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Model-based plantwide optimization of large scale lignocellulosic bioethanol plants

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Abstract

Second generation biorefineries transform lignocellulosic biomass into chemicals with higher added value following a conversion mechanism that consists of: pretreatment, enzymatic hydrolysis, fermentation and purification. The objective of this study is to identify the optimal operational point with respect to maximum economic profit of a large scale biorefinery plant using a systematic model-based plantwide optimization methodology. The following key process parameters are identified as decision variables: pretreatment temperature, enzyme dosage in enzymatic hydrolysis, and yeast loading per batch in fermentation. The plant is treated in an integrated manner taking into account the interactions and trade-offs between the conversion steps. A sensitivity and uncertainty analysis follows at the optimal solution considering both model and feed parameters.

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It is found that the optimal point is more sensitive to feedstock composition than to model parameters, and that the optimization supervisory layer as part of a plantwide automation system has the following benefits: (1) increases the economical profit, (2) flattens the objective function allowing a wider range of operation without negative impact on profit, and (3) reduces considerably the uncertainty on profit.

**Keywords:** Second generation bioethanol plant; Nonlinear model-based optimization; Uncertainty and sensitivity analysis; Steam pretreatment; Enzymatic hydrolysis; C5 and C6 co-fermentation.

---

**Nomenclature**

- $\beta_k$: The $\beta$ coefficient in global sensitivity analysis.
- $\delta_k$: Non-dimensional local differential sensitivity measure of cost function $c$ with respect to parameter $\theta_k$.
- $\dot{x}$: Vector of state derivatives used in dynamic modelling.
- $\dot{x}_f$: Vector of state derivatives used in fermentation dynamic model.
- $R_F$: Correlation matrix for fermentation model parameters.
- $R_L$: Correlation matrix for liquefaction model parameters.
- $R_P$: Correlation matrix for pretreatment model parameters.
- $R$: Correlation matrix for the entire integrated process.
- $\sigma_y$: Standard deviation of the objective function.
- $\sigma_{\theta_k}$: Standard deviation of parameter $\theta_k$.
- $\theta_k$: Model parameter $k$. See Table 1 from the supplementary material for a full list of model parameters.
$c(x, u)$ Cost function as a relation of model states $x$ and decision variables $u$.

$C_b$ Feedstock composition in [g/kg].

$c_f$ Cost of fermentation.

$c_o$ Value of cost function at the optimal solution.

$C_{AC_S}$ Acetlys concentration [g/kg].

$C_{CS}$ Cellulose concentration [g/kg].

$c_{eh}$ Cost of enzymatic hydrolysis.

$C_{Eth}$ Concentration of ethanol [g/kg].

$c_{ssk}$ Cost value in steady-state when varying parameter $k$.

$E_{HMF}$ 5-HMF activation energy.

$f(x, u)$ Nonlinear process model of states $x$ and inputs $u$ formulated as equality constraints.

$F_b$ Feedstock flow rate [kg/h].

$F_e$ Enzyme dosage [kg/h].

$F_s$ Steam flow rate [kg/h].

$g(x, u)$ Inequality constraints used as ranges for decision variables.

$h(x_f, u_f)$ Dynamic model for C5-C6 co-fermentation.

$K_2$ Cellulose to glucose reaction constant.

$M_y$ Yeast seed [kg].

$M_{Eth}$ Mass of ethanol [kg].

$P_b$ Feedstock price [unitcost/(kg/h)].
$P_e$ Enzyme price $[\text{unitcost}/(\text{kg}/\text{h})]$.

$P_s$ Steam price $[\text{unitcost}/(\text{kg}/\text{h})]$.

$P_y$ Yeast price $[\text{unitcost}/\text{kg}]$.

$P_{MX}$ Ethanol inhibition on xylose uptake.

$q_{Max_Ac}$ Maximum acetate uptake rate.

$R_B$ Severity factor dependency.

$t_f$ Final time in fermentation $[\text{h}]$.

$T_{tr}$ Thermal reactor temperature $[^\circ\text{C}]$.

$u$ Vector with all decision variables.

$u_f$ Input variables in fermentation.

$x_f$ Process states in fermentation.

$Y_{Cell_G}$ Biomass growth on glucose.

$Y_{Eth_G}$ Ethanol production from glucose uptake.

$Y_{Eth_X}$ Ethanol production from xylose uptake.

$z_i$ Initial guess for the optimization problem.

$z_o$ Optimal solution.

1. Introduction

Second generation lignocellulosic biorefineries reached commercial reality in 2012 [1], and several large scale plants are in operation nowadays including Beta Renewables, Abengoa Bioenergy, GranBio and POET-DSM [2]. Most biorefineries produce bioethanol, but the drop in oil price reduced the demand on the biofuel. However, plant upgrades for chemicals with higher-added values are pursued making biorefineries still competitive in an oil dependent environment [3].
This study deals with optimizing the daily operation of a large scale second
generation biorefinery with a well established conversion route for bioethanol
production. The focus is not a techno-economic assessment of alternative biomass
conversion technologies but rather assumes a plant has already been built and is
in operation.

The latest developments in biorefinery technology show that integrating the
facility with a nearby power plant following the Integrated Biomass Utilization
System (IBUS) \[4\] has a major impact on cost efficiency. E.g. the Inbicon
plant is integrated with Asnæsværket situated in Kalundborg Denmark and they
are both owned by the same company DONG Energy. The symbiosis between
the biorefinery and the power plant allows the exchange of by-products for
consumables, e.g. lignin bio-pellets for steam.

Modeling and simulation are used in this study as enabling technology to
analyze plant performance as basis for an overall optimization. The objective of
the optimization problem is to maximize the plant economical profit, considering
prices for the most important consumables and end products of the process:
biomass, enzymes, yeast and ethanol.

The conversion route from lignocellulosic material to products with higher
added value consists of: pretreatment, enzymatic hydrolysis, fermentation, and
purification \[1,4\]. Lignocellulosic biomass contains cellulose, hemicellulose
(xylan and arabinan), lignin, ash, and other residues \[5\]. The scope of the
pretreatment process is to open the biomatrix, relocate lignin and partially
hydrolyze the hemicellulose such that cellulose would become more accessible
to the downstream process of enzymatic hydrolysis \[4\]. During pretreatment,
inhibitors such as organic acids, furfural, and 5-Hydroxymethylfurfural (5-HMF)
are also created due to sugar degradation. Organic acids change the pH of
medium, but can be automatically neutralized by a pH controller for ensuring
optimal enzymatic conditions \[7\]. Furfural, 5-HMF, and acetate are fermentation
inhibitors \[8\], while the remaining hemicellulose fraction leads to xylooligomers
and xylene formation in the enzymatic hydrolysis process, which strongly inhibit
the enzymatic activity \[9\].
There are trade-offs between the conversion steps. Too little biomass pretreatment would reduce the exposed cellulose to enzymes, and also increases the amount of hemicellulose for enzymatic hydrolysis, which would eventually decrease the glucose yield due to xylose and xylooligomers inhibition. On the other hand, too much biomass pretreatment would increase the amount of fermentation inhibitors leading to a lower ethanol yield.

