	0.373	0.054	0.006	-0.145	0.121	0.058	0.084	0.223	1		
Par28	- 0.215	- 0.115	- 0.299	- 0.154	- 0.015	0.053	-0.017	0.019	0.221	0.123	0.01	-0.205	0.154	-0.023	-0.442	0.415	0.15	1	
Par29	0.053	0.079	0.025	0.075	- 0.003	-0.047	0.066	0.105	0.757	- 0.049	0	-0.511	0.426	0.019	0.447	0.192	0.314	0.18	1

LH2Table S3.5. Estimated values et $T_{ref}=392.15K$ and statistical data for LH2.

Parameters		Units	Estimates	HDP%
Par1	$k_{BL_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	3.61E-06	23.96
Par2	Ea_{BL_hyd}	$J.mol^{-1}$	3.17E+04	10.55
Par3	$K_{H2.}$	$m^3.mol^{-1}$	8.38E-03	241.80
Par4	$K_{BL}^{\wedge}$	$m^3.mol^{-1}$	3.99E-04	19.29
Par5	$K_{BHP}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	4.27E-05	48.01
Par7	$Ea'_{BHP_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	s^{-1}	6.41E-05	32.40
Par9	Ea_{BHP_noncat}	$J.mol^{-1}$	7.66E+04	39.69
Par10	$k_{LA_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	7.96E-06	23.99
Par11	Ea_{LA_hyd}	$J.mol^{-1}$	5.45E+04	7.00
Par12	$K_{LA}^{\wedge}$	$m^3.mol^{-1}$	1.47E-04	47.35
Par13	$K_{HPA}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	5.17E-04	53.05
Par15	$Ea'_{HPA_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	s^{-1}	5.20E-07	246.56
Par17	Ea_{HPA_noncat}	$J.mol^{-1}$	4.58E+05	39.27
Par18	$K_{BHP-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par20	$k_{BHP_RuC}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ RuC^{-1}$	2.20E-05	20.94
Par21	Ea_{BHP_RuC}	$J.mol^{-1}$	Fixed to zero	-

Par22	$k_{HPA_{RuC}}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	7.03E-05	8.46
Par23	$Ea_{HPA_{RuC}}$	$J.mol^{-1}$	Fixed to zero	-
Par24	$K_{\blacksquare BHP}$	$m^3.mol^{-1}$	Fixed to zero	-
Par25	$K_{\blacksquare HPA}$	$m^3.mol^{-1}$	Fixed to zero	-
Par26	K_i	$m^{9/2}.mol^{-3/2}$	Fixed to zero	-
Par27	K_{i2}	$m^{9/2}.mol^{-3/2}$	Fixed to zero	-
Par28	$k_{BHP_{diss}}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}$	1.42E-06	15.78
Par29	$Ea_{BHP_{diss}}$	$J.mol^{-1}$	1.05E+05	12.82
Par30	$k_{HPA_{diss}}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}$	3.76E-06	7.03
Par31	$Ea_{HPA_{diss}}$	$J.mol^{-1}$	7.43E+04	7.82

Table S3.6 Normalized parameter covariance matrix for LH2.

	Par1	Par2	Par3	Par4	Par6	Par8	Par9	Par10	Par11	Par12	Par14	Par16	Par17	Par20	Par22	Par28	Par29	Par30	Par31
Par1	1																		
Par2	-0.011	1																	
Par3	0.942	-0.064	1																
Par4	0.735	0.295	0.521	1															
Par6	0.025	0.007	0.021	0.025	1														
Par8	-0.004	0.115	-0.033	0.03	0.007	1.00													
Par9	-0.017	-0.204	0.016	-0.083	-0.007	-0.937	1												
Par10	0.97	-0.072	0.927	0.682	0.019	0.01	-0.022	1											
Par11	-0.023	0.026	-0.019	-0.004	-0.005	-0.083	0.116	0.039	1										
Par12	0.466	-0.52	0.474	-0.012	-0.004	0.052	-0.004	0.563	-0.116	1									
Par14	0.01	0.005	0.012	0.007	0	0	-0.001	0.014	0.005	-0.003	1								
Par16	-0.008	0.002	-0.017	0.011	0.002	0.026	-0.039	-0.041	-0.35	0.009	-0.002	1							
Par17	0.024	-0.005	0.036	-0.004	-0.002	-0.023	0.033	0.05	0.284	0.002	0.002	-0.968	1						
Par20	0.044	0.015	0.059	0.041	-0.002	-0.77	0.647	0.057	0.019	0.017	0.001	-0.009	0.008	1					
Par22	0.222	0.042	0.17	0.258	0.007	-0.027	0.022	0.174	0.216	0.048	-0.006	-0.357	0.316	0.018	1				

Par28	-0.05	-0.206	-0.118	-0.049	-0.036	-0.139	0.161	0.033	0.306	0.142	0.008	-0.122	0.093	-0.099	-0.065	1			
Par29	0.121	0.263	0.099	0.15	-0.007	0.233	-0.351	0.186	0.369	0.061	0.006	-0.14	0.114	0.05	0.085	0.219	1		
Par30	-0.267	-0.108	-0.309	-0.214	-0.014	0.052	-0.017	-0.122	0.219	0.079	0.01	-0.198	0.142	-0.023	-0.442	0.413	0.146	1	
Par31	0.053	0.086	0.034	0.082	-0.002	-0.048	0.066	0.086	0.753	-0.044	0.001	-0.502	0.407	0.02	0.447	0.189	0.312	0.177	1

ER1Table S3.7. Estimated values et $T_{ref}=392.15K$ and statistical data for ER1.

