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Dynamic optimisation of the aroma production in brewing

fermentation

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Abstract

Several key compounds for the final beer flavour (higher alcohols, esters, vicinal diketones) are produced during

the alcoholic fermentation phase. The paper demonstrates the possibility of obtaining various desired final aroma

profiles and reducing the total process time using dynamic optimisation of three control variables: temperature,

top pressure and initial yeast concentration in the fermentation tank. The optimisation is based on a sequential

quadratic programming algorithm, on a dynamic model of the alcoholic fermentation and on an aroma

- production model. The robustness of the optimal control profile with respect to model uncertainty is discussed.
-

Keywords

Beer, aroma compounds, constrained non-linear optimal control, model uncertainty

Introduction

 The alcoholic fermentation is an important stage in the beer production process. During this phase, fermentable sugars present in the brewing wort are transformed to ethanol and several aroma compounds important for the final beer flavour are produced. The contribution of the alcoholic fermentation phase to the final beer flavour depends on the wort composition, on the yeast strain and on the operating conditions. Industrial operating conditions for most existing beer brands were determined empirically and are confidential. The aim of this work is to demonstrate the possibility of optimising the fermentation temperature profile, the top pressure profile and the initial yeast concentration based on organoleptic and economic criteria. The organoleptic criterion takes into account target concentrations of the following compounds believed to be important for the final beer flavour [1]: two higher alcohols (isoamyl alcohol and phenyl ethanol), three esters (ethyl acetate, ethyl hexanoate and isoamyl acetate) and one vicinal diketone (diacetyl). The economic criterion is based on process time minimisation.

 Previous work on the optimisation of the brewing process was concerned with the computation of the temperature profile alone. Gee and Ramirez [2] used Pontryaghin's minimum principle for the temperature profile optimisation which insured a minimum fermentation time and a maximum ethanol production. Not surprisingly, they found that the fermentation temperature take its maximum possible value at any time. This purely economic optimum could be anticipated from the existing knowledge of the alcoholic fermentation process but is not used in practice because of its undesired effect on the aroma composition. Andres-Toro *et al*. [3] used an optimisation technique based on genetic algorithms and introduced the idea of aroma targets (for ethyl acetate and diacetyl) in addition to fermentation time minimisation. They obtained a non-trivial temperature profile where high temperatures were still favoured, however.

 The present contribution is an extension and a generalisation of the results of Titica *et al*. [4]. Dynamic optimisation is used to compute flexible control profiles instead of two-stage controls considered previously. Limitations of the process model that made it unsuitable for simulation in time-varying operating conditions were removed. The possibility of reducing the fermentation time substantially without altering the final aroma profile was demonstrated for a beer previously obtained in constant operating conditions. The feasibility (using a single yeast strain and wort composition) of four commercial beer aroma profiles is discussed and the limitations of the aroma control using the three considered operating conditions alone are pointed out. Finally, the robustness of the final aroma concentrations and of the optimal operating conditions with respect to unavoidable modelling errors is analysed.

Materials and methods

Experimental

 The alcoholic fermentation and the aroma production models were built and validated on data coming from nine laboratory scale experiments. Experiments were carried out in a 15 L, 0.5 m high, stainless steel bioreactor (LSL Biolafitte, France) filled with 12 L of wort, under gentle agitation (100 rpm). Preliminary experiments showed that mechanical agitation was needed to compensate the absence of the natural agitation that occurs in large scale 54 brewing (10 m high tanks or higher) due to $CO₂$ release. The lager wort and the industrial yeast strain, *Saccharomyces cerevisiae* var*. uvarum*, were provided by the Institut Français de Brasserie et Malterie (IFBM, 56 France). The conditions of the experimental runs R1 to R4 and R6 to R9 were selected according to a $2³$ experimental design. The three factors were the fermentation temperature (10 and 16°C), the top pressure (50 and 800 mbar) and the initial yeast concentration (5 and 20 million cells/mL). The run R5 was performed in intermediate operating conditions (13°C, 450 mbar and 10 million cells/mL). The concentrations of ethanol, isoamyl alcohol, phenyl ethanol, ethyl acetate, ethyl hexanoate and isoamyl acetate were determined by gas chromatography [5] coupled with mass spectrometry in the case of diacetyl [6]. The evolved carbon dioxide was recorded with a gas meter (Schlumberger, France), with a resolution better than 0.5% of the total amount of gas produced in each experiment. The measurements describing the alcoholic 64 fermentation (ethanol, wort density, CO₂ production, refractive index and fermentable sugar concentration) were reconciled using well-established stoichiometric relationships [7]. The yeast cell concentration was determined with a particle counter (Coulter Z1, Coultronics, France). Three counts were performed at 3 and 3.5 µm and the

logarithmic average of the six counts was taken.