Most existing studies focus on operational optimization conducting small scale experiments in the laboratory for finding the best pretreatment conditions such that ethanol yield is maximized \[10-13\]. The traditional focus is on one unit at a time (pretreatment versus enzymatic hydrolysis versus fermentation) but the entire process is rarely considered although the biomass conversion steps are inherently dependent and integrated. The single step methods are suboptimal from an economic point of view as they do not focus on overall process economics. Furthermore, in existing studies, the enzymatic hydrolysis and fermentation processes are usually conducted following a fixed recipe, i.e. no correction action or feedback is taken to counteract the effects of inhibitors. For example, one could increase the enzyme dosage when xylooligomers and xylose inhibit glucose production, or adjust the yeast seed in fermentation to compensate for inhibitors.

Therefore the focus of this paper is on systematic methods and tools to facilitate the further process optimization and daily operation of second generation bioethanol plants. The paper shows how overall optimization can be achieved and how sensitivity and uncertainty can be assessed with respect to feedstock composition and kinetic parameters. A Monte Carlo technique with Latin Hypercube Sampling and correlation control is used for the uncertainty analysis following the methodology from \[14, 15\].

This paper is structured as follows: the methods section revises the methodology for building the optimization layer for plantwide operation, along with the theoretical part of the sensitivity and uncertainty analysis. The results and discussion follow where the profit curve, costs, and optimal solutions are presented along with their uncertainty bounds. The paper concludes with a summary of all
important findings.

2. Methods

2.1. Second Generation Bioethanol Plant

Figure 1 illustrates a generic large scale second generation biorefinery concept for bioethanol production. The pretreatment process consists of a continuous thermal reactor and a separation press, which were modeled and analyzed in [16][17]. The thermal reactor is equipped with temperature control for adjusting the reaction temperature $T_{tr}$ [18]. When hemicellulose is hydrolyzed, it produces xylose and arabinose (C5 sugars). After separation, the liquid part containing the C5 sugars is directly pumped into fermentation reactors, bypassing the enzymatic hydrolysis reactors. Cellulose can also be degraded in the pretreatment process, but the produced glucose (C6 sugar) is not lost as it is added to fermentation along with the C5 sugars from the liquid fraction.

![Biorefinery diagram with assumed instrumentation. Pretreatment, enzymatic hydrolysis, and purification are continuous processes, while fermentation occurs in scheduled batch reactors. Feedstock composition is assumed to be known, and can be measured in reality with NIR equipment.](image)

The enzymatic hydrolysis process was thoroughly described and analyzed in [19]. It runs at a high dry matter content in a continuous mode and consists of a series of hydrolysis tanks. The first reactor is described in [20] followed
by conventional continuous stirred tank reactors in order to meet the necessary hydrolysis retention time of 140 h. The tanks are equipped with pH and temperature controllers in order to keep optimal conditions for the enzymatic activity: e.g. $pH = 5$, and $T = 50^\circ C$ [21]. Enzymes are added by a pump from a storage tank. The enzyme dosage $F_e$ can be adjusted accordingly and constitutes a degree of freedom in the optimization problem.

The fermentation process runs in scheduled batch mode in a number of tanks with a maximum holdup of $250 \text{ m}^3$. It is assumed that the size of the reactors has been optimally designed before construction to ensure high titer of ethanol. The fermentation tanks have pH and temperature controllers. Each batch consists of an inoculum phase, the fed-batch phase (which cannot be neglected because it takes 60 h to fill the tank, time when reactions already take place), the batch stage, and an unload step. These stages are illustrated in Figure 2a. In the inoculum phase, 10 t of hydrolyzed fibers rich in glucose are mixed with $M_y$ kilograms of yeast and diluted with water. The amount of yeast addition $M_y$ is one of the operation parameters. The fed-batch phase starts as soon as the inhibitors are removed, after about 10 h. The fermentation tank is filled up to 220 t with a constant feed rate calculated as the sum between the enzymatic hydrolysis outflow rate and the C5 liquid from the pretreatment process. Once the tank is filled, the batch phase begins where the C5 and C6 sugars are slowly depleted. The batch stage has a fixed duration set to 120 h.

A large scale biorefinery has several fermentation reactors running in parallel following a certain scheduling algorithm. Figure 2b shows the scheduling algorithm for 5 reactors such that the overall fermentation inflow and outflow rates have minimum interruptions. This is achieved by aligning in series the fed-batch phases from all tanks, and by synchronizing the unload stages.

The distillation and purification phase separates lignin and water from ethanol. Lignin is recovered as bio-pellets in a evaporation unit as a refinery by-product. The lignin bio-pellets are sent to a nearby power plant where they are co-burnt with coal for steam generation. Lignin bio-pellets are very important for the overall process economics because they contain enough energy
(a) Fermentation process consisting of 4 phases: inoculum (10 h), fed-batch (60 h), batch (120 h), and unload (70 h).

(b) An example of a fermentation process with 5 scheduled reactors. The reactors are scheduled such that the liquefied fiber inflow and ethanol outflow stay constant with minimum interrupts.

Figure 2: Fermentation process: sequential operation and scheduling.
to produce the required steam for ethanol recovery [22]. The exchange of bio-
pellets for steam illustrates the symbiosis between the biorefinery and the nearby
power plant following the Integrated Biomass Utilization System (IBUS) [4].

Bioethanol is the main product, which achieves a high concentration of 99.5 \% with the help of several molecular sieves.

The amount of total solids in each conversion step is measured and controlled. This is a key performance feature that enables constant residence times in the reactors and flattens distillation steam consumption.

One can change either the reaction time by modifying the retention time of each individual process, or adjusting the pretreatment temperature, enzyme dosage, and yeast seed to maximize ethanol yield. Since most second generation plants are new, the focus of this study is to maximize the daily operation efficiency of the plant assuming a constant throughput in the refinery, also defined as Mode I in [23]. This constraint translates to a fixed pretreatment time, i.e. 15 min, a constant enzymatic hydrolysis time of 140 h, and a fermentation time of 190 h per batch for the demonstration scale plant studied here. The degrees of freedom then become: the thermal reactor temperature $T_{tr}$, the enzyme dosage $F_{e}$, and yeast seed $M_{y}$, which are the key process parameters to use in the overall optimization.

For commercial scale plants, once the efficiency has been maximized, the optimization problem can be extended to maximize production or search for a maximum throughput by adding the raw material inflow as a degree of freedom in the optimization problem. Maximizing the throughput, defined as Mode II in [23], is not in focus in this study.

2.2. The Optimization Layer

This study aims to develop an optimization method for maximizing the daily operational efficiency that would be implemented as a supervisory layer at a large scale facility. Figure 3 shows the role of the optimization layer, and how it interacts with the system identification layer, the control system, and the real plant. The model identification layer utilizes real measurements to calibrate
the plant model such that predictions become more accurate. The control layer translates the optimal setpoints into actuator commands to ensure reference tracking for the key process states or variables. For pretreatment, temperature regulation is the key variable. For enzyme dosage, the flow of enzymes is controlled. Yeast seed control requires a mass estimator and control of added yeast. Common practice for a new plant is to design the control system to keep a constant throughput.

The dynamic models used in this study were validated against data that were collected at the Inbicon demonstration scale plant for a throughput of 1 t/h of raw biomass. This flow rate was chosen in order to minimize the impact of pretreatment disturbances on fiber composition. At higher throughputs vertical temperature gradients appear in the thermal reactor that create layers of different biomass composition.