Parameters		Units	Estimates	HDP%
Par1	$k_{BL_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	4.01E-06	22.86
Par2	Ea_{BL_hyd}	$J.mol^{-1}$	3.14E+04	10.88
Par3	$K_{H2.}$	$m^3.mol^{-1}$	5.81E-03	115.48
Par4	$K_{BL}^{\wedge}$	$m^3.mol^{-1}$	1.13E-03	31.80
Par5	$K_{BHP}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	4.29E-05	48.34
Par7	$Ea'_{BHP_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	s^{-1}	6.33E-05	32.79
Par9	Ea_{BHP_noncat}	$J.mol^{-1}$	7.91E+04	38.80
Par10	$k_{LA_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	8.88E-06	23.01
Par11	Ea_{LA_hyd}	$J.mol^{-1}$	5.49E+04	6.94
Par12	$K_{LA}^{\wedge}$	$m^3.mol^{-1}$	5.05E-04	53.91
Par13	$K_{HPA}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	5.20E-04	54.14
Par15	$Ea'_{HPA_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	s^{-1}	4.25E-07	271.12
Par17	Ea_{HPA_noncat}	$J.mol^{-1}$	4.73E+05	41.25
Par18	$K_{BHP-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par20	$k_{BHP_RuC}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ RuC^{-1}$	2.21E-05	20.69
Par21	Ea_{BHP_RuC}	$J.mol^{-1}$	Fixed to zero	-

Par22	$k_{HPA_{RuC}}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ RuC^{-1}$	6.95E-05	8.70
Par23	$Ea_{HPA_{RuC}}$	J.mol ⁻¹	Fixed to zero	-
Par24	$K_{\blacksquare BHP}$	m ³ .mol ⁻¹	Fixed to zero	-
Par25	$K_{\blacksquare HPA}$	m ³ .mol ⁻¹	Fixed to zero	-
Par26	$k_{BHP_{diss}}(T_{Ref})$	m ³ .mol ⁻¹ .s ⁻¹	1.37E-06	15.80
Par27	$Ea_{BHP_{diss}}$	J.mol ⁻¹	1.04E+05	12.92
Par28	$k_{HPA_{diss}}(T_{Ref})$	m ³ .mol ⁻¹ .s ⁻¹	3.74E-06	7.04
Par29	$Ea_{HPA_{diss}}$	J.mol ⁻¹	7.44E+04	7.89

Table S3.8 Normalized parameter covariance matrix for ER1.

	Par1	Par2	Par3	Par4	Par6	Par8	Par9	Par10	Par11	Par12	Par14	Par16	Par17	Par20	Par22	Par26	Par27	Par28	Par29
Par1	1																		
Par2	0.035	1																	
Par3	0.894	-0.047	1																
Par4	0.882	0.257	0.642	1															
Par6	0.027	0.007	0.022	0.026	1														
Par8	0	0.104	-0.046	0.019	0.005	1.00													
Par9	-0.021	-0.192	0.031	-0.065	-0.005	-0.938	1												
Par10	0.967	-0.033	0.878	0.835	0.02	0.014	-0.026	1											
Par11	-0.012	0.042	-0.01	0.008	-0.004	-0.077	0.108	0.05	1										
Par12	0.564	-0.476	0.575	0.237	0	0.048	-0.005	0.656	-0.118	1									
Par14	0.007	0.005	0.01	0.006	0	0	0	0.011	0.006	-0.002	1								
Par16	-0.006	0.006	-0.022	0.008	0.002	0.026	-0.039	-0.04	-0.352	0.006	-0.002	1							
Par17	0.019	-0.009	0.04	0	-0.001	-0.024	0.035	0.048	0.295	0.006	0.002	-0.973	1						
Par20	0.055	0.023	0.07	0.062	-0.001	-0.759	0.635	0.07	0.014	0.031	0.001	-0.007	0.007	1					
Par22	0.23	0.043	0.156	0.263	0.007	-0.025	0.021	0.181	0.216	0.076	-0.007	-0.359	0.325	0.018	1				
Par26	-0.006	-0.207	-0.088	-0.041	-0.035	-0.129	0.149	0.079	0.294	0.144	0.008	-0.121	0.097	-0.11	-0.055	1			
Par27	0.132	0.256	0.101	0.152	-0.007	0.226	-0.344	0.198	0.369	0.082	0.006	-0.138	0.115	0.064	0.081	0.225	1		
Par28	-0.22	-0.121	-0.276	-0.217	-0.014	0.058	-0.023	-0.067	0.213	0.058	0.011	-0.202	0.153	-0.023	-0.433	0.4	0.148	1	
Par29	0.052	0.079	0.027	0.073	-0.002	-0.045	0.064	0.086	0.753	-0.035	0	-0.506	0.423	0.016	0.442	0.188	0.304	0.185	1

NCLH21Table S3.9. Estimated values et $T_{ref}=392.15K$ and statistical data for NCLH21.