Alcoholic fermentation model

- The optimisation of the operating conditions with respect to the specified criterion is an iterative process. In each iteration candidate solutions are tested using a process model. The process model used in this work consisted in an alcoholic fermentation model coupled with an aroma production model.
- The alcoholic fermentation model is a modified version of a previously reported one [8]. The main modification
- 73 concerns the description of the CO₂ transfer between the wort and the headspace. In [8] it was assumed that $CO₂$
- was dissolved in the wort until saturation and released afterwards. This assumption was reasonable for constant
- operating conditions, but seems questionable in the dynamic optimisation context when variable temperature and
- 76 top pressure profiles are usually obtained. The stated assumption was replaced by a rigorous mass balance of the
- 77 CO₂ in the wort and in the headspace and a mass balance of the air in the headspace. Solubility of the air (N₂ and $78 \qquad O_2$) in the wort was found negligible as far as the mass balance was concerned.
- 79 The alcoholic fermentation can be described equivalently by the fermentable sugar consumption, ethanol
- 80 production, wort density decrease or CO_2 production [7]. The CO_2 was selected because it is the most convenient
- 81 variable to be measured on-line [9[\]\[10\].](#page-13-0) The model was constructed by analogy with classical microbial growth
- 82 kinetics with substrate limitation and product inhibition:

83
$$
\frac{dC_p(t)}{dt} = v(\theta(t), C_d(t)) \cdot \frac{S(t)}{K_s + S(t)} \cdot \frac{1}{1 + (E(t)/K_E)^2} \cdot (C_p(t) + K_X X_0)
$$
 (1)

$$
84 \t C_p(0) = 0 \t (2)
$$

$$
85 \t E(t) = Y_{E/C} C_p(t) \t (3)
$$

$$
S(t) = S(0) - Y_{S/C} C_p(t)
$$
\n(4)

The rate of the alcoholic fermentation was described by the rate of CO_2 production dC_p/dt . Simultaneously, 88 ethanol (*E*) is produced and fermentable sugars (*S*) are consumed, with constant yields ($Y_{E/C}$ and $Y_{S/C}$ 89 respectively). The initial fermentation rate, when $C_p = 0$, is proportional to the initial yeast concentration X_0 .

90 The "specific" fermentation rate ν was expressed as [11]:

91
$$
v(\theta, C_d) = K_v \exp(K_{v\theta}(\theta - \theta_{typ}) - K_{vC}(C_d - C_{d,typ}))
$$
 (5)

92 For small temperature variations $(\pm 3K)$ compared to the typical absolute fermentation temperature (286K), the 93 given relationship is a close approximation of the Arrhenius law. A similar dependence was assumed for the 94 dissolved CO_2 (C_d). The "typical" temperature and dissolved CO_2 values are those of the central point of the 95 experimental design (run R5) and correspond to usual values for lager beer making. For modelling purposes, it 96 was assumed that the produced CO_2 (C_p) is transferred into the solution, because of the large contact area 97 between the yeast cells and the wort. The mass transfer between the solution (C_d) and the headspace (C_h) , 98 depends on the partial CO₂ pressure in the headspace (p_c) and on the tank geometry and agitation through a 99 kinetic constant (7) :

100
$$
\frac{dC_d(t)}{dt} = \frac{dC_p(t)}{dt} - \frac{1}{\tau} [C_d(t) - C_{sat}(\theta(t), p_c(t))]
$$
(6)

$$
101 \t C_d(0) = 0 \t (7)
$$

102 C_{sat} is the equilibrium CO₂ concentration in the wort at a given temperature (θ) and at a given partial headspace 103 pressure (*pc*). It was determined from tables provided by Institut Français de Brasserie et Malterie and 104 approximated by the following empirical formula:

105
$$
C_{sat}(\theta, p_c) = K_C p_c \frac{\theta + \theta_{abs}}{\theta_{abs}} e^{-K_{C\theta}\theta}
$$
 (8)

106 The evolution of the CO_2 concentration in the headspace (C_h) is given by :

107
$$
\frac{dC_h(t)}{dt} = \frac{\gamma}{\tau} \Big[C_d(t) - C_{sat}(\theta(t), p_c(t)) \Big] - \frac{\gamma C_h(t)}{C_h(t) + C_a(t)} \Phi_g(t)
$$
\n(9)

$$
108 \t C_h(0) = 0 \t (10)
$$

109 where γ is the ratio of the wort volume to the headspace volume, C_a is the air concentration and $\Phi_{\rm g}$ the total gas

110 outflow rate $(CO_2 + air)$. The CO_2 outflow rate is proportional to the CO_2 mass fraction in the headspace. The air

111 concentration in the headspace diminishes with a rate proportional to the air mass fraction:

112
$$
\frac{dC_a(t)}{dt} = -\frac{\gamma C_a(t)}{C_h(t) + C_a(t)} \Phi_g(t)
$$
 (11)

$$
113 \t C_a(0) = \frac{p_{\text{atm}} M_a}{R(\theta(0) + \theta_{\text{abs}})}\tag{12}
$$

114 The partial CO2 (*pc*) and air (*pa*) pressures were calculated using the perfect gas law. The technologically 115 important variable is the total pressure (*p*):

116
$$
p_c = \frac{C_h}{M_c} R(\theta + \theta_{abs})
$$
 (13)

117
$$
p_a = \frac{C_a}{M_a} R(\theta + \theta_{abs})
$$
 (14)

$$
118 \qquad p = p_c + p_a \tag{15}
$$

119 where *R* is the perfect gas constant, M_c is the CO₂ molar mass and M_a is the equivalent molar mass of the air.

120 The parameter values of the alcoholic fermentation model are reported in Table 1. The differences with 121 previously reported values [8][12] are due to the way in which the dissolved carbon dioxide was estimated.

122 *Aroma production model*

 A mathematical model for some aroma compounds produced during the alcoholic fermentation, and considered important for the final beer flavour, was reported previously [11][12]. The same model was used for optimal control, with the following modifications: (i) In the new version, the operating conditions affect the yields versus CO₂ in a multiplicative, rather than additive way. (ii) The two-phase yield for the ester production was replaced by a single-phase, monotonically increasing yield. (iii) The model parameters were identified based on the whole pool of experiments, rather than separately for each experiment. This allowed a statistical analysis of the reliability of the parameter estimates. (iv) Parameters not significantly different from zero were removed from 130 the model, i.e. set to exactly zero.