The optimization layer is solved or updated either when the underlying models are recalibrated to fit new acquired data, when feedstock composition changes (e.g. due to different biomass type or variability in feedstock content,
which can be measured either offline or online with NIR equipment), or when prices change (e.g. ethanol price increases, enzymes price decreases, yeast can be grown at a lower price etc.). Model calibration and feedstock composition can change on a daily basis while prices are set through contracts with suppliers and remain constant over a much longer period.

The solution of the optimization problem provides setpoints for the pretreatment temperature, enzyme dosage, and yeast seed. The system constraints are formulated based on validated large scale models for: hydrothermal pretreatment with steam [17], enzymatic hydrolysis [19], and C5 and C6 co-fermentation [24]. The optimization problem is formulated as a constrained nonlinear programming problem (NLP) with the following generic formulation [25]:

$$\max_u c(x, u)$$
$$\text{subject to } f(x, u) = 0$$
$$g(x, u) \leq 0$$

where $u$ is a vector of the decision variables or degrees of freedom, $x$ represents the process variables, and $c(x, u)$ is the nonlinear objective function. $f(x, u) = 0$ and $g(x, u) \leq 0$ are equality and inequality nonlinear constraints as functions of process and decision variables. The solution of optimization problem (1) is found by a nonlinear programming solver (e.g. the optimization toolbox in Matlab contains function $\text{fmincon}$, which finds the minimum of a constrained nonlinear multivariate function).

### 2.3. Mathematical Models

The optimization layer uses a dynamic biorefinery simulator to calculate the stabilized or steady-state outputs for the continuous processes of pretreatment and enzymatic hydrolysis, and the final states at the end of the batch fermentation. The mathematical models are complex and nonlinear. Finding an analytical solution for steady-states might not be feasible. An alternative is to run a sufficiently long simulation until all outputs reach steady state. The fermentation model is a batch process and the outputs are collected after running a dynamic
simulation for 190 h, i.e. the end of the batch phase as shown in Figure 2a, which is a fixed amount of time by design due to setting a constant throughput of material in the biorefinery. The pretreatment and enzymatic hydrolysis models have already been published in [17] and [19]. The fermentation tank is modeled as a continuous stirred tank reactor (CSTR) with reaction kinetics derived from [24, 26]. The equations of all models used to simulate the integrated process are provided in the supplementary material.

The plant simulator is implemented in modules, one for each conversion step, i.e. pretreatment, enzymatic hydrolysis and fermentation. Each module has inputs, internal states, outputs and a set of parameters. The inputs and the outputs are data structures containing information on flow rate (in kg/h), composition (in g/kg) and enthalpy (in kJ/kg). E.g. the pretreatment block has 1 biomass input (i.e. the feedstock), which has a feed rate set to 1000 kg/h with the composition from Table 2, and 1 steam input. The thermal reactor is temperature controlled and the steam valve is adjusted accordingly to reach the desired pretreatment temperature. The pretreatment block has 2 outputs: 1 of fibers with high dry matter and 1 liquid stream rich in C5 sugars. The fibers output from the pretreatment block is connected to the input of the enzymatic hydrolysis block, while the output with C5 sugars is connected to one of the inputs of the fermentation block. The inputs and outputs of each block and their interconnection are represented with arrows in Figure 1.

Table 1 offers a summary of the integrated model complexity. The overall model accounts for 96 kinetic parameters, 580 states, 10 inputs and 25 outputs. The table also offers a split of the modular model based on the conversion step. The high number of states in pretreatment and enzymatic hydrolysis is due to the computational fluid dynamics tools (the convection equation discretized in space) used for modeling the thermal reactor and the first enzymatic hydrolysis tank. Nominal values for kinetic and feed parameters are given in Table 1 in the supplementary material.
Table 1: Model summary: number of parameters, states, inputs and outputs. Half of the pretreatment outputs (the pretreated fibers) are directed to the enzymatic hydrolysis process, while the other half (the C5 liquid) is connected to fermentation. The outputs from the enzymatic hydrolysis are connected to fermentation.

<table>
<thead>
<tr>
<th>Model</th>
<th>Parameters</th>
<th>States</th>
<th>Inputs</th>
<th>Outputs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pretreatment</td>
<td>17</td>
<td>298</td>
<td>10</td>
<td>36</td>
</tr>
<tr>
<td>Enzymatic Hydrolysis</td>
<td>46</td>
<td>257</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>Fermentation (1 tank)</td>
<td>33</td>
<td>25</td>
<td>37</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>96</td>
<td>580</td>
<td>10</td>
<td>25</td>
</tr>
</tbody>
</table>

2.4. Plantwide Optimization Methodology

The methodology steps for finding the optimal operational point of a plant are extended from [25]:

1. Select the objective or cost function;
2. Identify the decision variables;
3. Formulate process model constraints and set bounds for decision variables;
4. Formulate and solve the NLP optimization problem;
5. Sensitivity and uncertainty analysis of the optimal solution.

The optimal solution is analyzed from a sensitivity point of view using similar tools as in [14], [15], [17]. Mathematical models that describe complex systems are often over-parametrized. The sensitivity analysis quantifies the relation between the cost function and model parameters when the system runs at the optimal point. The aim is to rank all model parameters by their significance with respect to the profit value at the nominal operational point. Also a subset of relevant parameters can be extracted for calculating the uncertainty bounds.

A non-dimensional measure of local sensitivity suitable for steady-state signals is the differential sensitivity measure defined in [27] and [28]:

\[
\delta_k = \frac{\partial c_k}{\partial \theta_k} \frac{\theta_k}{c_{ss_k}}
\]
where $\partial c_k / \partial \theta_k$ is the variation in profit with respect to a model parameter, and is calculated based on finite differences. $\theta_k$ is the $k^{th}$ parameter, and $c_{ss_k}$ is the profit or the value of the cost function in steady-state. All model parameters are sorted in descending order with respect to $\delta_k$, and a subset is created with all parameters that have $\delta_k$ above a threshold. The reduced subset of model parameters reduces model complexity and is then used in the uncertainty analysis.

The propagation of uncertainty is analyzed with a Monte Carlo procedure as described in [14]:

1. Define input uncertainty;
2. Parameter sampling;
3. Monte Carlo simulations;
4. Output uncertainty.

The input uncertainty is defined with standard deviations and correlation matrices obtained from previous studies. Dealing with many parameters implies a large number of combinations of parameter values with high correlation between them. In order to reduce the number of parameter samples but preserve statistical meaning, a Latin Hypercube Sampling (LHS) technique with correlation control is utilized [29]. LHS generates less samples of parameters but is made statistically plausible with the help of a distribution function, standard deviation, and correlation matrix. For each set of samples, a simulation is then run and the output is collected. After all Monte-Carlo simulations are performed, enough output information is obtained to statistically compute the median and the 5th-95th percentile confidence intervals.

A global sensitivity analysis supplements the Monte-Carlo simulations results. The statistical distribution of both the model parameters and the biomass inflow composition are taken into account. The methodology includes two steps [15, 30]: (1) fitting a linear model from each variable of interest (model parameter and biomass composition) to the optimization objective function, which is the biorefinery profit calculated as in Equation (6) and generated through Monte Carlo simulations; (2) calculating the standardized regression coefficients as
a global sensitivity measure, which indicate how much of the output standard deviation is explained by the standard deviation in each input variable.