Parameters		Units	Estimates	HDP%
Par1	$k_{BL_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	3.25E-06	9.48
Par2	Ea_{BL_hyd}	$J.mol^{-1}$	3.02E+04	10.99
Par3	$K_{H2.}$	$m^{3/2}.mol^{-1/2}$	Fixed to zero	-
Par4	$K_{BL}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par5	$K_{BHP}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par6	$k'_{BHP_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	4.32E-05	47.89
Par7	$Ea'_{BHP_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par8	$k_{BHP_noncat}(T_{Ref})$	s^{-1}	6.32E-05	32.20
Par9	Ea_{BHP_noncat}	$J.mol^{-1}$	7.89E+04	38.44
Par10	$k_{LA_hyd}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}.kg_dry\ basis\ RuC^{-1}$	7.26E-06	10.27
Par11	Ea_{LA_hyd}	$J.mol^{-1}$	5.50E+04	6.87
Par12	$K_{LA}^{\wedge}$	$m^3.mol^{-1}$	5.41E-04	32.58
Par13	$K_{HPA}^{\wedge}$	$m^3.mol^{-1}$	Fixed to zero	-
Par14	$k'_{HPA_cat_Amb}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ Amb^{-1}$	5.32E-04	53.92
Par15	$Ea'_{HPA_cat_Amb}$	$J.mol^{-1}$	Fixed to zero	-
Par16	$k_{HPA_noncat}(T_{Ref})$	s^{-1}	2.67E-07	355.63
Par17	Ea_{HPA_noncat}	$J.mol^{-1}$	5.05E+05	44.16
Par18	$K_{BHP-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par19	$K_{HPA-SO3H}$	$m^3.mol^{-1}$	Fixed to zero	-
Par20	$k_{BHP_RuC}(T_{Ref})$	$s^{-1}.kg_dry\ basis\ RuC^{-1}$	2.23E-05	20.12
Par21	Ea_{BHP_RuC}	$J.mol^{-1}$	Fixed to zero	-

Par22	$k_{HPA_{RuC}}(T_{Ref})$	$s^{-1} \cdot kg_dry\ basis\ RuC^{-1}$	7.18E-05	8.63
Par23	$Ea_{HPA_{RuC}}$	$J.mol^{-1}$	Fixed to zero	-
Par24	$K_{\blacksquare BHP}$	$m^3.mol^{-1}$	Fixed to zero	-
Par25	$K_{\blacksquare HPA}$	$m^3.mol^{-1}$	Fixed to zero	-
Par26	K_c	$m^{9/2}.mol^{-3/2}$	1.43E-04	22.64
Par27	K_{c2}	$m^{9/2}.mol^{-3/2}$	Fixed to zero	-
Par28	$k_{BHP_{diss}}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}$	1.38E-06	15.88
Par29	$Ea_{BHP_{diss}}$	$J.mol^{-1}$	1.07E+05	12.61
Par30	$k_{HPA_{diss}}(T_{Ref})$	$m^3.mol^{-1}.s^{-1}$	3.70E-06	6.96
Par31	$Ea_{HPA_{diss}}$	$J.mol^{-1}$	7.60E+04	7.85

Table S3.10. Normalized parameter covariance matrix for NCLH21.

	Par1	Par2	Par6	Par8	Par9	Par10	Par11	Par12	Par14	Par16	Par17	Par20	Par22	Par26	Par28	Par29	Par30	Par31
Par1	1																	
Par2	0.094	1																
Par6	0.02	0.01	1															
Par8	0.08	0.099	0.006	1.00														
Par9	-0.097	-0.189	-0.007	-0.937	1													
Par10	0.825	-0.065	0.004	0.104	-0.098	1												
Par11	-0.031	0.019	-0.005	-0.08	0.11	0.116	1											
Par12	0.352	-0.49	-0.014	0.102	-0.046	0.588	-0.127	1										
Par14	0	0.008	0	0	-0.001	0.008	0.006	-0.012	1									
Par16	0.027	0.014	0.003	0.026	-0.041	-0.063	-0.378	0.02	-0.002	1								
Par17	-0.026	-0.013	-0.003	-0.024	0.037	0.053	0.335	-0.019	0.002	-0.982	1							
Par20	-0.005	0.035	-0.002	-0.753	0.626	0.029	0.015	-0.006	0	-0.006	0.005	1						
Par22	0.25	0.057	0.006	-0.019	0.012	0.112	0.217	0.017	-0.007	-0.364	0.339	0.014	1					