 Higher alcohols. Two higher alcohols were considered in this work, based on their organoleptic thresholds in beer [1]: the isoamyl alcohol (*IAL*) and the phenyl ethanol (*PHL*). Their production rate was related to the 133 alcoholic fermentation rate, with yields $(Y_{i/C})$ depending on the operating conditions: temperature (θ) , dissolved $CO_2(C_d)$ and initial yeast concentration in the wort (X_0) :

135
$$
\frac{dA_i(t)}{dt} = Y_{i/C}(\theta(t), C_d(t), X_0) \frac{dC_p(t)}{dt}
$$
 $i \in \{IAL, PHL\}$ (16)

136
$$
A_i(0) = 0
$$
 (17)

137
\n
$$
Y_{i/C}(\theta, C_d, X_0) = W_{i,1} \exp[W_{i,2}(\theta - \theta_{typ}) + W_{i,3}(C_d - C_{d,typ}) + W_{i,4}(X_0 - X_{0,app}) + W_{i,5}(\theta - \theta_{typ})(C_d - C_{d,typ}) + W_{i,6}(\theta - \theta_{typ})(X_0 - X_{0,app}) + W_{i,7}(C_d - C_{d,typ})(X_0 - X_{0,app}) + W_{i,8}(\theta - \theta_{app})(C_d - C_{d,app})(X_0 - X_{0,app})]
$$
\n(18)

- 138 **Esters**. The three selected esters were the ethyl acetate (*ETA*), the ethyl hexanoate (*ETX*) and the isoamyl acetate
- 139 (*IAA*). Their production rate was related to the alcoholic fermentation rate, as in the case of higher alcohols, but
- 140 the yield increased with fermentation progress:

141
$$
\frac{dA_i(t)}{dt} = Y_{i/C}(\theta(t), C_d(t), X_0) C_p(t) \frac{dC_p(t)}{dt} \qquad i \in \{ETA, ETX, IAA\}
$$
 (19)

$$
142 \t A_i(0) = 0 \t (20)
$$

143 The effect of the operating conditions on the yields $(Y_{i}/(\theta, p, X_0))$ had the same mathematical expression as in the 144 case of the higher alcohols (Equation 18).

 Vicinal diketones. The two important diketones in beer are the diacetyl and the pentanedione. In the considered experiments, the pentanedione concentration was always lower than the organoleptic threshold [12] and hence only the diacetyl model was considered for the optimal control. A detailed analysis of this model was given previously [12]. The diacetyl was simultaneously produced and reduced during the alcoholic fermentation:

149
$$
\frac{dA_{DL}}{dt}(t) = Y_{DL}/c(t) \frac{dC_p(t)}{dt} - W_{DL,1} \exp[W_{DL,2}(\theta - \theta_{typ})]
$$
 (21)

$$
150 \t A_{D1A}(0) = 0 \t (22)
$$

151 The production yield (*Y_{DIA/C}*) asymptotically decreased to zero. An empirical relationship describing this 152 behaviour was established:

153
$$
\frac{dY_{DLA/C}(t)}{dt} = -W_{DLA,3} \frac{dC_p(t)}{dt} Y_{DLA/C}(t)
$$
 (23)

$$
154 \t YDIA/C(0) = WDIA,4
$$
 (24)

 Statistical significance of the model coefficients. The significance test was based on the so-called Mallows statistic, which establishes a trade-off between the fit to experimental data and the model complexity, i.e. number of adjustable parameters [14]. This criterion says that a parameter should be retained in the model only if it reduces the residual variance sufficiently compared to the estimated measurement variance. For each aroma 159 compound (*i*), the Mallows statistic (I_i) was minimised over the various combinations of n_i nonzero parameters:

160
$$
I_i = \frac{1}{\sigma_i^2} \sum_k \left(A_i(t_k) - a_i(t_k) \right)^2 + 2n_i \qquad i \in \{ETA, \quad ETX, \quad IAA, \quad IAL, \quad PHL, \quad DIA\} (25)
$$

Here $a_i(t_k)$ are the measured concentrations and $A_i(t_k)$ are the concentrations predicted by the model with n_i 162 nonzero parameters. The measurement variance σ_i^2 was estimated as the residual variance of the complete 163 model, with all nonzero parameters.

164 The numeric values of the coefficients appearing in the aroma production model, together with their confidence 165 limits, are reported in the Table 2.

166 *Optimal control problem*

167 The optimal control of the fermentation process consisted in the selection of the operating conditions that 168 minimised an overall criterion *Q*, which reflected the desired product quality and plant operation mode. 169 Additional requirements were introduced via a set of constraints that admissible solutions must satisfy. The

-
- 170 operating conditions considered in this work are the wort temperature θ , controlled by the cooling rate Φ_{θ} , the

 top pressure *p*, controlled by the gas outflow rate ^Φ*g*, the initial yeast concentration in the wort *X*0, and the total 172 **fermentation time** t_f **. The optimal control problem is stated as:**

173
$$
\{\mathbf{r}_f, \boldsymbol{\varPhi}_{\theta}(t)_{t\in[0,t_f]}, \boldsymbol{\varPhi}_{g}(t)_{t\in[0,t_f]}, X_{0}\} = \arg\min Q
$$
 (26)

 The "arg min" symbol means the "the values which minimise". The optimisation criterion and the constraints are detailed below.