Step (1) finds coefficients $a$ and $b_k$ from Equation (3) using a least squares method. The linear model fits the objective function while the input parameters are contained in vector $\theta$, i.e. both model parameters $\theta_R$ and feedstock composition $C_b$.

$$y_{reg} = a + \sum_k b_k \theta_k$$

(3)

Step (2) uses Equation (4) to calculate the $\beta$ coefficients, which reflects how much of the output variation is explained by parameters uncertainty:

$$\beta_k = \frac{\sigma_\theta \theta_k}{\sigma_y} b_k$$

(4)

3. Results and Discussion

This section first presents values of model and feed parameters. The optimization problem is then formulated, it is shown how a solution is found and the properties of the results are discussed. A sensitivity analysis with respect to model kinetic parameters is then presented. A subset of sensitive parameters is identified, and Monte Carlo simulation is employed to quantify the uncertainty of the optimal solution. The costs and profit curves are also computed with uncertainty bounds. Furthermore, a global sensitivity analysis is made in order to identify bottlenecks in model predictions with respect to feed and model parameters. Uncertainty is then embedded in the formulation of a stochastic optimization problem. As a final result, dynamic simulations show the refinery operation at the optimal point.

3.1. Model Initialization

Table 2 shows the feed parameters, i.e. raw biomass inflow rate, composition and initial temperature. The inflow rate is set to 1000 kg/h, the throughput of a demonstration scale plant. The biomass composition resembles wheat straw with an initial dry matter of 89% [6].
Table 2: Biorefinery inputs: inflow rate, raw biomass composition, and initial temperature.

<table>
<thead>
<tr>
<th>Description</th>
<th>Value</th>
<th>Unit</th>
<th>% of dry matter</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Inflow rate</td>
<td>1000</td>
<td>kg/h</td>
<td></td>
</tr>
<tr>
<td>2 Cellulose</td>
<td>360</td>
<td>g/kg</td>
<td>40.45</td>
</tr>
<tr>
<td>3 Xylan</td>
<td>187</td>
<td>g/kg</td>
<td>21.01</td>
</tr>
<tr>
<td>4 Arabinan</td>
<td>23</td>
<td>g/kg</td>
<td>2.58</td>
</tr>
<tr>
<td>5 Lignin</td>
<td>200</td>
<td>g/kg</td>
<td>22.47</td>
</tr>
<tr>
<td>6 Acetyls</td>
<td>44</td>
<td>g/kg</td>
<td>4.94</td>
</tr>
<tr>
<td>7 Ash</td>
<td>26</td>
<td>g/kg</td>
<td>2.92</td>
</tr>
<tr>
<td>8 Water</td>
<td>110</td>
<td>g/kg</td>
<td>-</td>
</tr>
<tr>
<td>9 Other</td>
<td>50</td>
<td>g/kg</td>
<td>5.63</td>
</tr>
<tr>
<td>10 Temperature</td>
<td>15</td>
<td>ºC</td>
<td></td>
</tr>
</tbody>
</table>

Table 1 from the supplementary material indicates the values with units for all 96 model parameters. The table is split into pretreatment, enzymatic hydrolysis and fermentation. The model parameter values are taken from [17] (pretreatment), [19] (enzymatic hydrolysis) and [24] (C5 and C6 co-fermentation).

3.2. The Optimization Problem

The application of the optimization methodology is now highlighted for the case study:

1. Select the objective or cost function:

   The cost function from this study represents the profit for one fermentation batch defined as the difference between ethanol revenue and operating costs related to biomass, steam, enzymes, and yeast:

   \[
   c(M_{Eth}, F_b, F_s, F_c, M_y) = M_{Eth}(t_f)P_{Eth} - (F_bP_b + F_sP_s + F_cP_c + M_yP_y)
   \]  

   (5)

   Ethanol revenue is calculated as \( M_{Eth}(t_f)P_{Eth} \), i.e. mass of ethanol in kg at the end of the batch phase \( t_f \) times its price per kilogram \( P_{Eth} \). \( M_y \) is
the mass of yeast added to the fermentation tank in the inoculum phase. The operating costs are defined as flow rate or mass of utility times its price. The refinery consists of two continuous processes, i.e. pretreatment and enzymatic hydrolysis, and a batch process, i.e. fermentation. The weights \( P_b \) (cost of biomass), \( P_s \) (cost of steam), and \( P_e \) (cost of enzymes) are related to the continuous processes, i.e. pretreatment and liquefaction, and are measured in \( \text{unitcost}/(\text{kg/h}) \). \( P_y \) (cost of yeast) is measured in \( \text{unitcost}/\text{kg} \). The overall measuring unit of the cost function becomes the \( \text{unitcost} \), which can represent any currency.

Capital costs do not enter the objective function because they are not functions of the decision variables but rather a fixed amount. Capital costs diminish the potential profit by a constant and should be taken into consideration to give a full meaning to the profit solution from the optimization problem but they do not change the optimal solution.

The feedstock flow rate or refinery throughput \( F_b \) is kept constant. Product \( F_b P_b \) becomes a constant and its derivative with respect to any of the decision variables is 0. Therefore it can be dropped from the cost function. Distillation costs are neglected because the second generation biorefinery is considered to be integrated with a power plant exchanging lignin biopellets for steam.

The cost function then becomes:

\[
c(M_{Eth}, F_e, M_y) = M_{Eth}(t_f)P_{Eth} - (F_e P_e + M_y P_y)
\]  

(6)

Table 3 shows the weight values used in this optimization study. The weights from Table 3 are not fixed values but rather used as an example for the method. Changing the weights would naturally lead to a different solution, which is a matter of plant operation managers priority. The values depend on procurement contracts of feedstock and enzymes, and the relative importance may change.

2. Identify the decision variables:

The outcome of the pretreatment process is sensitive to the thermal reactor
Table 3: Cost function weights (prices).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P_\text{Eth}$</td>
<td>Ethanol</td>
<td>5 unitcost/kg</td>
</tr>
<tr>
<td>$P_e$</td>
<td>Enzymes</td>
<td>25 unitcost/(kg/h)</td>
</tr>
<tr>
<td>$P_y$</td>
<td>Yeast</td>
<td>50 unitcost/kg</td>
</tr>
</tbody>
</table>

Temperature and retention time [11]. When a constant throughput is required, the retention time is constant. The thermal reactor temperature then becomes the only degree of freedom in pretreatment.

The key parameters in enzymatic hydrolysis are: pH, temperature, and concentration of enzymes. The enzymatic activity is a function of pH and temperature, which resemble Gaussian curves with single peaks at pH of 5 units and temperature 50 °C [19]. Any deviations from the optimal point would reduce the enzymatic efficiency. Control loops keep the pH and temperature close to optimality [7] and it is not indicated to vary these variables. However, the concentration of enzymes can be adjusted by changing the inflow rate of enzymes $F_e$ and constitutes the only degree of freedom in enzymatic hydrolysis for the optimization problem.

The efficiency of the fermentation process is a function of pH, temperature, and yeast seed. The optimal pH level of the GMO yeast is relatively close to that of the enzymes, i.e. 5.5 units. The optimal fermentation temperature is 35 °C, which is different from the enzymatic optimal temperature. Controllers keep the pH and temperature conditions at the GMO yeast optimal levels throughout the entire fermentation process. The only degree of freedom considered in fermentation for the optimization problem is the yeast seed $M_y$ in the inoculum phase.