Par26	0.88	0.312	0.022	0.052	-0.105	0.698	-0.008	-0.003	0.005	0.028	-0.026	0.041	0.271	1				
Par28	0.143	-0.241	-0.035	-0.131	0.154	0.334	0.302	0.26	0.009	-0.14	0.123	-0.112	-0.054	-0.014	1			
Par29	0.118	0.263	-0.008	0.231	-0.35	0.261	0.38	0.057	0.006	-0.15	0.132	0.065	0.083	0.148	0.235	1		
Par30	-0.039	-0.182	-0.012	0.039	0	0.326	0.221	0.277	0.013	-0.227	0.191	-0.009	-0.442	-0.178	0.404	0.167	1	
Par31	0.065	0.065	-0.002	-0.047	0.065	0.138	0.759	-0.043	0.001	-0.528	0.465	0.017	0.45	0.076	0.194	0.314	0.18	1

NCLH1.2.

Table S3.11. Normalized parameter covariance matrix for Model NCLH1.2.

	Par1	Par2	Par3	Par4	Par6	Par8	Par9	Par10	Par11	Par12	Par14	Par16	Par17	Par20	Par22	Par26	Par27	Par28	Par29
Par1	1																		
Par2	0.17	1																	
Par3	0.453	0.069	1																
Par4	0.834	0.16	-0.072	1															
Par6	0.029	0.001	0.021	0.017	1														
Par8	-0.005	0.112	-0.07	0.035	0.005	1													
Par9	-0.028	-0.175	0.049	-0.066	-0.005	-0.932	1												
Par10	0.237	0.061	0.644	-0.127	0.002	-0.013	0.017	1											
Par11	0.013	0.095	0.039	-0.021	-0.006	-0.033	0.07	0.06	1										
Par12	-0.002	0.025	0.114	-0.081	-0.009	0.022	-0.011	0.783	-0.127	1									
Par14	0.006	0.001	0.016	-0.003	0	0	0	0.012	0.005	-0.006	1								
Par16	-0.009	-0.013	-0.033	0.012	0.002	0.012	-0.024	-0.019	-0.311	0.067	-0.002	1							
Par17	0.021	0.009	0.051	-0.008	0	-0.012	0.018	0.015	0.21	-0.06	0.002	-0.948	1						
Par20	0.061	0.074	0.052	0.066	0.002	-0.728	0.593	0.067	-0.014	0.059	0	0.006	-0.003	1					
Par22	0.096	0.029	0.187	0.002	0.005	-0.043	0.046	0.113	0.234	0.099	-0.005	-0.314	0.256	0.013	1				
Par26	0.102	-0.083	-0.123	0.14	-0.044	-0.085	0.084	0.17	0.261	0.166	0.006	-0.085	0.046	-0.123	-0.065	1			
Par27	0.112	0.327	0.088	0.096	-0.011	0.129	-0.245	0.217	0.392	0.13	0.005	-0.118	0.077	0.133	0.08	0.328	1		
Par28	-0.188	-0.006	-0.261	-0.082	-0.019	0.059	-0.031	0.305	0.189	0.377	0.005	-0.165	0.078	0.001	-0.368	0.381	0.187	1	
Par29	0.009	0.053	0.046	-0.027	-0.005	-0.03	0.061	0.14	0.764	0.042	0	-0.442	0.295	-0.004	0.398	0.207	0.31	0.292	1

NCLH2.2.

Table S3.12. Normalized parameter covariance matrix for Model NCLH2.2.

	Par1	Par2	Par6	Par8	Par9	Par10	Par11	Par12	Par14	Par16	Par17	Par20	Par22	Par26	Par28	Par29	Par30	Par31
Par1	1																	
Par2	0.19	1																
Par6	0.026	0.002	1															
Par8	0.113	0.165	0.011	1														
Par9	-0.163	-0.234	-0.011	-0.931	1													
Par10	-0.074	0.025	-0.017	0.035	-0.007	1												
Par11	-0.012	0.081	-0.008	-0.044	0.083	0.04	1											
Par12	-0.062	0.021	-0.013	0.025	-0.009	0.933	-0.139	1										
Par14	-0.002	0	0	0.001	-0.001	0.002	0.005	-0.008	1									
Par16	0.009	-0.007	0.003	0.014	-0.027	0.006	-0.3	0.073	-0.001	1								
Par17	-0.005	0.003	-0.001	-0.011	0.019	-0.026	0.196	-0.066	0.001	-0.946	1							
Par20	-0.02	0.019	-0.001	-0.768	0.635	0.044	-0.001	0.051	-0.001	0.002	-0.002	1						
Par22	-0.002	0.01	0.001	-0.035	0.042	-0.009	0.229	0.079	-0.008	-0.308	0.242	0.01	1					
Par26	0.97	0.209	0.023	0.117	-0.174	-0.102	-0.027	-0.077	-0.002	0.013	-0.007	0.006	0	1				
Par28	0.142	-0.11	-0.045	-0.106	0.109	0.342	0.273	0.193	0.008	-0.089	0.049	-0.106	-0.043	0.088	1			
Par29	0.137	0.356	-0.007	0.211	-0.329	0.207	0.375	0.121	0.003	-0.106	0.064	0.058	0.061	0.167	0.265	1		
Par30	-0.064	0.017	-0.016	0.036	-0.009	0.639	0.202	0.423	0.009	-0.176	0.087	0.015	-0.335	-0.09	0.376	0.214	1	
Par31	-0.024	0.039	-0.008	-0.04	0.074	0.139	0.762	0.03	-0.001	-0.431	0.277	0.007	0.394	-0.037	0.219	0.294	0.315	1