Optimisation criterion

 The considered problem is a multiobjective optimisation: approaching the five aroma targets as close as possible, reducing the final diacetyl concentration and the total processing time, as well as smoothing out the temperature and the top pressure profiles. The overall optimisation criterion is a weighted sum of the nine partial criteria. The weights specify the desired trade-offs between possibly conflicting objectives. In order to simplify the selection of the weights, all partial criteria were scaled by physically meaningful quantities. Thus, the partial criteria and the associated weights are dimensionless quantities of order of unity.

 Approaching the aroma targets. The final concentrations of the five aroma compounds, believed to be important for the final beer flavour, have to be as close as possible to their respective targets:

185
$$
Q_i = W_i \frac{\left(A_i(t_f) - A_{i,trg}\right)^2}{A_{i,tol}^2}
$$
, $i \in \{ETA, ETX, IAA, IAL, PHL\}$ (27)

Here Q_i is the partial criterion associated to the aroma compound *i*, W_i is the associated weight, $A_i(t_i)$ is the aroma

187 concentration at the end of the fermentation, $A_{i,trg}$ is the target concentration, and $A_{i,tol}$ is the accepted tolerance. Since there is no point in trying to approach the target closer than either the model accuracy or the perceived

difference, the tolerance was selected as:

190
$$
A_{i,tol}^2 = \sigma_i^2 + A_{i,org}^2
$$
 (28)

191 where σ_i is the model prediction accuracy (Table 2) and $A_{i,org}$ is the organoleptically significant difference in the concentration of the compound *i* taken as 10% of the organoleptic threshold of the considered compound in beer [\[1\].](#page-12-0) An alternative way of expressing the tolerance would be using the largest of the two values σ_i and $A_{i,org}$. The tolerance provides a natural scaling of the associated optimisation criterion.

 Minimising the diacetyl concentration. Diacetyl is an undesired compound in the finished beer. No target was defined for this compound because the lowest possible value is desired in practice (ideally zero). Minimisation of the diacetyl concentration at the end of the main fermentation is an economically important goal, since it may result in the reduction of the cost of the subsequent processing (dedicated "diacetyl rest" and/or lagering phases). Mathematically, this goal was expressed as:

200
$$
Q_{DIA} = W_{DIA} \frac{A_{DIA}(t_f)}{A_{DIA, typ}}
$$
 (29)

201 where $A_{DIA}(t_f)$ is the diacetyl concentration at the end of the alcoholic fermentation and $A_{DIA,typ}$ is a typical value of the final diacetyl concentration, used as a scaling constant.

 Minimising the fermentation time. The reduction of the processing time is desirable from an economic point of view. No target value was given for the fermentation time. The optimisation algorithm was free to select the shortest possible fermentation time compatible with the other requirements:

$$
206 \t\t Q_t = W_t \frac{t_f}{t_{\text{typ}}} \tag{30}
$$

207 where t_f is the final fermentation time and t_{typ} is a typical value for the final time, used for scaling.

 Smoothing out the temperature and top pressure profiles. Technologically, a smooth plant operation is usually desired. Smooth temperature profile requires a smaller peak cooling power. A smooth top pressure 210 profile avoids discontinuities in the $CO₂$ outflow, facilitating its recovery. Numerically, penalising zigzagging profiles favours robust convergence to a global optimum. The smoothing objective was expressed as a penalty of the curvature (second time derivative) of the temperature and top pressure profiles. For numerical computations, the profiles were sampled at equally spaced time points. The scaling factors were chosen to make the curvature criteria invariant with respect to the total fermentation time and to the range of the variables:

215
$$
Q_{\theta} = W_{\theta} \frac{1}{n_{t} - 2} \sum_{k=2}^{n_{t} - 1} \left(\frac{\theta_{k+1} - 2\theta_{k} + \theta_{k-1}}{\theta_{\text{max}} - \theta_{\text{min}}} \right)^{2}
$$
(31)

216
$$
Q_p = W_p \frac{1}{n_t - 2} \sum_{k=2}^{n_t - 1} \left(\frac{p_{k+1} - 2p_k + p_{k-1}}{p_{\text{max}} - p_{\text{atm}}} \right)^2
$$
(32)

Here n_t is the total number of sampling points, θ_k and p_k are wort temperature and total headspace pressure at the 218 sampling point number *k*.

219 **Overall optimisation criterion**. The overall optimisation criterion is the sum of the nine partial criteria. It 220 reflects the best possible trade-off between generally conflicting optimisation goals:

$$
221 \tQ = \sum_{i \in \{ETA, ETX, IAA, IAL, PHL, DIA, t, \theta, p\}} (33)
$$

222 *Technological constraints*

 The operating conditions (temperature, top pressure and initial yeast concentration) were limited to the model validity range, which is the range of the operating conditions of the experimental design that provided data for model identification and validation. Additionally, the initial top pressure must equal the atmospheric pressure p_{atm} , since it can be increased only by the $CO₂$ produced by fermentation:

$$
227 \qquad \theta(t) \in [\theta_{\min}, \ \theta_{\max}], \qquad \qquad t \in [0, \ t_f] \tag{34}
$$

$$
228 \t p(t) \in [p_{atm}, p_{max}], \t t \in [0, t_f]
$$
\n(35)

$$
p(\theta) = p_{atm} \tag{36}
$$

$$
230 \t X_{\theta} \in \begin{bmatrix} X_{\theta \min}, & X_{\theta \max} \end{bmatrix} \tag{37}
$$