In summary, the decision variables are: the pretreatment temperature $T_{tr}$ defined as the set-point for the thermal reactor temperature controller, the inflow rate of enzymes $F_e$ expressed in kg/h, and the yeast seed $M_y$ in kg.
as a set-point for the amount of yeast used to start the inoculum phase:

\[ u = [T_{tr} \ F_e \ M_y]^T \]  \hspace{1cm} (7)

3. Process model constraints, and bounds for decision variables:

The dynamic integrated models for pretreatment and enzymatic hydrolysis are detailed in the supplementary material, and are formulated as:

\[ \dot{x} = f(x, u) \]  \hspace{1cm} (8)

where \( f(x, u) \) is a nonlinear function of states \( x \) and inputs \( u \). The steady states are then found as the solution of \( \dot{x} = 0 \):

\[ 0 = f(x, u) \]  \hspace{1cm} (9)

Due to the model complexity and the nonlinear nature of \( f(x, u) \), an analytical solution to (9) cannot be easily found. As an alternative, the steady states are calculated by running a sufficiently long simulation till all states stabilize.

The dynamic model for fermentation is described by:

\[ \dot{x}_f = h(x_f, u_f) \]  \hspace{1cm} (10)

where \( h(x_f, u_f) \) represents a nonlinear complex model of states \( x_f \) and inputs \( u_f \). The final states at time 190 h are found by integrating the model numerically (dynamic simulation):

\[ x_f(t_f) = \int_0^{t_f} h(x_f, u_f)dt \]  \hspace{1cm} (11)

where \( t_f = 190 \) h, i.e. the end of the batch phase.

The decision variables are bounded as follows:

\[
\begin{align*}
150 & \leq T_{tr} \leq 210 ^\circ C \\
10 & \leq F_e \leq 1000 \text{ kg/h} \\
10 & \leq M_y \leq 1000 \text{ kg}
\end{align*}
\]  \hspace{1cm} (12)

which allows a wide range of operation for searching the optimal point.
4. Formulate and solve the overall NLP optimization problem:

\[
\max_{T_{tr}, F_e, M_y} \quad M_{Eth}(t_f) P_{Eth} - (F_e P_e + M_y P_y)
\]

subject to

\[
0 = f(x(t), u(t))
\]

\[
\dot{x}_f = h(x_f, u_f)
\]

\[
150 \leq T_{tr} \leq 210 \, ^\circ C
\]

\[
10 \leq F_e \leq 1000 \, \text{kg/h}
\]

\[
10 \leq M_y \leq 1000 \, \text{kg}
\]

(13)

The previous optimization problem is solved numerically by a constrained minimization solver (e.g. \texttt{fmincon} from Matlab) set to use a sequential quadratic programming algorithm (SQP) with scaled objective and constraints. The initial solution guess is picked to be feasible, and set to:

\[
z_i = \begin{bmatrix}
T_{tr} \\
F_e \\
M_y 
\end{bmatrix} = \begin{bmatrix}
170 \, ^\circ C \\
150 \, \text{kg/h} \\
200 \, \text{kg} 
\end{bmatrix}
\]

(14)

The initial states of all models are set to 0. The solver takes approximately 3 min to converge on a computer equipped with an Intel i7-5600U CPU. The following optimal point is reached:

\[
z_o = \begin{bmatrix}
T_{tr} \\
F_e \\
M_y 
\end{bmatrix} = \begin{bmatrix}
172 \, ^\circ C \\
110 \, \text{kg/h} \\
142 \, \text{kg} 
\end{bmatrix}
\]

(15)

The thermal reactor temperature should be set to 172\(^\circ\)C, the enzyme dosage is of about 110 kg/h, and the yeast seed is of 142 kg. This optimal set point gives a profit of:

\[
c_o = 7.6714 \times 10^4 \, \text{unitprofit}
\]

(16)

disregarding raw biomass, distillation and capital costs.

In order to gain process insight and to observe how pretreatment conditions affect the downstream processes, an iteration is created through pretreatment
temperatures between 160°C to 180°C with a step of 1°C. Each pretreatment temperature changes the fibers and C5 liquid composition. The enzymatic hydrolysis and fermentation processes are then optimized in an integrated manner for each pretreatment temperature:

\[
\max_{F_e, M_y} M_{Eth}P_{Eth} - (F_e P_e + M_y P_y)
\]

subject to

\[
0 = f(x(t), u(t))
\]
\[
\dot{x}_f = h(x_f, u_f)
\]

\(10 \leq F_e \leq 1000 \text{ kg/h}\)
\(10 \leq M_y \leq 1000 \text{ kg}\)  

(17)

In this way the pretreatment, liquefaction and fermentation costs, as well as refinery profit can be observed with respect to pretreatment conditions. The same methodology can be applied even if there are recycles between fermentation and liquefaction because these two processes are analyzed in an integrated manner in optimization problem (17). Algorithm 1 shows how to calculate the curves for profit, costs, and optimal solution as functions of pretreatment temperature. 

\(z\) is the optimal solution returned by the optimization problem solver, \(c_{eh}, c_f\) are the enzymatic hydrolysis, and fermentation costs, respectively. \(c\) is the value of the cost function or the profit. Algorithm 1 is used later in Algorithm 2 to determine the uncertainty bounds on profit, costs, and setpoints with respect to the pretreatment temperature.

3.3. Sensitivity Analysis of Profit Value at the Optimal Point

Figure 4 illustrates the sensitivity analysis of the profit curve with respect to all model parameters calculated as in Equation (2) at the optimal solution (15). A sensitivity threshold separates important parameters from unimportant ones. The sensitivity threshold is typically based on the context of application. E.g. 1% as a threshold states that any factors not contributing with 1% to the output variance are unimportant while the ones above that are important for further considerations (e.g. parameter estimation, optimization). For this particular application a value of 4.6% is chosen. A lower value would increase
Algorithm 1: Calculate optimal costs and profit with respect to pretreatment temperature given a fixed set of model parameters $\theta$ and feedstock composition $C_b$.

1: function $[z, c_{eh}, c_f, c] = \text{costs}(\theta, C_b)$
2:     Set a range of pretreatment temperatures $T_{tr} \leftarrow 160 \degree C : 1 \degree C : 180 \degree C$
3:     Set initial solution guess to $z_0 \leftarrow [100 \text{ kg/h} \quad 80 \text{ kg}]^T$
4:     for Each temperature in $T_{tr}$ do
5:         Run pretreatment process at temperature $T_{tr_i}$ and obtain composition of pretreated fibers and C5 liquid.
6:         $z_i \leftarrow$ Solution of optimization problem \((17)\) given the pretreated fibers composition and C5 liquid from previous step as inputs. Use as initial guess the solution from previous iteration $z_{i-1}$.
7:     Calculate mass of ethanol at final fermentation time: $M_{Eth_i} \leftarrow M(t_f)$.
8:     Calculate liquefaction cost: $c_{eh_i} \leftarrow F_{e_i} P_e$.
9:     Calculate fermentation cost: $c_f \leftarrow M_{y_i} P_y$.
10:    Calculate revenue: $r_i \leftarrow M_{Eth_i} P_{Eth}$.
11:    Calculate profit $c_i \leftarrow r_i - (c_{eh_i} + c_f)$.
12: end for
13: end function
the subset of important parameters but with little or negligible impact on the solution of the optimization problem.

Figure 4: Sensitivity measure $\delta_k$ of profit value with respect to model parameters. The model parameters set is reduced to 22 significant parameters out of 96.