S4. Parity plots for NCLH1.2 and NCLH2.2

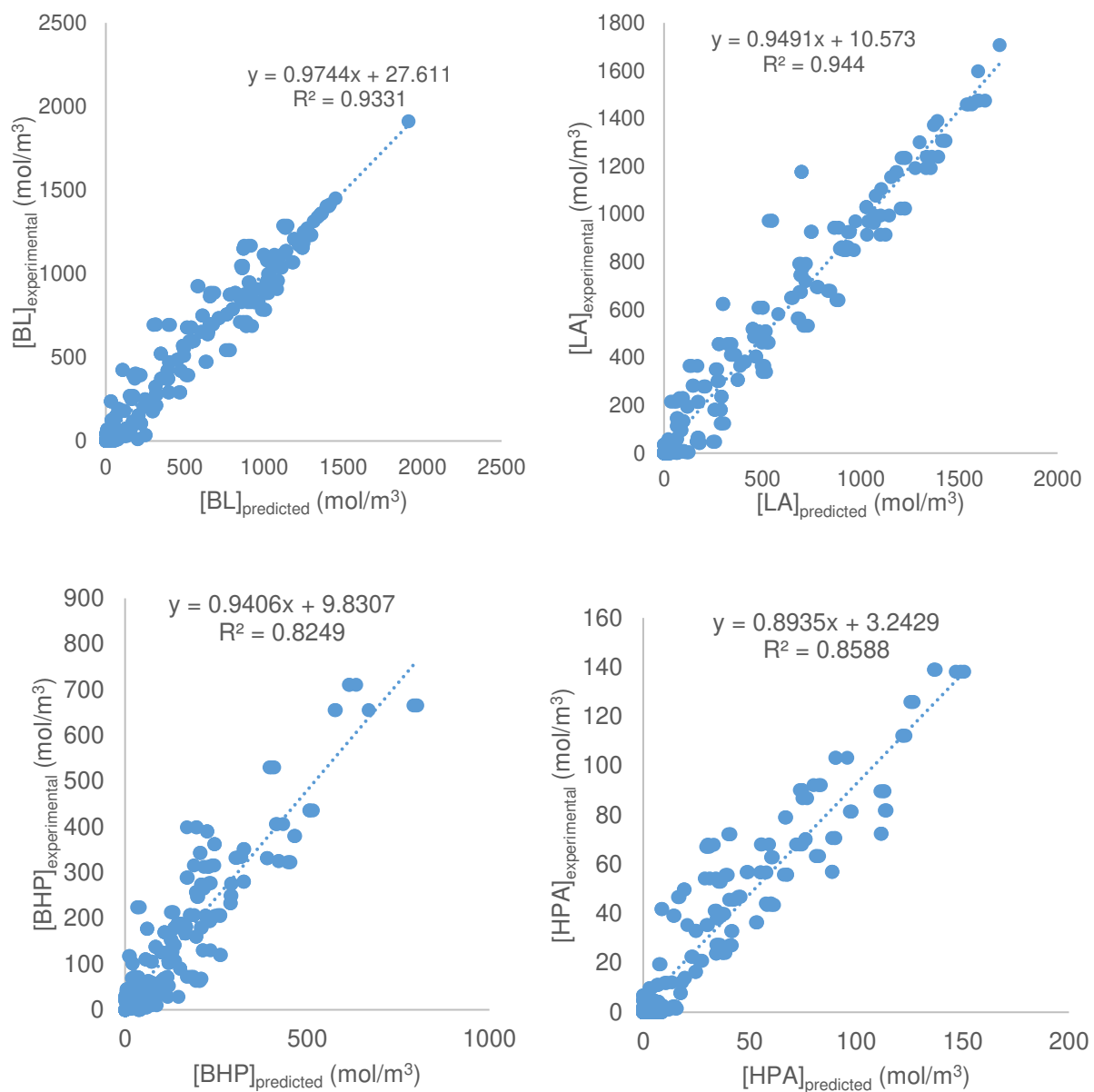


Fig. S4.1. Parity plots for Model NCLH1.2.