231 On most fermentation tanks, the temperature is controlled using a cooling jacket. The associated control variable, 232 namely the cooling rate Φ_{β} can only be positive since no heating device usually exists. Its amplitude is limited 233 by the maximum available cooling power or the maximum possible heat transfer rate:

234
$$
\boldsymbol{\Phi}_{\theta}(t) \in [0, \boldsymbol{\Phi}_{\theta \max}], \qquad t \in [0, t_f]
$$
 (38)

235 The top pressure is generally controlled by a valve. The corresponding control variable is the outflow gas rate ^Φ*^g* 236 which is also positive and limited by the valve diameter:

237
$$
\boldsymbol{\Phi}_g(t) \in \begin{bmatrix} 0, & \boldsymbol{\Phi}_{g \text{max}} \end{bmatrix}
$$
, $t \in \begin{bmatrix} 0, & t_f \end{bmatrix}$ (39)

- The alcoholic fermentation is considered finished when the residual fermentable sugar concentration reaches a
- specified value, corresponding to the so-called wort "attenuation" limit:

$$
240 \t S(t_f) = S_{res} \t (40)
$$

This constraint defines the fermentation end time t_f . The residual concentration S_{res} is generally known for a given wort and yeast strain.

- Numerical values of the optimal control problem parameters are reported in Table 3. The target aroma
- concentrations considered in this work are given in Table 4. The aroma profiles of the five sorts of beer listed in
- Table 4 are discussed in the "Results and discussion" section below.

Dynamic optimisation algorithm

- The considered optimal control problem is a dynamic one, since the unknown control variables are functions of time. The optimisation criterion contains linear and quadratic terms in the state variables. The constraints are linear in the state and control variables, except the top pressure constraint, which involves products of state 250 variables $(C_d(t)\theta(t)$ and $C_d(t)\theta(t)$). However, the main nonlinearity comes from the dynamic model of the fermentation process.
- Taking into account these characteristics of the control problem, a variant of the sequential quadratic programming (SQP) algorithm was used for solving it [15], coupled with a collocation (time discretisation) technique [16] and a safeguarded line search based on a "confidence region" method [17]. The main steps of the algorithm were:
- Step 0. **Initialisation**. A non-optimal but admissible (with respect to the constraints) solution was determined before entering the main algorithm. The initial fermentable sugar concentration is determined from the desired 258 ethanol concentration in the finished beer (E_{trg}) :

$$
259 \tS(0) = S_{res} + \frac{Y_{S/C}}{Y_{E/C}} E_{trg}
$$
\t(41)

 Random but admissible set-point profiles were generated for temperature and top pressure. A random admissible value was generated for the initial yeast concentration. The model equations were solved using simple proportional controllers to follow the set-points:

$$
263 \qquad \Phi_{\theta}(t) = K_{\rho\theta}(\theta(t) - \theta_{set}(t)) \tag{42}
$$

$$
264 \qquad \Phi_g(t) = K_{pp} \left(p(t) - p_{set}(t) \right) \tag{43}
$$

The fermentation end time (t_f) was determined as the moment when the fermentable sugar concentration fell below the specified limit (*Sres*).The resulting control variables (cooling rate and gas outflow rate) and the resulting fermentation end time were used as starting points for the optimisation.

268 Step 1. **Discretisation of the dynamic problem**. Let x be the vector of the $n_x = 12$ state variables and *u* the 269 vector of the $n_u = 2$ control variables:

270
$$
x = [C_p \ A_{ETA} \ A_{ETX} \ A_{LAA} \ A_{LAI} \ A_{PHL} \ A_{DLA} \ Y_{DLA/C} \ \theta \ C_d \ C_h \ C_a]^T
$$
 (44)
271 $u = [\Phi_\theta \ \Phi_g]^T$ (45)

- 272 The state and control variables were sampled at a finite number $(n_t = 49)$ of time points [16]. This reduced an infinite-dimensional problem to a finite-dimensional one, which could be handled by a numerical computer. The optimisation criterion, the constraints and the dynamic model equations were discretised on the same time grid.
- Step 2. **Construction of a local optimisation sub-problem**. The SQP method requires local approximation of the non-linear optimisation problem by a quadratic one with linear constraints, using limited Taylor series expansion. In the standard SQP method the quadratic approximation is made to the Lagrangean function. Its Hessian matrix includes curvature information on both the objective function and the constraints [15]. In this application the Hessian matrix of the objective function alone was used. The objective function being truly quadratic, its second order Taylor series expansion was exact. The discretised dynamic model equations were treated as additional constraints [15]. The limited Taylor series expansion being valid only locally, a limited search region around the current solution was established, using a set of "box" (min-max) constraints [17].
- Step 3. **Solution of the local optimisation sub-problem**. Commercial software, based on an "active constraint set" method was used to solve the quadratic optimisation problems with linear constraints [18]. This software did not exploit the special structure of the collocation equations and was unable to solve a problem of moderate size 286 $(2 + n_t(n_u + n_x) = 688)$. The number of variables was reduced to $2 + n_t n_u + n_x = 112$ by solving the linearized collocation equations explicitly. The reduction step involved a structurally well conditioned and sparse matrix, making the computation both fast and accurate.
- Step 4. **Convergence test**. The non-linear state equations were solved using the determined control variables. The value of the optimisation criterion was computed and the satisfaction of the constraints was checked. If the control variables, the state variables and the optimisation criterion were modified by less than a pre-specified amount, and if all constraints were satisfied, then calculations were halted. Else a new iteration was made, starting with step 1.
- 294 The algorithm was always run several (-10) times with various random initialisation in step 0. Robust convergence to the same optimum was observed. Occasionally, local optima in form of zigzagging control profiles were encountered. The control smoothing terms (Equations 31 and 32) were found very useful in avoiding these local optima. A detailed mathematical description of the algorithm is available from the authors on request.