The most sensitive parameter is $E_F$, i.e. the furfural formation activation energy. Furfural is a strong fermentation inhibitor produced during pretreatment, and ethanol yield is directly affected by the amount of furfural. The next sensitive parameter is $Y_{EthG}$, a yield parameter indicating the amount of ethanol in g produced per 1 g of glucose. $E_G$ or glucose activation energy follows indicating that cellulose degradation in pretreatment impacts the ethanol yield. Three more fermentation parameters with similar sensitivity follow, i.e. maximum acetate uptake parameter $q_{MaxAc}$, cell biomass yield on glucose $Y_{CellG}$, and ethanol yield on xylose $Y_{EthX}$. 5-HMF production during pretreatment has a relatively high sensitivity too as it influences both the glucose yield, by degrading it further, and also by inhibiting ethanol production in fermentation.

The first sensitive enzymatic hydrolysis parameter is $R_B$, i.e. the severity dependence of the enzymatic activity. $R_B$ shows the importance of biomatrix opening from the pretreatment process as a structural breakdown of the fiber, which affects cellulose accessibility for enzymes. Other important liquefaction parameters are $K_2$ and $K_7$, which indicate glucose production rate and enzyme deactivation in time.

The placement of pretreatment parameters such as $E_F$, $E_G$, $E_H$, and $E_{Ac}$
among the sensitive parameters shows how important pretreatment conditions are for downstream. Fermentation parameters are also numerous among the sensitive parameters. Fermentation with enhanced GMO yeast for bioethanol production is a key process in the biorefinery together with steam pretreatment. Liquefaction parameters have a lower importance because the overall hydrolysis time is long enough to compensate for any parameter uncertainties. The liquefaction process has a pure hydrolysis phase of 140 h followed by fermentation where enzymes are still active continuing cellulose degradation (simultaneous saccharification and fermentation).

The sensitivity threshold is set at 0.046, which reduces the parameters count to 22 out of 96 showing the importance of the sensitivity analysis in reducing model complexity. These parameters are then used in the following uncertainty analysis.

3.4. Uncertainty Analysis of Costs, Profit and Optimal Solution

The standard deviation and correlation matrix for pretreatment and enzymatic hydrolysis parameters is obtained from [17] and [18], respectively. Table 4 displays a list of all sensitive parameters with their assumed uncertainty as a normal distribution of a mean value and standard deviation. Regarding the model parameters for fermentation, there is no published real data that could be used for uncertainty characterization. Therefore, this study follows techniques from experimental design for system identification, and generates measurements through simulation, i.e. glucose, xylose and ethanol levels, which can be obtained in reality from sample based HPLC readings. Normally distributed measurement noise with 5% standard deviation is added, and then a parameter estimation procedure runs on the simulated data for estimating the standard deviation and correlation matrix for these parameters. The results are included in Table 4, where the standard deviation for fermentation parameters is obtained from simulated data as previously described.

The correlation matrix for the sensitive parameters presented in Table 4 is
Table 4: List with all sensitive parameters with their assumed distribution for uncertainty analysis. 

\(N(m, \sigma)\) stands for normally distributed with mean \(m\) and standard deviation \(\sigma\).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Distribution</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pretreatment</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(E_F)</td>
<td>N(327, 255, 285)</td>
<td>[17]</td>
</tr>
<tr>
<td>(E_G)</td>
<td>N(335, 616, 249)</td>
<td>[17]</td>
</tr>
<tr>
<td>(E_H)</td>
<td>N(299, 999, 2639)</td>
<td>[17]</td>
</tr>
<tr>
<td>(R_B)</td>
<td>N(2.915, 0.048)</td>
<td>This study</td>
</tr>
<tr>
<td>(E_{Ac})</td>
<td>N(242, 693, 174)</td>
<td>[17]</td>
</tr>
<tr>
<td>(E_{Xo})</td>
<td>N(298, 010, 98)</td>
<td>[17]</td>
</tr>
<tr>
<td><strong>Enzymatic Hydrolysis</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(K_2)</td>
<td>N(0.0065, 0.0001)</td>
<td>[18, 28]</td>
</tr>
<tr>
<td>(K_7)</td>
<td>N(2.5 \times 10^{-7}, 0.041 \times 10^{-7})</td>
<td>[18, 28]</td>
</tr>
<tr>
<td>(E_{MC}^E)</td>
<td>N(0.016, 0.00026)</td>
<td>[18, 28]</td>
</tr>
<tr>
<td>(I_{G_2})</td>
<td>N(0.067, 0.0011)</td>
<td>[18, 28]</td>
</tr>
<tr>
<td><strong>Fermentation</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Y_{EthG})</td>
<td>N(0.47, 6.0744 \times 10^{-3})</td>
<td>This study</td>
</tr>
<tr>
<td>(q_{Ac:Max})</td>
<td>N(1.23 \times 10^{-5}, 8.2395 \times 10^{-7})</td>
<td>This study</td>
</tr>
<tr>
<td>(Y_{EthX})</td>
<td>N(0.4, 1.3127 \times 10^{-2})</td>
<td>This study</td>
</tr>
<tr>
<td>(Y_{CellG})</td>
<td>N(0.115, 4.3453 \times 10^{-3})</td>
<td>This study</td>
</tr>
<tr>
<td>(P_{MPX})</td>
<td>N(100.2, 1.2852)</td>
<td>This study</td>
</tr>
<tr>
<td>(q_{MaxX})</td>
<td>N(0.8 \times 10^{-3}, 1.6115 \times 10^{-5})</td>
<td>This study</td>
</tr>
<tr>
<td>(q_{MaxG})</td>
<td>N(0.3 \times 10^{-3}, 6.7299 \times 10^{-6})</td>
<td>This study</td>
</tr>
<tr>
<td>(K_{AcS})</td>
<td>N(2.5, 5.7082 \times 10^{-2})</td>
<td>This study</td>
</tr>
<tr>
<td>(K_{I_{AcX}})</td>
<td>N(0.2, 8.9583 \times 10^{-3})</td>
<td>This study</td>
</tr>
<tr>
<td>(\gamma_X)</td>
<td>N(0.608, 4.0872 \times 10^{-2})</td>
<td>This study</td>
</tr>
<tr>
<td>(m_G)</td>
<td>N(2.6944 \times 10^{-5}, 8.6390 \times 10^{-7})</td>
<td>This study</td>
</tr>
<tr>
<td>(K_{I_{AcG}})</td>
<td>N(2.74, 4.2621 \times 10^{-2})</td>
<td>This study</td>
</tr>
</tbody>
</table>
built as a block diagonal matrix with the following structure:

\[
R = \begin{bmatrix}
R_P & 0 & 0 \\
0 & R_L & 0 \\
0 & 0 & R_F
\end{bmatrix}
\]  

where \( R_P, R_L \) and \( R_F \) are full matrices, and represent the correlation of the sensitive pretreatment, liquefaction and fermentation parameters grouped by refinery step. It is assumed that the refinery steps are not correlated with each other, e.g. pretreatment parameters are not correlated with liquefaction and fermentation parameters. Therefore, \( R \) has a block diagonal shape. The numerical values for \( R_P, R_L \) and \( R_F \) are included in the supplementary material. \( R_P \) and \( R_L \) are obtained from [17] and [28], respectively. The correlation matrix for fermentation parameters \( R_F \) is obtained in this study from simulated data.

Latin Hypercube Sampling (LHS) with correlation control can then be performed for all model parameters. Feedstock composition is sampled assuming uniform distribution with 5% variation in composition. \( N = 200 \) LHS samples are extracted for model and feed parameters. The Monte Carlo simulations are performed by running Algorithm 1 for each set of parameters. The simulation outputs are collected, and the 5th, 50th and 95th percentiles are then calculated for profit, costs, and optimal solution.