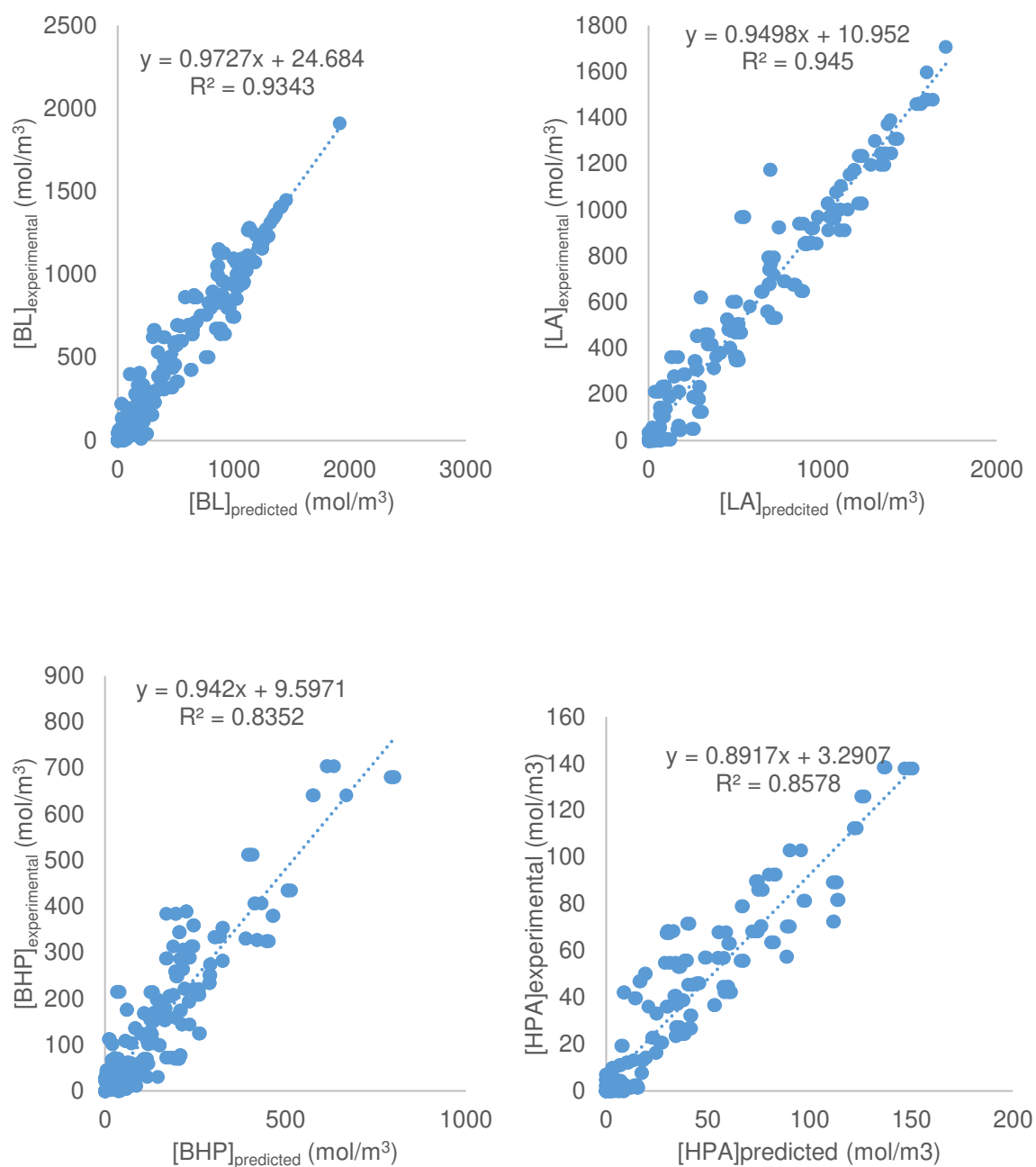
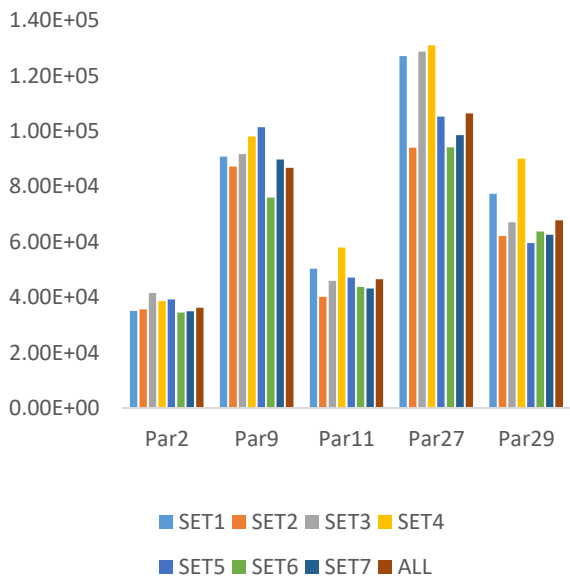
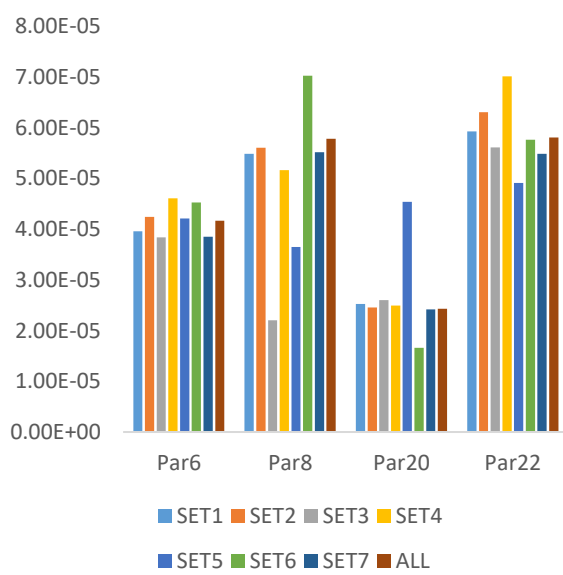