Results and discussion

Alcoholic fermentation model validation

 The main measured and simulated variables in the alcoholic fermentation model are presented in Figure 1 for the run R5, not used for the identification of the model coefficients. The carbon dioxide evolution rate is predicted 303 reasonably well. The evolution rate is zero for the first 24 hours: during this time period the produced $CO₂$ is partly dissolved in the wort and partly accumulated in the headspace increasing the top pressure. The top pressure increases until the set-point is reached and then remains constant being controlled by the outflow valve. The partial air pressure remains constant until the outflow valve is opened and then decreases, the air being 307 evacuated from the tank along with the $CO₂$. The partial $CO₂$ pressure equals the total one when the air was 308 evacuated completely. The ethanol production, proportional to the cumulated CO₂ production, is predicted almost perfectly in this run. It appears from the model simulation that the dissolved $CO₂$ exceeds its saturation

- 310 limit by about 50% during the most active fermentation period (between 50 and 100 hours). Dissolved $CO₂$
- could not be measured directly in the considered experiments but this value is consistent with data found in the
- 312 literature [19] and strongly depends on the tank geometry through the parameter τ in Equations 6 and 9. Taking
- 313 into account the CO₂ super-saturation is the main evolution of the model considered in this paper compared to
- previous work [8]. It has a moderate impact on the aroma concentrations predicted by the model, particularly in
- time-varying operating conditions when the top pressure is decreased quickly.

Aroma production model validation

 Measured and simulated concentrations of the aroma compound considered in this work are reported in Figure 2 for the experimental run R5, not used for model derivation. The concentrations of five out of the six compounds are predicted reasonably well taking into account the scatter of the experimental data. The concentration of isoamyl acetate is significantly overestimated, however. This was already the case with previous models [11]. The measured final isoamyl acetate concentration in run R5 is one of the lowest among all runs, while the operating conditions have intermediate values. This is probably due to anomalous measurements for this particular experiment: in later runs, carried out in conditions similar to run R5, final isoamyl acetate concentrations close to 1.5 mg/L were obtained. The diacetyl is simultaneously produced and reduced during the fermentation run. The production rate decreases gradually and approaches zero at 50 hours. The diacetyl concentration reaches a maximum when the reduction rate equals the production rate and declines exponentially when the production rate becomes negligible.

Producing an existing beer at a lower cost

 This section illustrates the reduction of the production cost for an existing sort of beer without altering its aroma profile. By "aroma profile" is meant here the set of final concentrations of the two higher alcohols and three esters considered in this work. The target aroma concentrations were those measured at the end of the run R5, that is at the central point of the experimental design. The simulated experiment corresponding to run R5 is shown in Figure 3(A). With constant temperature (13°C), constant top pressure (450 mbar above the atmospheric pressure) and a typical initial yeast concentration (10 million cells/mL) the target aroma profile is reproduced well, illustrating the adequacy of the model. The predicted fermentation time is 121 hours, in agreement with the experimental value (Figures 1 and 2).

 The result of applying the optimisation algorithm to the same target aroma profile is illustrated in Figure 3(B). The final predicted aroma profile matches the target equally well but the fermentation time is reduced to 81 hours, that is by 33%. This is achieved by increasing the average fermentation temperature and the initial yeast concentration. The aromatic equilibrium of the final beer is maintained, however, using variable operating conditions (slightly decreasing temperature and uniformly increasing top pressure) by taking advantage of the time-varying fermentation rate and of the time-varying ester yields. The final diacetyl concentration was the same as in run R5 (0.5 mg/L): the highest diacetyl degradation rate due to higher temperature was compensated by a shorter fermentation time.

Producing new beer flavours with the same yeast strain and wort

 The measured aroma concentrations in four existing beer brands (coded B1 to B4, Table 4) were given as targets to the optimal control algorithm. The results for the beers B2 and B3 are shown in Figure 4. Beer B2 has a relatively high concentration of higher alcohols and a low ester concentration, while the opposite is true for beer B3. Figure 4 shows that both aroma profiles can be achieved using the same wort and yeast strain but different operating conditions: Beer B2 needs a high initial yeast concentration, a high fermentation temperature (except for the final phase) and a bell-shaped top pressure profile, while the aroma profile of beer B3 requires roughly the opposite: low initial yeast concentration, V-shaped temperature profile and a low top pressure for most of the fermentation time. Note that beer B2 is produced much faster than beer B3 (91 hours instead of 152) mainly because of the higher average temperature and of the higher initial yeast concentration. This illustrates the fact that the selected weights in the optimisation criterion (Table 3) favour the reproduction of the desired target aroma profile compared to time minimisation, which is only a secondary objective: for beer B3, the fermentation time was increased as much as needed to obtain the desired final aroma composition.