The uncertainty analysis is carried out separately for feed, and then for model parameters in order to observe the contribution of each source of uncertainty. A last analysis combines the samples of feed and model parameters to find the overall effect of the uncertainty sources on the outputs.

The entire sensitivity and uncertainty analysis for combined model and feed parameters is summarized in Algorithm 2. To separate model and feed parameters step 11 from Algorithm 2 is modified either by keeping \( C_b \) or \( \theta \) constant.

The results of the uncertainty analysis are displayed in Figure 5 and commented below:

- The profit curve is drawn in Figure 5a, which is used to identify the optimal...
Algorithm 2 Sensitivity and uncertainty analysis with combined model and feed parameters.

1: Initialize model parameters $\theta$ and feed composition $C_b$.
2: Optimal deterministic solution: $[T_{tr}, F_e, M_y] \leftarrow (13)$.
3: Sensitivity analysis of the cost function in $[T_{tr}, F_e, M_y]$: calculate $\delta_k$ as in equation (2).
4: Rank all parameters with respect to $\delta_k$.
5: Select a subset $\theta_R$ such that $\theta_R$ is above a threshold.
6: Set standard deviations and correlation matrices for $\theta_R$.
7: $\theta \leftarrow$ LHS of $\theta_R$ with correlation control to generate N sets of model parameters.
8: Set bounds for feedstock composition.
9: $C_b \leftarrow$ Uniform LHS for feedstock composition to generate N sets of compositions.
10: for Each set of model and feed parameters do
11: $[z_i, c_{ehi}, c_{fi}, c_i] = \text{COSTS}(\theta_i, C_{bi})$
12: end for
13: Calculate the 5th, median and 95th percentile for profit, costs, and optimal solution.
(a) Potential profit per source of uncertainty for one fermentation batch. Comparison between traditional operation with a fixed recipe and optimized scenario.

(b) Refinery costs separated into liquefaction and fermentation. The costs are illustrated with respect to each source of uncertainty: feed, model, and combined parameters.

(c) Solution of the optimization problem with respect to each source of uncertainty as a function of pretreatment conditions.

Figure 5: Optimal costs, potential profit, and solution of the optimization problem with respect to pretreatment temperatures, and for each source of uncertainty: feed composition, model parameters and combined.
operational point. The traditional biorefinery operation is to follow a fixed recipe with little adjustments to pretreatment conditions. This traditional recipe is most often found by offline experiments on decoupled refinery steps that do not take into account the interactions between the conversion stages and utilities prices. The results show that the traditional operation is sub-optimal from an economic point of view. In contrast, the optimization layer is capable of adapting to pretreatment temperatures and finds the optimal operation by considering the integrated process. The optimized operation is superior to a traditional recipe with a higher median profit curve at any pretreatment temperature.

At low temperatures, most of the uncertainty is due to model parameters, but it shifts after $165 \, ^\circ \text{C}$ when feed uncertainty becomes dominant. The traditional operation is highly affected by feed uncertainty, while the optimized operation has a reduced uncertainty on the profit curve.

Another important result is that the optimized profit curve is flatter than the traditional curve allowing a wider range of operation with little impact on profit value. The optimal operational point can be picked as the maximum point on the median profit curve, and lies between $171 \, ^\circ \text{C}$ to $176 \, ^\circ \text{C}$. The optimal refinery operates at around 18% higher profit than a traditional plant without an optimization layer.

- Figure 5b shows the refinery costs split into liquefaction and fermentation as a function of pretreatment conditions. From left to right, the uncertainty analysis is carried with respect to separate feed and model parameters (left and center plots), and combined parameters (right plot). The pretreatment costs are only due to the steam used in the thermal reactor. The biorefinery is considered to be integrated with a local power plant, possibly owned by the same company following the IBUS principle [4]. Such a design lowers the cost of steam significantly and can be neglected. A higher pretreatment temperature demands more steam but the overall increase in cost for modifying the temperature from $160 \, ^\circ \text{C}$ to $180 \, ^\circ \text{C}$ is negligible.
compared to enzymatic hydrolysis or fermentation.

Liquefaction costs are high at low temperature because: (1) the biomatrix is not sufficiently opened to expose the whole cellulose, and (2) there is a large amount of unhydrolyzed hemicellulose, which leads to a high production of xylooligomers and xylose that inhibit the enzymatic hydrolysis further. In order to compensate for these negative effects, both the enzyme and yeast dosage are increased. The liquefaction costs decrease as the pretreatment temperature increases, which makes sense as the biomatrix opens significantly to expose cellulose, and also hemicellulose is partially removed from the enzymatic hydrolysis process.

Fermentation costs have the shape of a convex curve due to: (1) at low pretreatment temperatures a higher yeast seed could contribute to a faster digestion of sugars, which enhances the saccharification process from fermentation by reducing the C5 sugars inhibition leading to a higher ethanol yield; (2) at high temperatures the amount of inhibitors negatively affect fermentation but more yeast could compensate for the inhibitory effects of the pretreatment degradation products.

Feed uncertainty is rather constant through the entire temperature range. Uncertainty due to model parameters is high at low temperatures where the biomatrix opening highly affects the cost range. After 165 °C the model uncertainty is significantly reduced becoming lower than the feed. The combined model and feed uncertainty indicate high uncertainty at low temperatures when the pretreatment is insufficient.

- Figure 5c illustrates the optimal solution as a function of pretreatment temperatures. Enzyme dosage is expressed in kg/h, while yeast seed is given in kg. Uncertainty is higher at lower temperatures when the biomass is not sufficiently pretreated, and remains relatively constant once the biomatrix opens. Also, feed uncertainty has a higher impact than model parameters after 165 °C.

Increasing pretreatment temperatures is beneficial for enzymatic hydrolysis
as it lowers the necessary enzyme dosage, but is negative for fermentation as the amount of inhibitors rises with temperature. Also, around the optimal operational point, uncertainty due to feed parameters dominates that of model parameters.

In reality there are several factors that can degrade the performance of the optimization layer, and should be accounted for in real implementation. The feed rate in this study case was set to a low value, which does not allow inhibitors accumulation in the fermentation tank. However, at higher feed rates inhibitors accumulation becomes a bottleneck, which can be counteracted by calculating an optimal feed rate profile [31]. This study also disregards the temperature dependence of the yeast performance. In reality the enzymatic hydrolysis and fermentation processes run at different optimal temperatures. The solution is to calculate a temperature profile for finding the best trade-off between the two processes [32, 33].

3.5. Global Sensitivity Analysis at the Optimal Point

The $\beta$ coefficients are displayed in Table 5. The profit curve is mostly sensitive to feedstock composition, i.e. cellulose fiber and acetyls content $C_{CS}$ and $C_{ACS}$. 88% of profit curve variability is explained by cellulose content changes, while 31% is due to acetyls. A positive $\beta$ coefficient indicates that an increase in cellulose content would determine an increase in profit, while a negative value decreases the profit. Acetyls have a negative impact on profit because they are responsible for acid release in pretreatment and enzymatic hydrolysis, which inhibit both the liquefaction and fermentation processes. Yield performance to convert glucose $Y_{EthG}$ follows with 27%. It is also noted that unlike the local sensitivity analysis results presented above, only about 6 parameters are found to be globally influential on the profit curve (considering a threshold value for $\beta^2 >= 1\%$). This means that the profit curve is primarily influenced by feedstock composition.