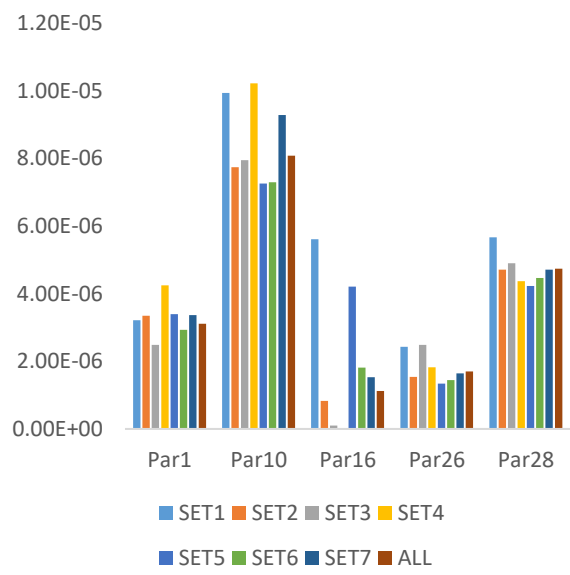
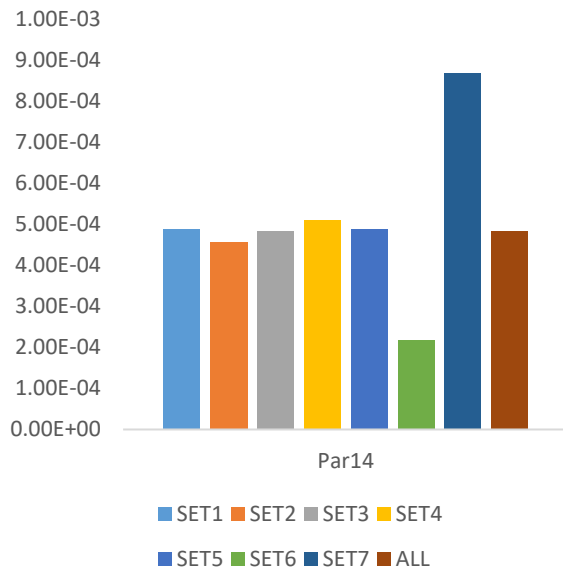