 It is expected, however, that not *all* aroma profiles can be reproduced with the given yeast strain and wort, even if the operating conditions are selected optimally in their admissible ranges. Two examples are given in Figure 5 for the aroma targets of beers B1 and B4 (Table 4). The difficulty of achieving the aroma profile B1 comes from the fact a low concentration of ethyl hexanoate can not be obtained simultaneously with a high concentration of 362 isoamyl acetate. The sensitivity of these compounds to the operating conditions (coefficients W_{i2} to W_{i8} , Table 2) are very similar, meaning that their concentrations can not be manipulated independently. The best possible solution found by the optimisation algorithm (Figure 5(A)) is a compromise where the final concentration of isoamyl acetate is lower than required and that of ethyl hexanoate is higher. The concentrations of ethyl acetate and isoamyl alcohol are reproduced correctly, however, and that of phenyl ethanol is in the tolerance domain. The situation is worse for the aroma profile B4 (Figure 5(B)) because the target concentration of ethyl hexanoate is even lower and the concentration of isoamyl acetate even higher. The only compound for which the target is achieved in this case is the isoamyl alcohol.

Final aroma profile robustness with respect to model uncertainty

 Due to the finite amount of experimental data available for model identification and to unavoidable measurement error the beer fermentation model parameters (Tables 2 and 3) can only be determined with limited accuracy. The uncertainty on the model parameters was expressed as a joint probability distribution and determined numerically (based on the so-called local Fisher information matrix) during the model fitting process [20]. The performance robustness was tested for the B2 beer by applying the pre-computed optimal operating conditions (temperature profile, top pressure profile and initial yeast concentration) to 20 models with parameters drawn at random from their respective probability distributions. The 95% confidence domain of the final aroma concentrations is shown in Figure 6(A). The confidence domain was determined as the range between the 2.5 and the 97.5 percentile of the calculated values. It appears from Figure 6 that the final aroma concentrations stay within their admissible tolerance limits despite model uncertainty. This is consistent with the selection of the tolerance limits that are always larger than the model accuracy (Equation 28).

Optimal control strategy robustness with respect to model uncertainty

 The sensitivity of the optimal control profile to the model uncertainty was investigated for the B3 beer by performing the optimal control calculation for 20 models with parameters drawn at random from their respective probability distributions. The 95% confidence domains of the final aroma concentrations and of the associated operating conditions (temperature, top pressure and initial yeast concentration) are shown in Figure 6(B). The confidence domain was determined as the range between the 2.5 and the 97.5 percentile of the calculated values. It appears that similar optimal control policies are obtained consistently despite model parameter variations. A V-shaped temperature profile, a bell-shaped top pressure profile and a low initial yeast concentration seem to be truly characteristic to the considered combination of yeast strain, wort composition and target aroma profile. The exact values of the operating conditions are not critical, however. For example, in a practical implementation, 392 $\pm 0.5^{\circ}$ C variations in the wort temperature, ± 200 mbar variations in the top pressure or ± 0.5 million cells/mL variations in the initial yeast concentration would not affect the final beer flavour significantly.

Conclusion

 A dynamic model of the beer fermentation process, including an alcoholic fermentation model and an aroma production model, was designed and validated. It is based on nine fermentation runs with operating conditions

397 (temperature, top pressure and initial yeast concentration) selected according to a $2³$ experimental design. The model was used for the dynamic optimal control of the beer fermentation process. A primary goal was to approach pre-specified final aroma targets (two higher alcohols and three esters simultaneously) as close as possible and a secondary objective was to reduce the fermentation time and the final concentration of an undesired aroma compound (diacetyl).

 An optimal time-varying control policy allowed the reduction of the fermentation time of an existing sort of beer, previously produced in constant operating conditions, by 33% while preserving the final aroma concentrations of the considered compounds. The optimal control strategy also allowed the reproduction of aroma profiles of two existing commercial beers (with different higher alcohols / esters ratios) by means of the operating conditions alone, i.e. using the same yeast strain and wort. Two other existing aroma profiles could not be reproduced satisfactorily (with the same yeast strain and wort) because some aroma compounds had similar sensitivities to the considered operating conditions and could not be manipulated independently.

 The sensitivity of the final aroma concentrations and of the optimal control policies with respect to modelling errors was explored numerically by generating random sets of model coefficients from their joint probability distributions estimated during the model identification phase. Achievable aroma profiles stayed within the tolerance limits and consistent optimal control profiles were obtained despite model uncertainty.