Measuring the inflow composition accurately would help to plan the production process in advance and reduce the uncertainty on the profit curve.
Table 5: Sensitivity measure at the optimal point \( z_o \). The linear model determination coefficient \( R^2 = 0.99 \).

<table>
<thead>
<tr>
<th>( \theta )</th>
<th>Description</th>
<th>( \beta )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>( C_{CS} )</td>
<td>Cellulose concentration in feedstock</td>
</tr>
<tr>
<td>2</td>
<td>( C_{ACS} )</td>
<td>Acetyls concentration in feedstock</td>
</tr>
<tr>
<td>3</td>
<td>( Y_{EthG} )</td>
<td>Ethanol production from glucose</td>
</tr>
<tr>
<td>4</td>
<td>( q_{MaxAc} )</td>
<td>Maximum acetate uptake rate</td>
</tr>
<tr>
<td>5</td>
<td>( Y_{CellG} )</td>
<td>Biomass growth on glucose</td>
</tr>
<tr>
<td>6</td>
<td>( Y_{EthX} )</td>
<td>Ethanol production from xylose</td>
</tr>
<tr>
<td>7</td>
<td>( P_{MPX} )</td>
<td>Ethanol inhibition on xylose uptake</td>
</tr>
<tr>
<td>8</td>
<td>( E_H )</td>
<td>5-HMF activation energy</td>
</tr>
<tr>
<td>9</td>
<td>( R_B )</td>
<td>Severity factor dependency</td>
</tr>
<tr>
<td>10</td>
<td>( K_2 )</td>
<td>Cellulose to glucose reaction constant</td>
</tr>
</tbody>
</table>

3.6. Stochastic Optimization Solution

Finding the optimal point by running the process through a wide range of pretreatment temperatures requires a long computational time, and is not feasible in an industrial application. A better way is to embed the feed and model parameters uncertainty into the objective function, and pick the mean cost value:

\[
\max_{T_{tr},F_e,M_y} \frac{1}{N} \sum_{i=1}^{N} \left[ M_{Eth}(t_f)P_{Eth} - (F_eP_e + M_yP_y) \right]
\]

subject to

\[
0 = f(x(t), u(t))
\]

\[
\dot{x}_f = h(x_f, u_f)
\]

\[
150 \leq T_{tr} \leq 210 \, ^\circ C
\]

\[
10 \leq F_e \leq 1000 \, \text{kg/h}
\]

\[
10 \leq M_y \leq 1000 \, \text{kg}
\]

This stochastic optimization is done using \( N = 200 \) as the number of random samples generated through LHS with correlation control. For each sample a simulation is run and the profit is assessed. After performing all \( N \) simulations the mean value of the profit is calculated, which becomes the cost function. The
following optimal solution and cost value are found:

\[ z_{so} = \begin{bmatrix} T_{tr} \\ F_c \\ M_y \end{bmatrix} = \begin{bmatrix} 171.5 \degree C \\ 113 \text{ kg/h} \\ 147 \text{ kg} \end{bmatrix} \quad c_{so} = 7.6020 \times 10^4 \text{ unit profit} \quad (20) \]

where \( z_{so} \) is the optimal solution in the stochastic optimization case, and \( c_{so} \) is the value of the cost function. Solution (20) is relatively close to the one found in the deterministic case from (15). A slightly lower profit value is found due to the nonlinear nature of the process. The model and feed parameters are assumed to be normally distributed but when propagating through a nonlinear process the outputs change their distribution type including the mean value.

3.7. Deterministic Simulations at the Optimal Point

As a final result, a deterministic simulation is run corresponding to the optimal operational point from solution (15). The pretreatment and the enzymatic hydrolysis are continuous processes and the steady state values at the optimal point are shown in Table 6. The pretreated fibers are rich in cellulose and have a dry matter content of about 35% as suggested by [34] for an efficient liquefaction process. Most solubles were separated from the fibers in the pretreatment process before liquefaction. The remaining hemicellulose continues to be degraded to sugars in the enzymatic hydrolysis tanks. When the level of C5 sugars increase, they strongly inhibit glucose production and a part of cellulosic fibers remain in solid state. This is why the liquefied fibers still contain cellulose before fermentation, i.e. 50 g/kg, approximately 30% of the initial cellulose content. The remaining cellulose continues conversion to glucose in the fermentation tank where enzymes are still active.

Figure 6 shows the fermentation batch process at the optimal point. The top plot illustrates C6 and C5 sugars depletion, ethanol production, and biomass growth. The bottom plot displays remaining cellulose and xylan conversion during simultaneous saccharification and fermentation. In the inoculum phase (first 10 h) the yeast concentration is high but as the fed-batch phase starts,
Table 6: Steady states for pretreatment and enzymatic hydrolysis at the optimal point.

<table>
<thead>
<tr>
<th>Flow/Mass</th>
<th>Pretreated fibers</th>
<th>C5 liquid</th>
<th>Liquefied fibers</th>
<th>Fermentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Content</td>
<td>2316 kg/h</td>
<td>628 kg/h</td>
<td>2487 kg/h</td>
<td>220 t</td>
</tr>
<tr>
<td>Composition</td>
<td>1000 g/kg</td>
<td>1000 g/kg</td>
<td>1000 g/kg</td>
<td>1000 g/kg</td>
</tr>
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biomass concentration is diluted in liquefied fibers from the enzymatic hydrolysis and C5 liquid from the pretreatment process. Ethanol production has several stages: (1) formation on glucose consumption till around 100 h, (2) production based on xylose consumption till 170 h, (3) as xylose is depleted, its inhibition on enzymatic hydrolysis disappears and glucose production from simultaneous saccharification and fermentation is accelerated in the last 20 h.

Figure 6: Simultaneous saccharification and C5-C6 co-fermentation.
4. Conclusions

This paper has presented a study on economical optimization of a large scale second generation biorefinery in a simulated environment. The proposed optimization method advises the operators of the plant at what temperature to pretreat the biomass, how many enzymes to add in the enzymatic hydrolysis tanks, and what the yeast seed should be in fermentation such that economical profit is kept high. Adjustments on these key figures need to be made on a daily basis in operation due to biomass composition and type variability, changes in consumables prices, and also due to any other disturbances that can reduce ethanol yield.

The optimization procedure is based on steady-state models (pretreatment and enzymatic hydrolysis), and dynamic fermentation model. It was found that increasing pretreatment temperature is positive for the performance of the enzymatic hydrolysis while negative for ethanol yield. Uncertainties in kinetics of pretreatment, liquefaction and fermentation were found to be negligible on the economic objective function around the optimal operational point. The main source of uncertainty was found to be the inflow feed composition.

The optimization layer reduced the uncertainty and flattened the profit curve allowing a wider range of operation with higher profit making the plant operation more robust to disturbances. The overall improvement of the optimization layer was found to be 18% higher than the traditional plant operation.

In addition to these specific results, the paper has contributed by suggesting a generic method for biorefinery operation optimization and with quantification of sensitivity and uncertainty on earnings.

References


[34] H. Jørgensen, J. Vibe-Pedersen, J. Larsen, C. Felby, Liquefaction of lignocellulose at high solids concentrations, Biotechnology and Bioengineering 96 (2007) 862–70.