Fig. S4.2. Parity plots for Model NCLH2.2.

S5. Cross validation



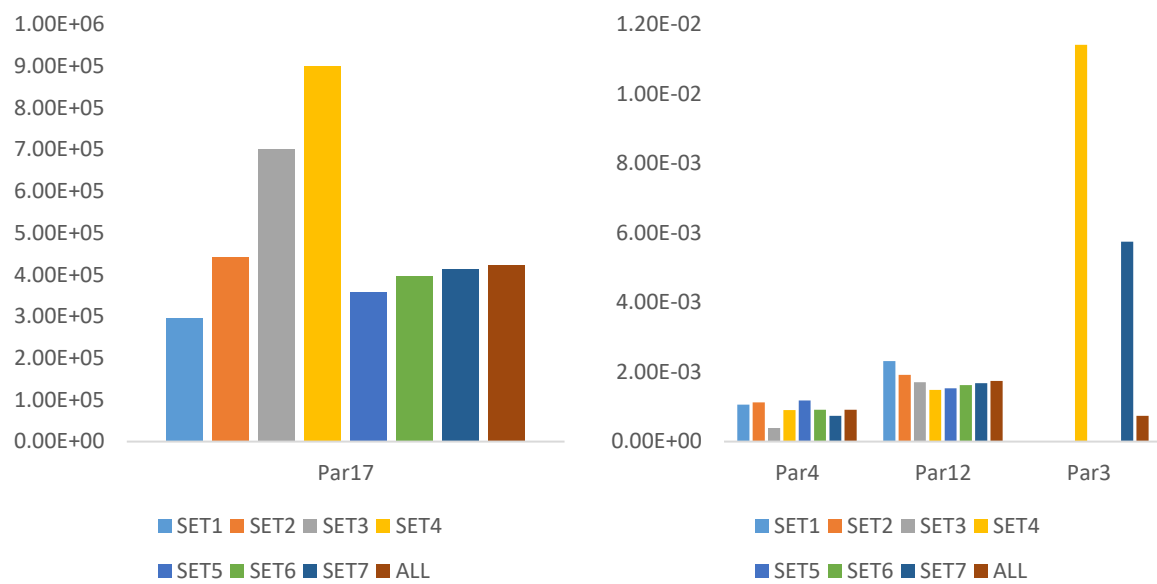


Fig. S5.1. Values of the estimated values for each set and for the whole experiments for NCLH1.2

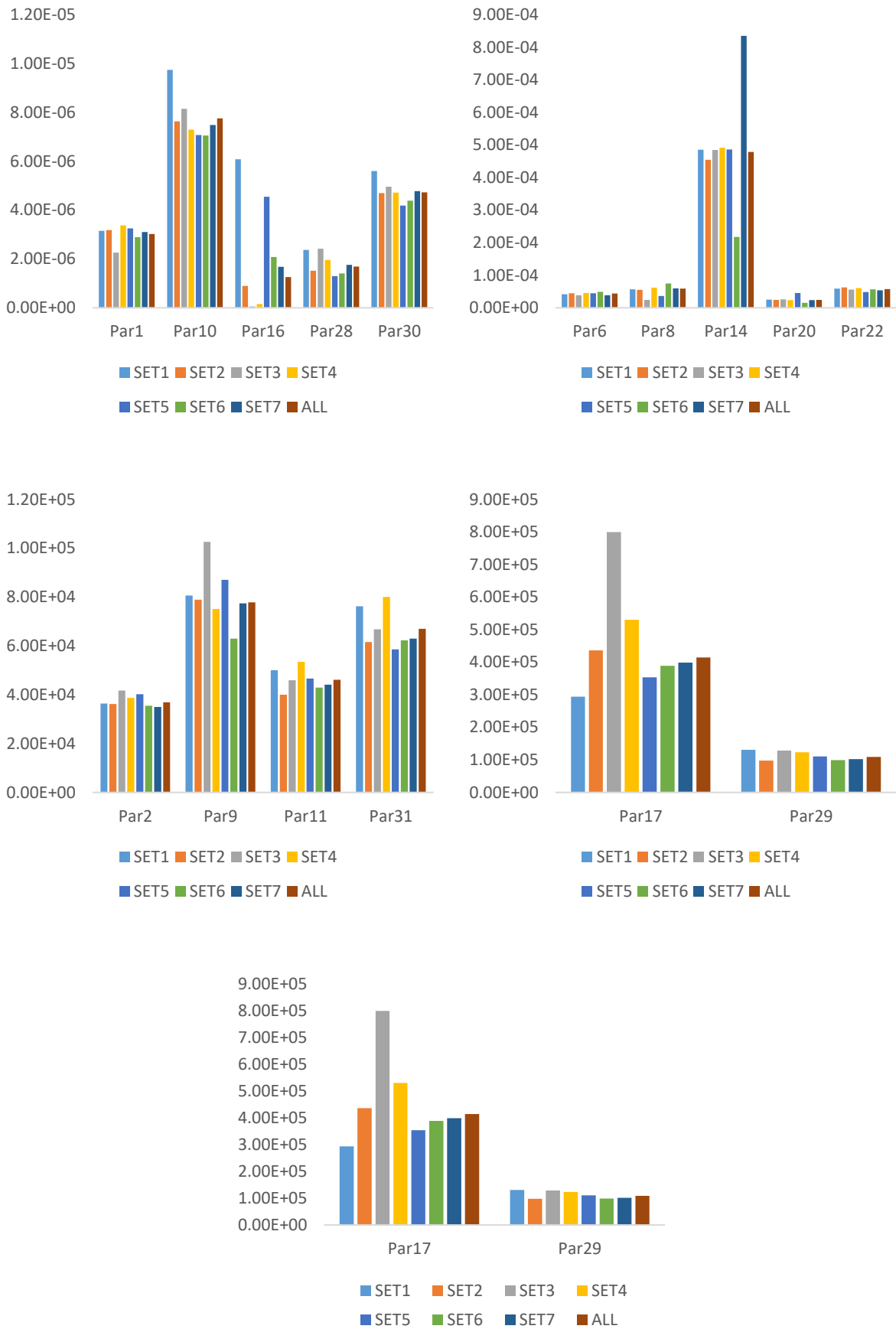


Fig. S5.2. Values of the estimated values for each set and for the whole experiments for NCLH2.2