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Nomenclature

Para	Units	Value (95% confidence limits)	Source		
meter					
θ_{abs}	K	273.16	Unit conversion: degrees Celsius to Kelvin		
θ_{typ}	$\rm ^{\circ}C$	13	Central point of the experimental design		
τ	\boldsymbol{h}	3	Separate experiment		
γ	$L L^{-1}$	$\overline{4}$	Experimental protocol		
$C_{d,typ}$	$g L^{-1}$	2.76	Central point of the experimental design, Equation (8)		
K_v	h^{-1}	0.0446(0.04130.0481)	Maximum likelihood, runs R1-R4 and R6-R9		
$K_{\nu\theta}$	$\mathrm{^{\circ}C^{\text{-}1}}$	0.132(0.1270.137)	Maximum likelihood, runs R1-R4 and R6-R9		
$K_{\nu C}$	g^{-1} L	0.074(0.0340.113)	Maximum likelihood, runs R1-R4 and R6-R9		
K_C	$g L^{-1}$	0.0283	Tables supplied by professional brewer association		
$K_{C\theta}$	$\mathrm{^{\circ}C^{\text{-}1}}$	-0.0335	Tables supplied by professional brewer association		
K_E	$g L^{-1}$	25.5(22.229.2)	Maximum likelihood, runs R1-R4 and R6-R9		
K_S	$g L^{-1}$	3	Reference [11]		
K_X	$g L^{-1} (10^6)$	0.145(0.1220.172)	Maximum likelihood, runs R1-R4 and R6-R9		
	cells mL^{-1}) ⁻¹				
M_a	$g \text{ mol}^{-1}$	28.8	$20\% O_2 + 80\% N_2$ by volume		
M_c	g mol ⁻¹	44	Chemical formula of $CO2$		
p_{atm}	mbar	1013	760 mm Hg		
\boldsymbol{R}	mbar $L \text{ mol}^{-1}$	83.1	Perfect gas constant		
	K^{-1}				
$Y_{E/C}$	$g g^{-1}$	1.028(1.0131.043)	Reference ^[7]		
$Y_{S/C}$	$g g^{-1}$	1.884(1.8341.934)	Reference ^[7]		

Table 1. Numerical values of the parameters in the alcoholic fermentation model

Table 2. Numerical values of the parameters in the aroma production model

(In parenthesis: 95% confidence limits)

Source: maximum likelihood estimation based on data from runs R1-R4 and R6-R9

NA = not applicable

Parameter	Units	Value	Parameter	Units	Value	Parameter	Units	Value
θ_{max}	$^{\circ}\mathrm{C}$	16	$K_{p\theta}$	h^{-1}	5	W_{ETX}	none	1
θ_{min}	$^{\circ}\mathrm{C}$	10	K_{pp}	$g L^{-1} h^{-1} mbar^{-1}$	5×10^{-4}	W_{IAA}	none	1
$\Phi_{\theta max}$	$^{\circ}$ C h ⁻¹	0.5	n_t	none	49	W_{IAL}	none	1
$\Phi_{\rm g \, max}$	$g L^{-1} h^{-1}$	$\overline{2}$	p_{max}	mbar	1813	W_{PHL}	none	1
$A_{DIA, typ}$	$mg L^{-1}$	0.1	S_{res}	$g L^{-1}$	$\mathbf{1}$	W_p	none	2×10^{-4}
$A_{ETA,org}$	$mg L^{-1}$	2.5	t_{fmax}	$\boldsymbol{\text{h}}$	200	W_t	none	0.1
$A_{ETX,org}$	$mg L^{-1}$	0.02	t_{typ}	$\boldsymbol{\mathrm{h}}$	100	$X_{0 \ min}$	10^6 cells 5	
							mL^{-1}	
$A_{IA,org}$	$mg L^{-1}$	0.20	W_{θ}	none	2×10^{-4}	$X_{0 \text{ max}}$	10^6 cells 20	
							mL^{-1}	
$A_{IAL,org}$	$mg L^{-1}$	5	W_{DIA}	none	0.1			
$A_{PHL,org}$	$mg L^{-1}$	$\overline{4}$	W_{ETA}	none				

Table 3. Numerical values of the optimal control problem parameters.

Table 4. Target ethanol and aroma compound concentrations

^a Final ethanol and aroma concentrations measured in run R5

^b Ethanol and aroma concentrations measured in commercial beers

Figure legends

Figure 1. Alcoholic fermentation model validation. Data from run R5, not used for model identification. Measured values (o) and simulated values $(-)$.

Figure 2. Aroma production model validation. Data from run R5, not used for model identification. Measured values (o) and simulated values $(-)$.

Figure 3. Fermentation time reduction of an existing beer without changing the final aroma profile. (**A**) Constant operating conditions corresponding to run R5. (**B**) Time-varying operating condition determined by the dynamic optimisation algorithm. Top: aroma concentrations at the end of the alcoholic fermentation. Bottom: operating conditions for the alcoholic fermentation process. Target values (o), admissible range (⋅ ⋅ ⋅) at ± 2*Ai,tol* and predicted values $(-)$.

Figure 4. Different final aroma profiles obtained with the same yeast strain and wort. (**A**) Beer B2. (**B**) Beer B3. Top: aroma concentrations at the end of the alcoholic fermentation. Bottom: operating conditions for the alcoholic fermentation process. Target values (o), admissible range (···) ±2 $A_{i,tol}$ and predicted values (—).

Figure 5. Aroma profiles impossible to achieve with the considered yeast strain and wort. (**A**) Beer B1. (**B**) Beer B4. Top: aroma concentrations at the end of the alcoholic fermentation. Bottom: operating conditions for the alcoholic fermentation process. Target values (o), admissible range (⋅ ⋅ ⋅) at ±2 *A_{i,tol}* and predicted values (--).

Figure 6. Sensitivity of the final aroma concentration and of the optimal control strategy to modelling errors. (**A**) The optimal control policy for beer B2 was applied without change to 20 possible model parameter sets. (**B**) The optimal control policy for beer B3 was recomputed for 20 possible model parameter sets. Top: aroma concentrations at the end of the alcoholic fermentation. Bottom: operating conditions for the alcoholic fermentation process. Target values (o), admissible range (⋅ ⋅ ⋅) at ±2 *Ai,tol* and 95% confidence intervals for the predicted values (\blacksquare) .

Figure 1

Figure 2

Figure 3.

Figure 5.

